



Discovery of Fungus-Specific Targets and Inhibitors Using Chemical Phenotyping of Pathogenic Spore Germination

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ABSTRACT There is a critical need for new antifungal drugs; however, the lack of available fungus-specific targets is a major hurdle in the development of antifungal therapeutics. Spore germination is a differentiation process absent in humans that could harbor uncharacterized fungus-specific targets. To capitalize on this possibility, we developed novel phenotypic assays to identify and characterize inhibitors of spore germination of the human fungal pathogen Cryptococcus. Using these assays, we carried out a high-throughput screen of \sim 75,000 drug-like small molecules and identified and characterized 191 novel inhibitors of spore germination, many of which also inhibited yeast replication and demonstrated low cytotoxicity against mammalian cells. Using an automated, microscopy-based, quantitative germination assay (QGA), we discovered that germinating spore populations can exhibit unique phenotypes in response to chemical inhibitors. Through the characterization of these spore population dynamics in the presence of the newly identified inhibitors, we classified 6 distinct phenotypes based on differences in germination synchronicity, germination rates, and overall population behavior. Similar chemical phenotypes were induced by inhibitors that targeted the same cellular function or had shared substructures. Leveraging these features, we used QGAs to identify outliers among compounds that fell into similar structural groups and thus refined relevant structural moieties, facilitating target identification. This approach led to the identification of complex II of the electron transport chain as the putative target of a promising structural cluster of germination inhibitory compounds. These inhibitors showed high potency against Cryptococcus spore germination while maintaining low cytotoxicity against mammalian cells, making them prime candidates for development into novel antifungal therapeutics.

IMPORTANCE Fungal pathogens cause 1.5 million deaths annually, and there is a critical need for new antifungal drugs. However, humans and fungi are very similar on a molecular level, and so many drugs that kill fungi also damage human cells, leading to extreme side effects, including death. The lack of fungus-specific targets is a major hurdle in the development of antifungal therapeutics. Spore germination is a process absent in humans that could harbor fungus-specific targets. To capitalize on this possibility, we developed new assays to identify and characterize inhibitors of spore germination of the human fungal pathogen *Cryptococcus*. Using these assays, we identified and characterized 191 novel inhibitors of spore germination while maintaining low cytotoxicity against mammalian cells, making them prime candidates for development into novel antifungal therapeutics.

KEYWORDS *Cryptococcus*, antifungal agents, fungi, germination, spores

uman fungal pathogens are an unmitigated problem causing \sim 1.5 million deaths a year worldwide (1). One of the biggest hurdles in the treatment of invasive fungal diseases is the lack of available therapeutics. There are three primary classes of

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Received 19 June 2021 **Accepted** 25 June 2021 **Published** 27 July 2021 antifungal drugs, all of which are suboptimal due to properties ranging from high toxicity to humans to rapid microbial resistance development (2–5). These classes target cell membranes or cell wall components, which have been the canonical targets for antifungal development (2, 6). While these cellular structures provide fungus-specific targets, the deficiency of novel antifungal agents indicates that new fungus-specific targets need to be identified and exploited. However, due to the eukaryotic nature of fungi and resulting conservation of molecular moieties between humans and fungi, the identification of fungus-specific pathways has been difficult.

One proposed solution is to target the process of spore germination (7). Spores are dormant stress-resistant cell types formed by many organisms to survive harsh environmental conditions and/or spread to new environments, and spores are infectious particles for most invasive human fungal pathogens (8, 9). To cause disease, fungal spores must escape dormancy through the process of germination, a process that appears unlike any in humans, and grow vegetatively in the host. Due to its specialized nature, spore germination may involve fungal pathways distinct from those in humans. We hypothesized that the process of spore germination would harbor new fungus-specific targets, and compounds that inhibit germination would therefore be less toxic to mammalian cells. Thus, germination inhibitors would be prime candidates for development into antifungal drugs for the prevention and/or treatment of many invasive fungal diseases.

The development of new antifungal drugs has been slow relative to that of other antimicrobial agents, such as those against bacteria and viruses, despite significant screening efforts (10). Traditionally, the primary method for identifying antifungal compounds was based on tracking changes in a biologically relevant readout such as fungal growth (phenotypic drug discovery). While this approach yielded many antifungal compounds over the years, most were also toxic to mammalian cells. As molecular techniques advanced, researchers in many fields moved toward targeted drug discovery, which relies on identification of inhibitors of a specific known molecular target. This approach proved to be useful in some arenas; however, it largely failed for antifungal drug development presumably because of a lack of identified fungus-specific targets. As a result, no new classes of antifungal therapeutics have come to the market in the last 20 years, and there is renewed interest in phenotypic drug discovery (2, 11). However, because growth inhibition screening alone has been used exhaustively as a target phenotype in fungi, for the promise of phenotypic drug discovery to be fully realized, the field must improve upon traditional growth screening methods and/or couple them with novel assays (10).

