1 Puf4 Mediates Post-transcriptional Regulation of Caspofungin Resistance in

2 Cryptococcus neoformans

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24 Abstract

25 Echinocandins have been on the market for 20 years, yet they are the newest class of 26 antifungal drugs. The human fungal pathogen Cryptococcus neoformans is intrinsically resistant 27 to the echinocandin antifungal drug caspofungin, which targets the β -1.3-glucan synthase 28 encoded by the *FKS1*. Analysis of a *C. neoformans puf4*∆ mutant, lacking the pumilio/FBF RNA 29 binding protein family member Puf4, revealed exacerbated caspofungin resistance. In contrast, 30 overexpression of PUF4 resulted in caspofungin sensitivity. The FKS1 mRNA contains three 31 Puf4-binding elements (PBEs) in its 5' untranslated region. Puf4 binds with specificity to this 32 region of the *FKS1*. The *FKS1* mRNA was destabilized in the *puf4* Δ mutant, and the abundance 33 of the FKS1 mRNA was reduced compared to wild type, suggesting that Puf4 is a positive 34 regulator FKS1 mRNA stability. In addition to FKS1, the abundance of additional cell wall 35 biosynthesis genes, including chitin synthases (CHS3, CHS4, CHS6) and deacetylases (CDA1, 36 CDA2, CDA3) as well as a β -1,6-glucan synthase gene (SKN1) was regulated by Puf4 during a 37 caspofungin time course. The use of fluorescent dyes to quantify cell wall components revealed that the $puf4\Delta$ mutant had increased chitin content, suggesting a cell wall composition that is 38 39 less reliant on β -1,3-glucan. Overall, our findings suggest a mechanism by which caspofungin 40 resistance, and more broadly, cell wall biogenesis, is regulated post-transcriptionally by Puf4. 41

42 Importance

Cryptococcus neoformans is an environmental fungus that causes pulmonary and central
nervous system infections. It is also responsible for 15% of AIDS-related deaths. A major
contributor to the high morbidity and mortality statistics is the lack of safe and effective
antifungal therapies, especially in resource-poor settings. Yet, antifungal drug development has
stalled in the pharmaceutical industry. Therefore, it is of importance to understand the

48 mechanism by which *C. neoformans* is resistant to caspofungin in order to design adjunctive
49 therapies to potentiate its activity toward this important pathogen.

50

51 Introduction

52 Invasive deep mycoses primarily impact immunocompromised populations causing high rates of 53 morbidity and mortality (1, 2). The pathogenic fungus Cryptococcus neoformans is the causative 54 agent of fatal meningitis most often in patients with defects in cell mediated immunity, including 55 transplant recipients and those living with HIV/AIDS (3–6). C. neoformans is responsible of the 56 15% of AIDS-related deaths (6). Treatment of cryptococcosis is difficult and therapeutic options 57 are limited. Even the best combination treatment using Amphotericin B with 5-fluorocytosine (5-58 FC) is not well tolerated, and 5-FC is often unavailable in resource-poor areas (7). Some of the 59 largest clinal challenges to invasive fungal infections are poor efficacy of the drugs, emerging 60 resistance issues, narrow selection and availability of antifungals especially in the areas where 61 needed the most (8, 9).

62

63 Another antifungal agent, fluconazole, is largely ineffective as the first-line therapy since 64 it lacks effective fungicidal activity against C. neoformans in vivo even at high concentrations 65 and presents resistance issues (10, 11). Echinocandins (such as caspofungin, micafungin and 66 anidulafungin) are the latest class of antifungal drugs approved by the Food and Drug 67 Administration (FDA) that target cell wall biosynthesis. The echinocandins are ineffective 68 against C. neoformans due to a high level of intrinsic resistance (12). Echinocandins 69 specifically target the β -1,3-glucan synthase encoded by *FKS1*, and *C. neoformans* Fks1 is 70 sensitive to inhibition by the echinocandins, suggesting that the mechanism of intrinsic 71 resistance is not related to biochemical differences in the target itself (13). In other pathogenic 72 fungi, such as Aspergillus and Candida species, resistance to caspofungin is manifested due to 73 mutations in FKS1, but this is not observed in Cryptococcus (14, 15). Another mechanism that is 74 discussed regarding caspofungin resistance in pathogenic fungi involves the cell wall 75 remodeling and integrity pathways (14–16). It has been shown that increased cell wall chitin 76 content contributes to caspofungin resistance (17, 18). Additionally, a defect in intracellular drug 77 concentration maintenance due to drug influx and efflux imbalance has been proposed to be a 78 potential mechanism to explain the intrinsic resistance phenomenon (19, 20). Discovering and 79 targeting the regulatory components behind the pathways involved in the intrinsic resistance 80 may result in a combination therapy that potentiates the antifungal activity of caspofungin 81 toward C. neoformans.

82

83 Calcineurin signaling plays a distinct role in intrinsic caspofungin resistance (21, 22). 84 Caspofungin synergizes with the calcineurin inhibitors FK506 and Cyclosporin A (23). Both the 85 A and B subunits of Calcineurin regulate tolerance to caspofungin (19). Crz1, the transcription 86 factor that is activated through dephosphorylation by calcineurin, translocates to the nucleus 87 following treatment with caspofungin, yet the $crz1\Delta$ mutant exhibits wild type sensitivity to the 88 drug, suggesting caspofungin resistance is Crz1-independent. (19). Calcineurin functions at the 89 intersection of multiple signaling pathways and interacts with a diversity of proteins involved in 90 calcium signaling, RNA processing, protein synthesis and vesicular trafficking among others 91 (24, 25). Since caspofungin resistance is calcineurin-dependent, yet Crz1-independent, RNA 92 processing targets of calcineurin that may be involved in resistance to caspofungin through 93 post-transcriptional modulation of gene expression are especially of interest. Tight control of not 94 only transcriptional but also post-transcriptional gene regulatory networks in drug resistance 95 phenotypes is underappreciated in fungal pathogens, yet may represent targets for adjunctive 96 therapies to improve the efficacy of drugs.

97

98 One of the targets of calcineurin involved in RNA processing is a pumilio-domain and 99 FBF (PUF) domain-containing RNA-binding protein - Puf4 (24). *C. neoformans* Puf4 is

100	homologous to the Saccharomyces cerevisiae paralogs Puf4 and Mpt5 (26). Our previous work
101	demonstrated that C. neoformans Puf4 recognizes the Mpt5 binding element in its mRNA
102	targets (27). It has been hypothesized that Puf4 may play a role in the regulation of cell wall
103	biosynthesis since the <i>puf4</i> Δ mutant is resistant to lysing enzymes, is temperature sensitive at
104	both 37°C and 39°C, and is sensitive to Congo red (28). Our group has previously shown that
105	Puf4 regulates endoplasmic reticulum (ER)-stress through controlling the splicing of a major ER-
106	stress related transcript, HXL1, and plays a role in the unfolded protein response pathway of C.
107	<i>neoformans.</i> Puf4 also contributes to virulence, as the <i>puf4</i> Δ mutant has attenuated virulence
108	compared to wild type in an intravenous murine competition model of cryptococcosis (27).
109	
110	In this study we demonstrated that Puf4 contributes to the intrinsic caspofungin
111	resistance of C. neoformans through post-transcriptional regulation of the mRNA encoding the
112	drug target, Fks1. Puf4 also regulated a number of the cell wall biosynthesis-related mRNAs.
113	This regulation is primarily at the level of mRNA stability and has functional consequences in
114	maintaining cell wall composition.
115	
116	<u>Results</u>
117	The <i>puf4</i> ∆ mutant is resistant to caspofungin.
118	Our previous work has implicated the pumilio family RNA binding protein Puf4 in the regulation of
119	ER stress in C. neoformans (27). Puf4 is an effector of calcineurin signaling, a pathway known to
120	regulate thermotolerance and cell integrity (24, 28). Given the connection of Puf4 to cell integrity
121	signaling, we assessed the sensitivity of $puf4\Delta$ mutant to the cell wall perturbing drug caspofungin.
122	We measured caspofungin sensitivity by spot plate analyses, and found that the $puf4\Delta$ mutant is
123	resistant to caspofungin above the published minimum inhibitory concentration of 16 μ g/ml (Fig.
124	1A). This result suggests that Puf4 is a negative regulator of caspofungin resistance in C.

overexpressed with a FLAG-tag (3-copies determined by Southern blot analysis), the hyper-126 127 resistance was not only suppressed, but the strain was more sensitive to caspofungin. Single-128 copy FLAG-Puf4 complementation of the $puf4\Delta$ mutant restored a wild type resistance phenotype 129 (Fig. 1A). In addition to the spot plate analyses, growth analysis using liquid cultures in a kinetic 130 plate reader assay in the presence of 8 µg/ml caspofungin showed similar trends to the 131 phenotypes observed in spot plates (Fig. 1B). Growth was not permissible in the presence of 16 132 µg/ml caspofungin in liquid culture for both wild type and the mutant cells suggesting that the 133 action of caspofungin may be influenced by the environmental constraints in different culture 134 conditions (data not shown).

