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Beauvericin counteracted multi-drug resistant Candida albicans by blocking ABC transporters

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A B S T R A C T

Multi-drug resistance of pathogenic microorganisms is becoming a serious threat, particularly to immunocompromised populations. The high mortality of systemic fungal infections necessitates novel antifungal drugs and therapies. Unfortunately, with traditional drug discovery approaches, only echinocandins was approved by FDA as a new class of antifungals in the past two decades. Drug efflux is one of the major contributors to multi-drug resistance, the modulator of drug efflux pumps is considered as one of the keys to conquer multi-drug resistance. In this study, we combined structure-based virtual screening and whole-cell based mechanism study, identified a natural product, beauvericin (BEA) as a drug efflux pump modulator, which can reverse the multi-drug resistant phenotype of Candida albicans by specifically blocking the ATP-binding cassette (ABC) transporters; meantime, BEA alone has fungicidal activity in vitro by elevating intracellular calcium and reactive oxygen species (ROS). It was further demonstrated by histopathological study that BEA synergizes with a sub-therapeutic dose of ketoconazole (KTC) and could cure the murine model of disseminated candidiasis. Toxicity evaluation of BEA, including acute toxicity test, Ames test, and hERG (human ether-à-go-go-related gene) test promised that BEA can be harnessed for treatment of candidiasis, especially the candidiasis caused by ABC over-expressed multi-drug resistant C. albicans.

1. Introduction

The opportunistic fungal pathogen Candida albicans is one of the major causes of systemic severe infections with a high mortality. It represents an emerging global health threat, especially among the immunocompromised populations. Unfortunately, only a limited number of effective classes of antifungals are available in the...
2. Materials and methods

2.1. Experimental subjects, chemicals, primers, and culture media

Cancer cell lines (A549, H1299, H157, H460, H358, H522, H1650, H1792, H226, Calu-1), C. parapsilosis (ATCC22019), and C. albicans SC5314 were maintained in our laboratory. Clinical isolates of C. albicans, C. krusei, and C. tropicalis were provided by a local hospital. All strains were maintained in 25% (v/v) glycerol at −80 °C. Specific pathogen-free ICR mice (white, 18−22 g, half females), were obtained from B & K Universal Group Limited, Shanghai, China.

RPMI 1640 (Invitrogen) was used according to the manufacturer’s protocol. YEPD liquid medium consisted of: yeast extract 1% (w/v), peptone 2% (w/v), dextrose 2% (w/v), 2% (w/v) agar will be added for plates.

The complete list of microbes used in this study is provided in Table S1. All primers used in this study are listed in Table S2. All chemicals were used in this study according to the manufacturer’s directions.

2.2. Virtual screening

The input files of receptor were prepared by AutoDock/Vina plugin of PyMOL, the sdf formatted ligands were downloaded from ZINC15 3D Tranches, under the parameters of “React.: Standard”, “Purch.: Wait OK”, “pH: Ref Mid”, and “Charge: −2 −1 0 +1 +2”. All tranches were imported into PyRx by Open Babel, then the molecules were processed by energy minimization and then all of them were converted to AutoDock Ligands. The virtual screening was carried out by Vina Wizard, the grid boxes were manually adjusted to center_x=21.55, center_y=65.64, center_z=13.71, size_x=62.13, size_y=50.11, size_z=52.25 for Cdr1; while center_x=21.06, center_y=65.66, center_z=14.23, size_x=62.51, size_y=50.60, size_z=52.23 for Cdr2. The exhaustiveness for both Cdr1 and Cdr2 is 8 (up to 9 poses).

2.3. Antifungal susceptibility and synergistic antifungal testing

Antifungal and synergistic antifungal tests were carried out as described previously, using a broth microdilution protocol modified from the Clinical and Laboratory Standards Institute M-27A3 methods. All minimal inhibitory concentration (MIC) tests were undertaken in triplicate. MICs were determined as the concentration of drugs that inhibited microbial growth by 90% relative to the corresponding drug-free growth control.

2.4. Rhodamine 6G efflux assay

Fungal strains were cultivated in YEPD liquid medium at 200 rpm (30 °C) for 16 hours. The harvested cells were washed twice with ice-cold glucose-free PBS. Cells were suspended in ice-cold glucose-free phosphate buffered saline (PBS) and incubated at 200 rpm (30 °C) for 4 hours under starvation conditions to reduce ABC pump activity. Cells were then washed twice and diluted to 10^6 cells/ml in ice-cold glucose-free PBS, as determined with a hemocytometer. BEA at a concentration of 16 μg/ml (about 2 × MIC) was added when necessary, while PBS was added as the negative control. All samples were incubated for another 2 hours at 200 rpm (30 °C). Then, 10 μM (final concentration) rhodamine 6G was added, and cells were incubated for further 1.5 hours at 200 rpm (30 °C). The external rhodamine 6G was then removed by washing with glucose-free PBS and glucose was added to the samples (to a final concentration of 3 mM) to reactivate the ABC efflux pumps, with PBS as the negative control. Cell samples (1 ml) were taken at designated time points, centrifuged and 100 μl of each supernatant was transferred into black 96-well microtiter plate with clear bottoms (Greiner, Germany). Rhodamine 6G fluorescence was measured with a Multilabel Plate Reader (Perkin Elmer, USA) at 510 nm excitation/535 nm emission wavelengths. Experiments were carried out at least in triplicate.
2.5. Intracellular calcium and ROS measurement