To address this need, we used fundamental biological discoveries of pathogenic spore biology to drive the development of two new phenotypic assays that use spore germination as a readout. The first is a luciferase-based assay for high-throughput screening of compounds to identify germination inhibitors, and the second is an automated, quantitative, microscopy-based germination assay for high-resolution evaluation of large populations of germinating spores. We developed these tools for use with the spores of the invasive human fungal pathogen Cryptococcus. This environmental budding yeast is the leading cause of fatal fungal disease worldwide, causing several hundred thousand deaths per year, particularly among people with compromised immune systems (5). The Cryptococcus system is known among human fungal pathogens to be well developed with many molecular and genetic tools. In addition, Cryptococcus spores germinate synchronously under nutritionally favorable conditions and do so largely independent of spore density (12). These unique properties facilitated the development of the phenotypic germination assays that we used to identify and characterize 191 novel fungal germination inhibitors. We discovered that population-level dynamics could be used to classify distinct chemical phenotypes, and we used those classifications to show that compounds with similar substructures demonstrated similar chemical phenotypes. This process led to the rapid identification of phenotypic outliers and facilitated target identification, resulting in the discovery of a novel set of fungus-specific electron transport chain inhibitors that are prime candidates for development into a new class of antifungal drugs for use in the prevention of fatal fungal diseases.

RESULTS

Eukaryotic translation inhibitors prevent initiation of *Cryptococcus* **spore germination.** Prior studies of *Cryptococcus* spore germination showed that different conditions, mutants, and drugs alter the behavior of spore populations during germination (7, 12). Based on these findings, we hypothesized that characterizing the behaviors of spore populations under different conditions would facilitate the identification of specific cellular processes required for spore germination. Because new protein synthesis is known to be required for successful germination in many fungi (13), we treated populations of spores under germinating conditions with the eukaryotic ribosome inhibitor cycloheximide and evaluated their responses using our quantitative germination assay (QGA).

In this automated microscopy-based assay, germination progression of spores is monitored as a function of changes in cell morphology over time (spores are small and oval; yeast are large and circular). Individual spores in a population ($\sim 1 \times 10^4$ per sample) are measured to determine size (area) and shape (aspect ratio) from the onset of germination, and the data are collected for each cell over the time of germination (Fig. 1A). Using the QGA, we determined that a concentration of $10 \,\mu$ M cycloheximide fully prevented spores from initiating any changes in size or shape, indicating full inhibition of germination and suggesting that new protein synthesis is required for *Cryptococcus* spores to germinate (Fig. 1B).

As the concentration of cycloheximide decreased, the amount of germination increased, exhibiting concentration-dependent inhibition of germination with a 50% inhibitory concentration (IC₅₀) of 1.7 μ M (Fig. 1C). In addition, we observed that even as the germination of the population of spores was slowing down due to inhibition, all of the spores responded in a similar manner and maintained their synchronous response (Fig. 1B). While the overall rate of germination for the population changed in response to cycloheximide, other properties were unchanged (population synchronicity, pattern of morphological changes, and integrity of individual spores), which facilitated the determination of specific rates of germination (i.e., transition out of the spore state) at each concentration tested (Fig. 1D). We observed that as the concentration of cycloheximide increased, germination rates decreased, causing a "slow-down" phenotype across the population.

From these data, we concluded that new protein synthesis was likely required very early in the germination process. We further surmised that if the cycloheximide phenotype was specific to the inhibition of protein translation (as opposed to off-target effects), other inhibitors of eukaryotic protein translation would produce the same phenotype. To test this hypothesis, we evaluated 3 structurally distinct inhibitors of eukaryotic protein translation (Geneticin, anisomycin, and puromycin) and determined their effects in QGAs (Fig. 1E). Although the inhibitors showed different potencies against germination (i.e., different concentrations were required to achieve similar effects), all three inhibitors caused the slow-down phenotype that maintained population synchronicity, mimicking the effect of cycloheximide.

Together, these data show that for each inhibitor, the QGA was an effective method for determining a concentration-dependent phenotype, determining precise inhibitory concentrations, and quantitating changes in germination rates. Furthermore, the consistent phenotypes across translation inhibitors suggested that inhibitors targeting the same cellular function generate similar phenotypes in the QGA. These findings indicated that QGAs would be a powerful tool in the validation, prioritization, and characterization of diverse germination inhibitors with unknown targets.