135

136 **Puf4 directly binds and stabilizes the FKS1 mRNA.**

137 The target of caspofungin, β -1,3-glucan synthase, is encoded by the *FKS1* gene. We searched 138 the FKS1 mRNA sequence for a potential Puf4 binding element. Our previous work suggests 139 that the Puf4 binding element in C. neoformans is homologous to that of S. cerevisiae Mpt5, 140 including the invariant initiating UGUGA followed by a four-nucleotide spacer sequence and 141 terminating UA. We found that the FKS1 mRNA contains two Puf4 binding elements in its 5' 142 UTR (Fig 2A). To determine if Puf4 can directly interact with its consensus element(s) in the 143 5'UTR of the FKS1 mRNA we performed electrophoretic mobility shift assay (EMSA). We 144 synthesized a 50-base long fluorescently labeled (TYE705 infrared label) RNA oligonucleotide 145 that span the FKS1 5' UTR containing the Puf4-binding elements (Table 2). Incubation of the 146 fluorescent oligonucleotide with the wild type cell lysate resulted a shift that was competed with 147 the unlabeled oligonucleotide. A mutant competitor, in which the UGUANNNNUA motif was 148 replaced by adenines, was unable to compete for binding to the wild type fluorescent 149 oligonucleotide. This suggests that Puf4 binds to the FKS1 mRNA through sequence-specific 150 recognition of the Puf4 binding elements in the 5' UTR (Fig. 2B). Incubation of the fluorescent 151 oligonucleotide with the puf4 Δ mutant lysate has the absence a shift that was observed with the 152 Puf4-FLAG cell lysate (Fig 2B, arrow). Because, we detected an interaction between Puf4 and the 5'UTR sequence of FKS1, we went on to investigate if loss of Puf4 would alter the 153 154 abundance or stability of FKS1 mRNA. FKS1 mRNA abundance in mid-log grown cultures of 155 $puf4\Delta$ cells were decreased 20% compared to wild type grown in parallel (Fig. 2C). PUF 156 proteins are known mRNA stability regulators, and so we asked if the reduction in FKS1 mRNA 157 in the $puf4\Delta$ mutant was due to destabilization. We performed an mRNA stability time-course 158 following transcription shut-off and found that the FKS1 mRNA was destabilized in the absence 159 of Puf4 compared to the wild type (Fig 2D). 160

161 **Puf4 protein expression is decreased following caspofungin treatment.**

162 Because Puf4 is a regulator of the FKS1 mRNA, we asked if inhibition of Fks1 by caspofungin 163 treatment would alter the abundance of PUF4 mRNA or Puf4 protein. First, we performed a caspofungin time course in which we treated wild type cultures grown to mid-log phase with 16 164 165 µg/ml caspofungin for 60 minutes and collected cells every 15 minutes. We guantified the PUF4 166 transcript levels, and found out that the PUF4 transcript levels are unchanged throughout time-167 course (Fig 3A). Then we utilized the Puf4-FLAG strain to investigate if Puf4 protein levels are 168 changed following treatment with caspofungin using immunoblotting. We found that Puf4 protein 169 levels are drastically decreased following treatment with caspofungin (Fig 3B and 3C). Since the 170 absence of Puf4 causes a hyper-resistant phenotype (as shown in Fig. 1A), we speculate that 171 the downregulation of Puf4 protein levels may be a contributing event to the intrinsic resistance.

172

173Puf4-dependent *FKS1* regulation correlates with reduced β-1,3-glucan staining in the *puf4* Δ 174mutant.

We next investigated the *FKS1* mRNA levels following treatment with caspofungin to determine if inhibition of the enzyme activates a feedback mechanism to upregulate expression of *FKS1*.,

177 The *FKS1* mRNA abundance is upregulated in wild type cells at 45 minutes post-caspofungin 178 treatment compared to 30 minutes (Fig 4A). Conversely, the *puf4* Δ mutant cells have 179 decreased levels of *FKS1* at 30 minutes post-caspofungin treatment, which is restored to the 180 basal levels at later time points (Fig. 4A). These different trends in transcript abundance 181 following caspofungin treatment may suggest that the cell wall β -1,3-glucan levels may differ 182 between wild type and the *puf4* Δ cells.

183

184 To investigate the functional consequences of the trends we observed at the transcript 185 level, we utilized aniline blue staining to investigate the β -1,3-glucan levels in the cell wall of the 186 wild type and the *puf4* Δ cells. Aniline blue specifically binds to β -1,3-glucan (29). Following 187 arowth to mid-log phase, both strains were treated with caspofungin for 60 minutes and stained 188 with aniline blue. Fluorescence microscopy revealed that aniline blue staining mainly localized 189 to the cell wall in the mid-logarithmic stage cells. In addition to the cell wall staining pattern, 190 staining pattern of the caspofungin treated cells also contained an intracellular punctate pattern 191 (Fig. 4B). Quantification of the microscopy images showed that the puf4 Δ cells had 30% less β -192 1.3-glucan compared to the wild type cells both at the mid-logarithmic stage and following 193 treatment with caspofungin (Fig. 4C). We concluded that the post-transcriptional regulation of 194 the *FKS1* mRNA by Puf4 directly affects cellular β -1,3-glucan levels.

195

196 **Deletion of** *PUF4* **leads to dysregulation of cell wall biosynthesis genes.**

A recent study revealed that multiple cell wall genes are influenced by caspofungin treatment
(19). We assessed the same panel of mRNAs for the presence of a putative Puf4 binding
element, and found that several caspofungin-sensitive genes contain Puf4 binding elements
(Table 1). These include genes that encode chitin synthases, chitin deacetylases, α-glucan and
β-glucan synthases.

202

We next asked if the caspofungin-responsiveness of these genes was dependent on 203 204 Puf4. Following growth to the mid-log stage, we challenged both wild type and the $puf4\Delta$ cells 205 with 16 µg/ml caspofungin over a 60-minute time course and investigated the changes in the transcript abundance of genes involved in cell wall biosynthesis. We found that CHS3 (Chitin 206 207 Synthase 3) is downregulated following caspofungin treatment in the puf4 Δ cells compared to 208 wild type (Fig 5A). Conversely, CHS4 and CHS6 are found to be upregulated in the $puf4\Delta$ 209 mutant compared to wild type (Fig 5B and 5C). Elevated cell wall chitin content is shown to 210 reduce susceptibility to caspofungin in Candida species (18). Therefore, upregulation of chitin 211 synthase genes in the puf4 Δ mutant may also contribute to the resistance phenotype in 212 Cryptococcus. The synthesis of chitosan from chitin is catalyzed by the chitin deacetylases and 213 chitosan is necessary for the integrity of the cell wall (30). During the caspofungin time course, 214 we found out that CDA1 (chitin deacetylase 1) is upregulated in the puf4 Δ mutant. On the 215 contrary, CDA2 and CDA3 were found to be downregulated in the puf4^Δ mutant (Fig 5D-F). The 216 CDA3 is the only chitin deacetylase gene that contain a PBE (Table 1). Of note, Cda1 is the 217 major chitin deacetylase and the only chitin deacetylase that is necessary for virulence (31). 218

