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# Exploring medium and long arm extensions of 1,2,4-triazole derivatives as Candida albicans $14\alpha$ -demethylase (CYP51) inhibitors†

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Fungal infections have been described as a silent crisis affecting more than one billion people each year. At least 150 million of these cases involve severe and life threatening invasive fungal infections, accounting for approximately 1.7 million deaths annually. 1,2,4-Trizoles such as fluconazole and posaconazole are widely used antifungal agents, but azole resistance is an increasing problem requiring further study. 1,2,4-Triazole derivatives with medium and long arm extensions designed to bind within the *Candida albicans* CYP51 (CaCYP51) access channel were synthesised to study their inhibition of CaCYP51 (IC<sub>50</sub>, MIC) and binding affinity ( $K_d$ ). A long arm extension using the amide linker was found to be most effective (e.g. 13), giving an antifungal profile vs. wild-type and resistant model fungal strains comparable with posaconazole.

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#### Introduction

Candida albicans (C. albicans), the predominant cause of candidiasis, is a commensal that resides in different parts of the human body including the gastrointestinal tract, genitourinary tract, oropharynx, and the skin. Under certain conditions C. albicans can cause infections that range from superficial to invasive, life-threatening, infections such as candidemia. Worldwide C. albicans is one of the most common healthcare associated bloodstream infections in intensive care units with a mortality of 30–50% for candidemia.

According to the Centre for Disease Control and prevention (CDC), many patients with severe COVID-19

symptoms in 2020 developed invasive candidemia from *C. albicans* as well as resistance to antifungal treatment owing to immune suppression caused by the virus.<sup>4–6</sup> This emphasises the need for improved antifungal treatments as the current range of effective antifungal agents is limited and susceptible to drug resistance.

Ergosterol is a fungal-specific and major sterol component of the fungal cell membrane that affects its physiochemical characteristics including fluidity. A key cytochrome P450 (CYP) enzyme involved in the synthesis of ergosterol is sterol 14 $\alpha$ -demethylase (CYP51, Erg11). The azole antifungals are the largest class of drugs that act by inhibiting CYP51. They are type II inhibitors that bind directly with the haem iron. The incidence of drug resistance to the available azole antifungals in yeast and other fungi is increasing and is attributed to the long treatments and prophylactic use in the clinic, and widespread use of agricultural azole fungicides to protect crops.  $^{9,10}$ 

Fluconazole (FLZ, Fig. 1) is one of the most commonly used antifungal agents and significant target-based FLZ resistance has been found in clinical isolates of *C. albicans*. <sup>10-14</sup> FLZ resistance has been attributed to single and double mutations within CYP51, with common single amino acid mutations involving Y132F and K143R located within the haem active site, and a G450E mutation within the electron transfer area that interacts with the NADPH-cytochrome P450 reductase redox partner. Y132 forms a key H-bonding interaction with azole antifungals *via* the tertiary hydroxy group of triazoles such as FLZ and voriconazole (VCZ, Fig. 1). Double mutations such as *C. albicans* CYP51

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Fig. 1 Structure of clinically used antifungal agents.

(CaCYP51) Y132H/K143R and Y132F/F145L further reduce the binding affinity of FLZ, increase the IC<sub>50</sub> value, 10 and probably account for increased MIC values found in C. mutations.11-14 strains containing these Conformational changes induced by such mutations are also likely to affect the binding between the azole nitrogen and the haem iron as well as the haem propionates with Y118 and Y132, e.g. by affecting H-bonding interactions of haem and by distorting the ionic interaction with K143. 9,10,15-18

Azole antifungals, especially the triazoles, are often a firstchoice treatment due to their high oral bioavailability, low toxicity, selectivity, tolerability by patient and the broad spectrum of activity. There are some limitations for the use of these azoles. For example, while VCZ (Fig. 1) has a broad spectrum of activity against most Candida species and fungal pathogens compared with FLZ, it has more adverse effects and drug-drug interactions than FLZ.19 Itraconazole (ITZ, Fig. 1) and posaconazole (PCZ, Fig. 1) have broad spectrum activities against candidiasis, but their low bioavailability has limited their clinical use.20

Recent studies on azole antifungals21 have focussed on azole-hybrid structures e.g. FLZ-COX inhibitor hybrids, 22 to explore dual activity and tetrazole derivatives e.g. VT-1161 (oteseconazole),<sup>23</sup> to improve CYP selectivity, althoughVT-1161 is susceptible to azole resistance.24 The increased incidence of resistant clinical isolates highlights the importance of developing new antifungals with activity against such strains, with azoles the focus of the research presented here. Our goal was to develop mid-sized and extended arm azoles targeting the CaCYP51 enzyme by using FLZ as a pharmacophore and applying the promising results obtained in our previous published study<sup>17</sup> to enhance activity, in particular against resistant

