Azole resistance is mediated by integration of sterol gene regulation and membrane transporter production by the zinc cluster-containing transcription factor Upc2A in *Candida glabrata*

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Running title: Upc2A global transcriptional regulation

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Draft: May 7, 2021
The most commonly used antifungal drugs are the azole compounds that interfere with biosynthesis of the fungal-specific sterol: ergosterol. The pathogenic yeast *Candida glabrata* commonly acquires resistance to azole drugs like fluconazole via mutations in a gene encoding a transcription factor called *PDR1*. These *PDR1* mutations lead to overproduction of drug transporter proteins like the ATP-binding cassette transporter Cdr1. In other *Candida* species, mutant forms of a transcription factor called Upc2 are associated with azole resistance, owing to the important role of this protein in control of expression of genes encoding enzymes involved in the ergosterol biosynthetic pathway. Recently, the *C. glabrata* Upc2A factor was demonstrated to be required for normal azole resistance, even in the presence of a hyperactive mutant form of *PDR1*. Using genome-scale approaches, we define the network of genes bound and regulated by Upc2A. By analogy to a previously described hyperactive *UPC2* mutation found in *Saccharomyces cerevisiae*, we generated a similar form of Upc2A in *C. glabrata* called G898D Upc2A. Chromatin immunoprecipitation coupled with Next Generation Sequencing (ChIP-seq) demonstrated that wild-type Upc2A binding to target genes was strongly induced by fluconazole while G898D Upc2A bound similarly, irrespective of drug treatment. We also carried out RNA-seq analysis to determine the genes that were direct or indirect targets of Upc2A transcriptional control. In addition to the well-described *ERG* genes as Upc2A transcriptional targets, we found a large group of genes encoding components of the translational apparatus along with membrane proteins. These Upc2A-regulated membrane protein-encoding genes are often targets of the Pdr1 transcription factor, demonstrating the high degree of overlap between these
two regulatory networks. Finally, we provide evidence that Upc2A impacts the Pdr1-Cdr1 system during the anaerobic response and also modulates resistance to caspofungin. These studies provide a new perspective of Upc2A as a master regulator of lipid and membrane protein biosynthesis.

Author summary (200 words)

In the pathogenic yeast *Candida glabrata*, expression of the genes encoding enzymes in the ergosterol biosynthetic pathway is controlled by the transcription factor Upc2A. *C. glabrata* has a low intrinsic susceptibility to azole therapy and acquires fluconazole resistance at high frequency. These azole resistant mutants typically contain substitution mutations in a gene encoding the transcription factor Pdr1. Pdr1 does not appear to regulate ergosterol genes and instead induces expression of genes encoding drug transport proteins like *CDR1*. Here we establish that extensive overlap exists between the regulatory networks defined by Upc2A and Pdr1. Genomic approaches are used to describe the hundreds of genes regulated by Upc2A that far exceed the well-described impact of this factor on genes involved in ergosterol biosynthesis. The overlap between Upc2A and Pdr1 is primarily described by co-regulation of genes encoding membrane transporters like *CDR1*. We provide evidence that Upc2A impacts the transcriptional control of the *FKS1* gene, producing a target of a second major class of antifungal drugs, the echinocandins. Our data are consistent with Upc2A playing a role as a master regulator coordinating the synthesis of membrane structural components, both at the level of lipids and proteins, to produce properly functional biological membranes.
Introduction

An almost inescapable problem for chemotherapy of microbes is the development of resistance. This problem is especially acute in the case of pathogenic fungi for which only 3 different drug classes exist for use in treatment of infections (reviewed in (1, 2)). The most commonly used drug class is the azole compounds, chief among these is the well-tolerated fluconazole (reviewed in (3)). Fluconazole targets ergosterol biosynthesis and has been used successfully since the 1980s but this success has led to the development of resistant organisms (recently discussed in (4)). The prevalence of fluconazole as an anti-Candidal therapy has likely contributed to the changing epidemiology of candidemias with the frequency of these fungal infections being increasingly associated with *Candida glabrata*; an increase that correlates with the introduction of fluconazole as an antifungal drug (5).

*C. glabrata* exhibits two features that complicate its control by fluconazole. First, this pathogen has a high intrinsic resistance to fluconazole (6). Second, high level resistant isolates easily arise that contain gain-of-function (GOF) mutations in a transcription factor-encoding gene called *PDR1* (7-9). The GOF *PDR1* alleles exhibit high levels of target gene expression and drive robust fluconazole resistance primarily through induction of expression of the ATP-binding cassette transporter-encoding gene *CDR1* (10, 11).

The primary species associated with candidemias is *Candida albicans* which can also acquire fluconazole resistance (recently discussed in (12)). Interestingly, the range of genes in which mutations are observed to associate with fluconazole resistance in *C.
*albicans* is much wider than in *C. glabrata*. Along with mutant forms of the genes encoding the well-described transcription factors Tac1 and Mrr1 (13, 14), two additional genes in which fluconazole resistant alleles can emerge in *C. albicans* are *ERG11* (15), that encodes the enzymatic target of azole drugs, and Upc2, the primary transcriptional activator of *ERG11* and other ergosterol biosynthetic pathway genes (16). Mutations in the cognate genes for these proteins have not been found in *C. glabrata*.

Two important observations have recently linked *C. glabrata* Pdr1 with the ergosterol biosynthetic pathway in this yeast. First, loss of *UPC2A* (*C. glabrata* homologue of *C. albicans* *UPC2*) was sufficient to strongly reduce fluconazole resistance of a GOF *PDR1* mutant allele (17). Second, genetic means of reducing the flux through the ergosterol pathway led to induction of the Pdr pathway in a Upc2A-dependent manner (18). Together, these data indicated that fluconazole resistance in *C. glabrata* was likely to involve coordination of the Pdr1- and Upc2A-dependent transcriptional circuits.

Given that Upc2A interfaced with the *PDR1* and *CDR1* promoters, we wanted to determine the full spectrum of genes bound and regulated by this factor. This was accomplished using chromatin immunoprecipitation coupled with Next Generation Sequencing (ChIP-seq). We also performed RNA-seq studies to identify the Upc2A-dependent transcriptome. Using the strong sequence conservation between *S. cerevisiae* Upc2 and Upc2A, we constructed a GOF form of Upc2A in *C. glabrata* based on an allele described for its *S. cerevisiae* homologue (19). This mutant Upc2A drove elevated fluconazole resistance and behaved like the hyperactive *S. cerevisiae* factor. ChIP-seq data indicated Upc2A bound to roughly 1000 genes and that this binding was
highly induced by fluconazole. Comparison of the genes bound by Upc2A with those we previously found to be associated with Pdr1 indicated a high degree of overlap between these two target gene suites. Upc2A-mediated transcription of \textit{PDR1} and \textit{CDR1} was linked with the response to anaerobic growth. Transcription of the \textit{FKS1} gene, encoding a \(\beta\)-glucan synthase protein was also found to be responsive to Upc2A, consistent with \textit{upc2A}\Delta strains being hypersensitive to caspofungin which is thought to act as a \(\beta\)-glucan synthase inhibitor (reviewed in (20)). Our data provide a new view of the global importance of Upc2A-mediated transcriptional activation as extending far beyond its well-appreciated control of \textit{ERG} gene expression. Upc2A appears to serve as a key coordinator of the biosynthesis of membrane lipids and proteins that are destined to function in this membrane environment.

Results

A gain-of-function form of Upc2A confers elevated fluconazole resistance.

Fluconazole resistance is most commonly caused in \textit{C. glabrata} by substitution mutations within the \textit{PDR1} gene (See (6, 21) for a review). These mutant transcription factors exhibit enhanced target gene expression when compared to the wild-type allele. Although mutations in the \textit{UPC2} gene in both \textit{Saccharomyces cerevisiae} and \textit{Candida albicans} have been isolated that drive fluconazole resistance (22-24), there are no gain-of-function (GOF) forms currently known for \textit{UPC2A}. To determine if a GOF allele of \textit{UPC2A} could be produced, we constructed an allele based on analogy with a mutation first found in \textit{Saccharomyces cerevisiae \textit{UPC2}} (ScUpc2) which caused the enhanced
function of this transcriptional regulator (25). The relevant mutation (*upc2-1*) (19) is a change of a glycine to an aspartate residue located at position (G888D) in the carboxy-terminus of ScUlp2. Alignment of *C. glabrata* Upc2A and ScUlp2 indicated that G898 was the analogous position in Upc2A. This residue was replaced with an aspartate to form the G898D *UPC2A* form of the gene. The resulting mutant allele was tagged with a 3X hemagglutinin (3X HA) epitope at its amino terminus as we have previously done for the wild-type *UPC2A* gene and both these forms of *UPC2A* were integrated into an otherwise wild-type *C. glabrata* strain. These tagged strains were then grown to mid-log phase along with isogenic wild-type and *upc2A*Δ cells. Serial dilutions of each culture were placed on rich medium containing the indicated concentrations of fluconazole (Figure 1A).