Combined HTS and QGA analysis identified 191 novel germination inhibitors. To identify potential inhibitors of spore germination, a nanoluciferase (NL)-based highthroughput screening (HTS) assay was developed, and compounds from three libraries



FIG 1 Eukaryotic translation inhibitors prevent initiation of *Cryptococcus* spore germination. (A) Diagram of the quantitative germination assay (QGA). Microscopy image data are processed using ImageJ and represented in two-dimensional histograms that show the numbers of cells in (Continued on next page)

of structurally diverse, drug-like small molecules (LifeChem 1 to 3) were screened (see Fig. S1 in the supplemental material). For the assay, a previously identified protein (CNK01510) was fused to the nanoluciferase protein (Promega) and introduced into its endogenous locus in the Cryptococcus genome (14). Strains harboring the integrated NL protein fusion were crossed under sexual development conditions to produce spores. Spores from the NL strains yielded very little NL enzyme activity; however, yeast from those strains produced a robust NL signal. Most importantly, the amount of NL signal correlated with the germination state. As spores germinated into yeast, the levels of NL signal increased, resulting in a robust signal over baseline at the end of germination (\sim 14-fold) (Fig. S1). Inhibitors of germination (e.g., cycloheximide) caused low levels of NL signal, and solvent-only controls (e.g., dimethyl sulfoxide [DMSO]) affected neither germination nor the NL signal (Fig. S1). Of the ~75,000 compounds screened, \sim 2,100 compounds caused a \geq 20% decrease in NL signal relative to that for the solvent-only control and were rescreened in duplicates. Compounds that showed \geq 50% inhibition of the NL control signal were selected, resulting in 238 putative germination inhibitors. Secondary screens were performed on these hits to determine effects on yeast replication, cytotoxicity against mammalian fibroblasts, and direct inhibition of the NL enzyme (see Data Set S1).

To confirm that the 238 hits from the high-throughput screen were bona fide inhibitors of germination, each compound was tested in the QGA. One hundred ninety-one of the 238 HTS hits showed inhibition of germination at a single relatively high concentration (80 μ M) in this assay (see Data Set S2). The majority of these confirmed germination inhibitors (121/191) showed low cytotoxicity to mammalian cells (<25% decrease in cell viability), and 167 of 191 caused at least 10% inhibition of yeast growth at 10 μ M. Six of the 191 germination inhibitors were also inhibitors of the NL enzyme assay. Overall, the QGA was highly effective for the validation of HTS hits and provided a high-confidence library of 191 novel germination inhibitors, resulting in the largest discovery of novel, confirmed fungal germination inhibitors in any system. Because the majority of these inhibitors exhibited low preliminary cytotoxicity against mammalian cells, the data supported the idea that germination could serve as a reservoir of fungus-specific drug targets.

Upon evaluation of the structures of the 191 confirmed inhibitors, we discovered that 76 of the compounds fell into 8 distinct groups with shared substructures, each of which contained 4 or more compounds (Fig. 2). The identification of multiple groups of similarly structured compounds from a library of diverse small molecules is advantageous, because similarly structured compounds are likely to have shared molecular targets (15, 16). Compounds that had shared substructures with other inhibitors, showed potent inhibition of both spore germination and yeast growth, and exhibited low mammalian cell toxicity (Fig. 2) were prioritized for further investigation.

QGA titrations of germination inhibitors identified 6 discernible phenotypes. Because our QGA data with translation inhibitors showed that inhibiting molecular targets within a specific cellular function resulted in a shared phenotype (Fig. 1), we hypothesized that compounds with shared substructures would induce shared phenotypes. To determine potential "chemical phenotypes," we titrated 86 confirmed germination inhibitors in the QGA. We discovered six different phenotypes, and they were distinguished from one another on the basis of differences in germination synchronicity, germination rates, and overall population behavior. These phenotypes fell into two general categories: "homogeneous germination" and "heterogeneous germination" (Fig. 3A).

Homogeneous germination phenotypes occurred when spores germinated syn-

FIG 1 Legend (Continued)

each position as a function of pixel intensity. In all two-dimensional histograms presented, the *x* axis represents area (μ m²), and *y* axis represents the aspect ratio (width/length). Spores (small oval cells) populate the lower left quadrant, and yeast (large circular cells) populate the upper right quadrant. (B) Germination profiles of ~6,000 spores in the presence of 0, 0.63, 1.25, 2.5, 5, or 10 μ M cycloheximide. (C) Dose-response curve of spores transitioned at 16 h relative to the control in the presence of 0.078 to 80 μ M cycloheximide (error bars represent standard deviations determined across 9 fields of view from 3 independent replicates). (D) Germination rates of spores at different time points in the presence of 0, 0.63, 1.25, 2.5, or 5 μ M cycloheximide (error bars represent standard deviations determined across 9 fields of view from 3 independent replicates). (E) Germination profiles of ~6,000 spores in the presence of protein translation inhibitors.

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FIG 2 Seventy-six of 191 germination inhibitors with shared substructures defined 8 groups. Diagram of each substructure with alphabetic assignments followed by their PubChem chemical identifiers (CIDs) for ease of identification (columns 1 and 2). Heat maps represent level of relative spore germination (at 80μ M), level of yeast growth inhibition (at 10μ M), and level of relative fibroblast vitality (at 10μ M) (columns 3 to 5).



