Identification of a New Class of Antifungals Targeting the Synthesis of Fungal Sphingolipids

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ABSTRACT Recent estimates suggest that >300 million people are afflicted by serious fungal infections worldwide. Current antifungal drugs are static and toxic and/or have a narrow spectrum of activity. Thus, there is an urgent need for the development of new antifungal drugs. The fungal sphingolipid glucosylceramide (GlCer) is critical in promoting virulence of a variety of human-pathogenic fungi. In this study, we screened a synthetic drug library for compounds that target the synthesis of GlCer, but not mammalian, GlCer and found two compounds [(N’-(3-bromo-4-hydroxybenzylidene)-2-methylbenzohydrazide (BHBM) and its derivative, 3-bromo-N’-(3-bromo-4-hydroxybenzylidene) benzohydrazide (D0) that were highly effective in vitro and in vivo against several pathogenic fungi. BHBM and D0 were well tolerated in animals and are highly synergistic or additive to current antifungals. BHBM and D0 significantly affected fungal cell morphology and resulted in the accumulation of intracellular vesicles. Deep-sequencing analysis of drug-resistant mutants revealed that four protein products, encoded by genes APL5, COS111, MKKI, and STE2, are involved in vesicular transport and cell cycle progression, are targeted by BHBM.

IMPORTANCE Fungal infections are a significant cause of morbidity and mortality worldwide. Current antifungal drugs suffer from various drawbacks, including toxicity, drug resistance, and narrow spectrum of activity. In this study, we have demonstrated that pharmaceutical inhibition of fungal glucosylerceramide presents a new opportunity to treat cryptococcosis and various other fungal infections. In addition to being effective against pathogenic fungi, the compounds discovered in this study were well tolerated by animals and additive to current antifungals. These findings suggest that these drugs might pave the way for the development of a new class of antifungals.

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Recent estimates suggest that >300 million people are afflicted by serious fungal infections worldwide, with the most common and life-threatening invasive mycoses being cryptococcosis, candidiasis, aspergillosis, and pneumocystosis (1, 2). The number of fungal infections is rising in part due to the increasing number of individuals with immunocompromising medical conditions such as AIDS or those using immunosuppressive medical treatment, such as some cancer therapies (3). Three classes of antifungal drugs are currently available: azoles (e.g., fluconazole), polyenes (e.g., amphotericin B), and echinocandins (e.g., caspofungin). However, these drugs are static and toxic, have a narrow spectrum of activity,
and interact with other drugs such as chemotherapy agents and immunosuppressants (4).

Due to the limited number of antifungals available on the market, their increasing use stimulates the development of drug resistance, and thus, new classes of antifungals are desperately needed. The fungal sphingolipid glucosylceramide (GlcCer) is critical in virulence, and thus, new classes of antifungals are desperately needed. The fungal sphingolipid glucosylceramide (GlcCer) is critical in promoting virulence of several plant- and human-pathogenic fungi, including Cryptococcus neoformans, Candida albicans, and Aspergillus fumigatus (5–10).

GlcCer regulates fungal cell replication in environments of neutral or alkaline pH (5, 11). When fungal cells lacking GlcCer are exposed to neutral/alkaline pH, they cannot progress through the cell cycle, and cytokinesis does not occur (5, 9, 12). GlcCer is most likely a panfungal virulence factor required during infection to promote fungal growth at neutral/alkaline environments in the host (e.g., alveolar spaces, cerebrospinal fluid, and bloodstream); as such, it is a promising novel drug target.

Antibodies to fungal GlcCer exert antifungal effects in vitro and in vivo (13); however, inhibitors that block fungal, but not mammalian, GlcCer synthesis are not available. In this study, we screened a synthetic drug library for compounds that specifically target the synthesis of fungal GlcCer. Two such compounds were found [N’-(3-bromo-4-hydroxybenzylidene)-2-methylbenzohydrazide (BHBM) and 3-bromo-N’-(3-bromo-4-hydroxybenzylidene) benzohydrazide (D0)] and were highly effective in vitro and in vivo against a series of pathogenic fungi. These compounds are safe and well tolerated in animals, demonstrate good pharmacokinetic properties, and are highly synergistic or additive to current antifungals. The compounds target the transport of fungal vesicles, which is how fungal ceramide is transported for the synthesis of GlcCer. Thus, we have identified a new class of antifungals that could potentially be used alone or in combination with existing antifungal drugs.

RESULTS

BHBM and D0 inhibit the synthesis of GlcCer. GlcCer regulates the growth of C. neoformans in neutral and alkaline pH (14). This phenotype was utilized for screening of compounds that can inhibit fungal GlcCer. A ChemBridge DIVERSet-CL library containing 49,120 compounds was screened to identify molecules that inhibit the growth of C. neoformans at neutral and alkaline pH. A total of 220 compounds with a MIC of ≤1 μg/ml were screened in acidic (4.0) pH. The 18 compounds that were inactive (MIC > 32 μg/ml) were subjected to an in vivo labeling assay for the inhibition of GlcCer synthesis. Two compounds, identified as N’-(3-bromo-4-hydroxybenzylidene)-2-methylbenzohydrazide (BHBM) and 3-bromo-N’-(3-bromo-4-hydroxybenzylidene) benzohydrazide (D0), a derivative of BHBM, significantly inhibited the synthesis of GlcCer in C. neoformans but not mammalian (1774.16) cells (Fig. 1). Both compounds were fungicidal in vitro with minimum fungicidal concentrations (MFC) of 4 μg/ml for BHBM and 1.2 μg/ml for D0. Because of their structure, we designated this class of compounds “hydrazycins.”

BHBM and D0 exert antifungal activity. The antifungal activity of BHBM was examined against a variety of clinically relevant fungi (Table 1). BHBM showed good in vitro activity against various Cryptococcus species with the MIC ranging between 0.25 μg/ml and 8 μg/ml. Importantly, a fluconazole-resistant C. neoformans strain (MIC > 64 μg/ml) and various virulent Cryptococcus gattii strains were highly susceptible to BHBM (MICs of 1 to 2 μg/ml and 0.5 to 2 μg/ml, respectively). Other fungi against which BHBM was highly active included Rhizopus oryzae, Blastomyces dermatitidis, Histoplasma capsulatum, Pneumocystis carinii, and Pneumocystis jirovecii (Table 2). BHBM showed moderate activity (MIC range of 2 to 32 μg/ml) against Candida krusei, Candida glabrata (depending on the strain), Candida guilliermondii, Candida parapsilosis, A. fumigatus, and Coccidioides spp. and poor activity (MIC > 32 μg/ml) against C. albicans, Candida parapsilosis, and Paeococales variotii (Table 1). In addition, both BHBM and D0 were highly synergistic when combined with fluconazole and amphotericin B and additive when combined with caspofungin (see Table S1 in the supplemental mate-
BHBM was synergistic with tunicamycin. In contrast to fluconazole, C. neoformans cells did not develop resistance to either BHBM or D0 after 15 days of in vitro passages (120 generations) with a population size of 10⁶ cells/ml (data not shown).

**BHBM and D0 are fungicidal against intracellular C. neoformans.** Since both BHBM and D0 exhibited fungicidal activity against C. neoformans, we examined their killing activity using a time-kill assay. BHBM showed concentration-dependent killing (Fig. 2A), whereas D0 showed time-dependent killing (Fig. 2B). BHBM is less inhibitory toward GlcCer synthesis compared to D0, and it is only at higher concentrations that its inhibitory action becomes comparable to that of D0 (Fig. 1). Thus, the dose response associated with BHBM might be directly related to its ability to inhibit GlcCer synthesis, resulting in increased fungicidal activity with increased GlcCer inhibition. C. neoformans is a facultative intracellular pathogen and is able to replicate within phagocytic cells, such as macrophages. Thus, the ability of BHBM to kill phagocytosed C. neoformans cells was also examined. The J774.16 macrophage cell line was infected with C. neoformans in the presence of opsonins. The opsonins and extracellular fungal cells were then washed off, and BHBM was added to test its activity against intracellular fungal cells. Analysis of CFU revealed that BHBM significantly decreased the intracellular replication of C. neoformans inside macrophages. This effect was dose dependent, particularly after 12 and 24 h of incubation (Fig. 2C). Drugs showed little cytotoxic activity against the J774 cells with a 50% effective concentration (EC₅₀) of 50 µg/ml for both drugs (see Fig. S1 in the supplemental material), leading to a favorable EC₅₀/MIC₈₀ selectivity index (50 for BHBM and >100 for D0). In addition, toxicity was also tested in A549 and L2 mammalian cell lines, and EC₅₀ of 112 µg/ml and 72 µg/ml were found for the two cell lines, respectively. These EC₅₀ concentrations did not inhibit mammalian GlcCer synthesis (data not shown), suggesting that drug toxicity at these concentrations is not due to the inhibition of mammalian GlcCer.