219 Lastly, we looked at the regulation of α -glucan and β -glucan genes during the 220 caspofungin time-course. We have found that AGS1 (α -glucan synthase 1) was present at a 221 slightly higher abundance in the $puf4\Delta$ compared to the wild type at the basal levels. Which then 222 decreased significantly at the 30 minutes time point. The AGS1 contains a PBE at its 5'UTR, 223 and may be a Puf4 target (Table 1). The β -1,6-glucan synthase genes *KRE6* and *SKN1* showed 224 opposite trends. Both wild type and the $puf4\Delta$ cells had comparable levels of KRE6 at t=0 225 minutes, yet the puf4 Δ cells had a significantly decreasing trend of KRE6 abundance during the 226 caspofungin time-course. Another β -1,6-glucan synthase gene SKN1 was upregulated in the

227	<i>puf4</i> Δ compared to wild type and remained upregulated throughout the 60 minutes (Fig 5G-I).
228	The PBE element that is present in the SKN1 3' UTR compared to KRE6, which does not have
229	a PBE, may explain the opposite trends in post-transcriptional gene regulation.
230	
231	Our quantitative analysis of the mRNAs involved in cell wall biosynthesis showed that
232	Puf4 plays a regulatory role in the fate of these mRNAs and modulate their abundances during
233	caspofungin treatment.
234	
235	Puf4 stabilizes cell wall biosynthesis genes involved in chitin and α -glucan synthesis.
236	To gain more mechanistic insight on how Puf4 may control the cell wall biosynthesis related
237	transcript abundances, we investigated the mRNA stability of the same transcripts. mRNA
238	stability is a crucial step in transcriptome remodeling to adapt to various environmental and
239	compound stressors (32). Therefore, we hypothesized that Puf4 may modulate cell wall
240	biosynthesis genes post-transcriptionally at the mRNA stability level.
241	
242	We found that CHS3 and CHS4 were destabilized to a great degree in the $puf4\Delta$ mutant,
243	whereas CDA3, AGS1 and CDA1 exhibited a slight reduction in stability in the absence of Puf4.
244	Unlike other genes we investigated, CDA1 does not contain a PBE and was included as a
245	negative control. Even though we expected CDA1 stability to be like wild type, it too was slightly
246	destabilized (Fig. 6). Our results show that Puf4-mediated post-transcriptional gene regulation in
247	mRNA stability may be crucial for cell wall remodeling that contributes to the caspofungin
248	resistance.
249	
250	Caspofungin treatment leads to increased cell wall chitin which is exacerbated in the
251	<i>puf4</i> ∆ mutant.

252 Since we have shown that Puf4 regulates cell wall biosynthesis related transcript abundances 253 and their mRNA stability, we further investigated the functional consequences of this Puf4 loss 254 by assessing cell wall chitin and chitooligomer content using calcofluor and wheat germ 255 agglutinin staining, respectively. Microscopy (Fig 7A) and flow cytometry analysis (Fig 7B-C) of 256 the chitin content using calcofluor dye showed that the cell wall chitin content increases 257 following caspofungin treatment in both wild type and the puf4 Δ cells. Importantly, the puf4 Δ 258 cells had significantly more chitin following caspofungin treatment compared to that of wild type 259 cells. In other pathogenic fungi such as Candida species and Aspergillus fumigatus, increased 260 chitin content is protective against caspofungin (17, 18).

261

Microbial cultures show molecular and phenotypical heterogeneity that may be important within the scope of antimicrobial resistance (33). Histogram graphs of the calcofluor staining showed that the increase in chitin content is at a sub-population level (Fig 7B). While the majority of the *puf4* Δ cell population shifted to a high chitin phenotype, wild type cells showed a more heterogeneous population. Lastly, we looked at the exposed chitooligomers as another form of chitin-derived structure, and saw a modest increase compared in the *puf4* Δ cells compared to the wild type.

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In this study, we have demonstrated that the caspofungin resistance of *C. neoformans* is regulated at the post-transcriptional level through the direct interaction of the mRNA encoding its target, *FKS1*, as well as through the regulation of multiple genes that regulate cell wall composition. Post-transcriptional regulation of cell wall remodeling genes by Puf4 has functional consequences, as the absence of Puf4 results to massive remodeling of the *C. neoformans* cell wall components. Future work will investigate the effect of Puf4 on the translation of these target mRNAs, as well as the mechanism by which Puf4 itself is regulated.

277

278 Discussion

279 The search for novel antifungal therapies is an ongoing battle in medical mycology, especially 280 with the growing number of fungal outbreaks and emerging drug resistance issues (33, 34). The 281 latest class of antifungals approved by the FDA are the echinocandins, and Cryptococcus is 282 intrinsically resistant to this class of antifungals (9, 14, 15, 36). Even though it is crucial to 283 design new therapies, it is also imperative to understand the resistance mechanisms to the 284 existing antifungals to avoid similar scenarios and to design adjunctive therapies to remedy the 285 current resistance issues. In this study, we elucidated the role of post-transcriptional gene 286 regulation in the molecular mechanism of action behind the caspofungin resistance. We 287 discovered that Puf4, a Pumilio domain-containing RNA binding protein, plays a role in the 288 resistance phenotype by stabilizing the mRNAs encoding cell wall biosynthesis genes post-289 transcriptionally. The functional consequence of this interaction is a change in cell wall 290 composition to a state that is more favorable during caspofungin challenge (Fig. 8). The intrinsic 291 resistance of C. neoformans to caspofungin involves multiple signaling pathways (13, 35). For 292 the first time, we have implicated post-transcriptional regulation of cell wall biosynthesis 293 mRNAs, including the mRNA encoding the target of caspofungin, in this intrinsic resistance.

294

295 PUF proteins, in addition to their interactions with other signaling proteins, alters mRNA 296 function, and this is often secondary to the translational repression or the inhibition of mRNA 297 decay (37, 38). Binding by Puf4 and other PUF proteins orchestrate mRNA fate-determining 298 processes including stability, splicing, localization and translatability (39, 40). For example, S. 299 cerevisiae Puf4p stabilizes the transcripts involved in rRNA processing, and deletion of PUF4 in 300 S. cerevisiae causes defects in translation. Additionally, Puf4p plays a role in the recruitment of 301 mRNAs to the translational machinery (41). In C. neoformans, Puf4 appears to play both 302 positive and negative regulatory roles. Puf4 is a positive regulator of the unconventional splicing 303 of the ER-stress transcription factor HXL1. In contrast, Puf4 is a negative regulator of the ALG7

mRNA, which is stabilized in the *puf4*∆ mutant (27). Interestingly, the Puf4 elements in the *HXL1*mRNA, like *FKS1*, are in the 5' UTR, which may suggest that 5' UTR Puf4-binding elements
exert positive regulatory activity. Stabilization of the 5' UTR Puf4 binding element containing *CHS3* and *AGS1* in the absence of Puf4 support this claim (Table 1 and Fig. 6). Caspofungin
challenge likely requires a reprogramming of gene regulatory networks for adaptation, and Puf4
and other related RNA-binding proteins may be involved in transforming the translating mRNA
pool to best respond to the pharmacologically induced stress by caspofungin.