FIG 3 Discovery of six unique germination inhibition phenotypes. (A) Two-dimensional histograms and diagrams of distinct phenotypes caused by representative compounds among 191 germination inhibitors (PubChem CID and effective concentration provided for each). Homogeneous germination includes (I) slow-down, (II) slow-start, and (III) slow-end phenotypes. Heterogeneous germination includes (IV) asynchrony, (V) bimodal, and (VI) morphological variability phenotypes. Each plot represents \sim 6,000 spores. (B) Germination rates of spores at different time points during homogeneous germination in panel A (error bars represent standard deviations determined from 6 fields of view from two independent replicates). (C) Population morphological spread at 0 and 16 h of germination for heterogeneous germination phenotypes shown in panel A.

chronously as a population but at a lower rate, and individual spores were affected equally throughout the population. In this category, there were three phenotypes, which were distinguished by the time point at which germination was most inhibited and were quantified by changes in germination rates. The most common of these (~30% of all inhibitors tested) was the (I) slow-down phenotype, in which the population of spores germinated synchronously but at a lower rate over 16 h of germination (Fig. 3A, rows 1 to 3). The (II) slow-start phenotype occurred with a single inhibitor at the beginning of germination. After initial inhibition (occurring between 0 and ~8 h), the rate of germination increased rapidly as spores overcame the inhibition. The (III) slow-end phenotype occurred with a few inhibitors, in which the population of spores initially germinated at a normal rate but exhibited a reduced rate later in germination (after ~8 h). This inhibition was observed primarily at the point in germination (~8 h) when spores that had germinated into small circular cells became uniformly larger (via isotropic growth). All three of these phenotypes showed lower rates of germination, but the points of inhibition and rates of germination varied among them (Fig. 3B).

Heterogeneous germination phenotypes occurred when spores no longer germinated synchronously as a population, and individual spores were affected differently. In this category, there were three phenotypes, all related to how a lack of synchrony manifested across a population. The most common of these inhibition phenotypes $(\sim 50\%$ of inhibitors tested) was the (IV) asynchrony phenotype, in which a population of spores lost synchronous germination and different levels of inhibition were observed across the population, resulting in large variations in cell shapes and sizes (Fig. 3A, rows 4 to 6). A (V) bimodal phenotype occurred with several inhibitors when the population split in two groups, with part of the population experiencing complete inhibition and the other germinating normally, resulting in a bimodal distribution. The (VI) morphological variability phenotype occurred with only two inhibitors, when spores germinated into cells with a variety of sizes and shapes, resulting in a higher proportion of more elongated and/or very large cells. This led to a more variable morphology in yeast once fully germinated. These three phenotypes demonstrate that spores in a population can respond differently to germination conditions and be distinguished by population-level and individual morphological differences (Fig. 3C). While the phenotypes observed were not necessarily a comprehensive accounting of all germination phenotypes, they provided an opportunity to further parse the 191 inhibitors into groups and evaluate structure-function relationships.

Similarly structured compounds elicit the same germination phenotypes. To test the hypothesis that compounds with shared substructures would induce shared phenotypes, we evaluated structural groups A and B in more detail. If similarly structured compounds showed similar phenotypes, it would support the idea that they share the same target. Groups A and B were chosen for comprehensive analysis because they (i) contained a relatively large number of compounds with similar substructures (8 and 10, respectively), (ii) displayed generally strong inhibition of germination at 80 μ M, (iii) were able to inhibit yeast growth to various degrees, and (iv) exhibited low cytotoxicity to mammalian cells.

Group A is composed of 8 compounds with a thieno[2,3-*d*]pyrimidin-4-amine substructure. To identify the phenotypes of inhibition for group A, we carried out titrations of each at concentrations from 2.5 μ M to 80 μ M (Fig. 4A and B). The majority of group A compounds (A1 to A7) demonstrated clear slow-down germination phenotypes; however, A8 showed the asynchrony phenotype. This phenotypic discrepancy suggests that A8 interacts differently with spores and may have a different cellular target. For this reason, A8 likely does not belong in this grouping of compounds and was considered an outlier during further characterization of group A. To determine the mammalian cytotoxicity of these inhibitors, dose-response cytotoxicity assays were performed on fibroblasts with A1 to A7 at concentrations from 0.5 μ M to 80 μ M (Fig. 4C). These assays showed that group A compounds exhibited relatively low cytotoxicity against mammalian cells at relevant inhibitory concentrations. Notably, the potency of



FIG 4 All but one group A compound caused the slow-down phenotype. (A) Germination profiles of \sim 6,000 spores in the presence of group A compounds at phenotypic concentrations. (B) Dose-response curves of compounds A1 to A7 at concentrations from 2.5 to 80 μ M (error bars represent standard deviations determined from 6 fields of view from 2 independent replicates). Data points for inhibitors that precipitated at higher concentrations were excluded. (C) Dose-response curves showing levels of cytotoxicity against human fibroblasts (error bars represent standard deviations from 2 independent replicates).

spore germination inhibition was not related to the levels of mammalian cytotoxicity, as exemplified by the strongest inhibitor in this group (A5) showing >80% germination inhibition at 5 μ M and <20% cytotoxicity at concentrations as high as 80 μ M. This suggests that structural moieties in this group could be altered to maximize antifungal activity while minimizing cytotoxicity. Together, these data support the hypotheses that similarly structured compounds demonstrate similar phenotypes of inhibition and that phenotypic characterization can identify outliers in structural groups. These results further support that group A compounds target the same biological process, have low cytotoxicity, and are promising candidates for antifungal development and target identification.