*C. albicans* SC5314 cells were cultivated in YEPD liquid medium at 200 rpm (30 °C) for about 16 hours. The harvested cells were washed twice with ice-cold PBS and diluted to 5 × 10^5 cells/ml with ice-cold PBS, as determined with a hemocytometer and confirmed by cfu counting. For calcium measurement, designated concentrations of each drug were added to 1 ml samples, and co-incubated at 200 rpm (30 °C) for 4 hours to get dose-dependent data. For the time-dependent assay, 64 μg/ml of each drug was added to 1 ml samples, and incubation was stopped at specific time points. For ROS detection, after co-incubation with each drug at 200 rpm (30 °C) for 4 hours, DCFH-DA (2',7’-dichlorofluorescein diacetate) was added to a final concentration of 10 μM and cells were incubated for another 30 min at 200 rpm (30 °C). The cells were then washed to remove external drugs, resuspended in Cal NW working solution (according to DiscoverX HitHunter™ Calcium No WashPLUS Assay Kit protocol) and incubated at 37 °C, 200 rpm for 2 hours. Then 100 μl of each sample was transferred into black 96-well microtiter plate with clear bottoms (Greiner, Germany). The plate was equilibrated at room temperature for 6 hours, but this step was not necessary for ROS detection. The fluorescence was measured with a Multilabel Plate Reader (Perkin Elmer, USA) at 485 nm excitation/510 nm emission wavelengths for calcium measurement, and 485 nm excitation/525 nm emission wavelengths for ROS detection. Experiments were carried out at least in triplicate.

2.6. Mouse model and ethics statement

Animal related experiments were carried out in Guangdong Laboratory Animal Monitoring Institute, which has earned AALAC accreditation (001469).

Specific-pathogen-free female ICR (Crl: CD-1) mice (female, white, 20–22 g) were used in this study. Experiments were carried out as described previously.14,20 Animal use protocols were reviewed and approved by IACUC of Guangdong Laboratory Animal Monitoring Institute in accordance with the Guide for the Care and Use of Laboratory Animals. Animals were bred in negative pressure isolation cages in an animal negative pressure facility with an isolators at 24 ± 2 °C on mice that had been starved for the previous 12 hours. BEA was dissolved in 0.5% sodium carboxymethyl cellulose at the concentration of 0.05 g/ml. Oral administration of 0.4 ml/10g.w BEA (equivalent the dose of 2 g/kg) was performed by gavage. For 14 days, the mice were weighed thrice daily and observed for signs of drug-related morbidity or mortality. After the observation period, animals were killed by CO2 exposure followed by cervical dislocation.

2.10. Ames test and hERG test

The Ames test and the hERG test were done by Shanghai InnoStar Bio-Tech Co., Ltd.

2.11. BEA-resistant mutants screening

To select BEA-resistant mutants, we used a rapid selection regime in which SC5314 cells were plated onto YEPD containing a high concentration of BEA (512 μg/ml). Only colonies of the largest size (>1.6 mm²) that had acquired robust, reproducible resistance with MIC increasing more than 10 fold were chosen.

3. Results

3.1. Homology modeling of C. albicans ABC transporters and virtual screening for inhibitors

The homology models of the *C. albicans* ABC transporters, Cdr1 and Cdr2 were built using Alignment Mode algorithm of Swiss-Model server,21–23 taking a known mouse P-glycoprotein crystal structure (PDB code: 3G60) as the template, which shares 29% sequence identity and 50% similarity to Cdr1 and 31% sequence identity and 51% similarity to Cdr2, respectively. As co-crystallization showed QZ59-RRR (cyclic-tris-(R)-valinolezalene, Mol. wt. 687.42 Daltons) and QZ59-RRS (cyclic-tris-(S)-valinolezalene, Mol. wt. 687.42 Daltons) can be poly-specifically bond to the P-gp internal cavity to block its efflux function.14 Based on the molecular weight of QZ59, we decided to screen the 3D ZINC15 database13 with >500 Daltons virtually. Meantime, QZ59-RRR and QZ59-RRS were added into our customized database manually as positive control. The virtual screening was carried out using the Vina Wizard of PyRx.17 No surprise that QZ59-RRR docked nicely to both Cdr1 and Cdr2 in a reasonable cavity position with the binding affinity of −9.0 kcal/mol and −8.3 kcal/mol, respectively; while the
binding affinity for QZ59-SSS is $8.3$ kcal/mol and $8.5$ kcal/mol, respectively. In total, 912 compounds with more than 500 Daltons that have available 3D information from ZINC15 database were used for virtual screening. Each compound was screened for up to 9 poses for each receptor, the binding affinity of those around 16200 poses varied from $13.1$ kcal/mol to $5.1$ kcal/mol for Cdr1, and from $13.8$ kcal/mol to $5.2$ kcal/mol for Cdr2 (Table S3). By manually checking the reasonable docking poses with the UCSF Chimera, we interestingly found that 8 BEA compounds have even better binding affinities to the internal cavity of both Cdr1 and Cdr2 than QZ59 compounds do (Table 1). BEA was previously reported to have highly effective synergistic antifungal activity against diverse fungal pathogens, but its precise molecular mechanism of synergy is yet unclear.