**BHBM and D0 have potent antifungal activity against cryptococcosis.** The efficacy of BHBM and D0 was tested in mouse models of cryptococcosis. Treatment was initiated the same day that mice were infected intranasally with 5 × 10⁵ C. neoformans cells with a drug concentration of 1.2 mg/kg of body weight/day. Most (90%) BHBM-treated mice and 70% of D0-treated mice survived C. neoformans infection for up to 60 days, whereas 100% of untreated mice died within 33 days (P < 0.001 for both BHBM- and D0-treated mice versus untreated mice) (Fig. 3A). The fungal burden in the lungs and brain were studied during the course of infection. Drug treatment significantly reduced the fungal burden in both organs, and no cells were detected in the brain tissue at the end of the study (see Fig. S2 in the supplemental material). It is possible that the drugs were able to act in the lungs or the bloodstream and prevent fungal dissemination to the brain. However, the total absence of fungal cells in the brain at later time points (Fig. S2B) suggests that perhaps the drugs may act directly in the brain tissue. Drugs did not show significant in vivo toxicity. Blood work of BHBM- and D0-treated mice after 60 days of treatment showed a slight but insignificant increase of liver aspartate amino-

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**TABLE 1** In vitro antifungal activities of N’-(3-bromo-4-hydroxybenzylidene)-2-methylbenzohydrazide (BHBM) determined by the MICs against several fungal clinical isolates and reference strains

<table>
<thead>
<tr>
<th>Species (strain) (n)</th>
<th>MIC range (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryptococcus neoformans (FTL strains) (13)</td>
<td>0.25–8</td>
</tr>
<tr>
<td>Cryptococcus neoformans (MDPL strains and S isolates) (8)</td>
<td>0.5–2</td>
</tr>
<tr>
<td>Cryptococcus neoformans (MDPL strains and R isolates) (4)</td>
<td>1–2</td>
</tr>
<tr>
<td>Candida albicans (FTL strains) (3)</td>
<td>0.5–2</td>
</tr>
<tr>
<td>Candida albicans (MDPL strains) (5)</td>
<td>&gt;32</td>
</tr>
<tr>
<td>Candida krusei (MDPL strain) (1)</td>
<td>32</td>
</tr>
<tr>
<td>Candida krusei ATCC 6258 (MDPL strain)</td>
<td>16</td>
</tr>
<tr>
<td>Candida krusei QC FTL strains</td>
<td>8</td>
</tr>
<tr>
<td>Candida glabrata (FTL strains) (10)</td>
<td>0.125–2</td>
</tr>
<tr>
<td>Candida glabrata (MDPL strains) (3)</td>
<td>4–32</td>
</tr>
<tr>
<td>Candida parapsilosis (MDPL strains) (3)</td>
<td>&gt;32</td>
</tr>
<tr>
<td>Candida parapsilosis QC FTL strains</td>
<td>&gt;16</td>
</tr>
<tr>
<td>Candida guilliermondii (FTL strains) (3)</td>
<td>2–16</td>
</tr>
<tr>
<td>Aspergillus fumigatus (MDPL strain) (1)</td>
<td>8</td>
</tr>
<tr>
<td>Aspergillus fumigatus (MDPL strains) (3)</td>
<td>&gt;16</td>
</tr>
<tr>
<td>Rhizopus oryzae (FTL strains) (3)</td>
<td>2</td>
</tr>
<tr>
<td>Blastomyces dermatitidis (FTL strains) (10)</td>
<td>0.5–1</td>
</tr>
<tr>
<td>Histoplasma capsulatum (FTL strains) (10)</td>
<td>0.125–1</td>
</tr>
<tr>
<td>Coccioides spp. (FTL strains) (10)</td>
<td>8–16</td>
</tr>
<tr>
<td>Pneumocystis carinii FTL strains</td>
<td>&gt;16</td>
</tr>
</tbody>
</table>

TABLE 2  **In vitro** antifungal activities of BHBM determined by the percentage inhibition of ATP (IC₅₀) against *Pneumocystis murina* and *Pneumocystis jirovecii*.

<table>
<thead>
<tr>
<th><em>P. murina</em> HA isolates</th>
<th>24 h</th>
<th>48 h</th>
<th>72 h</th>
<th>96 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>BHBM IC₅₀ (µg/ml) after exposure for:</td>
<td>1.02</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>2.02</td>
</tr>
<tr>
<td><em>P. jirovecii</em> VM isolates</td>
<td>0.912</td>
<td>0.159</td>
<td>0.074</td>
<td>0.072</td>
</tr>
</tbody>
</table>

*IC₅₀, 50% inhibitory concentration.*
analysis was performed using analysis of variance (ANOVA). Data were compiled from three independent experiments.

4 treated mice succumbed 11.1 protected the mice from infection (Fig. 3B). BHBM- and D0- compared with the control, although none of the drugs completely postinfection. All drugs prolonged survival significantly com-

transaminase (AST) (125 ± 75 U/liter) in BHBM-treated mice com-
pared to untreated mice (50 ± 9 U/liter). All other blood parameters for liver and kidney function were normal, as were the number of erythrocytes, thrombocytes, and leukocytes.

Pharmacokinetic studies following BHBM treatment were performed in immunocompetent control healthy mice and immuno-
suppressed/infected (IS/IN) mice. In immunocompetent mice, the half-life of high-dose intravenous (i.v.) BHBM was 1.03 h. The half-lives of BHBM after low-dose and high-dose administration were 1.70 and 1.43 h, respectively (see Table S2 in the supplemental material). In infected mice, the half-lives of BHBM after low-dose and high-dose administration were 0.79 h and 0.84 h, respectively. The changes in the half-lives of the drugs between immunocompetent and immunosuppressed/infected animals are due to interactions between the drugs and corticosteroids used for immunosuppression; we have found that these drugs do not in-
teract with another immunosuppressant, cyclophosphamide (data not shown).

A second mouse survival study was performed to test the efficacy of BHBM and D0 against invasive cryptococcal infection of the central nervous system (CNS). Fluconazole and amphotericin B were also included in these studies to compare the efficacies of these compounds to those of commonly used drugs. All mice in the vehicle-treated control group succumbed 7.0 ± 1.1 days postinfection. All drugs prolonged survival significantly com-
pared with the control, although none of the drugs completely protected the mice from infection (Fig. 3B). BHBM- and D0-
treated mice succumbed 11.1 ± 3.7 (P < 0.05 versus untreated mice) and 10.5 ± 4.2 days (P < 0.05 versus untreated mice) postinfection, respectively. This was similar to the survival pattern observed with fluconazole, with an average survival of 10 ± 2 days postinfection (P < 0.05 versus untreated mice). Mice treated with amphotericin B succumbed at an average of 23.5 ± 6.5 days postinfection (P < 0.005 versus untreated mice).

**FIG 2** Killing activity of BHBM and D0. Killing activity was determined using an in vitro killing assay in which the compounds were added to C. neoformans cells, which were then incubated at 35°C, 5% CO2, and pH 7.4. The number of CFU were counted during the 96-h incubation. (A and B) Both BHBM (A) and D0 (B) showed fungicidal activity. BHBM showed concentration-dependent killing, whereas D0 showed time-dependent killing. D0 is more effective in killing C. neoformans (100% dead cells within 24 h), and the killing activity does not occur earlier than 24 h with higher doses. BHBM kills more slowly, requiring at least 72 h of incubation to kill about 50% of the cells. Values that are significantly different are indicated as follows: *, P < 0.05 comparing treated cells versus untreated cells (no drug); #, P < 0.001 comparing treated cells versus untreated cells (no drug). (C) Intracellular activity of BHBM was assessed by incubating macrophages with internalized C. neoformans cells with different concentrations of BHBM in the absence of opsonins. Values that are significantly different are indicated as follows: *, P < 0.05 comparing extracellular or intracellular treated (0.25, 1, or 4 μg/ml) versus extracellular or intracellular untreated (0 μg/ml), respectively. Statistical analysis was performed using analysis of variance (ANOVA). Data were compiled from three independent experiments.
13 days of T/S treatment versus the negative control and versus BHBM groups (Fig. S2C and S2D).