Group B is composed of 10 compounds with a 2,5-dimethylfuran-3-carboxamide substructure. To identify phenotypes of inhibition for group B, we titrated these compounds at concentrations from 2.5 μ M to 80 μ M (Fig. 5A and B). While the 10 inhibitors showed various abilities to inhibit germination, all 10 demonstrated a clear asynchrony germination phenotype. Cytotoxicity assays were performed at concentrations from 0.5 μ M to 80 μ M (Fig. 5C) and showed that group B compounds exhibit relatively low cytotoxicity against mammalian cells at relevant inhibitory concentrations. Again, the potency of spore germination inhibition was not related to the levels of mammalian cytotoxicity. For example, the strongest inhibitor in this group (B2) showed >75% germination inhibition at 5 μ M and \leq 25% cytotoxicity at concentrations as high as 80 μ M. As for group A, these data suggest that all group B compounds target the same biological process, providing another group of promising candidates for further development.

In addition to evaluating all of the compounds in groups A and B, we evaluated a random subset of compounds from the largest groups (groups D, E, F, and G) to further test the hypothesis that similarly structured compounds would generate similar germination inhibition phenotypes. We discovered that all of the 12 compounds tested from group D caused a slow-down phenotype (see Fig. S2), and all of the 6 compounds tested from group E caused an asynchrony phenotype (see Fig. S3). In contrast, the 6 compounds tested from group F and the 6 compounds tested from group G showed different phenotypes within their groups, with group F compounds generating three different phenotypes, and group G compounds generating four different phenotypes (see Fig. S4 and S5). While these findings break with the pattern for groups A, B, D, and E, a closer examination of the structures in groups F and G revealed much more overall structural variation than in groups A, B, D, and E. These data indicate that the substructures initially identified for groups F and G were not predictive of phenotypes for their entire groups; however, compounds within groups F and G that shared phenotypes were more structurally similar to one another than to the phenotypic outliers. Together, these data support the hypothesis that similarly structured compounds elicit similar germination phenotypes, suggesting that phenotypes could be used to refine the relevant features of substructures conferring activity.

Complex II of the electron transport chain is the likely target of group B compounds. The relevant substructure in all the group B compounds is a furan carboxamide. This structure is found in known carboxamide fungicides. These fungicides have historically been used against plant pathogens and are part of the succinate dehydrogenase inhibitor (SDHI) class of fungicides, which target complex II of the electron transport chain (ETC) (17). SDHIs are a large class of fungicides that includes compounds of diverse structures that vary in their specificity for different plant fungal pathogens (17). Group B shows strong structural similarity with one SDHI in particular, furcarbanil, which has a structure nearly identical to compound B1, with only a single ethyl group differentiating the two molecules. Due to this shared similarity, we hypothesized that furcarbanil would inhibit germination at a level akin to that of B1 (the weakest group B compound). In fact, furcarbanil demonstrated the asynchrony phenotype at the same phenotypic concentration as B1 (80 μ M), showing a nearly identical profile that was weaker than most group B compounds (Fig. 5A and 6A).

Given these data, we hypothesized that other inhibitors of the ETC would result in the same germination phenotype as group B compounds and furcarbanil. To test this, we determined the inhibition phenotypes of rotenone, thenoyltrifluoroacetone (TTFA),

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FIG 5 All group B compounds caused the asynchrony phenotype. (A) Germination profiles of ~6,000 spores in the presence of group B compounds at phenotypic concentrations. (B) Dose-response curves of compounds B1 to B10 at concentrations from 2.5 to 80 µM (error bars represent standard deviations determined from 6 fields of view from 2 independent replicates). (C) Dose-response curves showing levels of cytotoxicity against human fibroblasts (error bars represent standard deviations from 2 independent replicates).



FIG 6 Group B compounds inhibit oxygen consumption by the electron transport chain. (A) Germination profiles of \sim 6,000 spores at phenotypic concentrations of the inhibitors furcarbanil (80 μ M), rotenone (10 μ M), TTFA (200 μ M), and antimycin A (2.5 μ M). (B) Oxygen (Continued on next page)