The virtual screening results here indicated that BEA can block the internal cavity of ABC transporter, and serve as potential inhibitors. We used PyMOL together with UCSF Chimera to display the docking models of BEA with Cdr1 and Cdr2, respectively (Fig. 1).

### Table 1

| ZINC Id     | Chemical name | Binding affinity kcal/mol | rmsd/ub | rmsd/lb |
|-------------|---------------|---------------------------|---------|---------|
|             |               | Cdr1 | Cdr2 | Cdr1 | Cdr2 | Cdr1 | Cdr2 |
| ZINC85643633 | Beauvericin   | $-9.8$ | $-8.4$ | 0.000 | 3.036 | 0.000 | 5.568 |
| ZINC95540658 | Beauvericin A | $-10.3$ | $-10.1$ | 0.000 | 0.000 | 0.000 | 0.000 |
| ZINC95607714 | Beauvericin H1| $-9.1$ | $-9.5$ | 5.899 | 0.000 | 9.572 | 0.000 |
| ZINC95542396 | Beauvericin H2| $-9.6$ | $-10.7$ | 16.142 | 0.000 | 20.455 | 0.000 |
| ZINC95613104 | Beauvericin H3| $-10.5$ | $-10.4$ | 0.000 | 0.000 | 0.000 | 0.000 |
| ZINC95607711 | Beauvericin G1| $-11.0$ | $-8.8$ | 0.000 | 21.689 | 0.000 | 25.254 |
| ZINC95613105 | Beauvericin G2| $-10.3$ | $-9.6$ | 0.000 | 0.000 | 0.000 | 0.000 |
| ZINC28974061 | Beauvericin G3| $-10.3$ | $-9.0$ | 0.000 | 10.572 | 0.000 | 14.905 |
| QZ59-RRR    |               | $-9.0$ | $-8.3$ | 21.093 | 17.523 | 23.909 | 20.773 |
| QZ59-SSS    |               | $-8.8$ | $-8.5$ | 14.237 | 24.328 | 19.356 | 28.751 |

#### 3.2. BEA indeed blocks the efflux function of *C. albicans* ABC transporters

In order to validate the virtual screening results, we used rhodamine 6G, a specific substrate of Cdr1 and Cdr2, as an indicator to study the efflux function of *C. albicans* ABC transporters. All testing *C. albicans* strains were starved for 4 hours in glucose-free PBS, and then 3 mM (final concentration) glucose was added to reactivate the ABC transporters when specified. When SC5314 was treated with 16 mg/ml of BEA, twice the MIC, the efflux of rhodamine 6G by ABC transporters was totally inhibited, even after 3 mM glucose was added (Fig. 2a, 2b). To confirm this observation, *C. albicans* strains with the following genes were knocked-out: CDR1 (strain 448), CDR2 (strain 653) and both CDR1 and CDR2 (strain 654) were subjected to the assay. Strain 653 behaved just like SC5314 did, while the efflux of rhodamine 6G in strain 448 was also sharply inhibited, but not as much as in strain 653, because the efflux function of Cdr1 is stronger than that of Cdr2 as reported in Ref. 28. Strain 654 did not show the inhibition since CDR1 and CDR2...
Fig. 2. BEA inhibits rhodamine 6G efflux in C. albicans CDR null mutants and in S. cerevisiae overexpressed CDRs. Error bars indicate standard deviation.
CDR2 were both knocked out (Fig. 2c–e).

To further investigate the effect of BEA on individual drug efflux pumps, *Saccharomyces cerevisiae* strains independently over-expressing *C. albicans CDR1* (strain CDR1), *C. albicans CDR2* (strain CDR2), and *C. albicans MDR1* (strain MDR1) were used to carry out the rhodamine 6G efflux assay. We found that BEA could significantly inhibit rhodamine 6G efflux in strain CDR1 and strain CDR2 (Fig. 2g, 2h). No difference of rhodamine 6G efflux was observed in strain AD1-8u- and strain MDR1, and both served as negative controls (rhodamine 6G is not a substrate of *C. albicans Mdr1*), with or without glucose (3 mM) (Fig. 2f, and Fig. S1f). Additionally, no inhibition of Nile Red efflux in strain MDR1 by BEA was observed, which is a substrate of *C. albicans Mdr1*.

