In silico Prediction of Protein–Protein Interaction Network Induced by Manganese II in Meyerozyma guilliermondii

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Recently, there has been an increasing interest in the use of yeast to produce biosorbent materials, because yeast is economical to use, adaptable to a variety of conditions, and amenable to morphological manipulations to yield better raw biomaterials. Previous studies from our laboratory have shown that Meyerozyma guilliermondii, a non-pathogenic haploid yeast (ascomycete), exhibits excellent biosorption capacity for Mn$^{2+}$, as demonstrated by kinetic analyses. Shotgun/bottom-up analyses of soluble fractions revealed a total of 1257 identified molecules, with 117 proteins expressed in the absence of Mn$^{2+}$ and 69 expressed only in the presence of Mn$^{2+}$. In this article, we describe the first in silico prediction and screening of protein–protein interactions (PPIs) in M. guilliermondii using experimental data from shotgun/bottom-up analyses. We also present the categorization of biological processes (BPs), molecular functions (MFs), and metabolic pathways of 71 proteins upregulated in the M. guilliermondii proteome in response to stress caused by an excess of Mn$^{2+}$ ions. Most of the annotated proteins were related to oxidation–reduction processes, metabolism, and response to oxidative stress. We identified seven functional enrichments and 42 metabolic pathways; most proteins belonged to pathways related to metabolic pathways (19 proteins) followed by the biosynthesis of secondary metabolites (10 proteins) in the presence of Mn$^{2+}$. Using our data, it is possible to infer that defense mechanisms minimize the impact of Mn$^{2+}$ via the expression of antioxidant proteins, thus allowing adjustment during the defense response. Previous studies have not considered protein interactions in this genus in a manner that permits comparisons. Consequently, the findings of the current study are innovative, highly relevant, and provide a description of interactive complexes and networks that yield insight into the cellular processes of M. guilliermondii. Collectively, our data will allow researchers to explore the biotechnological potential of M. guilliermondii in future bioremediation processes.

Keywords: Meyerozyma guilliermondii, manganese, bioremediation, proteome, protein–protein interactions, metabolic pathways
INTRODUCTION

Heavy metal pollution represents one of the most serious global environmental problems. Although methods exist to minimize the environmental impact caused by these elements, traditional decontamination methods are known to release toxic products into the environment. The development of new ecological strategies for the efficient treatment of contaminated water is therefore a key target for researchers. Such strategies should be based on biological removal, or bioremediation, which is safer and more economically viable than traditional methods (Azubuike et al., 2016; Bahafid et al., 2017; Amorim et al., 2018).

Manganese (Mn) is a heavy metal that is soluble in water. Under normal circumstances, Mn is essential for growth, development, and homeostasis; however, serious health problems can arise when excessive amounts of Mn accumulate in the body (Erikson et al., 2005; Johnson and Hallberg, 2005; Zhao et al., 2010; Kafaei et al., 2017; Kornblihit et al., 2018). In the state of Minas Gerais, Mn is naturally present in the soil and considered a natural constituent of the waters that drain from this region. However, Mn concentrations above the limits set by environmental resolutions (Conselho Nacional do Meio Ambiente-Conama, 2005, 2011; Instituto Mineiro de Gestão das Águas [IGAM], 2012, 2014) have been found in this region. It is generally thought that this increase in Mn concentration is related to activities in the mining and metallurgical sectors. Indeed, Mn has been detected in the effluents and drains of almost all Brazilian mining sites (U.S. Department of the Interior and U.S. Geological Survey, 2012; Departamento Nacional de Produção Mineral [DNPM], 2017). A more significant increase in the levels of Mn was detected after the rupture of a dam in the city of Mariana-MG in 2015; dam residues were shown to contain large amounts of manganese. More recently, another dam broke, resulting in the further release of waste into the environment (Departamento Nacional de Produção Mineral [DNPM], 2017).

Traditionally, the removal of Mn is accomplished by the addition of strong oxidizing agents such as chlorine (Cl₂), ozone (O₃), and potassium permanganate (KMnO₄); however, these agents generate toxic by-products. A potential alternative to these traditional processes is the use of biological technologies, such as bioremediation, that are more economically viable (Patil et al., 2016; Bahafid et al., 2017). The capacity of an organism for bioremediation, along with its tolerance or adaptability, is defined by molecular responses that are unique to each organism. Molecular techniques, such as proteomics and interactomics, allow us to identify and understand the metabolic potential of such organisms (Pandey and Mann, 2000; Mallick and Kuster, 2010). However, little is known about the molecular and biochemical mechanisms responsible for the accumulation and degradation of metals by yeast. In particular, the genetic and molecular basis of tolerance to heavy metals remains poorly understood. Consequently, there is an urgent need to identify the functional genes involved in tolerance and detoxification, and to elucidate the expression of key genes and proteins involved in the accumulation of heavy metals, along with their molecular interactions (Pieper and Reineke, 2000; Wase and Forster, 2003; Dixit et al., 2015; Bahafid et al., 2017).