311

312 Cell wall maintenance and perturbations in response to drug induced stress have a 313 broad appreciation by the medical mycology (42). For example, the mar1∆ mutant exhibits a 314 defect in intracellular trafficking of cell wall synthases and therefore exhibits a cell wall 315 composition that contains elevated exposed chitin and decreased glucan levels. These changes 316 in the cell wall composition and exposure of different carbohydrates play meaningful roles in the 317 immune recognition by the host and activate various signaling events in the host system (43). 318 Another example is the enhanced recognition of the $ccr4\Delta$ mutant by alveolar macrophages due 319 to increased unmasking of the β -1,3-glucan (44). The importance of Ccr4, an mRNA 320 deadenylase, in glucan masking is further evidence that post-transcriptional processes are 321 essential for adaptation to a number of stressors, including caspofungin treatment. In this study, 322 we show that Puf4 is a major regulator of cell wall biogenesis. We report that cell wall 323 biosynthesis genes showed different trends of expression in the $puf4\Delta$ compared to wild type in 324 caspofungin time-course experiments. We have also shown that FKS1, CHS3 and CHS4 325 mRNAs were destabilized in the absence of Puf4. This regulation of certain cell wall biogenesis 326 genes by Puf4 may be a necessary component of cell wall homeostasis under normal growth 327 conditions as well as facilitating rapid changes in cell wall gene expression during adaptation to 328 drug-induced stress. We have predicted that transcripts that have the PBE would have

329 enhanced decay in the $puf4\Delta$ mutant vet this was not true for all transcripts that carried PBEs. 330 We investigated the stability of CDA1 as a transcript that did not contain a PBE, yet we 331 observed a modest change in the mRNA stability. In that regard, it must be noted that other 332 RNA-binding proteins may play meaningful regulatory roles in conjunction with Puf4. 333 Additionally, the impact of Puf4 on its targets may be unique and this may be due to the location 334 of the PBE within the transcript, whether it is in the 3' or 5' UTR, introns or exons, and may yield 335 to variations in which Puf4 regulates transcript fate. Moving forward, screening additional RNA-336 binding proteins to understand the post-transcriptional regulatory network controlling cell wall 337 dynamics, as well as investigating the regulatory connections with the cell integrity pathway and 338 calcineurin signaling pathway will elucidate further the response of C. neoformans to 339 caspofungin that mitigate its toxicity and promote intrinsic resistance.

340

341 The complex cell wall structure of C. neoformans protects the cell from the extracellular 342 stressors including antifungals. A sturdier cell wall can also serve as a less permeable barrier to 343 antifungal drugs, making them less effective (45). We found that following treatment with 344 caspofungin wild type cells had an increase in the cell wall chitin. This increase was observed at 345 the sub-population level. This was an intriguing finding since sub-population impact of 346 antifungals are a neglected area of study, yet these heterogeneously resistant or tolerant 347 populations may play important roles in antifungal resistance (46-48). Recent studies show that 348 the effect of antifungals at subpopulation level is especially crucial to explain the growth of 349 Candida species at supra-MIC concentrations (33). This is mainly due to slow growth of 350 subpopulations compared to the rest of the cells. The heterogeneous nature of axenic microbial 351 cultures, and life in general, is an intricate phenomenon that may significantly contribute to our 352 understanding of antifungal resistance and tolerance. We hope that the single-cell genomics era 353 will enhance our understanding of the antifungal resistance and tolerance pathways in more 354 detail.

355

Puf4 is a *bona fide* downstream effector of the calcineurin pathway, as it was enriched in 356 357 a phosphoproteomics screen of the $cna1\Delta$ mutant (24). Calcineurin was portrayed to be a 358 longstanding player in the antifungal resistance of medically important fungi (49). Many groups 359 have shown that disruption of the calcineurin pathway, genetically or pharmacologically, using 360 novel or repurposed molecules, abolished the calcium homeostasis and led to death. This 361 specific inhibition of calcineurin suggested to us that the calcineurin pathway, especially the 362 downstream effectors, may contain potential targets which can be used as novel antifungal 363 targets (50). Caspofungin treatment causes the translocation of a calcineurin-dependent 364 transcription factor, Crz1. Translocation of Crz1 to the nucleus is an event that induces the 365 transcriptional changes in gene expression in response to caspofungin. Yet, this transcriptional 366 regulation does not contribute to the caspofungin resistance since the $crz1\Delta$ exhibits a similar 367 caspofungin sensitivity to that of wild type cells (19). On the other hand, the absence of Puf4 368 causes a hyper-resistant phenotype. While the absolute absence of Puf4 yields a resistance 369 phenotype, we demonstrated that Puf4 protein levels drastically decrease following caspofungin 370 treatment. Interestingly, this suggests that the downregulation of Puf4 may be necessary for the 371 paradoxical resistance, hence the hyper-resistant phenotype in the knockout. The converse was 372 true for the overexpression of Puf4 which showed hyper-sensitivity to caspofungin.

373

Further work is needed to determine if the phosphorylation state of Puf4 is governed by calcineurin, or if there is another post-translational modification, that is responsible for the rapid reduction in Puf4 protein abundance in response to caspofungin treatment. Inhibition of the mediator of Puf4 protein repression is another potential target to reverse the intrinsic resistance of *C. neoformans* to caspofungin. The post-transcriptional regulation of cell wall homeostasis by Puf4, a calcineurin-regulated RNA-binding protein, is another piece of the regulatory program that results in the intrinsic resistance of *C. neoformans* to caspofungin. Further elucidation of

this regulatory program may open new avenues to promote caspofungin sensitivity in C.

382 *neoformans* through adjunctive therapy.

383

384 Materials and Methods

385 Yeast strains and molecular cloning.

386 All strains used in this study were derived from C. neoformans var. grubii strain H99, a fully-387 virulent strain gifted from Peter Williamson (UIC, NIAID), which is derived from H99O gifted by 388 John Perfect (Duke University). Primers used to build the knockout and FLAG-tagged 389 complementation strains are included in Table 2. The plasmid construct to establish the PUF4-390 FLAG-(Hyg) mutant included native promoter and the terminator of the gene. Promoter and the 391 coding sequence were amplified as a single fragment using a forward primer that contained a 392 Notl cut site, and a reverse primer that contained a Sall cut site as well as the FLAG sequence. 393 Puf4 terminator fragment was amplified using forward and reverse primers that contained Sall 394 and BgIII sites, respectively. Following restriction digest with respective enzymes, these 395 fragments were cloned into pSL1180 containing the hygromycin B resistance cassette, as 396 described previously (51). Construct containing the PUF4-FLAG was introduced into the $puf4\Delta$ 397 mutant using biolistics transformation. Copy number were determined using southern blot 398 analysis. A strain that is a single copy tagged complement, and another strain that is a tagged 399 overexpression was established.

400

401 Growth analysis: Spot plates and plate reader assay.

402 Cells were grown overnight at 30°C in 5 mL cultures in YPD broth. Overnight cultures were
403 washed with sterile distilled water and the OD₆₀₀ was equaled to 1 in water. Adjusted cultures
404 were 1:10 serially diluted 5-times and 5 µl of each dilution spotted on YPD agar plates
405 containing indicated concentrations of caspofungin (Sigma). Plates were incubated at 30°C for 3
406 days and photographed. For the kinetic plate reader assay, overnight cultures were washed with

407 water once and then OD_{600} was equaled to 0.3 in YPD broth. 50 µl of YPD broth containing the 408 2x the final caspofungin concentration was placed in each well then 50 µl of the OD_{600} adjusted 409 cultures were placed in each well. Plate was incubated at 30°C for 20 hours while shaking in a 410 double orbital fashion and OD_{600} was measured every 10 minutes during this kinetic assay.

411

412 Electrophoretic mobility shift assay (EMSA).

413 EMSA reactions were set and analyzed as described previously (27). Briefly, all RNA binding 414 reactions contained 5 µg of total protein lysate, 0.5 pmol of the TYE705-labeled 415 oligonucleotide (IDT), and 4 µl 5x EMSA buffer (75 mM HEPES pH 7.4, 200 mM KCl, 25 416 mM MgCl₂, 25% glycerol, 5 mM dithiothreitol (DTT), and 0.5 mg/ml yeast tRNA) in a total 417 volume of 20 µl. For competition reactions, 5x, 10x, and 20x more unlabeled wild type or 418 mutant oligonucleotides added in addition to the TYE705-labeled oligonucleotide. Reaction 419 mixtures were incubated at room temperature for 20 minutes, and run on DNA retardation gel, then electrophoresed at 100V. Gels were imaged using a LiCor Odyssev imaging 420 421 system.

422

423 Motif Search - FIMO: Find Individual Motif Occurrences

424 Cell wall biosynthesis genes were scanned for the Puf4-binding-element using the FIMO tool on 425 the MEME-suite version 5.1.0 (52). RNA sequences of the cell wall biosynthesis genes in Table 426 1 were acquired from FungiDB and provided as the input. **UGUANNNNUA** motif was scanned 427 using the default settings. Only the given strand was searched and the p-value criterion was set 428 as p<0.0005 for significance cut-off. Results were manually curated to ensure accuracy in 429 detecting the desired motif.