Furthermore, we also observed that when BEA was added to the strains actively effluxing rhodamine 6G, the inhibition effect happened immediately (within 10 min) (Fig. S1). This result indicated that BEA acted on ABC transporters directly, rather than indirectly (through inhibition of gene transcription or/and translation). To confirm this observation, we first used quantitative real time PCR to verify the mRNA levels of CDR1 and CDR2 in SC5314 before and after BEA-treatment. No significant changes in gene transcription were observed. Moreover, we observed almost the same transcription level of ABC transporter genes in a laboratory acquired BEA-resistant strain BM-1 as that in its parental strain SC5314 (Fig. 3).

The results of rhodamine 6G efflux assay indicated that BEA can indeed specifically act on protein and inhibit the efflux function of *C. albicans* ABC transporters, but not MFS transporters. It is the mechanism of synergy we are looking for.

### 3.3. BEA has better synergistic antifungal effect with azoles against multi-drug resistant *C. albicans*

We hypothesized that as an efflux inhibitor, BEA might have better synergistic antifungal effect on those drug efflux pumps overexpressed multi-drug resistant strains. We tested the combination of BEA with KTC, and BEA with itraconazole (ICZ) against diverse Candida isolates, both drug sensitive and resistant (Table 2, Fig. 4). As expected, we found that BEA showed synergy with both azoles in a very low dosage on these yeast pathogens. Interestingly, the antifungal effect of BEA and azoles was indeed better in those drug efflux pumps overexpression strains than in drug sensitive strains, according to the fractional inhibitory concentration index (FICI) (Fig. 4).

### 3.4. BEA elevates intracellular calcium concentrations and triggers cell death in *C. albicans*

BEA alone showed some antifungal activity as well (Table 2). However, the inhibition of *C. albicans* ABC transporters is unlikely to be responsible for this antifungal activity, because these genes are not essential for the survival of *C. albicans*. Thus, there must be another target of BEA. Chemically, BEA is a cyclic hexadepsipeptide compound. Its ion-complexing capability might allow BEA to transport alkaline earth metal ions and alkaline metal ions across cell membranes. Previous studies had demonstrated that BEA is a potent calcium ionophore, can trigger apoptosis in lung cancer cells and human acute lymphoblastic leukemia. We want to see if there is similar effect in *C. albicans*. We compared the intracellular calcium concentrations of *C. albicans* SC5314 after treatment with BEA and A23187 (a known calcium ionophore), independently. Results showed that BEA could significantly (P < 0.001) increase the

### Table 2

Broad spectrum synergistic antifungal activity of BEA.

| Strain               | MIC (μg/ml) | FICI | MIC (μg/ml) | FICI |
|----------------------|-------------|------|-------------|------|
|                      | BEA[a]      | KTC[a] | KTC[c] | ICZ[a] | ICZ[c] |
| *Candida albicans* SC5314 | 8          | 0.008 | 0.002 | 0.5 | 0.032 | 0.004 | 0.375 |
| *Candida tropicalis*‡ | 16         | 1.6–3.2 | <0.064 | <0.145 | 0.032 | 0.008 | 0.5 |
| *Candida krusei*‡ | >32        | 0.8–1.6 | 0.025 | <0.28 | 0.5 | 0.064 | 0.378 |
| *Candida parapsilosis* ATCC14054 | 16        | 0.025 | <0.0064 | <0.5 | 0.25 | 0.004 | 0.266 |

- [a] Clinical isolates;
- [b] [a], alone, [c], combination with 1/4MIC of BEA;
- [c] FICI = (MICdrug A in combination/MICdrug A alone) + (MICdrug B in combination/MICdrug B alone); (FICI > 4: Antagonism; FICI < 0.5: Synergy; 0.5 < FICI < 4: Additive).
intracellular calcium concentration (Fig. 5a) in a clear dose- and time-dependent manner, but the different patterns between BEA and A23187 (Fig. S2) suggested that BEA and A23187 might have different modes of action. In our previous work, and A23187 (Fig. S2) suggested that BEA and A23187 might have time-dependent manner, but the different patterns between BEA intracellular calcium concentration (Fig. 5a) in a clear dose- and time-dependent manner.35,36 After 4 hours of BEA treatment (64 μg/ml), RTA2 was found up-regulated by increased intracellular calcium in a calcineurin-dependent manner.35,36 After 4 hours of BEA treatment (64 μg/ml), RTA2 transcription level indeed increased about 5-fold (P < 0.001) (Fig. 5b), which confirmed our observation that BEA can elevate the concentration of intracellular calcium. While the transcription level of calcineurin pathway-related genes CNA, CNB, and CRZ1 did not change significantly after BEA treatment, while the sensitivities of BEA to the calcineurin pathway mutants did not change either (Fig. 5c), indicating that BEA might target the upstream of calcineurin pathway (Fig. 5c).

Intracellular calcium, a secondary messenger, is a multi-functional molecule, and under some conditions, it could trigger programmed cell death.31,32 It has been demonstrated that reactive oxygen species (ROS) is one of the typical hallmarks of early apoptosis in C. albicans.37,38 Thus, we studied the effect of BEA on ROS generation. Not surprising, intracellular ROS was significantly induced (P < 0.001) by BEA; hydrogen peroxide was used as a positive control (Fig. 5d).