Introduction of the G898D mutation into *UPC2A* led to the resulting factor exhibiting elevated fluconazole resistance when compared to either the tagged or untagged version of the wild-type gene. Loss of *UPC2A* caused a dramatic increase in fluconazole susceptibility. This is consistent with G898D *UPC2A* behaving as a hyperactive transcriptional activator in *C. glabrata* as has previously been seen for G888D Ulp2 in *S. cerevisiae* (25).

To characterize the action of G898D Ulp2A in control of fluconazole resistance, the expression of a range of different genes involved in this phenotype was examined by RT-qPCR. Our previous experiments have identified Ulp2A as an inducer of expression of both the ATP-binding cassette transporter-encoding gene *CDR1* and the *PDR1* gene encoding a key transcriptional activator of *CDR1* (26). Levels of mRNA for
the ERG11 gene, encoding the enzymatic target of fluconazole, as well as for UPC2A itself were also evaluated in the presence and absence of azole drug challenge.

The presence of the G898D UPC2A gene led to a strong elevation of ERG11 transcription in the absence of fluconazole but had no significant effect on the other genes (Figure 1B). Treatment with fluconazole elevated transcription of all Upc2A target genes although the elevated ERG11 mRNA levels seen in the absence of drug were not further induced by fluconazole when G898D Upc2A was present. The fluconazole induction of both PDR1 and CDR1 was enhanced by the presence of wild-type UPC2A relative to the presence of the GOF allele. UPC2A transcription was approximately two-fold elevated by fluconazole, irrespective of the UPC2A allele tested.

These same strains were then used to compare expression of their protein products by western analysis with appropriate antibodies. All strains were grown in the absence or presence of fluconazole and whole cell protein extracts prepared. These were analyzed by western blotting using the indicated antibodies (Figure 1C and D).

The presence of the G899D UPC2A allele supported normal fluconazole induction of Cdr1 and showed a modest reduction in Pdr1 activation. The levels of the wild-type and G898D forms of Upc2A were not detectably different as shown by blotting with the anti-Upc2A polyclonal antiserum. Over the time course of fluconazole challenge (two hours), no differences in the levels of these two forms of Upc2A were seen. These data argue that the increased activation seen for ERG11 in the presence of the G898D UPC2A gene was due to increased function of Upc2A rather than a change in its expression compared to the wild-type factor.
Global analysis of Upc2A transcriptional targets. Having confirmed that the HA-tagged form of both wild-type and G898D Upc2A behaved as expected, we used these forms of Upc2A to carry out chromatin immunoprecipitation coupled with Next Generation Sequencing (ChIP-seq) to identify and compare genes that are direct transcriptional targets of Upc2A. Additionally, we assessed the effect of fluconazole induction on the suite of genes bound by either the wild-type or mutant forms of Upc2A. We also compared the binding sites found for Upc2A with those previously mapped for Pdr1 in ρ₀ cells (27) to determine the degree of overlap between these two transcriptional circuits involved in fluconazole resistance. The strains used above were grown to mid-log phase under control or fluconazole-treated conditions and fixed chromatin prepared and fragmented. Chromatin was immunoprecipitated with anti-HA antibody, purified and analyzed by Illumina sequencing. Reads were mapped and peaks called by use of the MACS2 algorithm (28). We first analyzed the peaks seen in these 4 different conditions (wild-type +/- fluconazole, mutant G898D +/- fluconazole) by determining the overlap of bound genes between them (Figure 2A).

The Venn diagram shown illustrates the extent of overlap between each different ChIP-seq reaction. The largest number of bound genes was seen for the wild-type factor in the presence of fluconazole (>1000 promoters bound). Interestingly, there were 565 of these promoters that only be bound Upc2A in the presence of fluconazole while others were bound in multiple different conditions. The next largest group of bound promoters (309) were found to be bound under all 4 different conditions. The third largest class of promoters (85) were only bound by Upc2A under conditions we consider induced (wild-type + fluconazole and G898D Upc2A +/- fluconazole). GO term
enrichment analysis of these three different classes of promoters indicated that very
different genes were associated with these patterns of Upc2A DNA-binding. The largest
class of genes was enriched for components of the translation apparatus
(Supplementary table 1A). The class of genes bound under all 4 different conditions
was most predominantly enriched for proteins associated with the plasma membrane
(Supplementary table 1B) while the inducible class of Upc2A-bound promoters showed
the strongest enrichment for genes involved in fatty acid biosynthesis (Supplementary
table 1C). While all these genes are Upc2A target loci, they exhibited unique patterns
of association with this factor.

We used the software MEME-ChIP to search the peaks associated with binding
of wild-type Upc2A in the presence of fluconazole for sequence motifs that were
enriched in this collection of binding sites. We chose this condition as it represented the
broadest collection of sterol response elements (SREs). We compared the MEME-ChIP
output to known binding sites for Upc2 in S. cerevisiae (ScUpc2) (29) and refer to the
binding elements for Upc2A in C. glabrata as SREs on the basis of their similarity to
those previously described in S. cerevisiae (30). This analysis is shown in Figure 2B.

The AR1b/c elements show the variation that is tolerated by Upc2 in S.
cerevisiae. These ScUpc2 SREs are most closely fit to the right-hand element of the
Upc2A SRE predicted by MEME-ChIP. The central CGTA sequence is conserved
between C. glabrata and S. cerevisiae, although the C. glabrata element has a nearby
conserved element (CACAGA) that shows a relatively constant spacing. It is important
to note that all analyses of S. cerevisiae Upc2 DNA binding were carried out before the
availability of global approaches like ChIP-seq which will impact predictions of
consensus elements as a more limited repertoire of regulated genes was considered. We examine the binding of Upc2A to its DNA target sites in detail below.

These data agree with the model proposed earlier that Upc2A appeared to accumulate inside the nucleus upon ergosterol limitation where the factor then binds to SRE-containing promoters (22). Since our previous work indicated overlap between Upc2A and Pdr1 target genes (26), we wanted to examine the degree of overlap between these different regulons. To make this comparison, we examined the shared Upc2A targets in cells treated with fluconazole (WTF) and the promoters bound by Pdr1 in ρ⁰ cells (27). We selected these two conditions for comparison since both involve a signal that activates the wild-type versions of Upc2A and/or Pdr1 (fluconazole or loss of the mitochondrial genome, respectively) (8, 31). Strikingly, more than 50% of Pdr1 target promoters were also associated with Upc2A binding (Figure 2C). The top 4 GO terms enriched in genes bound by both these Zn₂Cys₆-containing transcription factors were associated with transmembrane transport or integral membrane components (Supplementary table 2). As our earlier work had shown that both PDR1 and CDR1 were targets of Upc2A along with Pdr1 (18), these new data indicate that the overlap between these two transcriptional circuits extends well beyond the initial two genes.

Two different classes of Upc2A target promoters are shown in Figure 2D. The ERG11 gene is an example of a locus controlled by Upc2A but not Pdr1. Binding of wild-type Upc2A is represented by the read depth and can be seen to increase in the presence of fluconazole compared to in the absence of the drug. Binding of G898D Upc2A was constant, irrespective of the presence of the drug. Note that when the lack of a change in Upc2A expression is considered (see Figure 1C), these data support the
view that the DNA-binding activity of wild-type but not G898D Upc2A is increased by the presence of fluconazole, possibly by an increase in nuclear localization (22).

**CDR1** represents a Upc2A target gene that is also regulated by Pdr1. The bound regions for Pdr1 and Upc2A extensively overlap in the upstream region of **CDR1**. Pdr1 DNA-binding was strongly upregulated in $p^0$ cells, likely due in part to the large increase in **PDR1** expression in this background compared to wild-type cells (27). Upc2A DNA-binding to **CDR1** was regulated in a manner similar to that seen for **ERG11**.

