Supplemental Note and Figures for Yu et al. “Landscape of gene expression variation of natural isolates of *Cryptococcus neoformans* in response to biologically relevant stresses”

Supplemental Notes

- Lineage specific DEG on mating-type loci (MAT)
- Functional characterization of genes in the K-means clusters
- Differential expression between clinical and environmental pairs
- Supplementary references

Figure S1. Correlation matrix of all RNA-Seq samples.

Figure S2. Multi-dimensional scaling plots across lineages for each condition.

Figure S3. Number of lineage-specific DEGs identified in each condition and in the common set.

Figure S4. Percentage of GO terms for lineage DEGs in each condition.

Figure S5. Venn diagrams depicting overlap of DEGs between CSF and other conditions.
Supplemental Notes

Lineage-specific differentially expressed genes (DEGs) on mating-type loci (MAT)

To help understand if differences in gene expression between lineages correspond to gene presence/absence, we evaluated the presence of orthologs in Bt85 (VNBI) for each gene identified based on mapping to H99; 32 out of the 94 DEGs from the H99 comparison do not match an ortholog gene/group in Bt85 and were up-regulated in VNI strain. Most of these genes were located in five large regions, including Chromosome 1: 12789 - 19876 (CNAG_00002, CNAG_00003, and CNAG_00004), Chromosome 4: 1058232 - 1063107 (CNAG_07831, CNAG_05325, CNAG_05326), Chromosome 5: 7089 – 29713 (CNAG_06876, CNAG_06875, CNAG_06874, CNAG_06871, CNAG_06870, CNAG_06869), Chromosome 7: 384496 - 394513 (CNAG_06649, CNAG_06650, CNAG_06651, CNAG_06652, CNAG_06653), and Chromosome 8: 16125 - 32410 (CNAG_03086, CNAG_03087, CNAG_07707, CNAG_03090, CNAG_03092) (Table S5c). H99 and Bt85 have different mating types, MATα and MATa, respectively. As expected, three DEGs (CNAG_06812, CNAG_07015, and CNAG_07411) in the mating locus located on Chromosome 5 [1] were more highly expressed in H99 and down-regulated in Bt85 (Table S5b).

Functional characterization of genes in the K-means clusters

(1) Genes up-regulated during in vitro growth

The polysaccharide capsule is one of the main virulence factors of C. neoformans and plays a predominant role in the interaction with the host. This capsule, composed of glucuronoxylomannan and galactoxylomannan, not only provides resistance to stressful conditions but also has strong immunomodulatory properties to promote evasion and survival within the host [2, 3]. Many genes in k-means cluster 1 and cluster 5 are associated with cell wall and/or capsule formation (Table S8a and S8e). For instance, several genes are involved in the G protein-cAMP-PKA signaling pathway, which has been shown to promote mating and the production of virulence factors in C. neoformans [4, 5]. The PKAI gene
encodes the major cyclic AMP (cAMP)-dependent protein kinase catalytic subunit, and the \textit{PKRI} gene encodes the protein kinase A (PKA) regulatory subunit. The low-affinity phosphodiesterase, Pde1, is involved in the PKA1-dependent regulation and modulates the cAMP level [6]. The \textit{GPR4} gene encodes a G protein-coupled receptor that functions in sensing the amino acid methionine to activate cAMP-PKA signaling [7]. An adenylyl cyclase-associated protein encoded by the \textit{ACA1} gene and the Ga subunit encoded by the \textit{GPA1} gene cooperate to activate adenylyl cyclase, which plays a critical role in regulating melanin and capsule production, cell fusion, and filamentous growth [8]. Additional genes identified in this cluster include genes associated with acidification of vesicular compartments, growth at elevated temperatures, cell wall components, stress response, and capsule and melanin production. The vesicular (H+)-ATPase proton pump, encoded by the \textit{VPH1} gene, is important for the acidification of vesicular compartments, and disruption of \textit{VPH1} resulted in defective expression of polysaccharide capsule formation, growth at 37°C, and laccase and urease production [9]. The \textit{FPD1}, \textit{MP98}, and \textit{CDA3} genes encode polysaccharide deacetylases, including chitin deacetylases. Chitin deacetylases deacetylate chitin, converting it to chitosan. Both chitin and chitosan are associated with the cell wall integrity of \textit{C. neoformans} growing vegetatively [10] and are critical factors for the pathogenesis of \textit{C. neoformans} [11–13]. In addition, \textit{CSR1} encodes a chitin synthase regulator [13]. The Hex1 exochitinase, a hexosaminidase, degrades chitinoligomeric substrates to maintain cell wall plasticity [14]. Ssk1 and Ssk2 are involved in the stress-activated Pbs2-Hog1 mitogen-activated protein kinase (MAPK) signaling pathway, which governs several cellular events, including stress responses, sexual reproduction, drug sensitivity, and virulence [15]. The disruption of the \textit{SSK2} gene in H99 enhanced capsule and melanin biosynthesis, mating efficiency, and hypersensitivity to stresses [16]. \textit{KRE5} is a β-1,6-glucan synthesis-related gene that is vital for maintaining cell morphology and cell wall integrity and affects sensitivity to high temperature [17]. Other capsule or cell wall related genes are \textit{EGCrP1} [18], \textit{RIM20} [19], \textit{WSP1} [20], \textit{SCH9} [21], \textit{GIB2}, \textit{SSD1}, and \textit{ISP3} [10]. The \textit{MET3} gene encodes ATP sulfurylase and plays an important role in the methionine/cysteine biosynthetic pathway. Mutation of \textit{MET3} has effects on capsule and melanin formation, as well as on
growth rate, thermotolerance, and virulence [22]. \textit{LAC2} encodes a laccase important for melanin production and pathogenesis via \( \text{Ga protein/cyclic AMP (cAMP) pathway} \) [23]. Five other genes have also been found that when deleted are associated with melanization defects, including \textit{RTF1}, \textit{NUP75}, \textit{CSN4}, \textit{UBP14}, and \textit{LAG1} [24].