3.5. In vivo synergistic antifungal effect of BEA

To extend the in vitro observations, we evaluated the potential synergistic activity of BEA with a low dosage of KTC in a cyclophosphamide pre-treated immunocompromised murine model.14 Placebo-treated mice, infected with either C. albicans SC5314 or C. parapsilosis ATCC 14054, had demonstrated significant hyper trophy, bleeding, and fibrinoid necrosis in kidney, heart, spleen, lung, liver and brain tissues, yeast cells, and/or pseudohyphal forms, were visible (Fig. S3).

High dosage (50 mg/kg body weight) of KTC alone increased mean survival time (from 8.2 ± 0.4 d to 19.4 ± 2.0 d) of the treated mice, and caused a change in the relative number of histological foci of infection, but did not reduce the fungal burden in tissues comparing to the placebo-treated mice. Histopathologically, abscesses of renal body, and necrosis of glomerulus membrane, appeared with an absence of the renal capsule. Three types of foci were noticed: (1) abundant yeast cells and mycelia occupied the parenchyma which had been dissolved; (2) inflammatory cells infiltrated with fragmentation of necrotic cells; and (3) the boundary between the fungus and the host contained inflammatory, necrotic, and intumescent cells. Similar phenomena were observed in other organs. In this murine model, even the high dosage of KTC alone had limited therapeutic efficacy because KTC is fungistatic and the mice were pre-treated with immunosuppres sant, cyclophosphamide.

As one of the target organs in experimental candidiasis, kidney contains the greatest number of pathogens and has the most severe lesions, including diffuse foci of infection, acute inflammation, and necrosis. The cortical and medullar regions displayed acute diffuse glomerulonephritis; nephrons and proximal tubules were destroyed or even dissolved. Interestingly, kidneys from mice treated with 0.5 mg/kg BEA and KTC showed significant reduction in tissue damage and inflammatory cell infiltration as compared to those of placebo-treated mice control. Occasionally, abscesses of the renal body and granulomas of the glomerulus membrane were observed as well as limited necrosis and exfoliation of cells. Few fungal cells were observed and they were associated with only a limited number of inflammatory cells (mostly neutrophils, lymphocytes, monocytes, and fibroblasts). The periphery of these foci of infection contained numerous keratinocytes and fibrotic scars that formed peculiar immunological rings (Fig. 6). Even though there was limited tissue damage evident by microscopic evaluation. 0.5 mg/kg BEA and KTC had a significant therapeutic effect in the infected mice. By day 12 of post-infection, calculated by [log10 cfu (cell/g)], fungal burden of kidney was reduced by about 85%.14

3.6. Toxicity evaluation of BEA

Thus, BEA was proofed to be a high potential synergistic antifungal agent, and only a HepG2 toxicity evaluation14 is not sufficient to address that BEA is safe to use in clinic. An Ames test (Salmonella typhimurium TA98 and TA100)29,40 was adopted to evaluate the mutagenic potential of BEA. No compound related bacterial reverse mutagenicity was observed in both metabolic activation (+S9) and inactivation conditions (−S9) (Fig. 7a). Besides Ames test, ten lung cancer lines with different backgrounds were used to test the cytotoxicity of BEA, all IC50 are more than 20 μg/ml, indicating BEA only had limited cytotoxicity (Fig. 7b), but highly selective for Candida spp. Furthermore, we used a healthy murine model to test the acute toxicity of BEA. All testing mice survived.
Fig. 5. BEA elevates intracellular calcium and triggers the apoptosis pathway of C. albicans. The experiment was performed in triplicate. Error bars indicate standard deviation.

Fig. 6. Therapeutic effect of BEA synergizes with a low dosage of KTC on the infected mouse kidney.
without a toxic shock syndrome and observable signs of drug-related morbidity or mortality in a 2-week-observation period after a single-dose of BEA (2 g/kg body weight) gavage. Their body weight even increased (Fig. 7c). The effective therapeutic concentration of BEA is 0.5 mg/kg body weight. Therefore, the therapeutic index is very high. More and more structurally and functionally unrelated drugs have been found that can block the hERG potassium channel, prolong cardiac action potentials and lead to long QT syndrome, causing cardiac disease. So, the hERG test now is widely used as a predictor of cardiac risk in vivo,41,42 for drug discovery. The hERG test of BEA showed that 0.3 µM and higher concentration of BEA could inhibit the hERG potassium channel in a dose dependent manner; the IC_{50} of BEA is about 2.55 µM (Fig. 7d), which is considered controllable cardiac risk. All the toxicity tests indicated that BEA has good pharmacological potential with minimal toxicity.