The data above did not take into account a consideration of target gene expression. In order to link regulated Upc2A binding to changes in gene transcription of target genes, we carried out two additional analyses. First, we used software contained within the MACS2 algorithm called BDGdiff (28) that examines ChIP-seq data and identifies peaks that exhibit significant binding differences in the comparison of data from fluconazole-treated cells versus untreated cells. Secondly, RNA-seq assays were performed on isogenic wild-type and upc2AΔ cells in the presence and absence of fluconazole. RNA-seq data were processed and we focused on genes that had an adjusted P-value of <0.05 and significantly up-regulated by at least two-fold. These data are summarized in Figure 3 and included in supplementary table 3.

Figure 3A represents the union of all genes that are induced in the presence of fluconazole in either wild-type or upc2AΔ cells with genes that exhibited a significant increase in Upc2A DNA-binding when comparing ChIP-seq data from cells grown in the presence of fluconazole versus the absence of drug. Strikingly, only 53 genes required the presence of Upc2A DNA-binding to be induced by fluconazole while 274 were
induced either in the presence or the absence of the \textit{UPC2A} gene, although all these
genes were bound by Upc2A. The majority of fluconazole-induced genes (542) were
not dependent on the presence of Upc2A while 64 genes were fluconazole-induced,
required the presence of Upc2A but were not bound by this factor.

These data argue that the vast number of fluconazole-induced genes do not
depend on the presence of Upc2A. However, there are 117 genes that are only FLC-
inducible in the presence of Upc2A. The only GO term enriched in the 64 genes that
require the presence of Upc2A for FLC induction (yet are not bound by Upc2A)
represent loci involved in ergosterol biosynthesis. These Upc2A-dependent but indirect
targets include \textit{ERG4}, \textit{ERG8}, \textit{ERG9}, \textit{ERG24}, \textit{ERG26} and \textit{ERG27}. \textit{ERG8} is in the
earlier part of the ergosterol biosynthetic pathway while all other enzymes participate in
the conversion of farnesyl pyrophosphate to ergosterol (recently reviewed in (32)).

The simplest class of fluconazole-induced genes are represented by the 53 direct
target genes. GO terms enriched in this set of genes included ergosterol biosynthesis,
plasma membrane and cell wall biogenesis, membrane transport, sterol uptake and
RNA polymerase II core promoter-binding factors. The genes in this class of enriched
genes include \textit{UPC2B} and homologues of \textit{S. cerevisiae NRG2} and \textit{ADR1}. These
factors may be involved in regulation of genes that are controlled indirectly by Upc2A.

The other two categories of fluconazole-regulated Upc2A-bound genes were
either induced in both wild-type and \textit{upc2A}\textsuperscript{Δ} cells (201 genes) or only in the absence of
\textit{UPC2A} (73 genes). Only the group of 201 genes showed any significant GO term
enrichment with specific groups of genes involved in the response to stress and
glycogen catabolism being the top two categories.
The final and largest class of genes found to be bound by Upc2A were less than two-fold induced by fluconazole. GO term analysis of this class of genes indicated that the top three enriched categories were involved in translation and the ribosome (60 total). The next highest categories were plasma membrane or amino acid transmembrane transport and represented 74 genes. Together, these data strongly suggest that Upc2A impacts a wide range of cellular process as well as its well-described control of expression of genes involved in the ergosterol biosynthetic pathway.

To examine the range of expression of Upc2A-responsive genes, we compared the transcriptional response of a range of genes from the ERG pathway with loci that we have previously found to be targets of Pdr1 in C. glabrata (27). This comparison is presented in the form of a heat map (Figure 3B).

The majority of ERG genes showed at least two-fold induction by fluconazole in wild-type cells as long as these genes corresponded to steps later in the ergosterol biosynthetic pathway. ERG10, ERG13, HMG1 and ERG20, which all encode early steps in ergosterol biosynthesis, were not influenced by fluconazole challenge in wild-type cells although expression of these genes was strongly depressed under these same conditions in the absence of UPC2A. Genes encoding enzymes that function later in ergosterol biosynthesis (like ERG11) were induced by fluconazole at least two-fold in wild-type cells but depressed by at least two-fold in a upc2AΔ background.

There were two ERG genes that were exceptions to these general trends of regulation. ERG1, one of the genes showing the best induction by fluconazole in wild-type cells and most prominent peak of Upc2A ChIP-seq density, was similarly drug
induced in both wild-type and upc2AΔ cells. ERG8, encoding an enzyme that functions early in the ergosterol pathway, was also similarly fluconazole-induced in both gene backgrounds, irrespective of the presence of Upc2A. Strikingly, ERG8 was not seen to contain a detectable ChIP-seq peak for Upc2A binding. We interpret these complex responses to fluconazole treatment in C. glabrata as evidence for a multifactorial transcriptional network regulating ERG gene expression in which Upc2A participates as both a direct (later pathway genes) and indirect (early pathway genes) regulator.

Pdr1-regulated genes that were also associated with SREs exhibited less dependence on the presence of Upc2A for fluconazole induction than seen for most ERG genes. CDR1 fluconazole induction was reduced upon loss of Upc2A but PDR1 was similarly induced irrespective of the UPC2A background (Figure 3B). SNQ2 and PDH1 were reduced to approximately 50% of their normal drug-induced levels in the upc2AΔ strain while QDR2 and RTA1 showed higher levels of fluconazole induction in the same strain. Two other Pdr1 target genes (PDR16, RSB1) were repressed by the presence of fluconazole in wild-type cells and their expression lowered further when upc2AΔ cells were treated with fluconazole.

We also used RNA-seq to compare the gene expression profile of wild-type cells to an isogenic G898D UPC2A strain. These strains were grown to mid-log phase in the absence of fluconazole and then standard RNA-seq was carried out to determine the effects of this form of Upc2A on the transcriptome (Table 2).

The presence of the GOF form of UPC2A caused relatively small changes in gene transcription. There were only 11 genes observed to be elevated at least 1.4-fold. A striking feature shared by these genes was that nine of eleven encoded products that
were involved in the biosynthesis of ergosterol. Five of these 9 genes also contained SREs. As we have seen for ERG11 (Figure 1B and C), the presence of the G898D Upc2A protein led to enhanced expression of multiple genes involved in ergosterol biosynthesis. This coordinate up-regulation is likely responsible for the observed increase in fluconazole resistance caused by this allele.

Identification of functional SREs in Upc2A target genes. To confirm that the SREs present in the direct Upc2A target genes were required for normal in vivo functions, we both mapped the location of several SREs and prepared mutant versions of these sites that could not normally interact with Upc2A. DNase I protection assays with a recombinant form of the Upc2A DNA-binding domain were used to locate each SRE at nucleotide resolution. Radioactive probes were prepared from SREs contained in the ERG1, CDR1 and PDR1 promoters. These probes were used in a DNase I protection mapping experiment to locate the bounds of the region protected by Upc2A from nuclease digestion. The DNase I ladders were electrophoresed in parallel with chemical sequencing reactions on the same probe in order to locate the SRE. These data are shown in Figure 4.

The ERG1 promoter, which contains an everted pair of SREs (Figure 4A), showed the largest protected region of DNA and a strong DNase I hypersensitive site located immediately upstream of the SRE (Figure 4B). The CDR1 SRE exhibited two DNase I hypersensitive sites linked to Upc2A binding while the PDR1 SRE showed a clear protected region but no associated hypersensitive site.
Now that we could localize the SREs in each of these promoters to a relatively small segment of DNA, we mutagenized each to confirm its requirement for in vitro binding. To confirm that the predicted SREs were key for Upc2A binding, we used an electrophoretic mobility shift assay (EMSA) and prepared wild-type and mutant probes containing each SRE. Each probe was incubated with Upc2A and then resolved using non-denaturing electrophoresis. Bound and unbound probe was detected using a biotin moiety attached to the end of each probe. The results of this assay are shown in Figure 4C.

The wild-type ERG1 probe produced two different species of protein:DNA complex, possibly corresponding to either one or two binding sites being occupied with Upc2A. The mutant form of this SRE blocked formation of both complexes. CDR1 and PDR1 probes both formed primarily a single size of complex that was greatly diminished when the mutant SRE probe was used. These data argue that the SREs indicated in Figure 4A are likely required for Upc2A binding to each promoter.

Phenotypes caused by loss of SRE function. To validate the importance of the SREs identified in ERG1, CDR1 and PDR1, we prepared versions of these promoters that contained the mutations shown to block in vitro binding of Upc2A. These DNA-binding defective SREs (mSRE) were first introduced, along with their wild-type promoter, into a lacZ fusion plasmid to allow comparison of the expression supported by wild-type promoters to those that lacking Upc2A binding. These plasmids were transformed into wild-type cells, then grown to mid-log phase and challenged with or
without fluconazole. *C. glabrata* promoter-dependent β-galactosidase activity was then determined.