Other potentially important virulence genes induced \textit{in vitro} were also identified. \textit{URE1} encodes the urease apoenzyme, which has been shown to have a role in virulence, especially during brain invasion [25, 26]. Ure1 regulates the expression of transporters and catabolic enzymes in response to the availability and type of nitrogen source in the environment [27]. Glr1 is a glutathione reductase which responds to nitric oxide stress [28]. The thioredoxin proteins, Trx1 and Trx2 are important for nitrosative stress resistance and virulence of \textit{C. neoformans} [29]. In addition, we also found that more than 20 genes that encode V-type and F-type H\(^{+}\)-transporting ATPases, which are ATP-driven enzymes using the energy of ATP hydrolysis to pump protons across membranes, including plasma and mitochondrial inner membranes. In fungi, preservation of a proton gradient across membranes is critical for various cellular metabolisms, including receptor-mediated endocytosis, intracellular membrane trafficking, pro-hormone processing, protein degradation, uptake of small molecules, and storage and detoxification of metabolites and ions [30]. \textit{ATM1} encodes a mitochondrial ABC transporter that contributes to preservation of the homeostasis of cytosolic-nuclear Fe-S proteins during Cu stress [31].

(2) Genes up-regulated during \textit{in vivo} growth

2 (a) Inositol-related proteins

We found 21 genes in the \textit{in vivo} expression cluster are involved in inositol transport or catabolism, including myo-inositol transporters, inositol 2-dehydrogenases, inositol oxygenases, inositol/phosphatidylinositol kinases and phosphoinositide phospholipases (\textbf{Table S8c}). The human brain contains abundant inositol, which is a major osmolyte and plays a role in regulating normal neurological
responses [32]. If glucose availability is low in the brain, *Cryptococcus* may utilize inositol as a carbon source. Inositol in the brain has also been reported as a stimulator for promoting *Cryptococcus* penetration of the blood-brain barrier [33] and is required for virulence and mating [34, 35].

A large number of genes involved in DNA recombination, DNA repair, and DNA replication are also present in this cluster. These include DNA mismatch repair proteins (Pms2, Msh5, and Msh6), DNA polymerases, DNA primase, minichromosome maintenance proteins, DNA topoisomerases, exonucleases, endonucleases, and ATP-dependent DNA helicases. These results may reflect genome plasticity during the course of disease development within that host [36] and is already well-described with azole heteroresistance *in vivo* [2].