4. Discussion

Historically, infectious diseases were a huge threat to human health before antibiotics were discovered, but now, they are returning with a powerful weapon, multi-drug resistance, which is a worldwide threat.33,34 As eukaryotes, fungal pathogens share lots of features with human cells, which restrict the antifungals discovery. With prophylactic and excessive use of these limited antifungals, multi-drug resistance is rapidly emerging, fungal pathogens adopted intricate strategies to avoid the lethal effects of antifungals,4,5 and the most well-known and studied strategy is overexpression of ABC transporters, which is considered controllable cardiac risk. All the toxicity tests indicated that BEA has good pharmacological potential with minimal toxicity.

demonstrated that this strategy is doable. We virtually identified BEA as the potential inhibitor of C. albicans ABC transporters, which was further confirmed by real experiments. In fact, BEA can re-sensitize multi-drug resistance by blocking the drug efflux function of ABC transporters. This is the mechanism of synergistic antifungal activity of BEA. The more the drug efflux pumps, the better the synergistic antifungal activity of BEA. The synergistic antifungal activity of BEA and azoles was also evaluated in vivo.

After BEA + KTC treatment, in the kidney, one of the target organs of fungal infection, the amyloidosis and necrosis decreased, no broad hemorrhages were observed, and nephrons remained intact. Inflammatory cells were also noted, but to a lesser extent. Remarkably, rare fungal cells were besieged by the immunological rings formed by inflammatory cells, mostly neutrophils and lymphocytes. A few foci were infiltrated with monocytes and fibroblasts. The periphery of these foci was surrounded with keratinocytes and fibroblasts forming peculiar immunological rings. We assume that the synergistic treatment kills most of the pathogens, and the remaining pathogens probably will be taken care of by the recovering immune system, because the effect of cyclophosphamide can only last about 7 days after administration. On the other hand, whether BEA can enhance the immune response or not is under study.

It is interesting to note that enniatin, an analog of BEA, was found to inhibit S. cerevisiae Pdr5, a homolog of Cdr1, in vitro.45 We have determined which domains of Cdr1 and Cdr2 that enniatin binds to inhibit the function of C. albicans ABC in vitro.46 We also found that when CDR1, a homolog of KDR1 in Neurospora crassa, was knocked out, the synergistic effect of BEA and a low dosage of KTC was lost. These results indicate that BEA is an inhibitor of fungal ABC transporters. The same effect was observed in some cancer cells with P-glycoprotein, a Cdr1 and Cdr2 homolog.47 With all the knowledge we got, we believe that BEA can act as a special inhibitor of ABC transporters to reverse multi-drug resistance. This finding could also inspire us a new and effective strategy for cancer therapy. BEA alone is fungicidal. Previous studies demonstrated BEA as a potent calcium ionophore in some mammalian cells,33,34 In this study, BEA was also found to act as a calcium ionophore in C. albicans to elevate the intracellular calcium concentration. However, it is not clear whether the increased intracellular calcium was from the extracellular environment or from the intracellular calcium stores. The imbalance of intracellular calcium has been connected to apoptosis pathway. Mitochondria may act as buffers
of the intracellular calcium concentration, and regulate vital processes, including apoptosis.48 In our study, we found that one of the early apoptosis markers, ROS, was sharply induced after the intracellular calcium concentration increased. ROS has been considered to be responsible for the killing effects of antibiotics in some cases.39,50 However, the correlation between ROS and antibiotic-killing mechanisms has become controversial.31,52 In our case, BEA had multi–antifungal targets. ROS was partially responsible for its fungicidal activity. Antibiotics had largely contributed to human health, but during the wide usage, drug resistance has weakened the antibiotic effects, and made infections that were once easily curable very dangerous again, so we need to put more efforts to control those drug resistant pathogens, as antibiotics discovery is entering into the resistance era, and to find synergistic antibiotics is one way out of the looming antibiotic-resistance crisis.33 Another advantage of drug combination is it can take advantage of drug resistances to reverse them.33 BEA in this study is such a vivid example. A summary of the synergistic antifungal mechanisms of BEA is shown in Fig. S4. Based on our work, we believe that BEA has a great potential to be a new antifungal agent.

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Appendix. Supplementary material

Supplementary data to this article can be found online at doi: 10.1016/j.synbio.2016.10.001.