fungal strains, and binding affinity. The following design was proposed: one triazole moiety of FLZ was retained as the haem-binding group and the 2,4-difluorobenzene ring was replaced with the more lipophilic 4-chlorobenzene and 2,4-dichlorobenzene to enhance uptake across the lipophilic fungal cell wall and still fit in the small hydrophobic binding site of CaCYP51. The second triazole ring was replaced with either a medium or long arm extension (Fig. 2). The medium extensions contain an amide linked to either acetophenone or a 2-chlorothiazole ring identified through computational modelling to increase binding interactions. The long arm extensions contain a benzamide attached to a second benzene ring through either a thiourea, urea, amide or sulfonamide linker. The rationale for adding thiourea, urea, amide or sulfonamide linkers was to add H-bond acceptor and/or donor capability, which might form additional binding interactions with amino acid residues in the CaCYP51 access channel binding site, to compensate for the loss of the key H-bonding interaction between the azole tertiary hydroxy group and common active site mutations, such as Y132H/F (C. albicans) or Y140H/F (S. cerevisiae models) in resistant strains.

#### Results and discussion

#### Chemistry

The mid-sized acetyl (6a-b) and thiazole (7a-b) derivatives were obtained by a 5-step synthesis (Scheme 1) starting with introduction of the triazole ring by reaction of acetophenone (1) with 1,2,4-triazole and activated K2CO3 to give the 1-(substituted phenyl)-2-(1H-1,2,4-triazol-1-yl)ethan-1-one (2) derivatives.<sup>25</sup> Corey-Chaykovsky epoxidation reaction of the ethanone derivative (2) using TMSOI and 20% aqueous NaOH with heating at 60 °C for 6 h (ref. 26) gave the crude epoxides (3), which were directly converted

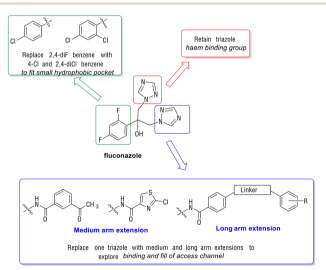


Fig. 2 Rationale for modifications to generate azole antifungals with medium and long arm extensions.

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Scheme 1 Reagents and conditions: (i) 1,2,4-triazole,  $K_2CO_3$ ,  $CH_3CN$ , 0–5 °C 30 min, r.t., 24 h or 80 °C, 4 h, 51–74% (ii) TMSOI, 2 M aq. NaOH, toluene, 60 °C, 6 h, quantitative crude yield (iii) NaN<sub>3</sub>, NH<sub>4</sub>Cl, DMF, 60 °C, 2 h, r.t., o/n, 65–66% (iv) (a) Ph<sub>3</sub>P, THF, r.t., 1 h (b) H<sub>2</sub>O, 60 °C, 4 h, 60–77% (v) CDI, DMF, r.t., o/n, 45–68% [a R<sup>1</sup> = Cl, R<sup>2</sup> = H; b R<sup>1</sup> = Cl, R<sup>2</sup> = Cl].

to the azides (4) on treatment with NaN3 and NH4Cl in good yields (4a 60% and 4b 63%) over two steps from 2. Azide reduction to the free amines (5) was performed using a Staudinger reaction,<sup>27</sup> with initial reaction of azides (4) with triphenylphosphine in THF at room temperature for 1 h followed by the addition of H<sub>2</sub>O and heating at 60 °C for 4 h. The mid-sized derivatives (6 and 7) were prepared by reaction of the amines (5) with either 3-acetylbenzoic acid 2-chlorothiazole-4-carboxylic using 1,1'carbonyldiimidazole (CDI) as the amide coupling reagent (Scheme 1). The mid-sized amides (6a-b and 7a-b) were obtained in yields ranging from 45-68% (Table 1) and structure and purity confirmed by <sup>1</sup>H/<sup>13</sup>C NMR and HPLC respectively.

Scheme 2 Reagents and conditions: (i) sat. aq. NaHCO<sub>3</sub>, CH<sub>2</sub>Cl<sub>2</sub>, r.t. o/n, 55-74% (ii) H<sub>2</sub>, Pd/C, EtOH, r.t., 3 h, 71-100% (iii) pyridine, r.t., o/n, 52-83% [a  $\mathbb{R}^1$  = Cl,  $\mathbb{R}^2$  = H; b  $\mathbb{R}^1$  = Cl,  $\mathbb{R}^2$  = Cl].

The nitro intermediates (9) for the preparation of the extended derivative, were prepared by reaction of 4-nitrobenzoyl chloride (8) with the free amines (5) at room temperature overnight, (Scheme 2). The nitro derivatives (9) were reduced to the amines (10) using 10% palladium on carbon in dry MeOH and H2 balloon for 3 h at room temperature. The synthesis of the final compounds (11-14) with the different linkers was achieved by adding 4-chlorophenyl derivatives to 4-amino-N-(2-(substituted phenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (12) in dry pyridine, and the reaction was left at room temperature overnight (Scheme 2). The extended compounds were obtained in good yields as white solids (Table 1), although lower yields were obtained for compounds 11b and 12b owing to more difficult purifications. Purity was determined either by elemental analysis or HPLC, while structure characterisation was performed by <sup>1</sup>H/<sup>13</sup>C NMR.