Introduction of the mSRE into *ERG1, CDR1* or *PDR1* promoters led to a reduction in the level of fluconazole-induced β-galactosidase activity produced by each respective fusion gene (Figure 5A). While some degree of fluconazole inducibility was retained in each mSRE-containing promoter, these data indicate that each SRE identified above is required for normal drug induced promoter activation.

To examine the effect of the loss of the SRE from the wild-type *CDR1* and *PDR1* genes, the mSRE mutations were introduced into otherwise wild-type versions of these two genes. Isogenic wild-type and mSRE versions of the *PDR1* locus were prepared by recombination into the normal chromosomal location of this gene in a strain containing a null allele of *CDR1*. Low-copy-number plasmids containing *CDR1* were constructed that varied only by the form of the SRE that was contained in the promoter region. These two different forms of *CDR1* were introduced into the wild-type and mSRE *PDR1 cdr1Δ* strain and transformants grown in the presence or absence of fluconazole. Whole cell protein extracts were prepared and examined for expression of proteins of interest using appropriate polyclonal antisera (Figure 5B).

Loss of the SRE from the *CDR1* promoter caused a significant drop in expression when cells were treated with fluconazole that was enhanced when combined with the mSRE version of the *PDR1* gene. Similar reductions in Cdr1 levels were seen in the absence of fluconazole, again with removal of the SRE from both *CDR1* and *PDR1* causing the largest reduction in Cdr1 levels. Expression of Pdr1 was not affected when the *CDR1* SRE was removed but fluconazole induction of Pdr1 was reduced when the
SRE was removed from the \textit{PDR1} promoter. Expression of Erg11 was unaffected in these backgrounds as these alterations were restricted to the \textit{CDR1} and \textit{PDR1} promoters.

These strains were also evaluated for their drug resistance phenotype using a serial dilution assay on fluconazole-containing media (Figure 5C). The major reduction in fluconazole resistance was caused by the presence of the mSRE-containing form of the \textit{CDR1} gene. This was modestly enhanced by the simultaneous loss of the SRE from the \textit{PDR1} gene. Together, these data demonstrate that the SREs present in \textit{CDR1} and \textit{PDR1} are required for normal expression of these genes and for full fluconazole resistance.

\textbf{Potential role for Pdr1/\textit{CDR1} in anaerobic growth.} The above data continue to support the notion that the Pdr pathway in \textit{C. glabrata} is linked to levels of ergosterol as we have argued previously (18). We wondered if a physiological parameter, key for ergosterol biosynthesis, might also involve the Pdr pathway. To test this possibility, we examined the effect of anaerobic growth on \textit{PDR1} and \textit{CDR1} expression. Oxygen is essential for normal ergosterol biosynthesis and anaerobic growth triggers rapid induction of \textit{ERG} gene expression in several fungi, including \textit{C. glabrata} (33). We also evaluated the requirement for the presence of \textit{PDR1} and \textit{CDR1} during anaerobic growth to test their contribution to this phenotype.

We validated our conditions of anaerobic growth by measuring the expression of several genes known from work in \textit{S. cerevisiae} to be induced under these conditions, along with others that were repressed in the absence of oxygen (34). Wild-type cells
were grown aerobically or anaerobically, total RNA prepared and levels of mRNAs assessed by RT-qPCR analyses.

Anaerobic growth led to 20-fold or higher induction of the genes corresponding to products important in sterol uptake such as the ABC transporter Aus1 and other proteins thought to be essential for this process (Dan1, Tir1) (Figure 6A). Anaerobic growth also repressed expression of genes encoding mitochondrial proteins involved in ATPase production (Atp3, Atp4) and an electron transport chain component (Sdh2).

Two different loci encoding enzyme involved in the tricarboxylic acid cycle were slightly reduced (KGD2) or unaffected (ACO1).

Having confirmed that the expected anaerobic gene regulation was seen under our growth conditions, we next tested expression of Cdr1, Pdr1 and Erg11 using the western blot assay described previously (Figure 6B). Both Cdr1 and Pdr1 were strongly induced, from 3- to 5-fold, while Erg11 was also induced albeit roughly 2-fold.

Since all these Upc2A target genes were induced in this assay, we wanted to determine if the SREs associated with CDR1 and PDR1 were required for this anaerobic activation. We used the strains described early in which either wild-type versions of CDR1 and PDR1 were present or these same genes containing mutations in their respective SREs were used. These isogenic strains were grown in the presence or absence of oxygen and levels of Cdr1, Pdr1 and Erg11 (as a control for anaerobic conditions) measured by western blotting.

Anaerobic induction of both Cdr1 and Pdr1 was diminished in the absence of the SRE motifs in their promoters (Figure 6C). As seen before for fluconazole challenge,
the SREs in the *CDR1* and *PDR1* promoters are required for normal induction. Erg11 was not affected as its SRE was unaltered in this experiment.

To test if the expression of *CDR1* and *PDR1* was involved in normal anaerobic growth, we carried out a competitive growth assay. Isogenic wild-type and double mutant *cdr1Δ pdr1Δ* strains were mixed in equal portions and then grown under either aerobic or anaerobic conditions. Once these mixed cultures had reached the end of log phase growth, aliquots were plated to determine if the percentage of the populations had changed.

The loss of *CDR1* and *PDR1* caused the resulting double mutant strain to exhibit a growth disadvantage when competed with the wild-type parent (Figure 6D) but only under anaerobic conditions. No change from the starting population was seen during aerobic growth. These data provide evidence that Upc2A-regulated expression of both *CDR1* and *PDR1* is required for normal anaerobic growth, linking the function of these two genes to this phenotype for the first time.

**Role for Upc2A in caspofungin resistance.** The ChIP-seq data predicted a potential SRE upstream of the *FKS1* and *FKS2* genes (Supplementary tables 1 and 2). To determine if these putative SREs had detectable roles in expression of the caspofungin resistance phenotype, we tested the ability of an isogenic set of strains varying in their *UPC2A* allele for the response to several different cell wall stress agents. Isogenic wild-type, *upc2AΔ* or epitope-tagged wild-type or G898D *UPC2A*-containing strains were tested for resistance to caspofungin, caffeine or high pH using a serial dilution assay. Caffeine and high pH are cell wall stresses and reflect general cell wall dysfunction (35).
Loss of *UPC2A* caused hypersensitivity to all these agents (Figure 7A) while both epitope-tagged alleles behaved like the wild-type strain. The finding of a caspofungin susceptible phenotype prompted us to examine expression of the three *FKS* genes in *C. glabrata* to determine if any of these showed a response to the G898D allele of *UPC2A*. None of these genes were altered in the presence of this gain-of-function form of *UPC2A* while both *ERG1* and *AUS1* were elevated (Figure 7B), confirming the functionality of this hypermorphic form of Upc2A.

To explain the observed caspofungin hypersensitivity of the *upc2A*Δ strain, levels of *FKS1* and *FKS2* mRNAs were measured using RT-qPCR in the presence or absence of caspofungin. Loss of *UPC2A* reduced basal expression of *FKS1* and had a modest effect on *FKS2* (Figure 7C). The addition of caspofungin strongly induced *FKS2* expression as expected (36) with this induction unaffected by the absence of Upc2A.

Since a defect was seen for *FKS1* expression, we prepared a DNA probe containing the putative SRE in this promoter for use in an EMSA to determine if recombinant Upc2A was able to recognize this element in vitro (Figure 7D). A mutant form of this SRE was also tested in this EMSA. The wild-type *FKS1* probe was strongly reduced in mobility when incubated with Upc2A while the mSRE-containing probe exhibited a band of reduced intensity upon loss of this sequence element.

To determine if the SRE was required for normal expression of *FKS1*, a *lacZ* translational fusion gene was prepared in which the *FKS1* regulatory region determined expression of β-galactosidase. Both the wild-type and mSRE-containing *FKS1* promoters were used and introduced on a low-copy-number plasmid into wild-type *C.*
glabrata cells. FKS1-dependent β-galactosidase activities were then determined in the presence or absence of caspofungin induction.

Loss of the SRE from the FKS1 promoter caused a significant reduction in FKS1-dependent expression of lacZ in the absence of caspofungin (Figure 7E). These data provide evidence that Upc2A-mediated gene activation is required for normal expression of FKS1 and wild-type caspofungin resistance.