BHBM and D0 have antifungal activity against invasive candidiasis. Most Candida spp. examined were resistant in vitro to both BHBM and D0. However, previous studies have shown that GlcCer is required for virulence through a mechanism other than facilitating growth at neutral/alkaline pH (8), which is the pH used in our library screening. In vitro treatment of C. albicans cells with BHBM inhibited GlcCer synthesis in a dose-dependent manner (data not shown). Thus, the efficacy of BHBM or D0 against invasive candidiasis was evaluated. Invasive candidiasis was established in CBA/J mice by intravenous injection with a lethal dose of $10^5$ C. albicans cells. Treatment started the same day with either BHBM or D0 using the same dose regimens used to treat cryptococcosis. After 21 days of infection, 75% of mice treated with D0 and 62.5% of mice treated with BHBM were still alive, whereas all untreated mice had succumbed by day 13 (Fig. 3D).

BHBM and D0 target metabolism and/or transport of certain ceramide species used for synthesis of GlcCer. Since the initial screening had demonstrated that BHBM and D0 affect the synthesis of GlcCer, biochemical studies were performed by thin-layer chromatography (TLC) and mass spectrometry to understand the effects of drugs on the sphingolipid pathway. Lipid profiling by TLC revealed that BHBM treatment led to a decrease in GlcCer levels and increased dihydroceramide, sphingosine, and sphingosine-1-phosphate (Fig. 4a). These changes were confirmed by mass spectrometry (Fig. 4b to e). Interestingly, the levels of C18 hydroxyceramide, the ceramide species that is a direct precursor of GlcCer, and its corresponding desaturated and methylated forms decreased upon BHBM (Fig. 4F to H) or D0 (data not shown) treatment, confirming a dysregulation of the ceramide synthesis upon treatment with BHBM. No significant changes in GlcCer levels were observed in mammalian cells after BHBM treatment at various time points (see Fig. S3 in the supplemental material).

BHBM affects fungal cell morphology and vesicular structure. The relationship between lipid composition and Golgi architecture is well established (15, 16). Thus, we set out to evaluate the effect of BHBM treatment on the Golgi accumulation of NBD C6-ceramide $\{N-[7-(4-nitrobenzo-2-oxa-1,3-diazole)]-6$-aminocaproyl-D-erythro-sphingosine$, which is a direct precursor to GlcCer$. The peripheral distribution reported for C. neoformans was observed for NBD C6-ceramide staining in control cells; however, a substantial accumulation of this compound was detected in BHBM-treated yeasts of C. neoformans as observed by microscopy and confirmed by fluorescence-activated cell sorting (FACS) analysis (Fig. 5A and B). The majority of BHBM-treated yeasts displayed a ring accumulation of NBD C6-ceramide. Similar results were observed when C. albicans cells were treated with BHBM (Fig. 5C).

FIG 3 Effects of BHBM and D0 on survival of mice upon infection with C. neoformans, P. murina, or C. albicans. (A) Cryptococcosis. Mice were infected intranasally with C. neoformans (Cn) and received 1.2 mg/kg/day of either BHBM or D0 intraperitoneally (10 mice per group). Values that are significantly different are indicated as follows: *, $P < 0.001$ comparing BHBM- or D0-treated mice versus untreated mice (no drug). (B) Cryptococcosis. Survival of mice infected intravenously with C. neoformans (8 mice per group) and treated intraperitoneally either with BHBM, D0, fluconazole (FLC), or amphotericin B (AMB). Values that are significantly different are indicated as follows: $^\wedge$, $P < 0.05$ comparing BHBM- or D0-treated mice versus untreated mice and AMB-treated mice versus untreated mice. (C) Pneumocystosis. Survival of corticosteroid-immunosuppressed (Cort) or CD4-depleted (CD4d) mice infected intranasally with P. murina after 13 days of intraperitoneal treatment with BHBM (3.2 mg/kg/day), D0 (1.25 mg/kg/day), or trimethoprim-sulfamethoxazole (T/S) (12 mice per group). Values that are significantly different are indicated as follows: $^\&$, $P < 0.05$ comparing BHBM- or D0-treated mice versus respective untreated controls. (D) Candidiasis. Survival of mice infected with C. albicans SC 5314 (Ca) intravenously received no drug or 1.2 mg/kg/day of either BHBM or D0 intraperitoneally (8 mice per group). Values that are significantly different are indicated as follows: $^\$, $P < 0.01$ comparing BHBM- or D0-treated mice versus untreated mice. Statistical analysis for survival studies was performed using Kruskal-Wallis test and by Student-Newman-Keuls t test for multiple comparisons using INSTAT.
were observed in C. albicans, confirming an altered distribution of NBD C6-ceramide after BHBM in hyphal forms. In addition, BHBM treatment significantly reduced the yeast-to-hypha differentiation in C. albicans (Fig. 5A). This defect in change of morphology might explain the loss of virulence in candidiasis (Fig. 3D), as Candida species that are locked in the yeast form are known to lose virulence (17). Additional fluorescent images are shown in Fig. S4 in the supplemental material.

Electron microscopy examination of drug-treated cells showed an accumulation of large vesicles inside the cells (Fig. 6A), while control cells maintained a regular organization and structure of intracellular vesicles (Fig. 6B). Vesicle accumulation in drug-treated cells was observed as single vesicles (Fig. 6A and D, arrows) and fused vesicles (Fig. 6A and D, arrowheads). Morphometric analysis revealed that there was a significant increase (P < 0.05) in the volume occupied by the intracellular vesicles in BHBM-treated cells (Fig. 6C). The numbers of vesicles per square micrometer of cell surface area were 0.84 ± 0.58 in BHBM-treated cells and 0.93 ± 0.54 in D0-treated cells compared to 0.33 ± 0.36 in untreated cells. The increase in both volume and number of intracellular vesicles suggests defects in secretion processes induced by the drug (Fig. 6C). In addition, plasma membrane-limited structures, which seemed to accumulate portions of the cytoplasm, were frequently seen accumulated in the periplasmic space (Fig. 6E). Transmission electron microscopy (TEM) images of untreated cells (Fig. 6C) and BHBM-treated cells (Fig. 6E) were frequently seen accumulated in the periplasmic space and other complex sphingolipids (19–21). These genes are conserved in C. neoformans, but they have not been characterized experimentally. The other six genes detected in the HIP-HOP assay comprising of essential genes as heterozygous diploids and nonessential genes as homozygous deletion profiling (18) was screened to find the potential pathways affected by the drug. Of this genome-wide collection of 5,900 mutants, 23 were >2-fold more sensitive to BHBM than the vehicle-treated control (see Table S3 in the supplemental material). Of these 23 genes, 8 were not considered further because they corresponded to dubious open reading frames or lacked any homology to C. neoformans. Among the remaining 15 significant sensitive genes, 9 genes (RET2, UBP3, SEC26, PEP7, SEC31, YML018C, SNP8, GOS1, and RET3) are involved in the regulation of vesicle sorting or transport between Golgi apparatus and endoplasmic reticulum (ER). This was in agreement with the observations of fungal cell morphology and vesicular structure after drug treatment (Fig. 5 and 6). These findings were also consistent with the decrease of GlcCer levels upon BHBM treatment, as vesicle transport is the main mechanism by which ceramide is transported between the ER and Golgi apparatus for the synthesis of GlcCer.