Interactomics allows researchers to identify all types of molecular interactions, such as protein–protein interactions (PPIs), and examine metabolic networks that are regulated under certain physiological conditions. Analysis of PPIs has been employed in previous research to predict the function of uncharacterized proteins based on the fact that interacting proteins have similar functions (Feng et al., 2014; Cafarelli et al., 2017). Proteins participate in complex networks of biochemical interactions, at both functional and physical levels, between DNA, RNA, other proteins, lipids, and small metabolites. Developing an understanding of cellular mechanisms requires the analysis of interactomic networks using data acquired from proteomics. Oxidative stress caused by excessive levels of metals triggers an increase in gene expression, metabolic factors, and signaling proteins related to the stress response. By specifically analyzing PPIs, it is possible to identify all protein interactions that are upregulated or downregulated in certain environmental scenarios, thus providing clues for the identification of groups of molecules activated during the process of metal resistance; such research could be vital in developing biotechnological processes to facilitate the removal of Mn (Goll and Uetz, 2006; Instituto Mineiro de Gestão das Águas [IGAM], 2012; Feng et al., 2014; López et al., 2016; Patil et al., 2016; Cafarelli et al., 2017; Wang et al., 2018; Ruas et al., 2019).

Meyerozyma guilliermondii is a haploid, osmotolerant, non-pathogenic yeast that is able to utilize various carbon sources for survival and growth. Some strains exhibit physiological characteristics that can be used for the bioremediation of environmental metal contamination (Kaszycki et al., 2004; Butler et al., 2009; Kurtzman et al., 2011; Defosse et al., 2014). In a previous publication, we demonstrated that this strain of yeast may be employed in bioremediation processes to remove Mn²⁺ ions from water contaminated by mines. We confirmed this biosorptive removal capability using kinetic equations (Amorim et al., 2018), and recently described the first soluble proteome for this genus and this particular species (Ruas et al., 2019).

In the present study, we aimed to create the first PPI map for M. guilliermondii and describe the interaction network of proteins that are upregulated under stress conditions caused by an excess of Mn²⁺ ions. The present finding provides new insights into the underlying molecular and biochemical mechanisms, particularly the expression of upregulated proteins and the activation of metabolic pathways, under stress conditions caused by the excessive accumulation of Mn. We consider that the present findings related to the molecular mechanisms and kinetics of Mn removal will facilitate the development of new strategies for the application of this Mn²⁺-resistant yeast in biotechnological processes (Schwikowski et al., 2000; McDermott et al., 2005; Feng et al., 2014).

MATERIALS AND METHODS

This study involved a dataset that was previously published by our group (Ruas et al., 2019) and deposited in the ProteomeXchange Consortium (Vizcaino et al., 2014) via the jPOST (Okuda et al., 2017) partner repository;
this dataset can be accessed using the following dataset identifiers: <PXD010049> and <PXD PXD010050>. This dataset was obtained from M. guilliermondii yeast grown, for 7 days, in Yeast Peptone Dextrose (YPD) medium containing 2% (w/v) glucose, 1% (w/v) yeast extract, and 2% (w/v) peptone (pH 7.4) and under two different conditions: the absence and presence 0.91 mM of Mn$^{2+}$ ions.