Table 1 Yields and melting points, <sup>1</sup>H and <sup>13</sup>C NMR of mid-sized (6 and 7) and linker (11 thiourea, 12 urea, 13 amide and 14 sulfonamide) of derivatives

$\mathrm{Cmpd}^a$	Yield (%)	MP (°C)	Amide/linker $^1$ H NMR $(\delta)^b$	Amide/linker $^{13}$ C NMR $(\delta)^b$
6a	52	Semisolid	NH 8.55	C=O 167.1
6b	45	160-162	NH 8.72	C=O 167.5
7 <b>a</b>	68	Semisolid	NH 8.10	C=O 160.1
7 <b>b</b>	68	Semisolid	NH 8.26	C=O 160.4
11a	68	146-148	2× NH 10.05, 10.00	C=S 180.0
11b	56	138-140	2× NH 10.06, 10.03	C=S 179.9
12a	63	226-228	2× NH 8.99, 8.90	C=O 152.63
12b	52	216-218	2× NH 9.06, 8.97	C=O 152.66
13a	80	222-224	NH 10.50	C=O 165.13
13b	76	190-192	NH 10.51	C=O 165.14
14a	77	236-238	NH 10.73	_
14b	83	228-230	NH 10.76	_

<sup>&</sup>lt;sup>a</sup> **a**  $R^1 = Cl$ ,  $R^2 = H$ ; **b**  $R^1 = Cl$ ,  $R^2 = Cl$ . <sup>b</sup> DMSO-d<sub>6</sub>.

#### In vitro biological assays

C. albicans susceptibility testing. Each novel compound was screened against C. albicans wild-type clinical isolate CA14 and C. albicans laboratory strain SC5314 using the standardised CLSI M27-S4 broth dilution method.<sup>28</sup> As shown in Table 2, the dichloro mid-sized compounds (6b and 7b, MIC < 0.03 μg mL<sup>-1</sup>) displayed promising antifungal activity against both strains compared with monochloro derivatives (6a and 7a) as well as with the control FLZ. This might be explained by the clog P data as the more lipophilic compounds should more readily penetrate the fungal membrane resulting in improved uptake or may relate to optimal fill of the small hydrophobic pocket leading to improved binding.

The dichloro with the thiourea, urea and sulfonamide linker of the extended derivatives (11b, 12b and 14b) were more effective at inhibiting C. albicans growth, for example: the dichloro thiourea 11b showed MIC 0.25 μg mL<sup>-1</sup> compared with the monochloro 11a MIC 4 µg mL<sup>-1</sup>. Even though the  $\log P$  of the amide inhibitors (13a,  $\log P$  3.53 and 13b, clog P 4.15) was more lipophilic than thiourea and sulfonamide (11a, clog P 3.13, 11b, clog P 3.73, 15a, clog P3.45 and 15b,  $\log P$  4.01) and less than urea (12a,  $\log P$  3.86 and 12b,  $\log P$  4.46), the mono and dichloro derivatives of the amide linker (13a and 13b) were the most effective over the three linkers as well as the standard FLZ with MIC < 0.03 μg mL<sup>-1</sup> (Table 2). This could be owing to the higher polarity of the thiourea and urea functional group compared with the amide, which may lead to a reduced uptake across the lipophilic fungal membrane, although other factors such as efflux and binding affinity may also play a role.

CaCYP51 IC<sub>50</sub>. Derivatives with a MIC  $<1 \mu g \text{ mL}^{-1}$  were chosen for IC<sub>50</sub> testing. In the mid-sized series (6 and 7) the dichloro derivatives of the acetyl 6b and the thiazole 7b showed inhibitory activity with IC50 values of 0.49 and 0.41 μM respectively compared with FLZ (IC<sub>50</sub> 0.31 μM) (Table 2). The extended derivatives of the amide linker (13) all showed

Table 2 MIC,  $IC_{50}$ ,  $K_d$  and clog P values for mid-sized and extended compounds

	MIC (μg mL <sup>-1</sup> )		$IC_{50}$		clog
Cmpd	CA14	SC5314	(μ <b>M</b> )	$K_{\rm d}$ (nM)	$P^a$
6a	0.25	0.25	1.30	$1445 \pm 266$	1.85
6b	< 0.03	0.06	0.49	$229 \pm 45$	2.46
7a	0.25	0.25	0.77	$635 \pm 181$	1.95
7 <b>b</b>	< 0.03	< 0.03	0.41	$106 \pm 15$	2.55
11a	4	4	_	_	3.13
11b	0.25	0.5	0.67	$125\pm41.0$	3.73
12a	1	2	_	_	3.86
12b	0.125	0.125	0.47	$137 \pm 14$	4.46
13a	< 0.03	< 0.03	0.56	$46 \pm 8$	3.54
13b	< 0.03	< 0.03	0.48	$166 \pm 39$	4.15
14a	1	1	_	_	3.45
14b	0.25	0.25	0.78	$94 \pm 24$	4.01
FLZ	0.125	0.125	0.31	$31 \pm 8$	0.86

<sup>&</sup>lt;sup>a</sup> clog P by Crippen's fragmentation.<sup>29</sup>

very good inhibition of CaCYP51 (IC<sub>50</sub> 0.56 μM for 13a and 0.48 µM for 13b) (Table 2). Of all derivatives, compounds 6b, 7b, 12b, 13a and 13b had MIC and IC<sub>50</sub> values comparable with or better than FLZ indicating that these compounds are promising candidates for further study.