To pinpoint the cellular targets of BHBM, a second approach, generation of BHBM-resistant mutants, was followed. The drug-sensitive S. cerevisiae RYO0622 strain was used for the generation of mutants (23). A prescreening study exposing this strain to var-

**FIG 4** Measurements of sphingolipids upon treatment with BHBM. (a) Thin-layer chromatography analysis of sphingolipids (see diagram) isolated from untreated or BHBM-treated C. neoformans cells after in vivo labeling with [3H]palmitate ([3H]-Palm). (b to h) Lipid analysis by LC-MS. (b) C18 hydroxy (Δ8) 9 methyl-glucosylceramide (GlcCer); (c) C18 dihydroceramide (C18 dhCer); (d) sphingosine and dihydrophosphogluco cerevisiae HIP-HOP (haploinsufficiency profiling-homozygous deletion profiling) assay comprising of essential genes as heterozygous diploids and nonessential genes as homozygous diploids (18) was screened to find the potential pathways affected by the drug. Of this genome-wide collection of 5,900 mutants, 23 were >2-fold more sensitive to BHBM than the vehicle-treated control (see Table S3 in the supplemental material). Of these 23 genes, 8 were not considered further because they corresponded to dubious open reading frames or lacked any homology to C. neoformans. Among the remaining 15 significant sensitive genes, 9 genes (RET2, UBP3, SEC26, PEP7, SEC31, YML018C, SNP8, GOS1, and RET3) are involved in the regulation of vesicle sorting or transport between Golgi apparatus and endoplasmic reticulum (ER). This was in agreement with the observations of fungal cell morphology and vesicular structure after drug treatment (Fig. 5 and 6). These findings were also consistent with the decrease of GlcCer levels upon BHBM treatment, as vesicle transport is the main mechanism by which ceramide is transported between the ER and Golgi apparatus for the synthesis of GlcCer and other complex sphingolipids (19–21). These genes are conserved in C. neoformans, but they have not been characterized experimentally. The other six genes detected in the HIP-HOP screen (MTW1, REG1, SPT4, SAP4, GLC7, and ACT1) regulate chromatin organization, cell cycle progression, or cell division, functions that are known to be regulated by GlcCer (14, 22).
ious concentrations of BHBM revealed that a drug concentration of 133 μg/ml completely inhibited yeast growth (100% inhibitory concentration [IC\textsubscript{100}]). Incubating 10\textsuperscript{6} cells of \textit{S. cerevisiae} RYO0622 strain with BHBM at the IC\textsubscript{100} resulted in seven resistant colonies, the genomes of which were sequenced and compared with the NCBI sacCer3 reference genome using the Genome Analysis Toolkit (GATK). This analysis led to the identification of mutations in four genes, \textit{APL5, COS111, MKK1}, and \textit{STE2}, that were present in all resistant mutants. The proteins encoded by these loci are known to be involved in vesicle trafficking, budding, and cell cycle progression (24–27), again in agreement with the observations of cell morphology and known phenotypes in the absence of GlcCer (14, 22). These findings also closely match the pathway proposed by the HIP-HOP analysis. Interestingly, these four genes each interact with UBI4 which encodes ubiquitin and which is conjugated to proteins to target them for degradation. UBI4 is a member of the endomembrane recycling pathways as defined by Finley et al. (28).

To confirm drug resistance, individual mutants along with the parent strain were grown in the presence of various concentrations of BHBM. Various concentrations of fluconazole and methyl methane sulfonate (MMS) were used as controls (Fig. 7). All individual mutants showed increased resistance to BHBM in the range of 11 to 92 μg/ml (Fig. 7), while all mutants showed similar susceptibility to fluconazole and MMS. The increased resistance of these mutants to BHBM treatment confirms that the above-mentioned genes are indeed the targets of BHBM, and the absence of these targets impairs the killing activity of the drug. It is worth noting that deep-sequencing analysis also revealed the presence of mutations in another gene, \textit{SLA2}, that is also involved in vesicular transport (29). Although the Δ\textit{sla2} mutant was not resistant to BHBM, it remains to be determined whether point mutations within the \textit{SLA2} open reading frame do in fact confer resistance.

**DISCUSSION**

In this study, we have identified and characterized a new class of antifungal compounds, the hydrazycins (specifically BHBM and D0), which decrease the synthesis of fungal, but not mammalian, GlcCer. These compounds were effective against cryptococcosis, candidiasis, and pneumocystosis in animal models and target four genes involved in vesicular trafficking and cell budding. The compounds were well tolerated by animals and can potentially pave the way for the development of a new class of antifungals.

Both BHBM and D0 are acylhydrazone compounds with the structure R\textsubscript{1}R\textsubscript{2}C = NNH\textsubscript{2} (Fig. 1) (30–32). Hydrazone molecules have been shown to exert diverse biological functions, including antibacterial, antifungal, antiviral, antiparasitic, anticancer, and antidepressant effects (33–36), but these activities are linked to specific structures of hydrazones, which we can readily modify.
Although previous literature exists on antimicrobial activity of hydrazone compounds, this is the first time that the ability of these compounds to block fungal GlcCer has been discovered. BHBM and D0 exhibited antifungal activity against a variety of clinically relevant fungi (Table 1) and acted synergistically with currently available antifungal drugs (see Table S1 in the supplemental material). The ability of these compounds to exert fungicidal activity on phagocytosed \textit{C. neoformans} cells is particularly important (Fig. 2). \textit{C. neoformans} is able to survive and replicate inside macrophages (37), and replication inside nonactivated macrophages is an important factor in its pathogenicity (38). Furthermore, BHBM and D0 exhibited low cytotoxicity in a macrophage cell line (see Fig. S1 in the supplemental material) and a good selectivity index (EC50/MIC ratio > 50), which is promising for their use as antifungals.

BHBM and D0 were effective against a variety of fungal infections \textit{in vivo}. In mouse models of cryptococcosis, the drugs significantly increased the survival of mice that were infected with a lethal dose of \textit{C. neoformans} via the intranasal route (Fig. 3A). Analysis of fungal burden demonstrated that drug treatment reduced the number of fungal cells in the brain throughout the course of the treatment, leading to the absence of fungal cells in the brain after 60 days (see Fig. S2 in the supplemental material). This observation indicates that drugs were able to reach the brain and clear the infection. This hypothesis is supported by the observa-

![FIG 6 Transmission electron microscopy examination of high-pressure frozen and freeze substituted \textit{C. neoformans} yeast cells. (A, D, and E) Cells treated with 4 \(\mu\)g/ml of BHBM for 6 h, showing accumulation of intracellular vesicles (arrows) and regions containing a "reticulum" presumably formed by the fusion of intracellular vesicles (arrowheads). M, mitochondrion. (B) Control cells showing a regular aspect. (C) Morphometric analysis results showing the volumetric density of the intracellular vesicles in control and treated cells. (E) Accumulation of cytoplasm-rich vesicles in the periplasmic space. Images and data are representative of the results of three separate experiments.](https://mbio.asm.org/)

![FIG 7 Effects of BHBM, fluconazole, and methyl methane sulfonate (MMS) on wild-type BY4741 and \Delta apl5, \Delta cos111, \Delta mkk1, and \Delta ste2 deletion strains. Relative growth inhibition was calculated by the average rate after normalizing the OD600 values in drug-treated wells against the DMSO control wells on each assay plate. The mutant strains show increased resistance to BHBM but not to fluconazole or MMS. Results are from two independent growth assays.](https://mbio.asm.org/)
tion that BHBM is readily found in brain tissue upon administration (data not shown), and this tissue distribution may explain the low serum half-life and the effect on cryptococcal meningitis. In an aggressive model of cryptococcosis, in which the infection was established via tail vein injection, the drugs performed similarly to fluconazole in prolonging survival in our murine model (Fig. 3B). The concentration of fluconazole was almost eight times higher than that of BHBM and D0, suggesting that these drugs could be more effective than fluconazole. While amphotericin B was the most effective drug in this mode of infection, this drug is associated with drawbacks such as adverse reactions and nephrotoxicity (39, 40). BHBM and D0 are well tolerated by animals; thus, it is possible to augment their dose regimen for improved antifungal effect.

In addition to cryptococcosis, both drugs were effective in in vivo infection models of pneumocystosis and candidiasis (Fig. 3C and D). In the case of pneumocytosis, BHBM was able to increase survival in corticosteroid-treated mice, while D0 was effective in improving the survival of CD4-depleted mice. Although T/S was the most effective drug, this drug has significant side effects, and relapse and recurrence of infections are often observed when using secondary therapies (41). Thus, BHBM and D0 can potentially provide new options for pneumocystosis treatment. Previous studies have shown antifungal activity of fluconazole against C. albicans in vivo despite in vitro resistance (42). A similar phenomenon was observed with our drug compounds. It has been shown that GlcCer-deficient mutants in C. albicans lack the ability to cause virulence in the mouse model despite showing no growth defect in vitro (8). Thus, inhibition of GlcCer by BHBM and D0 is the likely mechanism for partially protecting the mice against invasive candidiasis.