**LC-MS/MS Analysis**

All experiments were conducted in biological duplicates and techniques triplicates. Proteins were extracted, dosed, and subsequently processed separately; 20 μg aliquots of protein fractions were separated by electrophoresis short runs in order to concentrate the proteins and eliminate contaminants of low molecular mass such as secondary metabolites. Analysis involved 500 ng of peptides obtained from the enzymatic digestion of biological replicates that were separated by gradient elution using an Acclaim PepMap100 C18 Nano-Trap column (75 μm id × 2 cm, 3 μm, 100 Å, Thermo Scientific) with an Acclaim PepMap100 C18 capillary column (75 μm id × 15 cm, 2 μm, 100 Å, Thermo Scientific). Ultrahigh performance liquid chromatography (UHPLC) separation of previously digested tryptic peptides was performed using the Dionex UltiMate® 3000 UHPLC system (Thermo Scientific, Bremen). The spectra of biological duplicates were determined using a Q-Exactive™ mass spectrometry instrument (Thermo Scientific). The instrument was operated at 1.9 kV in the positive mode with a resolution of 70,000 at 300–1750 m/z, a maximum injection time of 120 ms and a target value for automatic gain control (AGC) of 1 and 6 ions. We also used a 2 m/z window, fragmented by high energy collisional dissociation (HCD) with a normalized collision energy of 28–30 V. MS/MS spectra were obtained at a resolution of 17,500, a maximum injection time of 60 ms, and an AGC target value of 5 and 5 ions. After each MS/MS, we applied a subsequent dynamic deletion of 30 s (Ruas et al., 2019).

**Processing and Storage of Data in a Repository**

Files containing raw data were subsequently searched by quantitative proteomics software package MaxQuant®, version 1.5.2.8 (Cox and Mann, 2008). Identified proteins were compared with prediction data in the Meyerozyma guilliermondii UniProt database, which contains 5,520 predicted sequences1 (accessed March 2017). In order to validate groups of proteins identified as being upregulated in the presence of an excess of Mn$^{2+}$ ions in the biological duplicates, we used PEAKS software. We also used MaxQuant® software to compare proteomic similarity and to determine unique proteins, as described previously (Kumar and Mann, 2009).

**Protein Interaction Datasets**

Analyses were carried out in *silico* using a range of bioinformatics tools, as described below. Protein annotation was performed using UniProt Knowledgebase (UniProtKB) and proteins identified as “uncharacterized” were subjected to a Basic Local Alignment Search Tool (BLAST) search using the UniProtKB and BLASTp databases and considering a minimum sequence identity of 60%. Analyses of biological process (BP) and molecular function (MF) through Blast2GO Software2 via Gene Ontology (GO)3 and KEGG. (Ogata et al., 1999) (Kyoto Encyclopedia of Genes and Genomes4). The PSORT program was used to predict subcellular protein localization5, and biochemical pathways involving differentially expressed proteins were analyzed using STRING6 (version 11.0), as described previously (Feng et al., 2014).

**RESULTS**

We successfully identified 1257 proteins in the total proteome of *M. guilliermondii*. The heatmap shown in Figure 1 demonstrates that proteins were downregulated and upregulated under the two different physiological conditions (the absence and presence of Mn$^{2+}$). Figure 2 and Supplementary Table S1 present for the first time the *in silico* prediction of the identified proteins, metabolic pathways, and the total proteome enrichment of the genus *Meyerozyma* sp. Results were obtained from the STRING database.

In total, 71 proteins were upregulated in the presence of an excess of Mn$^{2+}$ ions. These positively regulated proteins in the presence of Mn$^{2+}$ ions were classified according to the BPs in which they are involved (Figure 3) and their MFs (Figure 4). Most upregulated proteins relate to BPs associated with the cellular response to stress (32%), oxidation–reduction processes (11%), carbohydrate metabolic processes (5%), response to oxidative stress (4%), and phosphorylation (4%) (Figure 3). The most recurrent MFs in the differential proteome were transferase activity (19%), coenzyme binding (16%), oxidoreductase activity (16%), relative effect on the CH–OH group of donors (19%), NAD as an acceptor for the metal binding (14%), and ATP binding (12%) (Figure 4).

The first PPI networks of *M. guilliermondii* total and differential proteome *in silico* were performed using the STRING, KEGG, and UniProt database (Figure 2), as well as the first demonstration of stress-rich functional enrichment analysis caused by Mn$^{2+}$ (Figure 5). Seven functional enrichment analyses of the differentially caused by stress caused by Mn$^{2+}$ were found: metabolic pathways, glutathione metabolism, biosynthesis of secondary metabolites, biosynthesis of antibiotics, biosynthesis of amino acids, propanoate metabolism, and longevity regulating pathway—multiple species ($p = 5.29 \times 10^{-6}$) (Figure 5 and Supplementary Table S2). Of these enrichments,
Figure 1 | Heatmap representation of peptide quantification for the total proteome of *M. guilliermondii* based on the peak intensity of identified peptides from bottom-up proteomic data in the absence and presence of Mn$^{2+}$.
the most representative pathway was metabolic pathways, followed by biosynthesis of secondary metabolites, antibiotics, and amino acids.