Binding affinity  $(K_d)$ . The novel azole compounds (6, 7, 1)11-14) were designed as type II binding inhibitors i.e. bind directly to the Fe<sup>3+</sup> of the CaCYP51 haem prosthetic group by replacing the sixth axial water molecule via coordination with a triazole nitrogen atom.<sup>30</sup>

The binding affinity  $(K_d)$  values for FLZ and the novel inhibitors were determined using saturation curves for CaCYP51 that measured the change in the absorbance ( $\Delta A$ peak-trough) plotted against antifungal concentration (Fig. 3, S1 and S2†). For the mid-sized series, specifically the dichloro derivatives, the binding affinity for the thiazole (7b) compound was acceptable ( $K_d = 106 \pm 15 \text{ nM}$ ) compared with fluconazole ( $K_d = 31 \pm 8 \text{ nM}$ ), which showed tight binding with the haem iron. The binding affinity for the chloro and dichloro of the acetyl derivatives (6a-b) as well as the chloro of the thiazole (7a) derivative indicated weaker binding affinities (Table 2). For the extended series, the monochloro of the amide derivative (13a) showed the tightest binding ( $K_d$ = 46  $\pm$  8 nM) with CaCYP51 haem Fe<sup>3+</sup> compared with the mid-sized series as well as FLZ. Good binding affinity for the dichloro derivative of the thiourea (11b), urea (12b) and sulfonamide (14b) was observed (11b,  $K_d = 125 \pm 41$  nM, 12b,  $K_{\rm d}$  = 137 ± 14 nM and **14b**,  $K_{\rm d}$  = 94 ± 24 nM) (Table 2).

S. cerevisiae target and resistant mutant models. The most promising thiazole derivative 7b, and the extended amide derivatives 13a and 13b were evaluated against a panel of recombinant S. cerevisiae strains24,31,32 expressing control levels of wild-type CYP51 (ScCYP51), overexpressed wild-type ScCYP51, overexpressed azole-resistant ScCYP51 Y140F/H mutations (S. cerevisiae Y140 equivalent to C. albicans Y132),

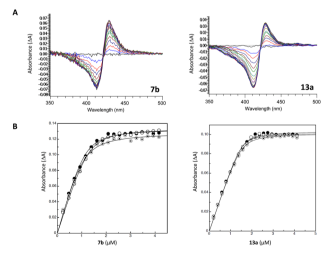


Fig. 3 (A) Absorbance difference spectra of exemplars 7b and 13a were measured during the titration of 2.5 μM native CaCYP51 with 7b and 13a (type II binding). (B) Saturation curve of 7b and 13a derived from type II difference spectra in (A) with 2.5 µM native CaCYP51. Azole titration was performed in triplicate.

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Compound	7 <b>b</b>	13a	13b	FLZ	PCZ
Strain			MIC (μg mL <sup>-1</sup> )		
S. cerevisiae mode	els				
Y2411 parent	0.125	0.25	< 0.03-0.06	4	< 0.03
Y2300 ScCYP51	0.5 - 1.0	1-2	0.25	16	0.06
Y2301 Y140F	4	1-2	0.25	32	0.06
Y2513 Y140H	2-4	1	0.125	32	< 0.03
C. albicans models					
Y525 CaMDR1a	2	0.125	< 0.03-0.06	32	< 0.03
Y570 CaCDR1B	>16	>16	>16	>128	8
H. sapiens model					
Y2760 HsCYP51	>16	>16	4-8	>128	8

Y2411 - azole sensitive control strain, Y2300 ScCYP51 overexpressed, Y2301 ScCYP51 Y140F mutant, Y2513 ScCYP51 Y140H mutant, Y525 MFS efflux pump CaMDR1a overexpressed, Y570 ABC efflux pump CaCDR1B overexpressed. FLZ fluconazole, PCZ posaconazole.

overexpressed C. albicans efflux pumps MDR1a or CDR1B and overexpressed H. sapiens wild-type CYP51 (HsCYP51) (Table 3) at 24 and 48 h with each assay performed in triplicate.