Discussion

These data provide important new appreciation for the expansive role of Upc2A in control of gene expression. Extensive previous work on Upc2A homologues in both S. cerevisiae and C. albicans was generally done prior to the availability of modern genomic approaches like ChIP- and RNA-seq (reviewed in (37)). Detailed analyses demonstrated the crucial role of these Upc2A-like factors in regulation of ERG gene biosynthesis (30, 38, 39)) but little was known about the full range of their target genes. A ChIP-chip experiment was carried out on C. albicans Upc2 and this factor was found to associate with the CDR1 gene in this species (40). To the best of our knowledge, there has been no follow-up linking C. albicans Upc2 with the Tac1 transcription factor (key regulator of CDR1 transcription) or S. cerevisiae Upc2 with either ScPDR1 or ScPDR3. This suggests the possibility that this C. glabrata connection between Upc2A and Pdr1 is a unique feature of this yeast and could help explain the high level of intrinsic azole resistance seen in this organism.

The large number of Upc2A target genes illustrates the breadth of processes that are transcriptionally influenced by this factor. Clearly, the ERG genes are an important
set of genetic targets but these are a small fraction of the whole. Upc2A appears to be coordinating a broad group of genes including a large number of plasma membrane-localized proteins (See supplementary table 1 and 3). Coupled with its control of ergosterol in this membrane, Upc2A appears to be a central determinant of the composition of this membrane compartment in *C. glabrata*. The regulation of plasma membrane constituents is of obvious importance in modulating the ability of substances to cross this barrier between the external and internal environments.

Comparing the members of the target gene sets defined by Upc2A and Pdr1 suggests a hierarchical relationship between these two transcription factors. Here we establish that a binding site for Upc2A lies upstream of *PDR1* and is required for normal activation of *PDR1* expression (Figure 6C) as well as many of the other genes controlled directly by Pdr1. We suggest Upc2A provides overarching control of both the Pdr1 regulon but also a variety of other genes that are not under Pdr1 control, serving to link these different classes of genes through this common transcriptional regulation.

While the full range of Upc2A target genes illustrate the global importance of this transcription factor, the *ERG* genes are especially sensitive to the level of activity of this regulator. The G898D *UPC2A* allele has a surprisingly limited effect on gene expression as this allele was seen to trigger significant transcriptional changes almost exclusively in genes associated with ergosterol biosynthesis (Table 2). It is possible that the transcriptional impact of the G898D Upc2A could be expanded if cells were treated with fluconazole as ergosterol limitation might be able to impact expression by regulatory inputs beyond Upc2A. Experiments to test this possibility are underway.
Construction and analysis of G898D Upc2A demonstrated that there is no particular prohibition on hyperactive alleles of \textit{UPC2A} existing in this pathogenic yeast. The \textit{S. cerevisiae UPC2-1} allele, from we derived the G898D \textit{UPC2A}, was originally isolated on the basis of permitting aerobic sterol uptake (19). Strikingly, no other \textit{S. cerevisiae UPC2} hypermorphic alleles are known. \textit{UPC2} mutations in \textit{C. albicans} have been found in multiple clinical strains and appear to be much more commonly isolated (23, 24). An interesting feature of the majority of \textit{C. albicans UPC2} GOF alleles is these all cluster with a region of the protein between residues 642 and 648 (16). This region shows strong sequence conservation with the 893-898 region of \textit{C. glabrata} Upc2A. The conserved location of these hypermorphic alleles suggests the possibility that a common function is being disrupted in both organisms.

Based on detailed structural and subcellular localization studies on \textit{S. cerevisiae} Upc2 (22), we propose that \textit{C. glabrata} Upc2A accumulates in the nucleus upon ergosterol limitation. Our data support this assertion in two ways. First, a large increase in ChIP-seq peaks is seen for Upc2A when azole-treated cells are compared to untreated cells. Second, the G898D Upc2A mutant protein shows constitutively high number of ChIP-seq peaks that is not significantly altered by azole challenge. These data are consistent with a model in which Upc2A nuclear accumulation is enhanced upon ergosterol limitation and this regulation requires the function of the region containing G898 in the C-terminus of this factor.

The finding of the interaction between Upc2A and the \textit{FKS1} gene provides an interesting connection between azole resistance, well-known to be impacted by Upc2A (41), and echinocandin resistance. These two antifungal drugs have been considered
to be defined by genetically separable pathways but here we provide evidence that

Upc2A may provide a link between them. Intervention in Upc2A-mediated
transcriptional activation may be able to cause reductions in resistance to both azole
drugs and the echinocandins.

Finally, our data also illuminate the complexity and interrelationship of expression
of genes involved in ergosterol biosynthesis with plasma membrane proteins and even
the cell wall. ERG gene regulation is an important task for Upc2A but this factor clearly
impacts transcription of a broad range of genes affecting multiple aspects of the plasma
membrane. Additionally, loss of Upc2A has clear phenotypes but is certainly not the
only regulator of ERG gene expression and fluconazole induction in C. glabrata. Loss
of UPC2A leads to a profound increase in fluconazole susceptibility, even in the
presence of a GOF form of PDR1 (17). However, only ~50 genes were both bound by
Upc2A and dependent on Upc2A for fluconazole induction while 880 were induced in
the presence of fluconazole independent of Upc2A (Figure 3A). This provides an
illustration of the overlapping modes of regulation controlling gene expression in
response to ergosterol limitation. The importance of ergosterol production and its
synchronization with biogenesis of membrane proteins in the plasma membrane is
central to a fungal cell producing a normally functioning membrane that can allow
growth. Understanding this regulatory circuitry will allow interventions to be developed
that can restore and potentially even enhance azole susceptibility, allowing the use of
this highly effective antifungal drug to be maintained.
Materials and Methods

**Strains and growth conditions.** *C. glabrata* was routinely grown in rich YPD medium [1% yeast extract, 2% peptone, 2% glucose] or under amino acid-selective conditions in complete supplemental medium (CSM) (Difco yeast nitrogen extract without amino acids, amino acid powder from Sunrise Science Products, 2% glucose). All solid media contained 1.5% agar. Nourseothricin (Jena Bioscience, Jena, Germany) was supplemented to YPD media at 50 μg/ml to select for strains containing the pBV133 vector (26) and its derivatives. All strains used in this study are listed in table 1.

**Plasmid construction and promoter mutagenesis.** All constructs used for homologous recombination into the chromosome were constructed in a pUC19 plasmid vector (New England Biolabs, Ipswich, MA). PCR was used to amplify DNA fragments and Gibson assembly cloning (New England Biolabs) employed to assemble fragments together. All isogenic deletion constructs were made by assembling the recyclable cassette from pBV65 (26) and fragments from the immediate upstream/downstream regions of the target genes. Eviction of the recyclable cassette left a single copy of loxP in place of the excised target gene coding region. Sequences of the repeated influenza hemagglutinin epitope tag (3X HA) was PCR amplified from BVGC3 background (26). This tag element was inserted before the start codon of *UPC2A* and G898D *UPC2A* with an addition of repeated 3X glycine-alanine linker sequence located between the 3X HA tag and the gene coding sequence. The G898D mutation in *UPC2A* was made by
Gibson assembly in which the overlapping primers contained the point mutation sequence.

Gene complementation constructs were made by Gibson assembling the fragments from the immediate upstream and downstream regions of the target genes [overlapping regions], coding region of the target genes, target gene terminators (about 250 base-pairs after the translation stop codon), and the recyclable cassette (located after the terminator). Eviction of the recyclable cassette in the complementation constructs left a single copy of loxP about 250 base-pairs downstream of the target gene stop codons. Complementation of LEU2 was done by PCR amplifying the LEU2 coding region and 500 base-pairs immediate upstream and downstream of the coding region from the CBS138 background. Linear DNA was then transformed into KKY2001 and the colonies were selected on CSM agar without Leucine.

All autonomous plasmids were derived from pBV133 (26) carrying nourseothricin marker. The lacZ gene encoding the E. coli β-galactosidase gene was amplified from pSK80 (42). ERG1, CDR1, PDR1 promoter fragments were amplified from the KKY2001 background. The CDR1 minimal promoter, which was fused to lacZ, contained the -1 to -1076 region (with the ATG of CDR1 considered as +1). The full CDR1 promoter, which was used in the complementing plasmid, contained the -1 to -1504 region. The PDR1 promoter contained the -1 to -847 region in all constructs. ERG1 promoter region consisted of the -1 to -916 region and the FKS1 promoter contained the -1 to -1795 region. SRE mutations in the target gene promoters were
done by modifying the SRE core sequence and 2 adjacent bases into a PacI restriction enzyme sequence with Gibson assembly in which the overlapping primers contained the PacI sequence. All constructs were verified by Sanger sequencing (University of Iowa Genomic Core).