2 (b) Cell wall and capsule formation

A total of 120 known virulence-associated genes were also identified within this *in vivo* cluster. At least 47 of 120 virulence-associated genes identified in this cluster are involved in capsule formation or cell wall integrity (Table S8c). *UGE1* and *UGD1*, which encode UDP-glucose epimerase and UDP-glucose dehydrogenase, respectively, are necessary for galactoxylomannan biosynthesis and virulence in *C. neoformans* [3, 37]. In addition, several proteins, Skn1 and Kre62 - Kre64, are involved in the synthesis of polysaccharide β-1,6-glucan, which is a major component of the cell wall of *C. neoformans* [17]. Genes encoding capsule-associated proteins (Cap4; Cap5; Cap6; Cap10; Cap60; Cas32; Cas41; Cas42; Cas9; Gmt2) involved in capsule biosynthesis were highly expressed [38–40]. The chitin synthases, Chs1, Chs3 – Chs8, the chitin synthase regulators (Csr3), and the endochitinase, Chi21 are involved in chitin metabolic and biosynthetic processes [14, 41]. We also found two signaling components, a high affinity cAMP phosphodiesterase (Pde2) and a PKA catalytic subunit (Pka2), related to the cAMP/PKA pathway which promotes mating and the production of melanin and capsule. Pde2 may be regulated by Pka2, which contributes to the regulation of virulence. The deletion of the *PDE2* gene, however, has been shown to have modest effects on mating or melanin or capsule production [6].
2 (c) Cation transporters

Two active sodium (Na+) efflux transporters, Ena1 and Nha1, cooperatively maintain the intracellular pH and homeostasis of several toxic cations, such as Na+, Li+, and K+ [42] (Table S8c). Nha1, a Na+/H+ antiporter, is required for short-term adaptation to high salt shock. ENA1 encodes a fungus-specific sodium or potassium P-type ATPase (Na(+)/ATPase), which is sensitive to alkaline pH conditions and essential for virulence in murine and rabbit models [43, 44]. Together, both genes play critical roles in cation homeostasis, pH regulation, membrane potential, and virulence in *C. neoformans* [42]. PMC1 encodes a calcium transporter and provides tolerance to high Ca2+ concentrations. Pmc1 has been described as a vacuolar calcium ATPase, which enables *C. neoformans* to penetrate the central nervous system and is critical for both the progression of pulmonary infection and brain colonization in a murine infection model [45, 46].

2 (d) Stress responses

Genes involved in *C. neoformans* stress responses were also up-regulated *in vivo*. Sre1, a sterol regulatory element-binding protein (SREBP), functions in an oxygen-sensing pathway. This gene is involved in oxygen sensing and sterol homeostasis and is important for adaptation and growth of *C. neoformans* in the brain, which has suboptimal conditions of oxygen concentration and nutrition for fungal growth [47, 48] (Table S8c). In addition, three genes encoding MAP kinases, Bck1 (MAPKKK), Mkk2 (MAPKK), and the presumed terminal kinase Mpk1 (MAPK) comprising the Bck1-Mkk2-Mpk1 signaling pathway, which is activated in response to thermal stress, were up-regulated [49, 50]. We also found a pH-response transcription factor, Rim101, in this cluster. The Rim signal transduction pathway is well known to sense and respond to changes in host pH. Rim101, an alkaline-responsive transcription factor, is required for the proper formation of the protective polysaccharide capsule as well as growth under stressful host conditions, such as elevated cation concentrations and alkaline pH [19]. Lastly, the known virulence-associated gene,
CNA1, encoding the calcineurin A catalytic subunit, confers sensitivity to temperature, CO₂, and pH [51, 52].

2 (e) Other potential virulence factors

LIV5 encodes a protein homologous to one required for fungal pathogenesis and the deletion of this gene resulted had reduced virulence in a murine inhalation model [24].