The most promising thiazole derivative 7b, and the extended amide derives 13a and 13b were all active against the susceptible strain Y2411 and S. cerevisiae CYP51 overexpressing strain Y2300. The extended amide derivatives (13a/b) retained good activity against the azole-resistant strains Y2301 and Y2513, expressing ScCYP51 Y140F and Y140H respectively and with 13b the most effective. Against the C. albicans efflux pump models, 13a and 13b retained activity against strain Y525, which overexpresses the C. albicans MFS transporter MDR1a. However inhibitory activity was significantly reduced against strain Y570, which overexpresses the C. albicans ABC transporter CDR1B, consistent with both FLZ and PCZ. Good selectivity was observed for ScCYP51 compared with HsCYP51 (Y2760 strain). All three derivatives were more effective than FLZ and 13b displayed a profile comparable with PCZ (Table 3) at both 24 and 48 h. Micafungin, a β-(1-3)-D-glucan synthase inhibitor, served as a negative control with MIC values of  $0.125 \mu g \text{ mL}^{-1}$  for all strains.

#### **Computational studies**

Protein-ligand complexes were prepared through docking of the R- and S-enantiomers of ligands 6a/b, 7a/b, 11a/b, 12a/b, 13a/b and 14a/b using Molecular Operating Environment (MOE) software<sup>33</sup> and the crystal structure of CaCYP51 cocrystallised with PCZ (pdb 5FSA).34 The protein-ligand complexes were subjected to 200 ns molecular dynamics simulations using the Desmond programme Schrödinger. 35,36 All complexes equilibrated, although fluctuations were observed in RMSD for some compounds due to changes in binding interactions over the 200 ns MD simulations (Fig. S3 and S4†). The ligands should be

positioned with the triazole above the haem Fe<sup>3+</sup> and perpendicular to the haem with a distance <3 Å for optimal binding. For nine of the ligands (6a, 6b, 7b, 11a, 12a, 12b, 13a, 13b and 14b) only the S-enantiomer was positioned optimally, for 11b only the R-enantiomer was optimally positioned and for 7a and 14a both enantiomers were optimally positioned for binding with the haem (Fig. S5†).

The ligands were positioned in contact with the haem, and within the small hydrophobic pocket and the access channel (Fig. 4). The small hydrophobic pocket accommodates the chlorobenzene (e.g. S-7b) or dichlorobenzene (e.g. S-13a) ring and the haem accommodates the triazole and amide linker (Fig. 4). The medium extended compounds (6 and 7) form  $\pi$ – $\pi$ stacking interactions with Tyr118, Tyr132 and Phe380 and H-bond interactions with Tyr132, Ser378 and Met508. The long extensions of compounds 11-14 form additional binding interactions along the access channel including Tyr64, Leu121, Phe233, Thr229, Ser506 and Ser507 (ligand interactions for each compound is provided in detail in Fig. S5 and S6†).

#### Conclusions

In general, the dichlorobenzene derivatives (b) displayed better inhibitory activity (MIC,  $IC_{50}$ ) and binding affinity ( $K_d$ ) compared with the monochlorobenzene derivatives (a). Among the medium arm extensions, the benzene acetyl derivative (6b) and the thiazole derivative (7b) had MIC and IC50 values comparable with FLZ (Table 2), however 7b had improved binding affinity ( $K_d = 106 \pm 15 \text{ nM}$ ) (Table 2). Comparison of the extended arm derivatives demonstrated the optimal linker with respect to antifungal activity was the amide (13), then urea (12), thiourea (11) and lastly sulfonamide (14), which was the least effective. The amide linker compounds (13a and 13b) were highly active against C. albicans strains CA14 and SC5314 with MIC  $< 0.03 \mu g \text{ mL}^{-1}$  and gave potent IC<sub>50</sub> values (13a IC<sub>50</sub> = 0.56  $\mu$ M, 13b IC<sub>50</sub> = 0.48  $\mu$ M), but only compound 13a showed very

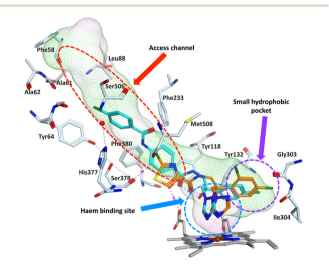


Fig. 4 Positioning of exemplar compounds S-7b (orange) and S-13a (cyan) in the CaCYP51 protein (for clarity not all amino acids are shown).

tight binding to CaCYP51 (13a  $K_d = 46 \pm 8$  nM) while a good binding affinity was noted for 13b ( $K_d = 166 \pm 39 \text{ nM}$ ,) compared with fluconazole (MIC =  $0.125 \mu g \text{ mL}^{-1}$ , IC<sub>50</sub> =  $0.31 \mu M$  and  $K_d$  =  $31 \pm 8 \text{ nM}$ ) (Table 2).

Evaluation (MIC) against S. cerevisiae models indicated the amide of the three lead compounds 7b, 13a and 13b all outperformed FLZ, with 13b optimal as it retained activity against mutant ScCYP51 Y140F/H mutant strains and the strain overexpressing C. albicans efflux pump MDR1a with a similar profile to PCZ. Computational studies suggest the S-enantiomers are generally preferred for optimal binding and positioning within the CaCYP51 active site, with the extended arm derivatives forming additional binding interactions in the access channel (Fig. 4 and S5†).