**C. glabrata transformation.** Cell transformations were performed using a lithium acetate method (43). After being heat shocked, cells were either directly plated onto selective CSM agar plates (for auxotrophic complementation) or grown at 30°C at 200 rpm overnight (for nourseothricin selection). Overnight cultures were then plated on YPD or CSM agar plates supplemented with 50 μg/ml of nourseothricin. Plates were incubated at 30°C for 24 to 48 h before individual colonies were isolated and screened by PCR for correct insertion of the targeted construct.

**Expression and purification of Upc2A DNA binding domain.** The DNA sequence corresponding to the first 150 amino acids of the N-terminus of Upc2A was amplified by PCR and cloned into pET28a+ vector [digested with NcoI and ScaI] with Gibson cloning. Correct clones were sequenced verified and transformed into the BL21 DE3 *E. coli* expression strain (Thermo Fisher, Waltham, MA). Mid-log phase cells were induced with 1 mM IPTG (Fisher Scientific, Hampton, NH) for 4 hours at 30°C. Collected cells were lysed using a French Press G-M high pressure disruptor (GlenMills, Clifton, NJ). The clarified lysate was subjected to Talon Metal Affinity column (Takara, Mountain View, CA) as per the manufacturer’s protocol. Purified protein was dialyzed with dialysis buffer.
and its concentration was quantified by Bradford assay (Bio-Rad, Des Plaines, IL).

**Quantification of transcript levels by RT-qPCR.** Total RNA was extracted from cells
by extraction using TRIzol (Invitrogen, Carlsbad, CA) and chloroform (Fisher Scientific, Hampton, NH) followed by purification with RNeasy minicolumns (Qiagen, Redwood City, CA). RNA was reverse-transcribed using an iScript cDNA synthesis kit (Bio-Rad, Des Plaines, IL). Assay of RNA via quantitative PCR [qPCR] was performed with iTaq universal SYBR green supermix (Bio-Rad). Target gene transcript levels were
normalized to transcript levels of 18S rRNA during fluconazole challenge and β-tubulin mRNA in other conditions. Primer sequences were listed in supplementary table 4.

**Spot test assay.** Cells were grown in YPD medium to mid-log-phase. Cultures were then 10-fold serially diluted and spotted onto YPD agar plates containing different concentrations [10 or 20 µg/ml] of fluconazole (LKT laboratories, St Paul, MN), caspofungin 100 ng/ml (Apexbio, Houston, TX), congo red 100 µg/ml (Sigma-Aldrich, St. Louis, MO), caffeine (Sigma-Aldrich). In some experiments, the YPD medium and agar was supplemented with 50 µg/ml nourseothricin to maintain plasmids derived from the pBV133 vector (26). All agar plates were incubated at 30°C for 24 to 48 h before imaging was performed. To adjust the pH level, 100 mM of MES (VWR, Radnor, PA) [pH 5.5], HEPES (RPI) [pH 7.0], and TAPS (Sigma-Aldrich) [pH 8.0] were added to the 2x YPD. Solutions were then filtered and mixed with autoclaved 3% agar to make YPD agar plates.
**Competitive growth assay.** The LEU2 coding region along with its immediate 500 bps upstream and downstream sequences were amplified from CBS138 genomic DNA. The product was then used to restore LEU2 in mSRE PDR1/mSRE CDR1 [KKY2001 background] and pdr1Δ/cdr1Δ at its native locus to generate mSRE PDR1/mSRE CDR1/LEU2 and pdr1Δ/cdr1Δ/LEU2. In the fluconazole competitive growth assay, mid-log growth culture [between OD of 1 and 2] was diluted to 0.5 O.D. in fresh YPD. 1:1 ratio of PDR1 CDR1 and mSRE PDR1/mSRE CDR1 cultures were mixed together, and treated either with fluconazole [20 μg/ml] or ethanol. At each time point, culture was collected, serially diluted, and plated on YPD and synthetic complete media without leucine for CFU determination.

In the anaerobic growth competitive assay, KKY2001 and its isogenic pdr1Δ cdr1Δ derivative were diluted to 0.01 O.D. and the mixed culture [1:1 ratio] was grown either aerobically [normoxic] or anaerobically [in a GasPak chamber (BD Biosciences, San Jose, CA)] for 24 hours. 1 mM Acetyl CoA (Sigma-Aldrich), 1% Tween80/Ethanol [1:1 ratio], 1 mM Squalene (VWR), 50 μM Lanosterol (Sigma-Aldrich), 50 μM Ergosterol (Sigma-Aldrich) were added to the mixed culture before the incubation. Acetyl CoA stock solution was made in sterile H₂O, and all sterol stock solutions were made in Tween80/Ethanol (1:1 ratio).

**Electrophoretic mobility shift assays (EMSA).** DNA probes were amplified by PCR with biotinylated primers (IDT, Coralville, IA) corresponding to the sequences listed in
S2 table. Fragments from the **ERG1** promoter -704 to -916, **CDR1** promoter -560 to -731, **PDR1** promoter -552 to -728, **FKS1** promoter -1489 to -1666, and **HO** promoter -787 to -957 regions were amplified. Reaction buffer [18 µl], containing 5 µg sheared salmon sperm DNA (Thermo Fisher Scientific, Waltham, MA), 5% Glycerol, 0.01% NP40, 0.1% bovine serum albumin (Thermo Fisher Scientific), and 2 µl of 10x binding buffer [100 mM Tris pH 7.5, 400 mM NaCl, 10 mM DTT and 100 µM ZnSO₄], was incubated with different concentrations of Upc2A-6X His or 1X binding buffer for 10 minutes at room temperature. Biotinylated probes [20 fmol] were added in a final reaction volume of 20 µl and incubated for additional 20 min at room temperature. Samples were immediately subjected to electrophoresis on 5% polyacrylamide Tris/Borate/EDTA [TBE] gel in 0.5x TBE running buffer at 4°C. Subsequently, samples were transferred into a nylon membrane (GE, Chicago, IL) in 0.5X TBE buffer at 4°C. Samples were then crosslinked on nylon membrane under UV light for 10 min. Membrane was blocked with Intercept blocking buffer (LI-COR Biosciences, Lincoln, NE) containing 1% SDS for 30 min before IRDye 680LT Streptavidin (LI-COR Biosciences) antibody was added at 1:20000 final dilution. After 35 min of incubation, the membrane was washed three times with phosphate buffer saline [PBS] containing 0.1% tween (RPI). Imaging was performed with Odyssey CLx Imaging System (LI-COR Biosciences) and analyzed by Image Studio Lite Software (LI-COR Biosciences).

**Chromatin immunoprecipitation-Next Generation Sequencing.** Overnight cultures were inoculated at 0.1 OD/ml in fresh YPD and allowed to grow to 0.4-0.5 OD/ml. Cells were treated with Fluconazole 20 µg/ml or ethanol for 2 hours. Cells were fixed with 1%
formaldehyde (Sigma-Aldrich) for 15 min at room temperature with mild shaking [120-150 rpm]. The fixing reaction was stopped with 250 mM glycine (RPI) for 15 min at room temperature with mild shaking [120-150 rpm]. Cells were centrifuged and washed once with PBS. The cell pellet was resuspended in lysis buffer [50mM Hepes pH 7.5, 140mM NaCl, 1mM EDTA, 1% Triton X-100, 0.1% Sodium deoxycholate, 1mM PMSF, 1x protease inhibitor cocktail (Roche Applied Science, Penzberg, Germany). Cells were then lysed with 1 ml glass beads (Scientific Industries Inc, Bohemia, NY) at 4°C for 10 min. Both cell lysate and debris were collected and subsequently transferred to an AFA fiber pre-slit snap-cap [6 x 15mm] microtube (Covaris, Woburn, MA) for additional cellular lysis and DNA shearing.