and antimycin A, which are well-characterized inhibitors of complexes I, II, and III, respectively (Fig. 6A). Each of these mitochondrial inhibitors demonstrated the asynchrony phenotype. Given the shared structural homology of group B to furan carboxamide SDHIs and the inhibitory phenotype of furcarbanil and other ETC inhibitors, we hypothesized that group B compounds hinder germination through ETC inhibition by targeting succinate dehydrogenase (complex II) specifically. To test the possibility that group B compounds inhibit the ETC, we performed oxygen consumption experiments to monitor the effects of these inhibitors on oxygen consumption rates of Cryptococcus yeast (Fig. 6B). We tested a weak (furcarbanil), an intermediate (B5), and a strong (B2) inhibitor of germination and found that all three compounds (at 10 μ M) were able to lower the oxygen consumption rate (OCR) of the cells. We anticipated that the ability to decrease the oxygen consumption rate would correlate with higher antifungal potency. Furcarbanil, which was the weakest of the three inhibitors, caused an initial substantial decrease in OCR followed by recovery, suggesting that the inhibition of oxygen consumption can easily be overcome. B5, which showed intermediate antifungal activity, caused a small decrease in OCR that was not recovered from, suggesting a modest but insurmountable inhibition of oxygen consumption. Finally, B2, which was the strongest inhibitor, showed a substantial decrease in OCR that was not recovered from, suggesting strong and insurmountable inhibition of oxygen consumption. These OCR inhibition patterns are consistent with the antifungal potencies of each inhibitor. These results support the idea that group B compounds inhibit germination by obstructing the ETC. While it is formally possible that these inhibitors could be altering OCR through pathways other than direct ETC inhibition, all the data presented here support the hypothesis that group B compounds are novel ETC inhibitors, likely targeting complex II. These novel inhibitors exhibit strong antifungal activity and low mammalian cytotoxicity, making them prime candidates for development into novel antifungal therapeutics.

DISCUSSION

In this study, we combined two new phenotypic assays that target fungal spore germination to identify, validate, and characterize 191 novel fungal germination inhibitors. Using QGAs, we identified 6 distinct chemical phenotypes distinguished from one another on the basis of differences in germination synchronicity, germination rates, and overall population behavior. Compounds that targeted the same cellular function or had shared substructures induced similar phenotypes. Thus, QGAs identified phenotypic outliers and facilitated target identification via comparisons between structurally similar compounds with the same phenotypes and compounds with known cellular targets. We identified a group of novel putative fungus-specific electron transport chain inhibitors that are promising candidates for antifungal development. Most importantly, this study supports the idea that the germination process holds fungusspecific pathways that could serve as targets for new antifungal drugs.

Chemical phenotyping can help overcome the hurdles of phenotypic drug discovery. The relative merits of targeted drug discovery versus those of phenotypic drug discovery have been debated across fields; however, in antifungal drug discovery, one of the key issues is the lack of known fungus-specific targets. This challenge supports using phenotypic drug discovery (PDD) approaches, but PDD presents other challenges such as (i) difficulties in validation of hits, (ii) an inability to establish structure-activity relationships, and (iii) difficulties in target identification (18). To overcome these limitations, new methods of phenotypic characterization have been used, such as molecular phenotyping in which transcriptome analysis was used as a secondary screening method (19). This approach facilitated the clustering of compounds based

FIG 6 Legend (Continued)

consumption rate (OCR) plots of *Cryptococcus* yeast with injections every 60 min with first glucose (20 mM) and then either (I) furcarbanil (10 μ M), (II) B5 (10 μ M), or (III) B2 (10 μ M), then rotenone/antimycin A (50 μ M), and finally 2-DG (100 mM) (error bars represent standard deviation across 3 independent replicates).

on shared profiles and helped identify their targets. Similarly, we used new phenotypic assays to both identify and characterize compounds and overcome the hurdles of PDD.

By using the NL-based high-throughput screen for inhibitors of germination (as opposed to growth), we increased the specificity of our initial screen, reducing the number of hits and increasing the likelihood that the hits would be fungus specific. Following the initial screen with QGAs enabled the identification of bona fide inhibitors of germination, eliminated false positives from the working pool of compounds, and addressed the first major challenge in PDD (validation). QGAs were also used to address the second major PDD challenge (establishing structure-function relationships) via the generation of chemical phenotypes for each compound of interest. We showed that inhibitors that target the same biological process share the same chemical phenotype. Thus, by characterizing the phenotypes of each compound in a structural group, we established structure-activity relationships and identified phenotypic outliers that could have targets that differ from the group overall. Finally, the use of chemical phenotyping also addressed the third challenge by lowering the barriers to target identification. Population dynamics of germinating spores vary in the presence of different inhibitors, stressors, nutrients, and mutations, all of which can be assessed using the QGA (7, 12). These provide an opportunity for comparative analyses to facilitate target identification. By combining the testing of inhibitors with alteration of nutrients, inhibition of known targets, or creation of knockout and overexpression constructs, we can mimic, intensify, alter, or eliminate a germination phenotype and thereby identify the target processes and pathways of specific inhibitors. For example, in a previous study, we found that disulfiram had a slow-end phenotype (7), indicating that it inhibits a target that is important for the isotropic growth phase of germination. The more we learn about the molecular programming of spore germination, the more we can gain from this type of phenotypic analysis.