In summary (Fig. 5), the dichlorobenzene ring was optimal for binding within the small hydrophobic pocket and, in combination with an extended arm (e.g. as seen in 13b) performed better in resistant strains than FLZ owing to additional binding opportunities along the access channel to stabilise ligand-protein interactions.

Consideration of the physicochemical properties of the most promising medium (7b) and extended (13a/b) arm derivatives compared with reference antifungal agents (Table 4), showed that the thiazole derivative 7b is comparable with FLZ, while the extended derivatives (13a/b) are intermediate between FLZ and PCZ. Compounds 13a and 13b are just over the ideal molecular weight resulting in one violation of Lipinski's Ro5, however this is still a significant improvement compared with PCZ with three violations. Improved physicochemical properties would make these compounds more suitable for oral administration, a major limitation of ITZ and PCZ, which combined with the addition of linkers and long arms capable of H-bonding interactions enables antifungal activity against resistant strains, in particular common mutations in the CYP51 active site.

## Experimental

#### General

All chemicals, reagents and solvents were purchased from Sigma-Aldrich, Alfa Aesar, VWR, Acros and Fluka. Solvents were dried over molecular sieves (4 Å) prior to use. For column chromatography, a glass column was slurry packed in the appropriate eluent with silica gel (Fluka Kieselgel 60). TLC was performed on pre-coated silica plates (ALUGRAM® SIL G/UV254) with visualisation via UV light (254 nm). Melting points were determined using an electrothermal instrument (Gallenkamp melting point apparatus) and were uncorrected. 1H and 13C NMR spectra were recorded on a Bruker Advance DP500 spectrometer operating at 500 MHz and 125 MHz, respectively. Chemical shifts are given in parts per million (ppm) relative to the internal standard tetramethylsilane (Me<sub>4</sub>Si). HPLC-HRMS or HPLC performed at the Department of Pharmacy and Pharmacology, University of Bath, Bath, UK (method A) or at Cardiff University (method B). Method A: was performed on a

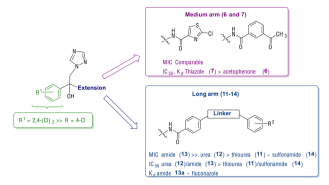


Fig. 5 Summary of antifungal (MIC), enzyme inhibition (IC50) and binding affinity ( $K_d$ ) of the medium (6 and 7) and long arm (11-14) extended compounds.

Zorbax Eclipse Plus C18 Rapid Resolution 2.1 × 50 mm, 1.8 μm particle size using a 7.5 minute gradient method 5:95 v/v water: methanol with 0.1% formic acid as additive; method B: was performed on a Shimadzu LC-2030C Plus C18 Rapid Resolution 250 × 4.6 mm, 5 μm particle size using a 7-10 min gradient method 5:95 v/v water: methanol. Details for the preparation of the intermediate amines (5 and 12) are provided in the ESI.†

#### Chemistry

General method for the synthesis of azoles (6, 7, 11–14). To a solution of 3-acetylbenzoic acid or 2-chlorothiazole-4carboxylic acid (1.5 eq.) in dry DMF (5 mL) was added CDI (1.5 eq.) and the reaction stirred at room temperature for 1 h. Then, a solution of 1-amino-2-(4-chlorophenyl)-3-(1H-1,2,4-triazol-1-yl)propan-2-ol (5) (1 eq.) in dry DMF (5 mL) was added and the reaction stirred at room temperature overnight. The reaction mixture was quenched with ice/cooled H<sub>2</sub>O (25 mL), the residue extracted with EtOAc (50 mL), washed with sat. aq. NaCl (25 mL × 2), and dried (MgSO<sub>4</sub>). The organic layer was evaporated under reduced pressure to give a light-yellow oil. Gradient chromatography was used to purify the compound, which eluted with CH2Cl2-MeOH to give a semi solid or white solid on drying (6 and 7). To prepare 11-14: to a cooled (0 °C, ice bath) solution of 4-amino-N-(2-(4-chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)

Table 4 Physicochemical properties of promising compounds and clinical antifungal agents

Cmpd	MW	$\operatorname{clog} P$	$n_{\mathrm{ON}}$	$n_{\mathrm{OHNH}}$	$n_{ m rotb}$	$n_{ m viol}$
7 <b>b</b>	399.81	2.55	7	2	6	0
13a	510.38	3.54	8	3	8	1
13b	544.83	4.15	8	3	8	1
FLZ	306.27	0.86	7	1	5	0
PCZ	700.77	5.74	12	1	12	3

MW = Molecular weight;  $n_{ON}$  = H-bond acceptor;  $n_{OHNH}$  = H-bond donor;  $n_{\text{rotb}}$  = number of rotable bonds;  $n_{\text{viol}}$  = number of Lipinsky violations.

benzamide (10) (1 eq.) in dry pyridine (5 mL) was added 4-chlorophenyl isothiocyanate or 4-chlorophenyl isocyanate or 4-chlorobenzoyl chloride or 4-chlorobenzenesulfonyl chloride (1.5 eq.) in portions and the reaction stirred at room temperature overnight. The solvent was evaporated, and the resulting oil extracted with EtOAc (50 mL), washed with 1 M aq. HCl (25 mL), H2O (2 × 25 mL), and dried (MgSO<sub>4</sub>). The organic layer was evaporated under reduced pressure to give a crude yellow-orange oil. The desired product was purified by gradient column chromatography using CH2Cl2-MeOH.