Both BHBM and D0 inhibit the synthesis of GlcCer and significantly affect the levels of various sphingolipids (Fig. 4). However, GlcCer is not the only target of these drugs. Loss of GlcCer has a static effect on fungal growth (6), whereas BHBM and D0 are fungicidal. Interestingly, sphingosine species accumulate in fungal cells upon BHBM or D0 treatment (Fig. 4), a phenomenon that is not observed when the glucosylceramide synthase gene (GCS1) is deleted (5), and these sphingolipids are highly toxic. These sphingolipids do not significantly accumulate in mammalian cells treated with BHBM or D0 (see Fig. S3 in the supplemental material), and this may also explain the lack of toxicity in this system.

The results of HIP-HOP analysis provided an indication that the drugs affect the genes involved in vesicular transport, cell budding, and cell cycle progression. Of the 23 genes identified in this analysis, 9 were involved in the regulation of vesicle sorting or transport between Golgi apparatus and ER, and endocytosis and another 6 genes were involved in cell budding and cell cycle progression (see Table S3 in the supplemental material). These observations suggested that vesicular transport, cell division, and cell cycle progression are targeted by the drugs. This provided a potential target pathway, which was supported by the electron microscopy images showing an accumulation of intracellular vesicles (Fig. 6).

Further understanding of the drug targets was provided by generation of mutants resistant to BHBM. Mutations in four genes, APL5, COS111, MKK1, and STE2, were found in all seven colonies that showed resistance to BHBM. The corresponding single deletion mutants of each of these genes (from the yeast deletion collections [43]) showed increased resistance to BHBM, but no difference in resistance to the unrelated drugs fluconazole or MMS (Fig. 7), confirming that these genes encode proteins that have a critical role in the cellular response to BHBM. Of these 4 genes, APL5 plays an important role in vesicular transport and has been shown to be involved in cargo inclusion into vesicles during the secretory and endocytic pathways (44, 45). Furthermore, the APL5 protein directly interacts with the VPS3 and GOS1-encoded proteins, which were identified by the HIP-HOP assay (46, 47). Similarly, MKK1 directly interacts with RET2 and RET3 identified by the HIP-HOP assay (48), COS111 and STE2 may also interact with genes identified by the HIP-HOP assay in a process mediated by ubiquitin (49–52). In fact, one of the genes identified by the HIP-HOP assay to be highly upregulated is UBP3, which encodes a ubiquitin-specific protease working together with UBI4 to regulate protein degradation and endocytosis. In fact, UBI4 physically interacts with all the genes identified by the HIP-HOP assay except SPT4, SAP4, SNF8, and VPS3. UBI4 also interacts with all proteins identified by the deep-sequencing analysis. UBI3 is also essential for the regulation of the COPII system (encoded by RET2, RET3, SEC26, and SEC31). Thus, UBI4 may be involved in the regulation of endocytosis and vesicle transport by interacting with RET2, RET3, UBP3, REG1, SEC26, SEC31, and PEP7, all of which are genes identified by the HIP-HOP analysis. UBI5 may be a hub between the proteins identified by the deep-sequencing analysis and those identified by the HIP-HOP assay. Vesicular transport is the main mechanism by which ceramide is transported from the ER to the Golgi apparatus for the synthesis of GlcCer, which in turn regulates cell cycle progression (5, 9, 12). The results of deep-sequencing analysis suggest that the drug compounds inhibit a protein product in the endocytic/vesicle transport pathway that leads to the formation of GlcCer; this will strip the drug of its main target, leading to increased resistance.

In summary, we have identified hydrazine molecules (BHBM and D0) that exhibit potent antifungal activity through a specific inhibition of fungal GlcCer. These molecules exerted antifungal activity in the mouse models of cryptococcosis, pneumocystis, and candidiasis and were well tolerated by animals. BHBM and D0 also affected fungal cell morphology and vesicular trafficking. These molecules target four genes that are involved in vesicular secretion and cell cycle progression. To the best of our knowledge, this study marks the first drugs that act based on the inhibition of fungal GlcCer. Given the antifungal potency of these drugs, further studies are warranted for their preclinical and eventually clinical development.

MATERIALS AND METHODS

Strains, media, and reagents. A series of fungal clinical isolates and reference strains were used in this study, including Cryptococcus neoformans (strain H99), Cryptococcus gattii, Candida albicans, Candida krusei, Candida glabrata, Candida parapsilosis, Candida guilliermondii, Aspergillus fumigatus, Rhizopus oryzae, Blastomyces dermatitidis, Histoplasma capsulatum, Coccidioides spp., Paeilomyces variotii, Pneumocystis murina, Pneumocystis jiroveci, and Saccharomyces cerevisiae. These strains were obtained from existing collections at the Fungal Testing Laboratory (University of Texas Health Science Center at San Antonio, San Antonio), Del Poeta’s laboratory (Stony Brook University), Cushion’s laboratory (University of Cincinnati), and Nilson’s laboratory (University of British Columbia). Yeast peptone dextrose (YPD), yeast nitrogen base (YNB), Luria-Bertani (LB), Roswell Park Memorial Institute (RPMI 1640), and Dulbecco’s modified Eagle’s medium (DMEM) were purchased from Invitrogen Life Technologies (Grand Island, NY). Fluconazole, amphotericin B, dexa-
methasone, cyclophosphamide, and tunicamycin were purchased from Sigma-Aldrich (St. Louis, MO). Caspofungin and posaconazole were obtained from Merck (Rahway, NJ). Voriconazole was obtained from Pfizer (Rey Brook, NY). N-‘(3-bromo-4-hydroxybenzylidene)-2-methylbenzohydrazide (BHBM) and 3-bromo-N-‘(3-bromo-4-hydroxybenzylidene) benzohydrazide (D0) were obtained from ChemBridge (San Diego, CA). Cryptococcal capsular monoclonal antibody MAb 18B7 was a gift from Arturo Casadevall’s laboratory (Johns Hopkins University).

**Library screening.** The DIVERSet-CL library was obtained from ChemBridge (San Diego, CA) in a 96-well plate format and contained 10 mM compound per well in 100% dimethyl sulfoxide (DMSO). In each well, 10 compounds were mixed together. The compounds were first diluted to 1 mM each (1:10 dilution with 10% DMSO) with YNB medium buffered with HEPES at pH 7.4 containing 2% glucose and subsequently diluted to 300 μM (1:3.3 dilution) with the same medium (3% DMSO). A 100-μl aliquot of this solution was placed into each well of a 96-well plate and stored at −20°C until use. Then, 4 × 10⁵ C. neoformans H99 cells in 100 μl of YNB medium buffered at pH 7.4 with HEPES were added to each well. The final concentration of the tested drugs was 150 μM in YNB medium containing 1.5% DMSO. The plates were incubated at 37°C in the presence of 5% CO₂ for 48 h. The optical density at 495 nm (OD₄⁹⁵) was recorded using the FilterMax 5 multimode microplate reader (Molecular Devices, Sunnyvale, CA). Compound cocktails showing an OD of <80% compared to the OD in the control well (1.5% DMSO but no drug) were selected for further studies.