(3) Genes up-regulated in CSF conditions, down-regulated in MP conditions

Many genes specifically up-regulated in CSF are tRNA/rRNA methyltransferases (Enzyme Commission (EC) number: 2.1.1.-) and tRNA pseudouridine synthases (EC 5.4.99.-). RNA methyltransferases catalyze the transfer of a methyl group on rRNA/tRNA and tRNA pseudouridine synthases catalyze the transition of uridine to pseudouridine on tRNA. In eukaryotes, these epigenetic modifications of rRNA nucleotides can alter ribosome function, which could allow modulation of translation and the cellular proteome in response to specific intracellular or environmental stresses [53, 54]. Six genes induced in CSF are linked to capsule formation, including NSTX, CPS1, CRG1, GMT1, PBX1, and SET302 [10, 38, 55]. Also in this cluster are genes in the heat shock protein associated pathways. We identified several heat-shock related protein family members (Hsp60/Hsp70/Hsp90/Hsp104), including CNAG_00305, CNAG_06443, CNAG_05199, CNAG_03899, CNAG_01727, CNAG_01750, CNAG_03459, CNAG_05619, CNAG_07558, CNAG_00100, CNAG_06150, CNAG_06208, CNAG_07347, CNAG_01568, CNAG_03459, CNAG_05252, CNAG_03944, and CNAG_06106. Native misfolding of protein and protein aggregation caused by protein denaturation in pH, oxidative, osmotic, or high temperature stresses may lead to the loss of protein functions and may promote apoptosis suggesting that the CSF is a stressful environment [56].
Differential expression between clinical and environmental pairs

In comparing clinical and environmental isolates from VNI, we found 68 genes up-regulated and 137 genes down-regulated during MP interaction. Some of these genes were associated with fungal defense in macrophages and the survival in the host. For instance, we found the up-regulated gene, quinone oxidoreductase, which is a homolog of S. cerevisiae ZTA1 gene (NADPH:quinone reductase). In C. albicans, quinone reductases promote resistance to oxidative stress and appear to be required for virulence in mouse infections studies [57]. The SIT1 siderophore iron transporter (CNAG_00815), is up-regulated in clinical isolates, and is associated with iron acquisition, melanin formation, and cAMP signaling in C. neoformans [58]. In Candida spp., the SIT1 gene is required for epithelial invasion and penetration and for its siderophore-mediated iron acquisition that is critical for enhancing yeast survival to the microbicidal activities of macrophages [59, 60].

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**Figure S1. Correlation matrix of all RNA-Seq samples.** Sample correlation matrix displays the overall similarity in expression profile for each sample from different conditions. Red color intensity indicates increasing sample correlation, whereas blue color intensity indicates decreasing sample correlation. The dendrogram clustering on the Y axis indicates the overall similarity of all samples. The samples are named with the strain name, the condition, and the experimental replicate.
Figure S2. Multi-dimensional scaling plots across lineages for each condition. These MDS plots show the clear separation among two lineages, VNB and VNI (including sublineages VNIa and VNIb), for each condition, including VNIa clinical (filled red triangle) and environmental (open red triangle) isolates; VNIb clinical (filled red square) and environmental (open red square) isolates; VNB clinical (filled blue circle) and environmental (open blue circle) isolates.
Figure S3. Number of lineage-specific differential expressed genes (DEGs) identified in each condition and in the common set. The significant differential expression genes are identified between VNI and VNB lineages for each condition by exact test at FDR $P$-value $< 0.05$. The red bar represents the number of DEGs that are up-regulated in VNI strains, compared to VNB strains. By contrast, the blue bar represents the number of VNI down-regulated DEGs. The common category is the intersection of lineage specific DEGs across all five conditions (94 genes).
Figure S4. Percentage of GO terms for lineage DEGs in each condition. The GO SLIM terms of DEGs were mapped by the FungiDB database. Genes without GO SLIM annotations were classified in two categories; Hypothetical Proteins do not have assigned names or annotated functions in FungiDb and Others have an assigned name or function although not an assigned GO SLIM term.
Figure S5. Venn diagrams depicting the overlap of DEGs between CSF and other conditions. The plots show the number of DEGs between CSF and each of the other conditions and the overlaps between these DEG sets. A. Up-regulated DEGs in CSF condition compared to the other conditions. B. Down-regulated DEGs in CSF condition compared to the other conditions. The total number of up-regulated (lower left panel) and down-regulated CSF (lower right panel) compared to the other conditions varies depending on the condition.