References

1 Brown GD, Denning DW, Levitz SM. Tackling human fungal infections. Science 2012;336:647.
2 Pfaller MA, Diekema DJ. Epidemiology of invasive candidiasis: a persistent public health problem. Clin Microbiol Rev 2007;20(1):131–63.
3 Cui J, Ren B, Tong Y, Dai H, Zhang L. Synergistic combinations of antifungals and anti-virulence agents to fight against Candida albicans. Virulence 2015;6:362–71.
4 Anderson JB. Evolution of antifungal-drug resistance: mechanisms and pathogen fitness. Nat Rev Microbiol 2005;3(7):457–56.
5 Cowen LE. The evolution of fungal drug resistance: modulating the trajectory from genotype to phenotype. Nat Rev Microbiol 2008;6(2):187–98.
6 Prasad R, Kapoor K. Multidrug resistance in yeast. J Rev Cytol 2005;242:215–48.
7 Prasad R, Gaur NA, Gaur M, Komath SS. Efflux pumps in drug resistance of Candida. Infect Disord Drug Targets 2006;6(2):69–83.
8 Sanglard D, Ischer F, Parkinsson T, Falconer D, Bille J. Candida albicans mutations in the ergosterol biosynthetic pathway and resistance to several antifungal agents. Antimicrob Agents Chemother 2003;47(8):2404–12.
9 Zhang L, Demann AL. Natural products: drug discovery and therapeutic medicine. Totowa (NJ): Humana Press; 2005.
10 Aller SG, Yu J, Ward A, Weng Y, Chittaboina S, Zhuo R, et al. Structure of P-glycoprotein reveals a molecular basis for poly-specific drug binding. Science 2009;323:1718–22.
11 Lionta E, Spyrou G, Vassiliadis DK, Couriia Z. Structure-based virtual screening for drug discovery: principles, applications and recent advances. Curr Top Med Chem 2014;14:1923–38.
12 Sterling T, Irwin JL. ZINC 15-ligand discovery for everyone. J Chem Inf Model 2015;55:3324–37.
13 Fukuda T, Ari A, Yamaguchi Y, Masuma R, Tomoda H, Obara S. New beavercins, potentiators of antifungal micafuconazole activity, produced by Beauveria sp FK-1366 – I. Taxonomy, fertilization, isolation and biological properties. J Antibiot 2004;57(2):110–6.
14 Zhang LX, Yan KZ, Zhang Y, Ruan J, Zheng CS, et al. High-throughput system screening identifying multiple microbial metabolite combination agents for the treatment of fungal infections. Proc Natl Acad Sci USA 2007;104(11):4606–11.
15 Mei L, Zhang L, Dai R. An inhibition study of beavercin on human and rat cytochrome P450 enzymes and its pharmacokinetics in rats. J Enzyme Inhib Med Chem 2009;24:753–62.
16 Trott O, Olson AJ. AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multithreading. J Comput Chem 2010;31:455–61.
17 Dallakyan S, Olson AJ. Small-molecule library screening by docking with PyRx. Methods Mol Biol 2015;1283:243–50.
18 Song F, Dai H, Tong Y, Ren B, Chen C, Sun N, et al. Trichodermaealexides A-D and 7-O-methylkoninginin D from the marine fungus Trichoderma koningii. J Nat Prod 2010;73:806–10.
19 Clinical and Laboratory Standards Institute. Reference method for broth dilution antifungal susceptibility testing of yeasts; Approved standard. 3rd ed, CLSI document M27-A3. Wayne, PA: National Committee for Clinical Laboratory Standards. 2008.
20 Wu Y, Min F, Pan J, Wang J, Yuan W, Zhang Y, et al. Systemic Candida parapsilosis infection model in immunosuppressed Kcr mice and assessing the antifungal efficiency of fluconazole. Vet Med Int 2015:2015. 370641.
21 Schwede T, Kopp J, Guex N, Peitsch MC. SWISS-MODEL: an automated protein homology-modelling server. Nucleic Acids Res 2003;31:3381–5.
22 Arnold K, Bordoli L, Kopp J, Schwede T. The SWISS-MODEL workspace: a web-based environment for protein structure homology modelling. Bioinformatics 2006;22:195–201.
23 Guex N, Peitsch MC. SWISS-MODEL: the Swiss-PdbViewer: an environment for comparative protein modeling. Electrophoresis 1997;18:2714–23.
24 Pettersen EF, Goddard TD, Huaye CC, Couch GS, Greenblatt DM, Meng EC, et al. UCSF chimera—a visualization system for exploratory research and analysis. J Comput Chem 2004;25:1505–12.
25 Schrodinger LLC. The PyMOL Molecular Graphics System. Version 1.5.0.4. 2010.
26 Sanglard D, Ischer F, Monod M, Bille J. Susceptibilities of Candida albicans multidrug transporter mutants to various antifungal agents and other metabolite inhibitors. Antimicrob Agents Chemother 1996;40:2300–5.
27 Sanglard D, Ischer F, Monod M, Bille J. Cloning of Candida albicans genes conferring resistance to azole antifungal agents: characterization of CDR2, a new multidrug ABC transporter gene. Microbiology 1997;143(3/4):205–16.