**Labeling of fungal cells with tritiated palmitate ([3H]palmitate).** C. neoformans cells were grown in YNB (pH 7.4) at 37°C in the presence of 5% CO₂ for 16 h. The cells were centrifuged for 10 min at 3000 rpm at room temperature. Supernatant was removed, and the cell pellet was suspended and counted. Next, 900 μl containing 5 × 10⁵ C. neoformans cells was placed into a 15-mI round-bottom Corning centrifuge tube. Then, 100-μl volumes of different concentrations of BHBM or D0 diluted in YNB containing 0.1% DMSO were added, resulting in final concentrations of 0.25, 1, and 4 μg/ml or 0.075, 0.3, and 1.2 μg/ml, respectively. The tubes were incubated in a shaker incubator at 225 rpm at 37°C in the presence of 5% CO₂ for 4 h. [3H]palmitate (30 μCi/ml) (PerkinElmer, Waltham, MA) was added to the culture and incubated for an additional 2 h. Cells without the drug were included as a negative control. The cells were then pelleted, washed once with distilled sterile water, and resuspended in 1.5 ml of Mandala lipid extraction buffer. Lipids were extracted by the methods of Mandala et al. (53) and Bligh and Dyer (54), followed by methanolic base hydrolysis as previously described. Extracted lipids were dried in an SPD210 SpeedVac system (Thermo Fisher Scientific). Dried lipids were resuspended in 30 μl of 1:1 methanol-chloroform and loaded on thin-layer chromatography (TLC) silica gel 60 plates (EMD Millipore, Billerica, MA). Glycosylceramide (GlcCer) standard from soybean (Avanti Polar Lipids, Alabaster, AL) was added in a separate lane as a control. The sample was resolved in a tank containing chloroform-methanol-water (65:25:4) as the mobile phase. The TLC plates were then dried and exposed to iodine vapor for the identification of the GlcCer standard band, which was marked. The TLC plate was then enhanced by spraying with Enhancer (PerkinElmer) and exposed to X-ray film at −80°C for 72 h, after which the film was developed.

**Labeling mammalian cells with tritiated palmitate ([3H]palmitate).** The murine macrophage cell line J774.16 (ATCC) was maintained in DMEM containing 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin (pen-strep) and were used until passage eight. Cells at a density of 5 × 10⁵ cells/ml were cultured in a six-well culture plate for 14 h to achieve adherence. BHBM or D0 at the same concentrations used for fungal cells (see above) was added to the plate for 4 h. Then, 30 μCi/ml of [3H]palmitic acid was added, and the plate was further incubated for 2 h. Labeled, untreated J774.16 cells were included as a control. The cells were harvested by adding 0.05% trypsin-EDTA and scraping with a cell scraper, and then the cells were washed once with phosphate-buffered saline (PBS) and dissolved in 2 ml methanol and 1 ml chloroform. Lipids were extracted by the method of Bligh and Dyer (54) followed by base hydrolysis. Lipid samples were dried in a SpeedVac vacuum concentrator, suspended in 30 μl of 1:1 methanol-chloroform, and loaded on a TLC plate with GlcCer as the standard.

**In vitro susceptibility testing.** MICs were determined following the methods of the Clinical and Laboratory Standards Institutes (CLSI) with modifications. RPMI 1640 or YNB medium (pH 7.0, 0.2% glucose) buffered with HEPES was used for MIC studies. HEPES was used instead of morpholinepropanesulfonic acid (MOPS), because MOPS was found to inhibit the activity of BHBM and D0. BHBM and D0 were serially diluted from 32 to 0.03 μg/ml or 19 to 0.02 μg/ml, respectively, in a 96-well plate with the respective medium. The inoculum was prepared as described in the CLSI protocol M27-A3 guidelines (55). The plates were incubated at 37°C with 5% CO₂ for 24 to 96 h. The MICs were determined as the lowest concentration of the drug that inhibited 50% of growth compared to the control. The minimum fungicidal concentration (MFC) was also determined by taking 100 μl of serial dilutions from each well, spreading it on YPD agar plates, and counting the CFU. MFC was defined as the lowest drug concentration that yielded three or fewer colonies (i.e., 99% of the inoculum was killed).

**In vitro killing assay.** C. neoformans cells from a culture grown overnight were washed in PBS and resuspended in YNB buffered with HEPES at pH 7.4. The cells were counted, and 2 × 10⁴ cells were incubated with 1, 2, or 4 μg/ml of BHBM or D0 in a final volume of 10 ml with a final concentration of 0.4% DMSO. The tubes were then incubated at 37°C with 5% CO₂ on a rotary shaker at 200 rpm. Aliquots were taken at time points and diluted, and 100-μl portions were plated onto YPD plates. YPD plates were incubated in a 30°C incubator and after 72 h, the numbers of CFU were counted and recorded.

**In vitro testing against P. murina and P. jiroveci.** A characterized P. carinii strain isolated from rat lung tissue was distributed into triplicate wells of 48-well plates with a final volume of 500 μl and a final concentration of 5 × 10⁷ nuclei/ml. The cells were incubated with either 0.1, 1, 10, or 100 μg/ml of BHBM or D0. The negative controls were media alone and ampicillin 10 μg/ml, and the positive control was pentamidine isethionate at 1 μg/ml. At 24, 48, and 72 h, 10% of the well volume was removed, and the ATP content was measured using PerkinElmer ATP-liteM luciferin-luciferase assay. The luminescence generated by the ATP content of the samples was measured by using a spectrophotometer (PolarStar Optima BMG, Ortenburg, Germany). Reduction in ATP consumption by P. jiroveci cells was considered a measure of drug efficacy. A control for toxicity, ATP consumption was also measured in A549 and L2 mammalian cells. A sample of each group was examined microscopically on the final day of the assay to rule out the presence of bacterial contamination.

**Intracellular effect of BHBM.** To assess whether BHBM would be effective against intracellular C. neoformans, J774.16 macrophage cells were incubated with C. neoformans cells at a 1:1 ratio in the presence of opsonins (complement and antibody MAb 18B7 against the cryptococcal capsular antigen). After 2 h of incubation, about 60 to 80% of macrophages had at least one C. neoformans cell internalized. At this time, cells were washed to remove extracellular fungal cells and fresh DMEM with different concentrations of BHBM was added to each well. The plates were incubated at 37°C with 5% CO₂. At 0, 6, 12, and 24 h, extracellular cells were collected by washing and plated onto YPD agar plates for counting the CFU of extracellular cells. Macrophages were also lysed and plated onto YPD agar plates for counting the CFU of intracellular fungal cells.

**Synergism assay.** Synergistic activity was assayed by calculating the fractional inhibitory index (FIC). Briefly, in a 96-well plate, drug A (either BHBM or D0) was serially diluted from 16 to 0.015 μg/ml (11 dilutions), whereas drug B (either fluconazole, amphotericin B, caspofungin, or tunicamycin) was serially diluted from 12 to 0.19 μg/ml, 5 to 0.078 μg/ml, 70 to 1.09 μg/ml, and 6 to 0.09 μg/ml (seven dilutions), respectively. The FIC was defined as (MIC combined/MIC drug A alone) + (MIC com-
bined (MIC drug B alone). Synergism was categorized as follows: strongly synergistic effect, FIC < 0.5; synergistic effect, FIC < 1; additive effect, FIC = 1; no effect, 1 < FIC < 2; antagonistic effect, FIC > 2.

**In vitro toxicity.** The murine macrophage cell line J774.16 was maintained in DMEM containing 10% FBS and 1% pen-strep. At passage 7, 10^6 cells/well in DMEM containing 10% FBS were transferred into 96-well plates and cultured for 14 h for the cells to adhere to the wells. BHBM or D0 was added to the cells at concentrations ranging from 0.1 to 100 μg/ml. The wells without the drug served as controls. The plate was incubated at 37°C with 5% CO2. After 24 or 48 h, the supernatant was removed, and 50 μl of 5-mg/ml 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) solution in PBS was added to each well. The plates were incubated for an additional 4 h. The formazan crystal formed inside the cell was dissolved by adding 50 μl of isopropanol containing 0.1 N HCl. The optical density was measured at 570 nm.

**Animal studies for cryptococcosis.** Survival studies were performed with two routes of infection, intranasal and intravenous (i.v.) (tail vein). For survival studies with intranasal infection, 4-week-old CBA/J female mice (Jackson Laboratory, Bar Harbor, ME) were used. Ten mice per treatment or control group were evaluated. Mice were infected by inoculation of 20 μl containing 5 × 10^5 cells of C. neoformans. Treated mice received an intraperitoneal (i.p.) injection of 1.2 mg/kg of body weight/day of either BHBM or D0 in a final volume of 100 μl of PBS containing 0.4% DMSO (PBS–0.4% DMSO) 2 h after receiving the inoculum. Untreated mice received 100 μl of PBS–0.4% DMSO. Mice were fed *ad libitum* and monitored closely for signs of discomfort and meningitis. Mice showing abnormal gait, lethargy, tremor, significant loss of body weight, or inability to reach water or food were sacrificed, and survival was counted until that day. At the end of the survival study, tissue burden culture was performed in mice that survived the infection. Mice were sacrificed and their organs were extracted and homogenized in 10 ml sterile PBS using a homogenizer (Stomacher80; Cole-Parmer, Vernon Hills, IL). Organ homogenates were serially diluted 1:10 in PBS, and 100-μl portions were plated on YPD agar plates and incubated at 30°C for 72 h for CFU count.