More specific analysis of the 71 upregulated proteins showed that they were associated with 42 metabolic pathways. The pathways associated with the largest number of proteins were metabolic pathways (19 proteins) followed by biosynthesis of secondary metabolites (10 proteins), biosynthesis of antibiotics (nine proteins), biosynthesis of amino acids (six proteins), glutathione metabolism (four proteins), and carbon metabolism (four proteins) (Table 2).

**DISCUSSION**

Previous studies from our group showed that *M. guilliermondii* possesses the ability to carry out manganese remediation via an efficient biosorption process. Due to these characteristics,
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FIGURE 3 | Description of the biological processes associated with upregulated proteins in the presence of Mn$^{2+}$. Sequences were analyzed by Blast2GO and Gene Ontology (GO).

this yeast can be effectively used for bioremediation of metal contamination in contaminated water. Results from the present study provide novel insights into molecular and biochemical mechanisms that allow this species of yeast to tolerate excessive concentrations of Mn$^{2+}$ (Eccles, 1999; Shakya et al., 2015; Amorim et al., 2018).

In a previous study, we isolated *M. guilliermondii* and demonstrated that this species of yeast removes Mn$^{2+}$ ions efficiently. Indeed, our experiments indicated that 100% of Mn$^{2+}$ ions were removed from water in only 6 days (0.91 mM Mn$^{2+}$); the removal kinetics were also indicative of efficient bioremediation (Amorim et al., 2018). Species that are capable of tolerating excessive concentrations of heavy metals may possess specific biochemical and molecular characteristics such as the expression of specific proteins and metabolic pathways that confer the ability to survive under heavy metal stress. We already know that proteins, their activities, and functions suffer interference from Mn$^{2+}$, presented in the differential expression of proteins in Figure 1, and that the organism described herein uses such mechanisms to overcome stress conditions to the metal, thus generate a response that allows it to survive under conditions of excess metal. This yeast has mechanisms of gene alteration, differential protein expression, and activation of metabolic pathways that allow to correct and protect cellular and genetic integrity (Stohs and Bagchi, 1995; Kotrba and Ruml, 2000; Chen and Shi, 2002; Ercal et al., 2005; Wysocki and Tamás, 2010; Ruas et al., 2019).

In our previous paper, we reported that the majority of proteins in the proteome of *M. guilliermondii* are involved in BPs related to genetic and metabolic mechanisms (Ruas et al., 2019). The presence of excess Mn$^{2+}$ ions caused stress to yeast, as expected, since most proteins that were upregulated in the presence of Mn$^{2+}$ ions are mainly involved in BPs associated with cellular stress response (32%) (Figure 3). BPs associated with reduction and oxidation processes (11%) and oxidative stress response (4%) (Figure 3) also stood out. In general, oxidation–reduction processes act as catalysts for BPs that require electron transfer, such as metabolism and signaling processes that govern gene regulation and expression in certain situations (Prabhulkar et al., 2012). The proteins related to these processes are the majority in the differential proteome (Table 1).

The key MFs of this expressed proteome are protein binding, transerase activity, and oxidoreductase activity (Figure 4). These results corroborate the number of BPs and demonstrate the capacity of the studied species under stressful conditions.
triggered by Mn$^{2+}$. It has become clear that this species expresses proteins that activate detoxification regulatory mechanisms which allowed it to tolerate considered Mn$^{2+}$ concentrations (Hohmann and Mager, 2003; Wysocki and Tamás, 2010; Ruas et al., 2019).

Stressful conditions in yeast triggered by the presence of Mn$^{2+}$ upregulated the expression of 71 proteins. Some of the upregulated proteins were directly related to protein biosynthesis, thus confirming the need for the synthesis of protective proteins in response to overexposure to the tested ions. Others, most of them, are related to oxidation–reduction, metabolism, and oxidative stress response processes. These results confirm that M. guilliermondii has molecular and biochemical mechanisms to overcome Mn$^{2+}$ toxicity. The effects of metal toxicity on metabolism and homeostasis in M. guilliermondii, particularly in response to Mn exposure, are not known, therefore, the need for this report.

We know that the exposure of cells to excessive concentrations of heavy metals causes an increase in the cellular levels of reactive oxygen species (ROS) that impair and interfere with various cellular components, such as nucleic acids and proteins in yeasts. Heavy metals can cause damage in yeast cells via increased levels of ROS, which lead to the abnormal regulation of redox-sensitive pathways, and through oxidative changes to essential biomolecules (Fridovich, 1998; Shanmuganathan et al., 2004; Ercal et al., 2005; Schrader and Fahimi, 2006; Bonekamp et al., 2009; Pan et al., 2017).