Electron transport chain as a fungus-specific target in antifungal development. The ETC has been suggested as a good target for antifungal drug development because of the role of respiration in regulating virulence traits and the existence of fungus-specific ETC elements. However, complex II/succinate dehydrogenase (SDH) has yet to be exploited (20). SDH could be a promising target, having been implicated in virulence of some human fungal pathogens. Specifically, SDH mRNA transcripts are overrepresented in Cryptococcus during murine pulmonary infections (21), and the SDH inhibitor thenoyltrifluoroacetone (TTFA) has been shown to prevent hyphal formation in Candida albicans, a key virulence trait (22). It is unclear whether there are fungus-specific properties of SDH, but some carboxamide SDHIs have shown narrowspectrum use against basidiomycete plant pathogens (17, 23), which suggests that fungal SDH is unique. Alternatively, some SDHIs have been shown to also inhibit complex III, implying more complex interactions (24). Nevertheless, it is promising that group B compounds in this study show high efficacy and low cytotoxicity, supporting the idea that their target(s) harbors fungus-specific features. This low cytotoxicity suggests that these compounds can inhibit the cryptococcal ETC while not significantly inhibiting fibroblast mitochondrial activity. While it is difficult to irrefutably conclude that group B inhibitors are targeting SDH, the data provided here strongly support this hypothesis. Future studies will be needed to fully characterize the mechanism of these inhibitors, to optimize them for increased antifungal potency and reduced mammalian cytotoxicity, and to test optimized inhibitors in murine models of invasive fungal infections. Overall, this group of compounds is extremely promising for further development into antifungal drugs.

Spore germination as a target reservoir for antifungal therapeutics. Spore germination is a process that appears to be distinct from any process in humans, making it a potential reservoir for fungus-specific targets for drug development. Here, we determined that 121 of the 191 germination inhibitors we identified showed preliminary low cytotoxicity against mammalian cells, suggesting that they may be targeting fungus-specific molecules. Additionally, both group A and B compounds showed

relatively low cytotoxicity at relevant inhibitory concentrations, and germination inhibition ability was not linked to cytotoxicity. This provides the opportunity to modify their structures to maximize antifungal activity while minimizing human cytotoxicity, thus increasing the difference between the effective dose and the toxic dose, leading to a higher therapeutic index. These data support the idea that targeting spore germination will result in the identification of low-toxicity antifungal drug candidates.

Targeting spore germination also provides an opportunity for the prevention of fungal disease, which is an area of disease management that is underexplored in the field of human fungal pathogenesis. Spores play an important role in disease progression in the majority of invasive human fungal pathogens, and spore germination is required for spores to cause disease (8, 9). Therefore, inhibiting spore germination (in addition to the subsequent vegetative replication) could provide a unique opportunity for antifungal prophylaxis in immunocompromised individuals to prevent fatal disease. The potential role of germination inhibitors in antifungal prophylaxis has been explored previously (7), and with the identification of novel inhibitors of both germination and growth, the development of preventative therapeutics can now be pursued.

MATERIALS AND METHODS

Strains and strain manipulation. *Cryptococcus neoformans* serotype D (*deneoformans*) strains JEC20, JEC21, CHY3833, and CHY3836 were handled using standard techniques and media as described previously (7, 25, 26). *Cryptococcus* spores were isolated from cultures as described previously (27). Briefly, yeast of both mating types (JEC20 and JEC21 or CHY3833 and CHY 3836) were grown on yeast-peptone-dextrose (YPD) medium for 2 days at 30°C, combined at a 1:1 ratio in $1 \times$ phosphate-buffered saline (PBS), and spotted onto V8 (pH 7) agar plates. Plates were incubated for 5 days at 25°C, and spots were resuspended in 75% Percoll in $1 \times$ PBS and subjected to gradient centrifugation. Spores were recovered, counted using a hemocytometer, and assessed for purity by visual inspection.

Nanoluciferase germination screen. Screening of the LifeChem libraries (Life Chemicals) was carried out with the assistance of the University of Wisconsin—Madison (UW-Madison) small-molecule screening facility. All NL screening was performed as described previously (7). Briefly, CHY3833 and CHY3836, reporter strains harboring a CNK01510-NL fusion construct, were used to produce spores for library screening. Screening was carried out with 1×10^4 spores incubated in 384-well screening plates in $10 \,\mu$ l of germination medium ($0.5 \times$ YPD) for 10 h at 30°C. Cells were then incubated with $10 \,\mu$ l of Nano-Glo luciferase assay reagent (Promega Corporation) prepared as suggested by the manufacturer at 22°C for 10 min and then read using a Perkin-Elmer EnSpire plate reader at 460 nm.

Secondary screens. (i) NL enzyme test. CHY3833 was grown overnight in liquid YPD, washed 3 times in $1 \times$ PBS, and resuspended to an optical density at 600 nm (OD₆₀₀) of 1.00. Cells (100 μ l) were added to 384-well plate wells containing 10 μ M each compound, incubated with 10 μ l of Nano-Glo luciferase assay reagent (Promega Corporation) at 22°C for 10 min, and then read using a Perkin-Elmer EnSpire plate reader at 460 nm. Compounds that caused a >50% decrease in luciferase signal were determined to be nanoluciferase enzyme assay inhibitors.