For survival studies using the intravenous injection of *C. neoformans*, 4-week-old CBA/J female mice (Jackson Laboratory, Bar Harbor, ME) were used. A total of 40 mice were infected by tail vein injection of 200 μl containing 10^6 cells of *C. neoformans* cells and were randomly separated into five groups (eight mice per group). The treated mice received an intraperitoneal injection of 1.2 mg/kg/day of BHBM, D0, or amphotericin B or 10 mg/kg/day of fluconazole in a final volume of 100 μl of PBS containing 0.4% DMSO a few hours after receiving the inoculum. Untreated mice received 100 μl of PBS–0.4% DMSO. Mice were fed *ad libitum* and monitored closely for signs of discomfort and meningitis. Mice showing abnormal gait, lethargy, tremor, significant loss of body weight, or inability to reach water or food were sacrificed, and survival was counted until that day.

**Animal studies for pneumocystosis.** For survival studies, 4- to 6-week-old C3H/HeN mice from the National Cancer Institute (Bethesda, MD) were used. Mice were infected with *P. murina* through exposure to mice with a fulminating *P. murina* infection (seed mice). These mice were immunosuppressed by the addition of 4 mg/liter dexamethasone to the drinking water. One milliliter of sulfuric acid was also added per liter of drinking water for disinfection. The seed mice were rotated within the cages for 2 weeks and then removed. After the mice had developed a moderate infection level (approximately 5 weeks), they were divided into groups containing 12 mice each: negative-control (control steroid), positive-control (trimethoprim-sulfamethoxazole [TS]), BHBM-treated, and D0-treated groups. BHBM or D0 was administered intraperitoneally or by oral gavage on a milligram/kilogram/day basis for up to 3 weeks. Treatment was initiated after the mice developed infection. The dose, route, and frequency of administration varied depending on the agent being tested. At the end of the treatment, mice were sacrificed and processed for analysis. Slides were made from the lung homogenates at different dilutions and stained with Diff-Quik to quantify the trophic forms and cresyl echt violet to quantify the asci. An additional group of 12 mice was selectively depleted of their CD4+ lymphocytes by antibody treatment with 300 μg of GK 1.5 antibody (Biovest International, Minneapolis, MN) administered intraperitoneally three times on days 1, 3, and 7. After this initial treatment, the mice were infected by exposure to *P. murina*-infected mice. Mice were then treated with 100 μg of GK 1.5 antibody intraperitoneally once a week for 6 weeks. After this, mice were treated with 1.25 or 12.5 mg/kg/day of D0 for 14 days while continuing the GK 1.5 treatment. Control mice received vehicle.

**Animal studies for candidiasis.** For survival studies, 8-week-old CBA/J female mice (Jackson Laboratory) were used. Eight mice per treatment or control group were evaluated. Mice were infected by intravenous inoculation of 100 μl containing 10^7 cells of *C. albicans* strain SC–5314. Treated mice received an intraperitoneal injection of 1.2 mg/kg/day of either BHBM or D0 in a final volume of 100 μl of PBS containing 0.4% DMSO immediately after receiving the *Candida* inoculum. Untreated mice received 100 μl of PBS–0.4% DMSO. Mice were fed *ad libitum* and monitored closely for signs of discomfort. At the end of the survival study, tissue burden culture was performed in mice that survived the infection. Mice were sacrificed, and their organs were extracted and homogenized in 10 ml sterile PBS using a homogenizer. Organ homogenates were diluted 10 times in PBS, and 100-μl portions were plated on YPD agar plates and incubated at 30°C for 72 h for CFU count.

**In vivo toxicity.** Mouse toxicity studies were performed using 4-week-old CBA/J female mice from Jackson Laboratory. Five mice received 1.2 mg/kg/day BHBM for 60 days. Three control mice received a solvent injection per day. At day 60, blood was collected in two tubes: one with K2EDTA and the other without K2EDTA to allow blood clotting. The blood clot was then centrifuged at 1,500 rpm for 10 min, and the serum was collected and analyzed for liver and kidney blood tests. The noncoagulated blood was used for hematocrit and blood cell analysis. These tests were done using MASCOt HEMAVET 950FS (Drew Scientific Group, Düsseldorf, Germany).

**Pharmacokinetics of BHBM.** BHBM was dissolved in a mixture of cremophore-ethanol (1:1) to prepare a 10 mg/ml stock solution. The stock solution was diluted in PBS to obtain 200 μg/ml and 400 μg/ml solutions for i.v. and i.p. administrations in C3H/HeN mice (n = 3). BHBM was administered to control (healthy) mice or immunocompromised mice infected with *P. murina* at doses of 0.8 mg/kg and 1.6 mg/kg via tail vein injection or intraperitoneal injection in a final volume of 100 μl. The mice were sacrificed, and blood samples were collected pre-dose and 0.5, 1, 2, 4, 8, 12, and 24 h after administration into K2EDTA-containing tubes. The samples were centrifuged immediately, and plasma was collected and stored at −80°C until analysis. Plasma samples were extracted using methylene chloride. Briefly, 50 μl of the plasma sample was taken into a glass vial, and 10 μl of internal standard N’-(3-bromobenzylidene)-4-hydroxybenzoylhydrazide was added. After the contents of the glass vials were mixed, 1 ml of methylene chloride was added to the vial, and the samples were vortex mixed for 30 s followed by centrifugation for 5 min. Then, 800 μl of supernatant was transferred to another tube and evaporated to dryness using a centrifugal evaporator. The residue was reconstituted in 100 μl acetonitrile-water (50:50) solution, mixed, and transferred to mass spectrometry vials. Separation was performed under isocratic reverse-phase chromatographic conditions using a Waters XBridge C18 column (3.5 μm; 2.1 × 100 mm) (Waters, Milford, MA), a Finnigan Surveyor MS pump (Thermo Fisher Scientific), and a Finnigan Micro AS autosampler (Thermo Fisher Scientific). The mobile phase consisted of water-acetonitrile with 0.1% formic acid (30:70) run at a flow rate of 200 μl/min. Then, 5 μl aliquots were analyzed using LTQ-FT liquid chromatography/tandem mass spectrometer (LC/MS/MS) with electrospray source in the positive ion mode (Thermo Fisher Scientific). The retention time of BHBM was 5.7 min. The lower limit of quantification (LLOQ) was 10 ng/ml. Systemic exposure of BHBM in mice was quantified by calculating the area under the concentration-time curve.
were grown in YNB (pH 7.4) at 37°C with 5% CO2 and treated for 6 h with the amount of time. For HPF, BHBM-treated and control cells were centrifuged at 3,000 rpm (1,700 x g) for 10 min, washed with PBS, and stained cell suspensions were mounted over glass slides as described above and analyzed using an Axioskop 2 microscope (Zeiss, Germany). Fluorescence intensity of yeasts labeled with NBD C6-ceramide staining was determined by analyzing dose response over the course of 16 h of treatment. Values were expressed as means ± standard error of the mean (SEM) and analyzed by using Student's t test.

**Lipid mass spectrometry.** For lipid analysis by mass spectrometry, fungal cells (C. neoformans H99 or C. albicans SC-5314) were grown in YNB and incubated with BHBM or D0 as explained above for in vivo labeling (except that tritiated palmitate was not added) for 6 h. Samples without drug were included as a control. Before lipid extraction, lipid internal standards (C17 ceramide and C17 sphingosine) were added. Lipids were then extracted, and one-fourth of the sample was aliquoted for determination of the inorganic phosphate. The remainder of the sample was subjected to base hydrolysis and then analyzed using LC-MS as previously described (6). Results were normalized using the inorganic phosphate levels.