430

431 **RNA stability time-course.**

432 Overnight cultures grown at 30°C were used to inoculate 35 mL of YPD broth at the OD₆₀₀ 433 between 0.15 - 0.2 in baffled erlenmeyer flasks. Cultures were grown in baffled flasks at 30 °C 434 while shaking at 250 rpm until they reach the mid-log stage $-OD_{600}$ between 0.6 and 0.7. Mid-435 log stage cultures were supplemented with 250 µg/mL of the transcriptional inhibitor 436 1,10-phenanthroline (Sigma). Then, 5-mL aliguots of each culture were transferred to snap cap 437 tubes and pelleted every 15 minutes for 60 minutes. 50 μ I RLT supplemented with 1% β -438 mercaptoethanol was added to each pellet prior to flash freezing in liquid nitrogen. Pellets were 439 stored at -80 °C until RNA extraction. Cells were lysed by bead beating using glass beads. RNA 440 was extracted from each sample using the RNeasy Mini Kit (QIAGEN) following manufacturer's 441 instructions. RNA was DNase digested on-column using the RNase free DNase Kit (QIAGEN) 442 or using the AMBION TURBO DNA-free Kit (ThermoFisher). cDNA for real time guantitative 443 PCR (RT-gPCR) was synthesized using the Applied Biosystems High Capacity cDNA Reverse 444 Transcription Kit (ThermoFisher) using random hexamers. 800 ng to 1000 ng RNA was used to 445 synthesize cDNA. Samples were quantified using the second-derivative-maximum method and 446 fitted to a standard curve of five 4-fold serial dilutions of cDNA. For experimental samples, 447 cDNA was diluted 1:5 in nuclease-free water. To make the reaction mixture, 5 µl of the 2X 448 SYBR Green Blue Mix (PCRbiosystems) was combined with 4 µl of 1.5 µM primers (970 µl 449 water + 15 µl forward +15 µl reverse). 9 µl reaction mixture was placed in each well and 1 µl of 450 either experimental samples or standards added to their respective wells. Samples from 3 451 biological samples in duplicate wells were tested. Primer sequences are listed in Table 2 in the 452 along with the gene IDs. Statistical differences were compared by determining the least squares 453 fit of one-phase exponential decay non-linear regression analysis with GraphPad Prism 454 software. Significance between curves was detected with the P-value cut-off of 0.05, which 455 determined that the data from two different curves create different regression lines therefore 456 yielding to different half-lives of the same transcript investigated in different mutants.

457 Caspofungin time-course.

458 Cells were grown to the mid-log stage as described in the previous section. At this stage, 459 cultures were supplemented with 16 µg/mL caspofungin and 5 mL aliquots were collected in 460 snap-cap tubes every 15 minutes for 60 minutes. 50 μl of buffer RLT supplemented with 1% β-461 mercaptoethanol was added to each pellet prior to flash freezing in liquid nitrogen. Pellets were 462 stored at -80 °C until RNA extraction. Cells were lysed by bead beating using glass beads. RNA 463 was extracted from each sample, cDNA was synthesized and transcript abundances were 464 calculated using RT-gPCR as described in the previous section. Primer sequences are listed in 465 Table 2. Statistical differences were determined using a two-way-ANOVA.

466

467 **Immunoblotting**.

468 Cells were grown to the mid-logarithmic stage and half of the culture were treated with 16 µg/mL 469 caspofungin for an hour while the other half was left untreated. Cell pellets were flash frozen in 470 liquid nitrogen and stored at -80 °C. At the time of lysis, 50 µl cold lysis buffer (50 mM Tris HCl 471 pH 7.4, 150 mM NaCl, 1 mM EDTA, 1% Triton X100, 10 µl/ml HALT protease and phosphatase 472 inhibitor [ThermoFisher]) was added and cells were lysed by bead beating using glass beads. 473 250 µl of the cold lysis buffer was added to the beads and lysate was extracted from the glass 474 beads. Lysate was centrifuged at 20,000 x g for 15 min, and supernatant was transferred to a 475 new tube. Protein quantities were measured using Pierce 660nm Protein Assav Kit 476 (ThermoFisher). 25 µg of protein was run per sample on 4-15% Mini-PROTEAN TGX stain-free 477 precast gels (BioRad) at 150V. Total protein was imaged using a BioRad Gel Documentation 478 System with the stain-free gel setting. Gels were transferred to nitrocellulose membrane and 479 blocked with LiCor Odyssey blocking buffer for an hour. Then, incubated overnight with the 480 mouse Anti-FLAG antibody (1:1000 in Tris-Buffered Saline-Tween20 [TBS-T] with 10% LiCor 481 Odyssey blocking buffer) at 4°C. The blot was washed three times with 15 minutes incubations 482 in TBS-T. Then, LiCor rabbit anti-mouse 800 secondary antibody (1:10000 in TBS-T with 10%

LiCor Odyssey blocking buffer) was added. The blot was incubated with the secondary antibody for an hour at room temperature, then washed with TBS-T and blot was imaged using a LiCor Odyssey imaging system.

486

487 Cell wall staining, microscopy and flow cytometry.

488 Cells were grown to the mid-log stage and treated with caspofungin as described previously.
489 Cells were prepared and cell wall components were stained for microscopic and flow cytometric
490 analyses as previously published (43, 53). Briefly, cells were pelleted and washed with 1X PBS
491 once. Cells were fixed with 3.7% formaldehyde for 5 minutes at room temperature, and washed
492 with 1X PBS twice.

493

494 Cells were stained with calcofluor and FITC-conjugated wheat germ agglutinin (WGA: 495 Molecular Probes) to visualize chitin. Calcofluor dye stains the total chitin while WGA only stains 496 the exposed chitooligomers. Cells were incubated in the dark with 100 µg/ml FITC-WGA for 35 497 minutes, then consecutively stained with 25 µg/ml calcofluor white for 15 minutes. Cells were 498 washed twice with 1X PBS before analysis. Stained cells were imaged using a Leica TCS SP8 499 Confocal Microscope. For microscopy, WGA was detected using the GFP settings and 500 calcofluor was detected using the DAPI settings. Images were taken using the 100X objective. 501 Representative images were shown. Flow cytometry data was acquired using a BD 502 LSRFortessa Cell Analyzer. WGA signal was detected using 488nm laser, and the calcofluor 503 was detected using the 405nm laser. Flow cytometry data were analyzed FlowJo v10.0 504 software. Representative histogram graphs were shown. 505

506 Cells were stained with Aniline Blue to detect the β-1,3-glucan levels. Unfixed mid-log
 507 stage cells in YPD, untreated and treated with 16 µg/ml caspofungin, were washed with 1X PBS

508	and stained with 0.05% Aniline Blue (Wako Chemicals, Japan) for 10 minutes, and then cells
509	were imaged using the DAPI channel on the Leica TCS SP8 Confocal Microscope. Microscopy
510	images were analyzed using the Fiji (Image J, NIH). Mean fluorescent intensity of at least 100
511	cells in 3-4 different fields were quantified and normalized. Representative images are shown.
512	
513	Statistical Analysis.
514	Data analysis was performed using the GraphPad Prism software version 6. Each figure legend
515	contains the statistical test information that is used to assess the statistical significance. Briefly,
516	we have utilized the one-phase exponential decay analysis to determine the half-life of the
517	mRNAs analyzed. Immunoblot data was analyzed using unpaired t-test with Welch's correction.
518	Gene expression and microscopy quantification data were analyzed using either one-way or
519	two-way-ANOVA followed by a post-hoc multiple comparison test. For all of the graphs, *:
520	p<0.05, *** p<0.001 and ****: p<0.0001. All error bars represent the SEM.
521	
522	Acknowledgements
523	We would like to acknowledge Dr. Amanda L. M. Bloom for advice and stimulating discussions.
524	This work was funded by NIH R21 AI133133 to JCP.
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690 Figure Legends