Thus, M. guilliermondii activates these oxidative control mechanisms. Most proteins found in the proteome are related to protein oxidoreductase activity, oxidative stress response, and mitochondrial activity and peroxisomes. This reduction activity is confirmed by the protein interaction network in Figure 5 and the pathway enrichment functional analysis (Table 2). We found upregulated acyl-CoA dehydrogenase (A5DBX1) that catalyzes the early stage of the mitochondrial fatty acid β-oxidation cycle (Colin and Kimt, 1995). Isocitrate dehydrogenase [NADP] peroxisomal (A5DHQ4), which acts on the krebs cycle by providing NADPH for maintenance of the reduced glutathione and peroxiroxin systems (Table 1 and Supplementary Table S2) (Smolková and Ježek, 2012). The Peroxiredoxin (Prx) (A5DE26) is involved in redox homeostasis and oxidative stress (Bonekamp et al., 2009; Murray et al., 2011; Nelson et al., 2011; Soito et al., 2011) and glutathione S-transferase (A5DBE9, A5DJ25) plays essential role in antioxidant activity (Stohs and Bagchi, 1995).

In studies with Saccharomyces cerevisiae, it was observed that metals and metalloids, such as cadmium (Cd), arsenic III (As III), among other metals, were neutralized. Neutralization triggers a mechanism of glutathione depletion (GSH), usually associated with metal toxicity. GSH depletion influenced the redox environment and impaired the activities of GSH dependent enzymes such as glutathione peroxidases, glutathione S-transferases, and glutaredoxins, affecting many cellular processes (Stohs and Bagchi, 1995). M. guilliermondii...
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FIGURE 5 | Network interactions of upregulated proteins in M. guilliermondii under excess Mn$^{2+}$ stress, using tools and databases that can predict STRING protein function (PPI enrichment p-value: $5.29 \times 10^{-6}$).

We identified using TabPath metabolic pathways of glycolysis/gluconeogenesis (UDP-glucose 4-epimerase—A5DRG2) and peroxisome (isocitrate dehydrogenase [NADP] peroxisomal—A5DHQ4; superoxide dismutase—A5DA67) (Table 2). Previous research has shown that heavy metal toxicity increases the activity of enzymes such as glucose-6-phosphate dehydrogenase and peroxidases. Heavy metals can also affect plasma membrane lipids by modifying membrane properties linked to permeability, fluidity, the modulation of membrane-bound ATPase activities, and interfere with proper protein folding (Niki, 2009; Jan et al., 2015).

ATPase activity was identified as one of the MFs that was significantly represented in the proteome of M. guilliermondii (10%) (Figure 4). Damage to cell membrane ATPase activity and increased lipid peroxidation are indication of increased...
### TABLE 1 | Differentially expressed proteins in the presence of Mn$^{2+}$ ions.