Genomic DNA was sheared with E220 focused-ultrasonicator (Covaris) [peak incident power: 75 W, duty factor: 10%, cycles per burst: 200, treatment time: 16 min, temperature: 10°C max, sample volume: 130 µl.] The sheared sample was centrifuged and the clear lysate was collected. Upc2A was immunoprecipitated with Dyna beads-protein G magnetic beads (Invitrogen, Carlsbad, CA) and anti-HA antibody (Invitrogen) [1:50 dilution] overnight at 4°C. Beads were then washed twice with lysis buffer, once with lysis buffer+ 500mM NaCl, once with LiCl buffer [10 mM Tris pH 8, 250 mM LiCl, 0.5% P-40, 0.5% Sodium deoxycholate, and 1 mM EDTA], and once with Tris-EDTA buffer. Beads were resuspended in TE and treated with RNAse A (Thermo Fisher Scientific) at 37°C for 30 min. Beads were then washed once with Tris-EDTA buffer, resuspended in Tris-EDTA buffer with 1% SDS, and incubate at 65°C for at least 5 hours to reverse crosslink. Eluted DNA was subsequently purified with mini elite clean-
up kit (Qiagen). Qubit fluorometric assay (Thermo Fisher Scientific) was used to analyze
the yield quantity and Agilent Bioanalyzer (Agilent Technology, Santa Clara, CA) was
used to determine the average sheared DNA size.

All ChIPed DNA libraries were generated with Accel-NGS plus DNA library kit (Swift
Biosciences, Ann Arbor, MI) according to the manufacturer’s instructions. Samples
were sequenced at the University of Iowa Institute for Human Genetics Genomics
Division using an Illumina NovaSeq 6000 instrument. Read quality was confirmed using
FastQC (Babraham Bioinformatics). The reads from duplicate experiments were
combined and mapped to the C. glabrata CBS138 genome using HISAT2 (44). Paired
reads with intervening fragments greater than 1000 bp were removed during the
mapping. The total number of mapped reads were reduced by randomly selecting 2.5%
from each bam file which were then sorted and indexed using Samtools. ChiP-seq
peak calling was done with the callpeak function of MACS2 using a false discovery rate
(q-value) cutoff of 0.001 and a maximum allowable gap between peaks of 100 bp (28).
Differential peak detection among the experimental conditions was done using the
bdgdiff function of MACS2. The default likelihood ratio cutoff of 1000 was used. Output
files from both callpeak and bdgdiff were annotated to identify candidate downstream
genes using the ChIPpeakAnno R package (45).

RNA-sequencing. A single colony of each C. glabrata strain was used to inoculate 2
ml of YPD, which was grown overnight at 30° C in an environmental shaking incubator.
Cell density was then adjusted to OD_{600}=0.1 in 10 ml YPD, and cultures were grown as
before for 6 hrs (mid-log phase). For the fluconazole-treated strains, either fluconazole (50 μg/ml final concentration) or DMSO (diluent control) was added to the 10 ml culture and grown for 6 hrs. Cells were collected by centrifugation, supernatants discarded, and cell pellets stored at -80° C. RNA was isolated from cell pellets via a hot phenol method as described previously (46). The quantity and purity of RNA were determined by spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE) and verified using a Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA). Library preparation and RNA sequencing analysis were performed as previously described (47). Transcript quantification of expression levels and analysis of differential expression were done using HISAT2 and Stringtie (48). Differential expression was analyzed using DESeq2 (49).

**DNase I protection assay.** DNA probes were generated by PCR. To generate a 5′ [γ-32P] singly end-labelled probe one of the PCR primers was modified [5 Amino-MC6 (Integrated DNA technologies, Coralville, IA)] at the 5′ end to prevent phosphorylation by polynucleotide kinase. Probes were end-labelled [1 pmol] using 10 μCi of [γ-32P]-ATP (PerkinElmer, Waltham, MA) and 10 U polynucleotide kinase (New England Biolabs, Beverly, MA) as instructed by the manufacturer. Unincorporated [γ-32P]-ATP was removed using a nucleotide removal column (Qiagen). The binding reaction was done as described in the EMSA section, and the sample was digested with DNase I (NEB [1:20 dilution]) for 30 seconds at room temperature. DNase I foot-printing and DNA sequencing reactions were performed as previously described (50).
β-galactosidase assay. Harvested cells were lysed with glass beads (Scientific Industries Inc) in breaking buffer [100 mM Tris pH8, 1 mM Dithiothreitol, and 20% Glycerol] at 4°C for 10 min. Lysate was collected and β-galactosidase reactions carried out in Z-buffer [60 mM Na₂HPO₄, 40 mM NaH₂PO₄, 10 mM KCl, 1 mM MgSO₄, 50 mM 2-Mercaptoethanol] with 650 µg/ml O-nitrophenyl-β-D-galactoside [ONPG]. Miller units were calculated based on the equation: [OD₄₂₀ x 1.7] / [0.0045 x total protein concentration x used extract volume x time]. The Bradford assay (Bio-Rad) was used to measure the total protein concentration in the lysate.

Western immunoblot. Cells were lysed with lysis buffer [1.85 M NaOH, 7.5% 2-Mercaptoethanol]. Proteins were precipitated with 50% Trichloroacetic acid and resuspended in Urea buffer [40 mM Tris pH8, 8.0 M Urea, 5% SDS, 1% 2-Mercaptoethanol]. Cdr1, Pdr1, and Upc2A rabbit polyclonal antibodies were previously described (18, 27). Mouse anti-HA monoclonal antibody was purchased from Invitrogen. Secondary antibodies were purchased from LI-COR Biosciences. Imaging was performed with Odyssey CLx Imaging System (LI-COR Biosciences) and analyzed by Image Studio Lite Software (LI-COR Biosciences). Detected target band fluorescence intensity was normalized against tubulin fluorescence intensity and compiled from 2 biological replicate experiments and 2 technical replicates in each experiment, giving 4 replicates in total.

Statistical analysis. Unpaired T-test was used to compare between isogenic deletion mutant and wildtype strains. Paired T-test was used to compare between the drug
treated and non-treated conditions. *, **, *** were designated for $P \leq 0.05, 0.01, 0.001$
respectively.

Acknowledgements

We thank Dr. Damian Krysan for helpful discussions.

References


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Figure Legends

Figure 1. **Characterization of UPC2A gain-of-function allele.** A. An isogenic series of strains was prepared that varied in their UPC2A allele: wild-type (CBS138), upc2AΔ, or strains containing an amino-terminally HA-tagged form of wild-type (3X HA-UPC2A) or the G898D form of UPC2A (3X HA-G898D UPC2A). These strains were grown to mid-log phase and then tested for their resistance to the indicated levels of fluconazole in rich medium. B. G898D Upc2A induced ERG11 mRNA under basal conditions. The strains described in A were grown in the presence or absence of fluconazole and total RNA prepared. Levels of mRNA were assessed by RT-qPCR. C. Western blot analysis of Upc2A and target gene-encoded proteins. The indicated strains were grown as described previously in the presence or absence of fluconazole, whole cell protein
extracts prepared and analyzed by western blotting using the antisera listed at the left side. Upc2A was detected using either an anti-Upc2A polyclonal antibody (α-Upc2A) or anti-HA (α-HA). Tubulin was used as a loading control along with Ponceau S staining of the membranes. Quantitation shown in the right hand panel was performed as described in Materials and Methods.

Figure 2. Chromatin immunoprecipitation-high throughput sequencing (ChIP-seq) analysis of UPC2A. A. 3X HA-tagged forms of either the wild-type (WT) or G898D (M) alleles of UPC2A were used for a standard ChIP-seq experiment. Strains were in the presence (WTF, MF) or absence (WT, M) of fluconazole, followed by ChIP-seq analysis as described earlier (27). A Venn diagram showing the overlap of genes detected in each sample is shown with largest number of genes color-coded from dark to light. B. MEME-ChIP analysis of sterol response element (SREs) shared in Upc2A peaks. A logo is shown representing the most commonly enriched element associated with Upc2A ChIP-seq peaks that was detected by MEME-ChIP analysis. AR1b and AR1c show the corresponding Upc2 consensus sites from Saccharomyces cerevisiae (29). C. Venn diagram showing the overlap between binding sites found for fluconazole-stressed 3X HA-UPC2A (WTF) and ρ0-induced wild-type Pdr1 (27). D. Comparison of Upc2A- and Pdr1-binding to the CDR1 and ERG11 promoters. Plots of relative ChIP-seq density are shown for ChIP reactions performed with either anti-Pdr1 or α-HA to detect epitope-tagged Upc2A. ChIP in samples treated with fluconazole is denoted +F while the corresponding no drug control is represented by -F. The bottom
line is a control ChIP reaction using wild-type cells that lack the HA-epitope tag (Upc2A). CDR1 is controlled by both factors while ERG11 only responds to Upc2A.