**NBD C6-ceramide staining.** The Golgi apparatus of C. neoformans H99 and C. albicans SC-5314 was stained with NBD C6-ceramide (Molecular Probes, Eugene, OR) in a final concentration of 1 μM. Cells were incubated with NBD C6-ceramide in PBS for 1 h at 37°C. The cells were examined in a Zeiss Axioskop microscope equipped with a Megaview III camera operating at 80 kV. For analysis of vesicle volumetric density, images of at least 30 cells from control and treated groups were acquired and measured. The total area of the cell and the area of intracellular vesicles were measured, and the percentage of cell volume occupied by these vesicles was estimated according to the Delesse principle. Statistical significance was determined by Student's t test. P values of <0.05 were considered statistically significant.

**HIP-HOP library screening.** The yeast deletion collection is comprised of approximately 5,900 individually bar-coded heterozygous diploid strains (HIP [haploinsufficiency profiling]) and ~4,800 homozygous diploid strains (HOP [homozygous deletion profiling]) (58). Pools of approximately equal strain abundance were generated by robotically pooling (S and P Robotics, Ontario, Canada) each strain (from frozen stocks) onto YPD agar plates as arrays of 384 strains/plate. After 2 days of growth at 30°C, colonies were collected from plates by flooding with YPD, and cells were adjusted to an optical density at 600 nm (OD600) of 2. The fitness of each strain in each experimental pool was assessed as described previously (58). The BHBM dose that resulted in 15% growth inhibition in S. cerevisiae BY4743 (the parent strain of the yeast deletion collection) was determined by analyzing dose response over the course of 16 h of growth at 30°C. Screens of the homozygous deletion collection were performed for 5 generations of growth in BHBM, and screens of the heterozygous deletion collection were collected after 20 generations of growth. Cells were processed as described previously (59). Genomic DNA was extracted from each sample and subjected to PCR to amplify the unique bar code identifiers. The abundance of each bar code was determined by quantifying the microarray signal as previously described. A ranked list of all genes in the genome was generated for each experiment and then compared using gene set enrichment analysis (GSEA).

**Generation of BHBM-resistant strains.** For the generation of BHBM-resistant strains, the drug-sensitive S. cerevisiae RYO0622 haploid strain was used (23). Prescreening studies were performed to determine the IC50 dose of BHBM for this strain (the 50% inhibitory concentration [IC50] at which 100% yeast cell growth is inhibited upon drug exposure). For this screening, 20 μl of RYO0622 cells (at an OD600 of ~10−4) were plated on solid synthetic complete (SC) medium alone or with DMSO or with various BHBM concentrations (67, 133, 266, 533, and 1,066 μg/ml) in a 48-well plate. The plates were incubated for 2 days at 30°C in the dark. These studies revealed an IC50 dose of 133 μg/ml of BHBM.

Screening for the BHBM-resistant mutants was performed by growing the RYO0622 cells to mid-log phase (OD600 of ~0.5) in liquid SC medium before adjusting the cell density to 1 x 106 cells/ml (equivalent to an OD600 of ~0.1). One milliliter of cells was plated on solid SC medium containing DMSO solvent control (0.26% [vol/vol]) or BHBM
(133 μg/ml IC_{50}, dose) and incubated at 30°C in the dark. A lawn of cells grew on the solvent control, while seven BHBM-resistant colonies were identified after 9 days. Longer incubation did not result in the appearance of further resistant colonies. To confirm BHBM resistance, single colonies isolated from the BHBM-containing SC medium were plated onto fresh solid SC medium containing 133 μg/ml BHBM and incubated for 2 days at 30°C in the dark. Robust BHBM-resistant cells were seen.

Next-generation sequencing of BHBM-resistant strains. Genomic DNA was extracted from RY00622 and BHBM-resistant cells using a standard yeast DNA extraction protocol (60). Genomic DNA samples were quantified using Qubit fluorometry (Life Technologies) and diluted for sequencing library preparation using a Nextera XT library preparation kit according to the manufacturer’s instructions (Illumina, San Diego, CA). For the initial round of sequencing, individual sequencing libraries were prepared for the parent and a single BHBM-resistant clone. These libraries were pooled and sequenced on a single MiSeq lane (Illumina), generating paired-end 150-bp reads. Further BHBM-resistant colonies were obtained in a second screen, and their DNAs were pooled at equal concentrations before preparation of a single sequencing library for the pool. This pool was sequenced alongside a new library for the parent strain on a single HiSeq 2500 lane (Illumina), generating paired-end 100-bp reads.

Mapping and variant calling. Raw FASTQ paired-end reads for the parent (RY00622) and the BHBM-resistant pool were independently aligned to the NCBI sacCer3 reference genome (genbank/genomes/eukaryotes/fungi/saccharomyces_cerevisiae/saccer_3r12011 at http://hgdownload.test.cse.ucsc.edu/goldenPath/sacCer3/bigZips/) using bwa mem v0.7.4-r385 (61) with the -M flag to mark shorter split hits as secondary for compatibility with Picard. Resultant SAM files were converted to BAM format using samtools v1.1 and sorted by coordinate using Picard v1.96 (SortSam) (http://picard.sourceforge.net). PCR duplicate reads were filtered out using Picard MarkDuplicates and indexed using Picard BuildBamIndex. To call single nucleotide variants (SNVs), the GATK Unified Genotyper v2.1-8 (62) was run with the NCBI sacCer3 reference genome, stand_call_conf=30, and stand_emit_conf=10 (63). The ploidy parameter was set at 1, since the parent and resistant strains are in haploid state. Realignment around known indels and quality score recalibration was not performed, since a database of known indels and known single nucleotide polymorphisms (SNPs) is not available.

Validation of BHBM-resistant yeast mutants. Four yeast genes (ALPS, COS111, MKKI, and STE2) were selected based on the high-quality variant calls present in the BHBM-resistant pool. To confirm BHBM resistance, the individual haploid Δnull, Δcos111, Δmkki1, and Δste2 deletion mutants were assayed for growth fitness after treatment with BHBM. Unrelated drug controls, including methyl methane sulfonate (MMS) (cytotoxic) and fluconazole (antifungal) were assayed in parallel. Strains were cultured to mid-log phase (OD_{600} of ~0.5) in liquid YPD medium before adjusting the cell density to an OD_{600} of 0.0625 with YPD medium. The cells were transferred to 96-well plates containing 100 μl of YPD with DMSO solvent control (2% [vol/vol]), BHBM (6 to 733 μg/ml), MMS (10 μg/ml to 625 μg/ml), or fluconazole (2 to 306 μg/ml) and incubated at 30°C for 24 h. The fitness of individual strains was calculated by the average rate after normalizing the growth inhibition was calculated by the average rate after normalizing the growth inhibition.

SUPPLEMENTAL MATERIAL

We thank Rao Movva for help with the HIP-HOP experiments. We thank the staff at the Transmission Electron Microscopy Facility in the Central Microscopy Imaging Center (C-MIC) at Stony Brook University, Stony Brook, NY, for their contributions to TEM preparation and data collection.

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REFERENCES


New Antifungals against Fungal Sphingolipids

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Erratum for Mor et al., “Identification of a New Class of Antifungals Targeting the Synthesis of Fungal Sphingolipids”


The compound that was used in our article was \( N'-(3\text{-bromo-6-hydroxybenzylidene})\)-2-methylbenzohydrazide (BHBM) and not \( N'-(3\text{-bromo-4-hydroxybenzylidene})\)-2-methylbenzohydrazide. All experiments described in the paper were done using BHBM \( [N'-(3\text{-bromo-6-hydroxybenzylidene})\]-2-methylbenzohydrazide\), and the structure noted in Fig. 1B of our original article should be as indicated in Fig. 1 here.

The structure of BHBM used in the mBio studies was confirmed by proton nuclear magnetic resonance (NMR), which indicated the following chemical shifts: \( ^1H\) NMR (500 MHz, dimethyl sulfoxide-d6) \( \delta \) 2.38 (s, 3H), 6.90 (d, 1H, \( J = 8.8 \) Hz), 7.28 to 7.32 (m, 3H), 7.37 to 7.43 (m, 2H), 7.47 (d, 1H, \( J = 7.5 \) Hz), 7.78 (s, 1H), 8.47 (s, 1H), 11.19 (s, 1H), 12.05 (s, 1H).
All findings illustrated in the paper are correct.