| Accession | Gene       | Description                                           | e-Value   | Sim mean |
|-----------|------------|-------------------------------------------------------|-----------|----------|
| A5DBF8    | PGU_00613  | Voltage-dependent anion channel protein 2             | 0.0       | 92.15    |
| A5DA0     | PGU_01951  | 40S ribosomal protein S20                              | 6.51E-73  | 94.34    |
| A5DQK9    | PGU_05660  | Uroporphyrinogen-III synthase                         | 0.0       | 71.25    |
| A5DCX8    | PGU_01133  | ATP synthase subunit alpha, mitochondrial              | 0.0       | 96.25    |
| A5DHA7    | PGU_02658  | 60S ribosomal protein L9-B                             | 3.89E-130 | 94.64    |
| A5DDN3    | PGU_01384  | ATP synthase subunit beta, mitochondrial               | 0.0       | 97.92    |
| A5DCC1    | PGU_00636  | Actin-related protein 2/3 complex subunit 1           | 0.0       | 84.71    |
| A5DG8     | PGU_02329  | Golgi vesicle protein                                  | 0.0       | 81.12    |
| A5DFK1    | PGU_02052  | Ornithine carbamoyltransferase                         | 0.0       | 85.34    |
| A5DQ9     | PGU_05410  | Eukaryotic translation initiation factor 2 subunit gamma| 0.0       | 94.51    |
| A5DR3     | PGU_05884  | Karyopherin functions in nuclear transport of protein | 2.02E-135 | 81.69    |
| A5DDQ0    | PGU_01401  | Eukaryotic translation initiation factor 1A           | 4.88E-70  | 98.69    |
| A5DEB0    | PGU_00604  | Glutathione S-transferase                             | 2.27E-167 | 77.01    |
| A5DBX1    | PGU_00776  | Acyl-CoA dehydrogenase                                 | 0.0       | 87.75    |
| A5DG40    | PGU_01205  | Isocitrate dehydrogenase [NADP] peroxisomal           | 0.0       | 94.86    |
| A5DQBS    | PGU_05464  | NAD(P)-binding protein                                 | 0.0       | 77.64    |
| A5DDQ5    | PGU_00805  | Inosine-5’-monophosphate dehydrogenase                 | 0.0       | 94.52    |
| A5DA9     | PGU_00384  | Ribosomal protein L4                                   | 1.18E-169 | 78.33    |
| A5DQS7    | PGU_05628  | Aspartate transcarbamoylase                            | 0.0       | 91.57    |
| A5DAY0    | PGU_00435  | Actin/actin-like protein                               | 0.0       | 94.62    |
| A5DGJ8    | PGU_02529  | Predicted protein                                      | 0.0       | 66.99    |
| A5DK41    | PGU_03642  | Opaque-phase-specific protein OP4                      | 2.64E-156 | 72.65    |
| A5DOQ8    | PGU_05649  | Putative vacuolar sorting protein                      | 4.53E-160 | 77.31    |
| A5DB30    | PGU_00485  | Eukaryotic translation initiation factor 2 subunit alpha| 0.0       | 93.07    |
| A5DFH3    | PGU_02024  | aarF domain-containing kinase                         | 0.0       | 83.19    |
| A5DBW2    | PGU_00767  | Glycoside hydrolase family 81 protein                  | 0.0       | 82.84    |
| A5DNL9    | PGU_03470  | P-loop containing nucleoside triphosphate hydrolase protein| 0.0     | 92.9     |
| A5DEV2    | PGU_01813  | U1 small nuclear ribonucleoprotein                     | 6.07E-129 | 67.83    |
| A5DAG2    | PGU_056863 | UDP-glucose 4-epimerase                                | 0.0       | 81.85    |
| A5DQ86    | PGU_05439  | Fe-S cluster assembly protein DRE2                     | 0.0       | 73.44    |
| A5DEL5    | PGU_01716  | D-3-phosphoglycerate dehydrogenase 1                  | 0.0       | 94.77    |
| A5DQ39    | PGU_05390  | Secreted protein                                       | 0.0       | 68.28    |
| A5DQ70    | PGU_05558  | Pyridoxal kinase                                       | 0.0       | 73.51    |
| A5DNY7    | PGU_04988  | Regulation of G-protein function                       | 0.0       | 73.48    |
| A5DBX4    | PGU_00779  | Actin/actin-like protein                               | 0.0       | 97.87    |
| A5DEQ2    | PGU_01753  | Predicted protein                                      | 0.0       | 69.69    |
| A5DB15    | PGU_00470  | Elongation factor 3                                    | 0.0       | 95.71    |
| A5DDM8    | PGU_01379  | Protein MET17                                         | 0.0       | 91.27    |
| A5DHJ1    | PGU_02742  | 3,4-Dihydroxy-2-butanone 4-phosphate synthase          | 4.37E-143 | 90.82    |
| A5DTK9    | PGU_03881  | WD40 repeat-like protein                               | 8.77E-128 | 79.25    |
| A5DFT8    | PGU_02139  | Predicted protein                                      | 2.92E-112 | 80.51    |
| A5DKE0    | PGU_03741  | Glyoxylate reductase 1                                 | 0.0       | 83.28    |
| A5DE26    | PGU_01527  | Peroxiredoxin HYR1                                    | 1.91E-122 | 88.86    |
| A5DE13    | PGU_01514  | CIP1 protein                                          | 0.0       | 70.44    |
| A5DNZ9    | PGU_05600  | Phosphoacetylglucosamine mutase                       | 0.0       | 81.25    |
| A5DR5     | PGU_05886  | Protein MNN4                                         | 0.0       | 74.75    |
| A5DG27    | PGU_05623  | Nicotinamidase                                        | 3.75E-160 | 74.04    |
| A5DR7     | PGU_05898  | Probable xylose and arabinose reductase                | 0.0       | 78.81    |
| A5DHN1    | PGU_02782  | HIT-like protein                                      | 3.69E-125 | 80.15    |
| A5DK4    | PGU_03905  | Cysteine hydrolase                                    | 1.02E-130 | 73.44    |
| A5DE42    | PGU_01543  | Lysine acetyltransferase (lysine N(6)-acetyltransferase) (LAT)| 0.0   | 67.62    |
| A5DO4     | PGU_02495  | FAD/NAD(P)-binding domain-containing protein          | 0.0       | 73.63    |
| A5DMR7    | PGU_04568  | Inositol-pentakisphosphate 2-kinase                    | 0.0       | 78.13    |