691

692 Figure 1. The *puf4* mutant is resistant to caspofungin. (A) Spot plate analysis. The 693 indicated strains were diluted to an OD₆₀₀ of 1.0 and five, 1:10 serial dilutions were spotted on 694 agar plates containing 0, 16, 32, 40, and 48 µg/ml caspofungin. Plates were incubated at 30°C 695 for 3 days and photographed. (B-C) Growth assay. The indicated strains were diluted to an 696 OD₆₀₀ of 0.3 and mixed 1:1 with either fresh YPD or YPD containing caspofungin in a 96 well-697 plate. Plates were incubated and at 30°C for 20 hours while shaking. OD₆₀₀ was measured every 698 10 minutes. 699 700 Figure 2. Puf4 directly binds and stabilizes the FKS1 mRNA. (A) The FKS1 mRNA contains 701 Puf4 binding elements (PBE) in its 5' UTR - UGUANNNNUA. (B) Puf4 binds to FKS1. 702 Electrophoretic mobility shift assay was performed using a fluorescently labeled synthetic RNA 703 oligonucleotide designed for the FKS1 5'UTR that contain the PBEs. Unlabeled mutant (contains mutated PBEs) and unlabeled wild type probes were used as controls for sequence 704 705 specificity. $puf4\Delta$ was included as a control to show binding is absent when Puf4 is not present. 706 A representative gel image is shown, n=3. (C) The *FKS1* is downregulated in the *puf4* Δ . The 707 FKS1 mRNA abundance in mid-log samples grown at 30°C was determined using RT-gPCR 708 with GPD1 as the normalization gene. 3 replicates were plotted and unpaired t-test with Welch's 709 correction was performed, *: p<0.05. (D) The FKS1 is destabilized in the puf4 Δ mutant. FKS1 710 abundance was determined using RT-gPCR following transcription shut-off to determine the 711 decay kinetics. GPD1 was utilized as the control for normalization. 3 replicates were plotted 712 and differences among two strains were analyzed using one-phase exponential decay analysis. 713 Error bars show SEM.

714

715 Figure 3. Puf4 protein expression is decreased following caspofungin treatment. (A)

716 PUF4 transcript levels are unchanged during a 60-minute caspofungin time course. Cells were 717 grown at 30°C and treated with caspofungin. The abundance of FKS1 mRNA was determined in 718 samples collected every 15 minutes using RT-gPCR with GPD1 as the normalization gene. 3 719 replicates were plotted and one-way-ANOVA using Tukey's test was performed, p=0.5814 (B) 720 Puf4 protein levels are decreased following 60-minute caspofungin treatment. Puf4-FLAG strain 721 (*puf4*^{\Delta} expressing *PUF4-FLAG* C-terminal fusion) was grown to mid-log and treated with 722 caspofungin for 1 hour. SDS-PAGE followed by immunoblotting using anti-flag antibody is 723 shown, (C) Anti-flag signal is normalized to total protein signal from the stain-free gel. 5 724 replicates were plotted and unpaired t-test with Welch's correction was performed, **: p<0.01. 725 Error bars show SEM.

726

727 Figure 4. Puf4-dependent *FKS1* regulation correlates with reduced β -1,3-glucan staining. 728 (A) FKS1 abundance is regulated by Puf4 during a 60-minute caspofungin time course. Cells 729 were grown at 30°C and treated with caspofungin. The abundance of FKS1 mRNA was 730 determined in samples collected every 15 minutes using RT-gPCR with GPD1 as the 731 normalization gene. 3 replicates were plotted and one-way-ANOVA using Tukey's test was 732 performed. (B) The *puf4* Δ mutant has decreased levels of β -1,3-glucan. Cells were grown to 733 mid-log at 30°C and stained with Aniline Blue to detect β -1,3-glucan. Representative Aniline 734 Blue staining images for each strain is shown to assess both levels and localization. (C) 735 Microscopy images were quantified on Fiji and Aniline Blue signal was normalized to DIC signal. 736 At least 5 fields from 3 biological replicates were plotted and one-way-ANOVA followed by Dunn's multiple comparisons test was performed. *: p<0.05, **: p<0.01 Error bars show SEM. 737 738

739 Figure 5. Deletion of *PUF4* leads to dysregulation of certain cell wall biosynthesis genes.

- 740 The mRNA abundance of select cell wall biosynthesis genes was determined during a 60-
- 741 minute caspofungin time course by collecting samples every 15 minutes and determining
- abundance by RT-qPCR using GPD1 as a normalization gene. (A) CHS3, (B) CHS4, (C) CHS6,
- (D) CDA1, (E) CDA2, (F) CDA3, (G) AGS1, (H) KRE6, and (I) SKN1. 3 biological replicates with
- 2 technical replicates were plotted and two-way ANOVA was used to determine statistical
- significance. '#' denotes comparison between wild type and *puf4*∆ while '*' denotes comparison
- 746 between indicated time points within a strain. #/*: p<0.05, ##/**: p<0.01, ###/*** p<0.001 and
- 747 ####/****: p<0.0001. Error bars show SEM.
- 748

749 Figure 6. Puf4 stabilizes cell wall biosynthesis genes involved in chitin and α-glucan

750 synthesis. (A) CHS3, (B) CHS4, (C) CDA3, (D) AGS1, (E) CDA1 transcript abundance were

751 determined using RT-qPCR following transcription-shutoff to determine the decay kinetics.

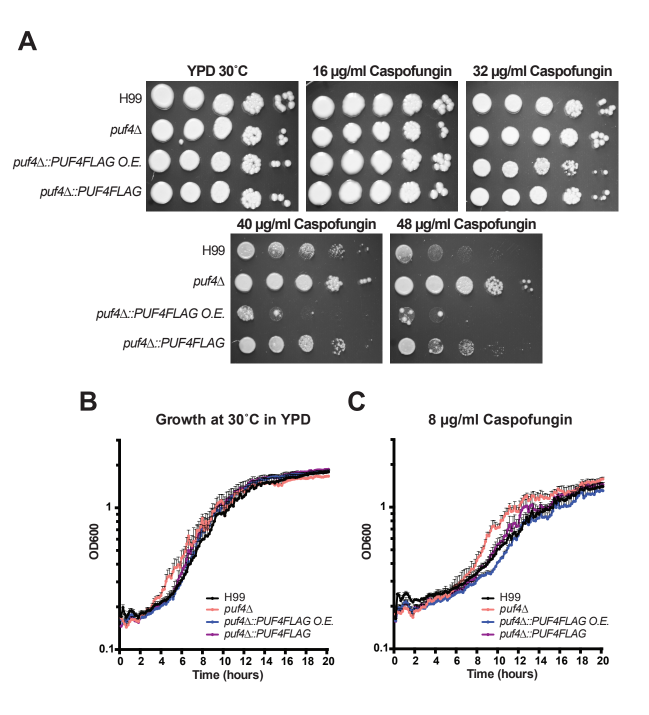
- 752 *GPD1* was utilized as the control for normalization. 15 minutes post-transcription shut-off was
- denoted as t=0. 3 replicates were plotted and differences among two strains were analyzed
- using one-phase exponential decay analysis. Error bars show SEM.

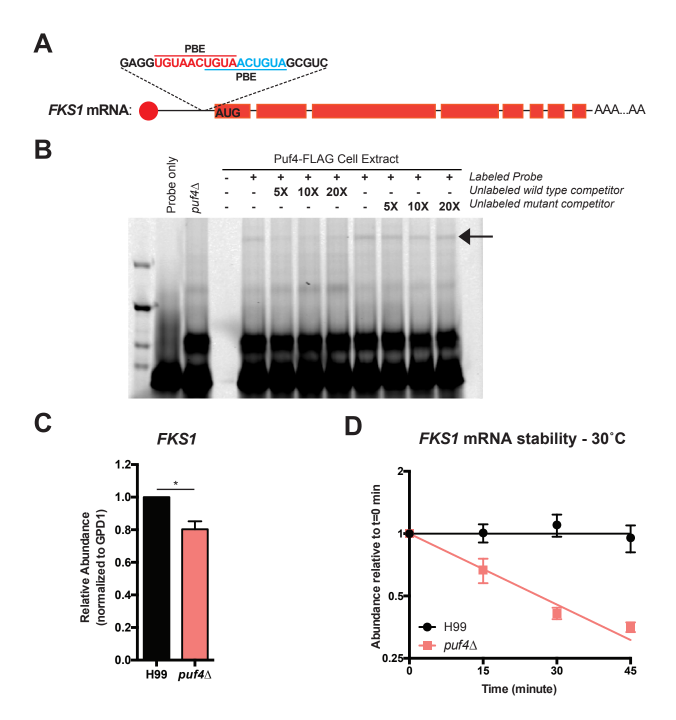
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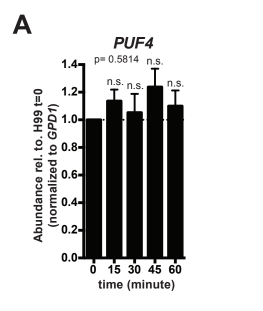
Figure 7. Caspofungin treatment leads to increased cell wall chitin which is exacerbated in the *puf4*∆ mutant. (A-C) Cell wall chitin levels are increased response to caspofungin. Cells were grown to mid-log at 30°C and stained with calcofluor white, then fluorescent intensity (A-B) and staining pattern (C) were determined using flow cytometry and fluorescence microscopy, respectively. (D-E) Levels of exposed chitooligomers did not change significantly. Exposed chitooligomers were stained using wheat germ agglutinin conjugated with FITC. Fluorescent intensity was determined using flow cytometry. For (A) and (D), 3 biological replicates were