28 Holmes AR, Lin YH, Niimi K, Lamping E, Keniya M, Niimi M, et al. ABC transporter Cdr1p contributes more than Cdr2p does to fluconazole efflux in fluconazole-resistant Candida albicans clinical isolates. Antimicrob Agents Chemother 2008;52:3581–62.
29 Decottignies A, Grant AM, Nichols JW, de Wet H, McIntosh DB, Goffeau A, ATPase and multidrug transporter activities of the overexpressed yeast ABC protein Yor1p. J Biol Chem 1998;273:12612–22.
30 Lamping E, Monck BC, Niimi K, Holmes AR, Tsao S, Tanabe K, et al. Characterization of three classes of membrane proteins involved in fungal azole resistance by functional hypersusceptirion in Saccharomyces cerevisiae. Eukaryot Cell 2007;6:1150–65.
31 Ivantysine-Steele I, Holmes AR, Lamping E, Monck BC, Ridd KL, Isolation of Nile red as a fluorescent substrate of the Candida albicans ATP-binding cassette transporters Cdr1p and Cdr2p and the major facilitator superfAMILY transporter Mdr1p. Anal Biochem 2009;384:87–91.
32 Eliopoulos GMM. In: Lorian V, editor. Antibiories in laboratory medicine. Baltimore (MD): Williams & Wilkins; 1991. p. 432–52.
33 Chen BF, Tsai MC, Jow GM. Induction of calcium influx from extracellular fluid by beauvericin in human leukemia cells. Biochem Biophys Res Commun 2006;340:134–9.
34 Jow GM, Chou CJ, Chen BF. Beauvericin induces cytotoxic effects in human acute lymphoblastic leukemia cells through cytochrome c release, caspase 3 activation: the causative role of calcium. Cancer Lett 2004;216:165–73.
35 Jia XM, Ma ZP, Jia Y, Gao PH, Zhang JD, Wang Y, et al. RTA2, a novel gene involved in azole resistance in Candida albicans. Biochem Biophys Res Commun 2008;367:331–6.
36 Jia XM, Wang Y, Jia Y, Gao PH, Xu YG, Wang L, et al. RTA2 is involved in calpain-mediated azole resistance and sphingolipid long-chain base resistance in Candida albicans. Cell Mol Life Sci 2009;66:122–34.
37 Dai BD, Cao YY, Huang S, Xu YG, Gao PH, Wang Y, et al. Baicalein induces programmed cell death in Candida albicans. J Microbiol Biotechnol 2009;19:801–8.
39 Ames BN, McCann J, Yamasaki E. Methods for detecting carcinogens and mu-
tagens with the Salmonella/mammalian-microsome mutagenicity test. Mutat Res 1975;31:347–64.
40 Flamand N, Meunier J, Meunier P, Agapakis-Causse C. Mini mutagenicity test: a
miniaturized version of the Ames test used in a prescreening assay for point
mutagenesis assessment. Toxicol In Vitro 2001;15:105–14.
41 Bowlby MR, Peri R, Zhang H, Dunlop J. hERG (KCNH2 or Kv11.1) K+ channels:
screening for cardiac arrhythmia risk. Curr Drug Metab 2008;9:965–70.
42 Priest BT, Bell IM, Garcia ML. Role of hERG potassium channel assays in drug
development. Channels (Austin) 2008;2:87–93.
43 Cosgrove SE, Carmeli Y. The impact of antimicrobial resistance on health and
economic outcomes. Clin Infect Dis 2003;36:1433–7.
44 Smith PA, Ronesberg FE. Combating bacteria and drug resistance by inhibiting
mechanisms of persistence and adaptation. Nat Chem Biol 2007;3(9):549–56.
45 Hiraga K, Yamamoto S, Fukuda H, Hamanaka N, Oda K. Enniatin has a new
function as an inhibitor of Pdr5p, one of the ABC transporters in Saccharomyces
cerevisiae. Biochem Biophys Res Commun 2005;328:1119–25.
46 Tanabe K, Lamping E, Nagi M, Okawada A, Holmes AR, Miyazaki Y, et al. Chi-
ermas of Candida albicans Cdr1p and Cdr2p reveal features of pleiotropic drug
resistance transporter structure and function. Mol Microbiol 2011;82(2):416–33.
47 Sharom FJ, Lu P, Liu R, Yu X. Linear and cyclic peptides as substrates and
modulators of P-glycoprotein: peptide binding and effects on drug transport
and accumulation. Biochem J 1998;333(Pt 3):621–30.
48 Rizzuto R, De Stafani D, Raffaello A, Mammucari C. Mitochondria as sensors and
regulators of calcium signalling. Nat Rev Mol Cell Biol 2012;13:566–78.
49 Kohanski MA, Dwyer DJ, Hayete B, Lawrence CA, Collins JJ. A common mech-
anism of cellular death induced by bactericidal antibiotics. Cell 2007;130:
797–810.
50 Dwyer DJ, Kohanski MA, Collins JJ. Role of reactive oxygen species in antibiotic
action and resistance. Curr Opin Microbiol 2009;12:482–8.
51 Liu Y, Imlay JA. Cell death from antibiotics without the involvement of reactive
oxygen species. Science 2013;339:1210–3.
52 Keren I, Wu Y, Inocêncio J, Mukahi LR, Lewis K. Killing by bactericidal antibi-
otics does not depend on reactive oxygen species. Science 2013;339:1213–6.
53 Brown ED, Wright GD. Antibacterial drug discovery in the resistance era. Nature
2016;529:336–43.
54 Baym M, Stone LK, Kishony R. Multidrug evolutionary strategies to reverse
antibiotic resistance. Science 2016;351:aae3202.