(Continued)
oxidative stress caused by metals (Niki, 2009). Adaptive response to lipid peroxidation requires the transcriptional regulation of antioxidant genes, including those encoding enzymes related to glutathione synthesis and metabolism (four proteins: A5DHQ4, A5D25, A5DJ25, A5DBE9) and two glutathione S-transferases (A5DJ25 and A5DBE9) that play a key role in redox homeostasis, cell signaling, and detoxification (Figure 5 and Table 2) (Niki, 2009; Zhang et al., 2013; Jan et al., 2015; Özaslan et al., 2017). The functional enrichment analysis results show the glutathione metabolism pathway as one of the highlights, just as PPI networks show four proteins related to the same functions including two glutathione S-transferase (A5DBE9 and A5DJ25), peroxiredoxin (A5DE26), and peroxisomal isocitrate dehydrogenase [NADP] (A5DHQ4).

Certain metabolic pathways are activated allowing stress to be overcome by detoxification or metal elimination mechanisms. Some induce the precipitation of heavy metals, including metal carbonates, phosphates, and sulfates. Other pathways allow stress to be overcome by mechanisms of detoxification and the elimination of metals. Species that have the ability to adapt or tolerate metals are of very important and could potentially be utilized in important biotechnological processes (Kotrba and Ruml, 2000; Ercal et al., 2005).

Meyerozyma guilliermondii upregulated proteins are involved in 42 metabolic pathways, according to TabPath analysis. Functional enrichment analysis demonstrated seven-way enrichment and that proteins have a network of interactions between proteins that confirm the results of analyzes of biological functions and processes. This pathway relates to several metabolic reactions (catabolism and anabolism) that are necessary for carrying out all processes for body maintenance, including glycolysis, pentose pathway, electron transported chain, and others. Ten identified proteins belong to biosynthesis of secondary metabolites, which in yeast involves many proteins and protein complexes that allow it to respond to various environmental stimuli such as stress to metals, just as M. guilliermondii was exposed (Fox and Howlett, 2008). Other pathways that have been observed as defense reactions and detoxification mechanisms in yeast due to oxidative damage induced by heavy metals were biosynthesis of amino acids, propanoate, and longevity regulating pathway (Figure 5 and Supplementary Table S2).

Species that possess the ability to remove heavy metals may exhibit adaptive characteristics such as specific metabolic pathways for the remediation of metals. Certain metabolic pathways mediate the precipitation of heavy metals into carbonates, phosphates, and metal sulfates with potential biotechnological applications (Kotrba and Ruml, 2000). In addition, some species can produce metal-binding compounds, such as bio-pigments (flavins, flavonoids, polyhydroxy anthraquinones, and tannin), which contribute to the stress response induced by exposure to metals by acting as the first barrier against a variety of metals. These binders are predominant in metal-binding sites, when dead biomass is used as a biosorbent material (Kotrba and Ruml, 2000; Farooq et al., 2010). Indeed, a recent study involving M. guilliermondii reported that dead biomass removed manganese ions more efficiently than live biomass (Amorim et al., 2018), suggesting that metal binding compounds may play a role in this process.

In addition to producing metal binding compounds, biosynthesis of secondary metabolites, amino acids, antibiotics, activation of metabolic pathways also contribute to success in a such stressful situation. Consequently, these factors are of vital importance for the decontamination of metals and thus present meaning for possible biotechnological applications.
TABLE 2 | Metabolic pathways of proteins related to manganese tolerance in differential proteomes.