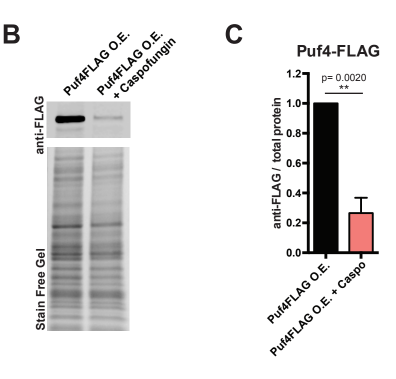
- 763 plotted and one-way-ANOVA followed by Dunn's multiple comparisons test was performed. *:
- 764 p<0.05, *** p<0.001 and ****: p<0.0001. Error bars show SEM.
- 765
- 766 Figure 8. Model: Post-transcriptional regulation of cell wall remodeling by Puf4 is a path
- to caspofungin resistance in *C. neoformans*. In the wild type cells, *FKS1* is stabilized by
- 768 Puf4. When wild type cells are treated with caspofungin, we observe increase of cell wall chitin
- and β -1,3-glucan staining intensity. In the *puf4* Δ , *FKS1* as well as other cell wall biosynthesis
- transcripts are destabilized. Therefore, we observed a different response to caspofungin in the
- puf4 Δ involving a robust increase in cell wall chitin content. Graphics modified from (9).
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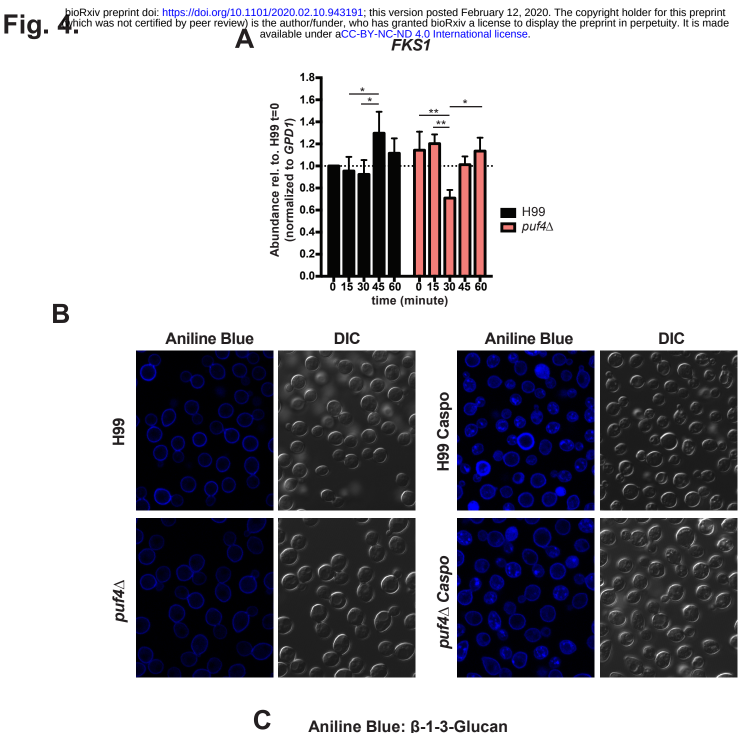
Fig.