(ii) Yeast replication. CHY3833 and CHY3836 were each grown in YPD liquid overnight at 30°C to saturation and then resuspended in $0.5 \times$ YPD at an OD₆₀₀ of 0.005. Strains were aliquoted into 384-well plates with inhibitors and grown for 12 h at 30°C at 3,000 rpm before OD₆₀₀ readings were taken. Compounds that caused a >10% decrease in growth were considered yeast growth inhibitors.

Fibroblast cytotoxicity. PrimaPure normal human dermal fibroblasts (NHDF) cells (Genlatis PH10605A) were cultured and passaged in fibroblast growth medium (Genlatis PM116500) according to the manufacturer's protocol. NHDF cells (1×10^3 per well) were plated in a 384-well plate in cell culture medium. The cells were incubated with each compound of interest at $10 \,\mu$ M concentration for 72 h at 37°C and 5% CO₂. Following treatment, CellTiter-Glo (Promega) reagent was used to assay ATP-dependent luminescence and thus provide a measure of cell viability. Compounds that resulted in >75% cell viability were considered low toxicity. For cytotoxicity dose-response curves (Fig. 4C and Fig. 5C), compounds were titrated from 80 μ M to 0.5 μ M in duplicates.

Quantitative germination assay. Germination assays were modified from those described by Barkal et al. to introduce automation, increase throughput, and refine assay consistency (12). Briefly, 384-well plates (Thermo Scientific 142762) were loaded with 10⁵ spores per well, and at 0 h, synthetic medium plus 2% dextrose (SD medium) containing the compounds of interest was added to the sample (final volume of 40 μ l). All compounds were tested initially at 80 μ M; however, concentrations were changed on a case-by-case basis for subsequent experiments. All assays and controls were performed with a final concentration of 0.8% DMSO (the compound solvent) unless specified otherwise. Spores were germinated at 30°C in a humidified chamber, and the same $\sim 5 \times 10^3$ cells were monitored every 2 h for 16 h. Imaging was performed on a Ti2 Nikon microscope, and each condition was visualized in a minimum of two individual wells with three fields of view acquired from each well. All images were analyzed as described previously based on cell shape and size using ImageJ. The population ratios of spores, intermediates, and yeast were determined. Error bars in plots are based on the variation among all fields of view acquired. Level of germination was determined by quantifying the decrease in the proportion of spores in a population, and rates were quantified by determining the change in this proportion over time. For all two-dimensional histograms, the x axis represents area from $0\,\mu\text{m}^2$ to $25\,\mu\text{m}^2$, and the y axis represents the aspect ratio from 0.4 to 1.

(i) Translation inhibitors and concentrations. Cycloheximide (0.078 μ M to 80 μ M) (Dot Scientific, Inc. DSC81040-1), G418 (1.6 mM) (Fisher Scientific AAJ6267106), and puromycin (20 mM) (Dot Scientific, Inc. DSP33020-0.025) were tested with no DMSO (all were water soluble), whereas anisomycin (62.5 μ M) (Sigma-Aldrich A9789-5MG) was tested in 2.5% DMSO.

(ii) Mitochondrial inhibitors and concentrations. Furcarbanil (80 μ M) (Sigma-Aldrich T313122), rotenone (10 μ M) (Fisher Scientific 501687383), and antimycin A (2.5 μ M) (Santa Cruz Biotechnology, Inc. sc-202467A) were all tested in 0.8% DMSO, and TTFA (200 μ M) (Sigma-Aldrich 88300-5G) was tested in 3% DMSO.

Oxygen consumption rate experiments. Oxygen consumption experiments were performed on a Seahorse Biosciences XFe96 extracellular flux analyzer, and the assays were modified from those described by Lev et al. 2020 (28). Briefly, cartridges were hydrated overnight in Agilent Seahorse XF calibrant. JEC21 yeast were grown overnight in YPD, washed with double-distilled water (ddH₂O) and resuspended to an OD₆₀₀ of 0.08 in Seahorse XF Dulbecco's modified Eagle medium (DMEM) at pH 7.4 (Agilent 103575-100). Cells (180 μ I) were loaded into each well with the exception of blank wells, which were filled with XF DMEM. Each injection solution was 10× the final volume in XF DMEM. The assay was carried out at 30°C with injections occurring every 60 min. OCR was read every 6 min during each hour interval, with 3 min of mixing and 3 min of measuring. The following final concentrations were achieved after each injection: (a) 20 mM dextrose, (b) 10 μ M chosen inhibitor (furcarbanil, B2, or B5) and 1% DMSO, (c) 50 μ M rotenone, 50 μ M antimycin A, and 1% DMSO, and (d) 100 mM 2-deoxy-D-glucose (2-DG; Sigma-Aldrich D8375-16). Each condition was tested in triplicates.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. DATA SET S1, XLSX file, 2.1 MB. DATA SET S2, PDF file, 4 MB. FIG S1, PDF file, 0.7 MB. FIG S2, PDF file, 3.1 MB. FIG S3, PDF file, 1.8 MB. FIG S4, PDF file, 1.8 MB. FIG S5, PDF file, 3.5 MB.

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