Figure 3. RNA-seq analysis of fluconazole induced genes in wild-type and upc2AΔ strains. A. Venn diagrams illustrating the union of genes that are at least two-fold induced in wild-type (2X Induced WTF) or isogenic upc2AΔ cells treated with fluconazole along with genes containing a Upc2A SRE that is exhibits fluconazole inducible binding compared to untreated samples (WTF over WT). B. Heat map of representative fluconazole induced genes is shown. The values refer to the log2 score of the ratio of RPKM of fluconazole treated over untreated samples. The scale for the heat map is indicated at the bottom and the presence of a SRE is denoted by the solid dot in the lefthand column.

Figure 4. DNA-binding of Upc2A to SREs in target genes. A. Locations of SREs from several different promoters are shown. ERG1, CDR1, PDR1 and FKS1 all correspond to the C. glabrata locus while ScERG2 is from S. cerevisiae and CaERG2 is from C. albicans. Note that ERG1 contains an everted SRE repeat indicated by the arrows. The single SRE is shown in black and white. The extent of DNA protected from cleavage by the DNase I mapping experiment (see below) is shown in gray. B. DNase I protection of the indicated C. glabrata promoters is shown. The position of each SRE is indicated by the bar at the righthand side and DNase I hypersensitive sites are noted by the arrows. AG refers to the purine-specific reaction of Maxam-Gilbert chemical sequencing and is carried out on the same radioactive DNA fragment used in the
DNase I reaction. Recombinant Upc2A was added (+) to the DNA probe or omitted (-) as indicated. C. Electrophoretic mobility shift assay (EMSA) analysis of Upc2A binding to wild-type and mutant SREs. Biotinylated probes were prepared from the indicated C. glabrata promoter regions containing either wild-type (wt) or mutant (mut) versions of each SRE. Sequences of these different SREs are shown at the bottom of the panel with the altered residues in lower case. The SRE repeats in the ERG1 promoter are shown by the divergent arrows at the top of the sequence. Position of the shifted protein:DNA complexes are shown by the arrows at the lefthand side of each image. The presence or absence of Upc2A protein is indicated by the (+) or (-), respectively.

Figure 5. Phenotypes of SRE mutations. A. Normal expression of lacZ gene fusions requires the presence of intact SREs in Upc2A target promoters. Low-copy-number plasmids containing translational fusions between ERG1, CDR1 and PDR1 5’ regulatory region and E. coli lacZ were generated containing either the wild-type version of each promoter or the same fragment with the SRE mutant (mSRE) shown to reduce in vitro binding in Figure 4. These plasmids were introduced into wild-type cells, grown in the absence or presence of fluconazole and then β-galactosidase activity determined. B. Western blot analysis of CDR1 and PDR1 expression upon loss of the wild-type SRE. All alleles of PDR1 were integrated into the chromosome while all alleles of CDR1 were carried on a low-copy-number plasmid. The presence of either the wild-type (wt) or mutant (m) SRE at each gene is indicated at the bottom of the panel. Each strain was grown in the presence or absence of fluconazole and levels of proteins of interest determined using western blotting with appropriate antibodies as described above.
Erg11 was detected using an anti-peptide antiserum. The right hand panel shows the quantitation as described in Figure 1. C. The strains described above were tested by serial dilution for their growth on rich medium (YPD) or the same medium containing fluconazole (FLC).

Figure 6. Role for \textit{CDR1} and \textit{PDR1} in anaerobic growth. A. Wild-type cells were grown under aerobic or anaerobic conditions and total RNA prepared. Levels of the mRNA corresponding to the indicated genes were determined using RT-qPCR. B. Western blot analysis of the indicated proteins was performed using appropriate antisera described earlier. Whole cell protein extracts were prepared after growth in the presence of oxygen (aerobic) or its absence (anaerobic). Quantitation is shown in the right hand panel. C. Expression of \textit{CDR1} and \textit{PDR1} in response to aerobic versus anaerobic growth was assessed by western blotting. Strains containing these genes under control of their wild-type promoter regions or an isogenic strain with both SREs removed from these promoters were used to prepare whole cell protein extracts. Quantitation of at least three western blot experiments is shown on the righthand side of the panel. D. Competitive growth assay between wild-type and \textit{cdr1}\textsuperscript{Δ} \textit{pdr1}\textsuperscript{Δ} double mutant strains was performed. These two strains were mixed at a starting proportion of 50:50 and then grown in rich medium under aerobic or anaerobic conditions. After growth, the final proportion was determined by plating cells and counting the number of leucine prototrophic colonies (specific for \textit{cdr1}\textsuperscript{Δ} \textit{pdr1}\textsuperscript{Δ} cells containing \textit{LEU2}).
Figure 7. **Upc2A transcriptionally regulates FKS1 expression.** A. An isogenic series of strains with the listed **UPC2A** genotypes was tested for resistance to the indicated stress agents that affect the cell wall. Strains were grown to mid-log phase and then serially diluted across each plate. B. Strains containing the different alleles of **UPC2A** were grown and analyzed for levels of the indicated RNAs using RT-qPCR. C. Isogenic wild-type and **upc2AΔ** strains were grown in the presence or absence of caspofungin and levels of **FKS1** and **FKS2** RNA assayed as above. D. A probe from the **FKS1** promoter containing the putative SRE shown in Figure 4A was used in a EMSA experiment. A version of this probe lacking the SRE (mSRE) was also compared for its behavior in this assay. Upc2A was either omitted (-) or added (+) to the reaction prior to electrophoresis. E. An **FKS1-lacZ** fusion gene containing either the wild-type or the mSRE version of the promoter was introduced into wild-type cells. Levels of **FKS1**-dependent β-galactosidase were determined in the presence or absence of caspofungin.
Table 1. Strains and relevant genotypes for C. glabrata cells used in this study.

<table>
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<tr>
<th>Name</th>
<th>Parent strain</th>
<th>Genotype</th>
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<tbody>
<tr>
<td>KKY2001</td>
<td>CBS138</td>
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<td>KKY2001</td>
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Table 2. **Genes induced by log2 of 0.5 or greater in the presence of the G898D UPC2A.** RNA-seq was used to compare the levels of transcripts produced in the presence of the G898D UPC2A allele and the ratio of these mRNA levels/those seen in the presence of wild-type UPC2A presented as log2 Fold Change.

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<tr>
<th>Gene</th>
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<th>SRE</th>
<th>Gene</th>
<th>Sc homologue</th>
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Figure 1. Vu, et al.

A.  

<table>
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<tr>
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<tr>
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<tr>
<td>3X HA-G898D UPC2A</td>
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B.  

![Graph showing gene expression levels for different conditions.](https://doi.org/10.1101/2021.05.07.443069)

C.  

<table>
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<tr>
<th></th>
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<tr>
<td><strong>3X HA-UPC2A</strong></td>
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<td><strong>3X HA-G898D UPC2A</strong></td>
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</table>
Figure 2. Vu, et al.

AR1c TCGTATA
AR1b TCGTTTA
Figure 3. Vu, et al.
Figure 4. Vu, et al.

A.  
ERG1: CGACTCTATCACGATCGTATACTGTTGCTACCAACCACCCAG  
CDR1: TACCCGAAACGATTAGCTCTGATATGCTGACACCAGGAC  
PDR1: TCTGTGCTTCATTTTCTACCTCGTAGATAGGTTACGTTAAAATTTT  
FKS1: CAGGGCTACTGCAATTTCTATCGTAAACGAAACTGAGAATTGAA  
ScERG2: GCAGAAATCGAACCTAGCGGTGTCAAGCGCAAGGGAAATACC  
CaERG2: ATCCGAATAGTTCTGATTGCTGTATAAAAAAATTTGTG

B.  
Upc2A

<table>
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<tr>
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<th>-</th>
<th>+</th>
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ERG1  
CDR1  
PDR1

C.  
ERG1  
CDR1  
PDR1

<table>
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<tr>
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<th>-</th>
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<th>mut</th>
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<tr>
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</table>
Figure 7. Vu, et al.

A.  

<table>
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<tr>
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</tr>
<tr>
<td>upc2AΔ</td>
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<tr>
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<td>3X HA-G898D UPC2A</td>
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</tbody>
</table>

B.  

Relative expression levels for FKS1, FKS2, FKS3, ERG1, and AUS1 under different conditions:

- 3X HA-UPC2A
- 3X HA-G898D UPC2A

C.  

Comparison of FKS1 and FKS2 expression levels under untreated and caspofungin-treated conditions:

- wt
- upc2AΔ

D.  

FKS1 expression in wild type (wt) and mSRE conditions.

E.  

Expression levels of FKS1-lacZ and mSRE FKS1-lacZ under untreated and caspofungin-treated conditions.