| Pathway                          | Accession |
|----------------------------------|-----------|
| Metabolic pathways               | A5DFK1    |
|                                  | A5DM8     |
|                                  | A5DX8     |
|                                  | A5DK7     |
| Biosynthesis of secondary metabolites | A5DFK1    |
|                                  | A5DM8     |
|                                  | A5DK9     |
| Biosynthesis of antibiotics      | A5DFK1    |
|                                  | A5DM8     |
| Biosynthesis of amino acids      | A5DFK1    |
|                                  | A5DM8     |
| Glutathione metabolism           | A5DQK9    |
|                                  | A5DQS7    |
| Carbon metabolism                | A5DQK7    |
|                                  | A5DQS7    |
| Propanoate metabolism            | A5DFK3    |
| Ribosome                         | A5DFA0    |
| RNA transport                    | A5DQS2    |
| Longevity regulating pathway     | A5DQS2    |
| Oxidative phosphorylation        | A5DX8     |
| Arginine biosynthesis            | A5DFK3    |
| Purine metabolism                | A5DFA0    |
| Glycine, serine, and threonine metabolism | A5DQK9    |
| Amino sugar and nucleotide sugar metabolism | A5DQK9    |
| Pyruvate metabolism              | A5DQK9    |
| MAPK signaling pathway           | A5DQK9    |
| Protein processing in endoplasmic reticulum | A5DQK9    |
| Peroxisome                       | A5DQK9    |
| Glycolysis/gluconeogenesis       | A5DQK9    |
| Citrate cycle (TCA cycle)        | A5DQK9    |
| Pentose and glucuronate interconversions | A5DQK9    |
| Galactose metabolism             | A5DQK9    |
| Fatty acid degradation           | A5DQK9    |
| Alanine, aspartate, and glutamate metabolism | A5DQK9    |
| Cysteine and methionine metabolism | A5DQK9    |
| Valine, leucine, and isoleucine degradation | A5DQK9    |
| Beta-alanine metabolism          | A5DQK9    |
| Glycerolipid metabolism          | A5DQK9    |
| Inositol phosphate metabolism    | A5DQK9    |
| Arachidonic acid metabolism      | A5DQK9    |
| Methane metabolism               | A5DQK9    |
| Riboflavin metabolism            | A5DQK9    |
| Vitamin B6 metabolism            | A5DQK9    |
| Nicotinate and nicotinamide metabolism | A5DQK9    |
| Porphyrin and chlorophyll metabolism | A5DQK9    |
| Sulfur metabolism                | A5DQK9    |
| 2-Oxocarboxylic acid metabolism  | A5DQK9    |
| Fatty acid metabolism            | A5DQK9    |
| mRNA surveillance pathway        | A5DQK9    |
| Phosphatidylinositol signaling system | A5DQK9    |
| Endocytosis                      | A5DQK9    |

(Holan and Volesky, 1995; Kotrba and Ruml, 2000; Ahemad and Kibret, 2001; Farooq et al., 2010). The possibility of altering the properties of living species used in heavy metal remediation and construction of chimeric organisms that have desirable characteristics using genetic engineering are now under study in many laboratories.
CONCLUSION

Heavy metal removal processes by bioremediation represent an economical alternative to physicochemical decontamination methods. In this paper, we describe the molecular interactions of a yeast strain, *M. guilliermondii*, with proven resistance to Mn$^{2+}$ ions. Collectively, our findings allow us to identify the molecular interactions associated with proteins that are upregulated in the presence of Mn$^{2+}$ and describe the metabolic pathways that play a role in the expression of these proteins. The majority of the upregulated proteins were related to oxidoreductase activity. Our findings indicate that *M. guilliermondii* is able to tolerate excessive Mn$^{2+}$ concentrations via the expression of antioxidant proteins, thus creating a more effective ecotoxicological defense response. Our data corroborate the findings of previous studies in that this species has significant potential for application in biotechnological processes used to remove Mn$^{2+}$ contamination from water. The results presented here, in addition to contributing to scientific knowledge about the species, may help to develop future methods for monitoring potentially hazardous and toxic manganese-contaminated areas or waters (rivers, dams). It may further contribute to the establishment of future standardized tests using the species *M. guilliermondii* or even employ as a suitable eukaryotic organism model for bioremediation processes.

DATA AVAILABILITY STATEMENT

Our mass spectrometry proteomics data have been deposited in the ProteomeXchange Consortium [49] via the jPOST [50] partner repository with the following dataset identifiers: <PXD010049> and <PXD010050>.

AUTHOR CONTRIBUTIONS

FR designed and performed Mn$^{2+}$ experiments with *M. guilliermondii*, processed the data, carried out bioinformatic analyses and statistical analysis, deposited the proteomic dataset in public repositories, and wrote and reviewed the manuscript. RG-S designed and supervised the study and reviewed the manuscript. Both authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.00236/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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