Research Article

3-Acetyl-N-(2-(4-chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1yl)propyl)benzamide (6a,  $R^1 = Cl$ ,  $R^2 = H$ ). Prepared from 1-amino-2-(4-chlorophenyl)-3-(1H-1,2,4-triazol-1-yl)propan-2-ol (5a) (0.15 g, 0.59 mmol) and 3-acetyl benzoic acid (0.14 g, 0.88 mmol). Purified by gradient column chromatography eluting with 3.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a semisolid, yield 0.12 g (52%). TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5: 0.5 v/v), R<sub>f</sub> 0.4. <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  8.55 (t, J = 5.7 Hz, 1H, NH), 8.27 (d, J = 7.2 Hz, 1H, Ar), 8.26 (s, 1H, triaz), 8.09 (d, J = 7.8 Hz, 1H, Ar), 7.98 (d, J = 8.3 Hz, 1H, Ar), 7.83 (s, 1H, triaz), 7.60 (d, J = 7.8 Hz, 1H,Ar), 7.45 (d, J = 8.8 Hz, 2H, Ar), 7.32 (d, J = 8.8 Hz, 2H, Ar), 6.07 (s, 1H, OH, ex), 4.61 (q, J = 14.4 Hz, 2H, CH<sub>2</sub>-triaz), 3.90 (dd, J =6.8, 14.0 Hz, 1H, CH<sub>2</sub>-NH), 3.65 (dd, J = 5.3, 14.0 Hz, 1H, CH<sub>2</sub>-NH), 2.61 (s, 3H, CH<sub>3</sub>).  $^{13}$ C NMR (DMSO-d<sub>6</sub>):  $\delta$  197.99 (C,  $CH_3C=O$ ), 167.08 (C, C=O, amide), 150.01 (CH, triaz), 145.45 (CH, triaz), 141.40 (C, Ar), 137.19 (C, Ar), 134.92 (C, Ar), 132.31 (CH, Ar), 132.17 (C, C-Cl), 131.41 (CH, Ar), 129.29 (CH, Ar), 128.37 (2× CH, Ar), 128.09 (2× CH, Ar), 127.29 (2× CH, Ar), 76.28 (C-OH), 57.01 (CH<sub>2</sub>-triaz), 48.24 (CH<sub>2</sub>-NH), 27.30 (CH<sub>3</sub>). HPLC (method A): 100%, RT = 4.37min. HRMS (ESI, m/z). Calcd for  $C_{20}H_{19}ClN_4O_3$  [M + H]<sup>+</sup>, 399.1224; found, 399.1225.

3-Acetyl-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4triazol-1yl)propyl)benzamide (6b,  $R^1 = Cl$ ,  $R^2 = Cl$ ). Prepared 1-amino-2-(2,4-dichlorophenyl)-3-(1*H*-1,2,4-triazol-1-yl) propan-2-ol (5b) (0.15 g, 0.52 mmol) and 3-acetyl benzoic acid (0.13 g, 0.78 mmol). Purified by gradient column chromatography eluting with 3.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.1 g (45%). M.p.: 160-162 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f = 0.45$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  8.72 (t, J = 6.0 Hz, 1H, NH), 8.34 (s, 1H, triaz), 8.30 (s, 1H, Ar), 8.10 (d, J = 7.8 Hz, 1H, Ar), 8.01 (d, J = 8.3Hz, 1H, Ar), 7.74 (s, 1H, triaz), 7.59 (m, 3H, Ar), 7.29 (dd, J =2.2, 8.6 Hz, 1H, Ar), 6.35 (s, 1H, OH, ex), 5.10 (d, J = 14.4 Hz, 1H, CH<sub>2</sub>-triaz), 4.70 (d, J = 14.4 Hz, 1H, CH<sub>2</sub>-triaz), 4.08 (dd, J= 5.7, 14.0 Hz, 1H,  $CH_2$ -NH), 3.97 (dd, J = 6.4, 14.0 Hz, 1H,  $CH_2$ -NH), 2.61 (s, 3H,  $CH_3$ ). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  198.00 (C, CH<sub>3</sub>C=O), 167.58 (C, C=O, amide), 151.02 (CH, triaz), 145.56 (CH, triaz), 138.11 (C, Ar), 137.20 (C, Ar), 134.71 (C, Ar), 133.29 (C, C-Cl), 132.40 (CH, Ar), 131.51 (CH, Ar), 131.50 (C, C-Cl), 130.43 (CH, Ar), 129.30 (2× CH, Ar), 127.39 (2× CH, Ar), 76.90 (C-OH), 54.09 (CH<sub>2</sub>-triaz), 45.95 (CH<sub>2</sub>-NH), 27.32 (CH<sub>3</sub>). HPLC (method A): 100%, RT = 4.49 min. HRMS (ESI, m/z), calcd for  $C_{20}H_{18}Cl_2N_4O_3$  [M + H]<sup>+</sup>, 433.0834; found, 433.0832.