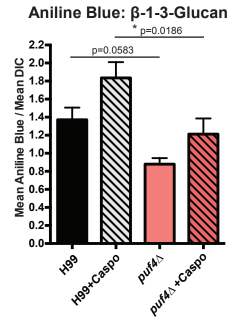




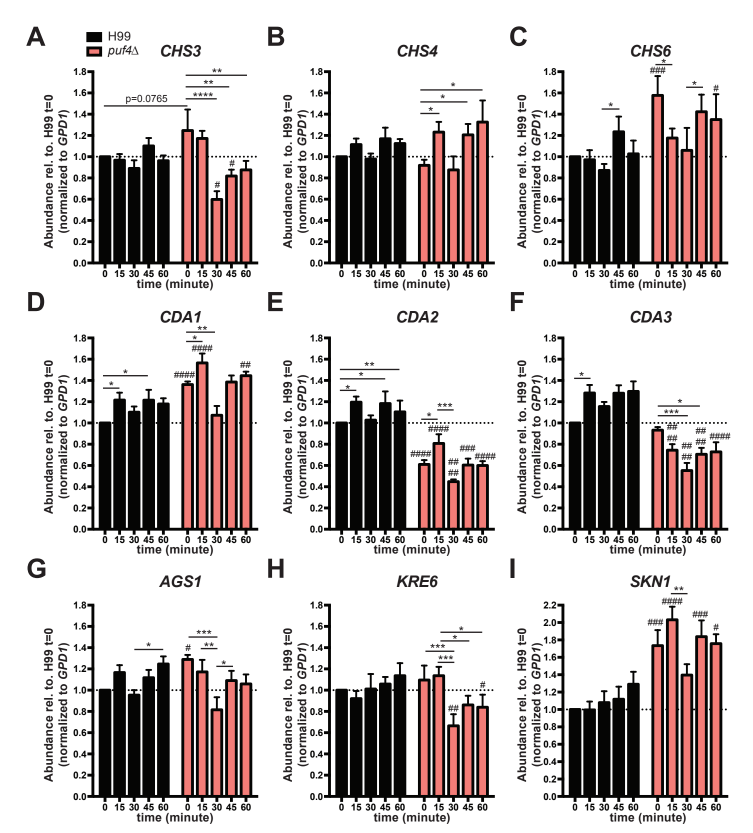








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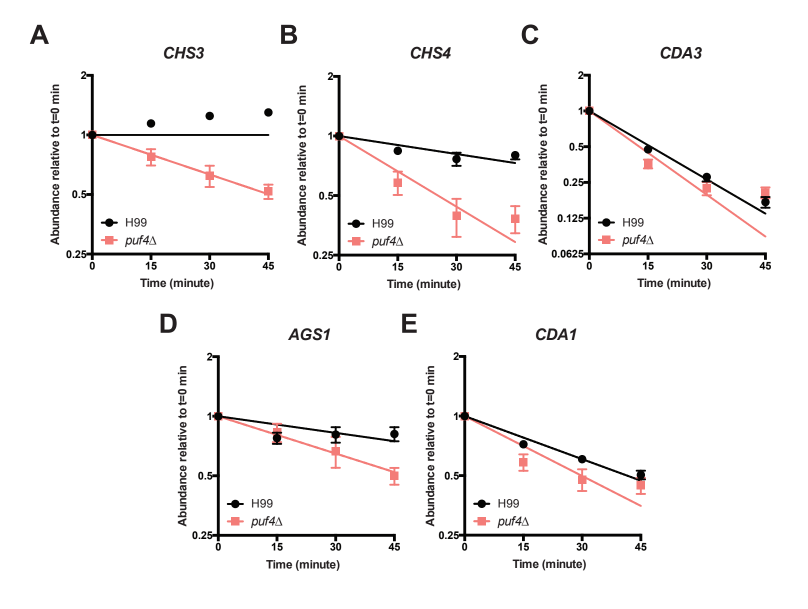
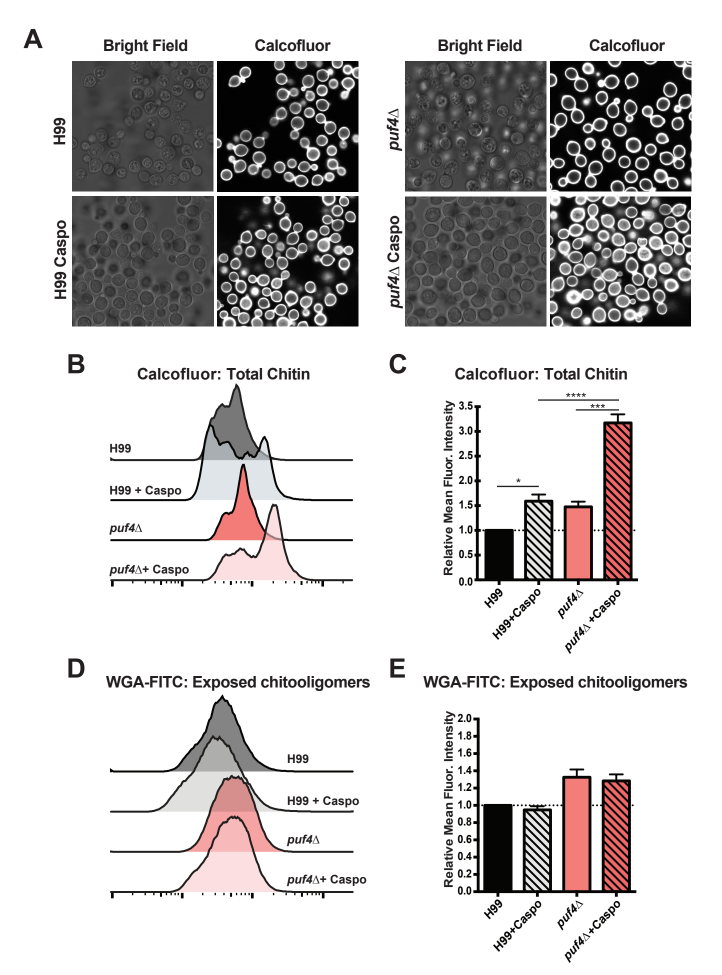


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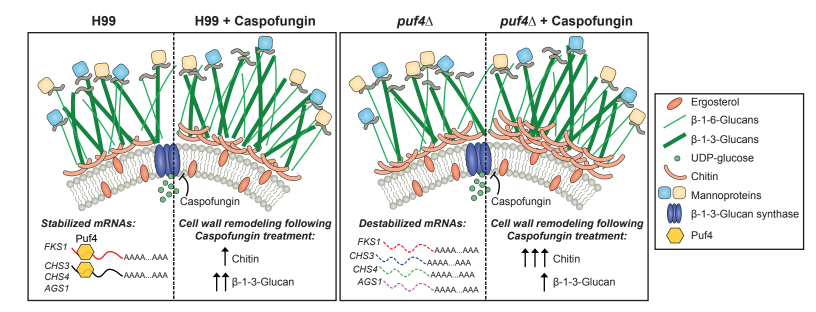


Table 1. List of Puf4 binding element containing cell wall biosynthesis related genes.
UGUANNNNUA motif was searched in target genes using FIMO (MEME-suite version 5.1.1.).
Results were manually confirmed and location of the motifs were identified.
* Indicates the genes selected for further mRNA stability analysis.

Gene	Gene ID	Start	End	p-value	Location
CHS3*	CNAG_05581	242	251	0.000262	5' UTR
CHS4*	CNAG_00546	3257	3266	0.00013	Exon
		3516	3525	0.000203	Exon
CHS7	CNAG_02217	88	97	6.59E-05	5' UTR
		3264	3273	6.59E-05	Exon
CHS8	CNAG_07499	1479	1488	7.96E-05	Exon
CDA3*	CNAG_01239	1916	1925	7.96E-05	3' UTR
FKS1*	CNAG_06508	281	290	7.96E-05	5' UTR
		307	316	0.000162	5' UTR
		313	322	0.000162	5' UTR
SKN1	CNAG_00897	2471	2480	0.000122	3' UTR
AGS1*	CNAG_03120	202	211	0.000107	5' UTR

Cloning			
Name	Primer Sequence	Restriction site	Referenc
HYG F	GGCTGCGAGGATGTGAGC		This study
HYG R	GCCACCAAGCGTTAAGGCC		This study
PUF4 5' Promoter F	TAATAAGCGGCCGCCGCGATGCTGGCAAGGC	Notl	This study
PUF4 +FLAG R	TAATAAGTCGACTCATTTATCATCATCCTTGTAATCCATCATCGGAACGTGGGAAGG	Sall	This study
PUF4 Terminator 5' end F	TAATAA GTCGAC CTGAAGAGCGGATAAAGTCCG	Sal1	This study
PUF4 Terminator 3' end R	TAATAAAGATCTGCATGTCGATGAGAATGTCAGG	BgIII	This study
EMSA			
Name	Sequence	Reference	
FKS1 5' UTR-TYE705	/5TYE705/rGrArCrGrUrGrUrArGrArGrCrUrArUrCrGrArCrArGrGrArArGrArGrArGrArGrGrUrGrUrArArCrUrGrUrArArCrUrGrUrArGrCrGrU	This study	
FKS1 5' UTR -WT-unlabeled	rGrArCrGrUrGrUrArGrArGrCrUrArUrCrGrArCrArGrGrArGrArGrArGrArGrArGrGrUrGrUrArArCrUrGrUrArArCrUrGrUrArGrCrGrU	This study	
FKS1 5' UTR-MUT-unlabeled	rGrArCrGrArArArArGrArGrCrUrArUrCrGrArCrArGrGrArArGrArGrArGrArGrArArArA	This study	
RT-q-PCR			
Gene	Primer Seguence	Reference	
PUF4 F	AAGACGACGATGTTGATGCG	This study	
PUF4 R	GCACCAGGAACAGCAGAAAA	This study	
CHS3 F	ACCCAGGTCTGGCATTCC	(Esher et. al., 2018)
CHS3 R	AGGATCAACATTGGAAGC	(Esher et. al., 2018)
CHS4 F	CGGTCTTCAGGCATTGATTT	(Esher et. al., 2018)
CHS4 R	TTCGGAGTGAAGTGATGCTG	(Esher et. al., 2018)
CHS6 F	TTGACCCTTGGCACATCT	(Esher et. al., 2018)
CHS6 R	GTTGGCATAAGTATCCTT	(Esher et. al., 2018)
CDA1 F	TCGAGCTATTGCTGCTCAGA	(Esher et. al., 2018)
CDA1 R	GCTGGTAGATGTCGTGCTCA	(Esher et. al., 2018)
CDA2 F	GTAACGAGGTCGTCTTTG	(Esher et. al., 2018)
CDA2 R	TGTAGTTGGTGAGCTCGT	(Esher et. al., 2018)
CDA3 F	ATGTGGCCGATGCTTTTAAC	(Esher et. al., 2018)
CDA3 R	GAAGTGAGAAGGCCTGTTGG	(Esher et. al., 2018)
AGS1 F	ATCCTTATCCGTTATTCC	(Esher et. al., 2018)
AGS1 R	AGCTGTTCCTCTAGCGAGC	(Esher et. al., 2018)
FKS1 F	TGGACTGGTGTTTGGTTCAA	(Esher et. al., 2018)
FKS1 R	GTACAAAAGACCGTACTTG	(Esher et. al., 2018)
KRE6 F	GTCTCGGAAGGCGACTCAT	(Esher et. al., 2018)
KRE6 R	TCAACTCATTCTTTGGGAAGG	(Esher et. al., 2018)
SKN 1 F	CTGGACAATGTATGCGGATG	(Esher et. al., 2018)
SKN 1 R	TCCGCAGTGGGATAATCTTC	(Esher et. al., 2018	/
GPD1 F	AGTATGACTCCACACATGGTCG	(Esher et. al., 2018	/
GPD1 R	AGACAAACATCGGAGCATCAGC	(Esher et. al., 2018	/