2-Chloro-N-(2-(4-chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)thiazole-4-carboxamide (7**a**,  $R^1 = Cl$ ,  $R^2 = H$ ). Prepared from 1-amino-2-(2,4-dichlorophenyl)-3-(1H-1,2,4triazol-1-yl)propan-2-ol (5a) (0.16 g, 0.63 mmol) and 2-chlorothiazole-4-carboxylic acid (0.15 g, 0.95 mmol). Purified by gradient column chromatography eluting with 3.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a semisolid, yield 0.17 g (68%). TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5: 0.5 v/v),  $R_f = 0.45$ .  $^{1}$ H NMR (DMSO-d<sub>6</sub>):  $\delta$  8.23 (s, 1H, thiazole), 8.22 (s, 1H, triaz), 8.10 (t, J = 5.2 Hz, 1H, NH), 7.84 (s, 1H, triaz), 7.41 (d, J = 8.8 Hz, 2H, Ar), 7.32 (d, J = 8.8 Hz, 2H, Ar), 6.12 (s, 1H, OH, The state of the staex), 4.53 (s, 2H,  $CH_2$ -triaz), 3.96 (dd, J = 7.3, 14.1 Hz, 1H,  $CH_2$ -NH), 3.58 (dd, J = 5.1, 14.1 Hz, 1H,  $CH_2$ -NH). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  160.06 (C, C=O, amide), 151.44 (C, C-Cl), 151.13 (CH, triaz), 147.72 (C, Ar), 145.45 (CH, triaz), 141.28 (C, thiazole), 132.28 (C, Ar), 128.30 (2× CH, Ar), 128.19 (2× CH, Ar), 128.03 (CH, thiazole), 75.72 (C-OH), 57.26 (CH<sub>2</sub>triaz), 47.22 (CH<sub>2</sub>-NH). HPLC (method A): 100%, RT = 4.52 min. HRMS (ESI, m/z). Calcd for  $C_{15}H_{13}Cl_2N_5O_2S$  [M + H]<sup>+</sup>, 398.0245; found, 398.0242.

2-Chloro-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4triazol-1-yl)propyl)thiazole-4-carboxamide (7b,  $R^1 = Cl$ ,  $R^2 = Cl$ ). Prepared from 1-amino-2-(2,4-dichlorophenyl)-3-(1H-1,2,4triazol-1-yl)propan-2-ol (5b) (0.15 g, 0.52 mmol) and 2-chlorothiazole-4-carboxylic acid (0.13 g, 0.78 mmol). Purified by gradient column chromatography eluting with 2.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a pale yellow wax, yield 0.15 g (68%). TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f$  = 0.55. <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  8.32 (s, 1H, thiazole), 8.26 (t, J =5.9 Hz, 1H, NH), 8.25 (s, 1H, triaz), 7.72 (s, 1H, triaz), 7.54 (d, J = 7.7 Hz, 2H, Ar), 7.29 (dd, J = 2.2, 8.6 Hz, 1H, Ar), 6.33 (s, 1H, OH, ex), 5.03 (d, J = 14.4 Hz, 1H, CH<sub>2</sub>-triaz), 4.64 (d, J =14.4 Hz, 1H,  $CH_2$ -triaz), 4.05 (dd, J = 6.7, 14.0 Hz, 1H,  $CH_2$ -NH), 3.99 (dd, J = 5.8, 14.0 Hz, 1H, CH<sub>2</sub>-NH). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  160.43 (C, C=O, amide), 151.45 (C, C-Cl), 151.03 (CH, triaz), 147.67 (C, Ar), 145.58 (CH, triaz), 137.98 (C, thiazole), 133.35 (C, C-Cl), 132.07 (C, C-Cl), 131.49 (CH, Ar), 130.47 (CH, Ar), 128.18 (CH, thiazole), 127.30 (CH, Ar), 76.27 (C-OH), 54.14 (CH<sub>2</sub>-triaz), 45.06 (CH<sub>2</sub>-NH). HPLC (method A): 100%, RT = 4.63 min. HRMS (ESI, m/z). Calcd for  $C_{15}H_{12}Cl_3N_5O_2S[M+H]^+$ , 431.9855; found, 433.9826.

N-(2-(4-Chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1yl)propyl)-4-(3-(4-chlorophenyl)thioureido)benzamide (11a,  $R^1$  = Cl,  $R^2 = H$ , X = S). Prepared from 4-amino-N-(2-(4chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10a) (0.16 g, 043 mmol) and 4-chlorophenyl isothiocyanate (0.11 g, 0.64 mmol). Purified by gradient column chromatography eluting with 2.5% MeOH in CH2Cl2. Product obtained as a white solid, yield 0.15 g (68%). M.p.: 146–148 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>–MeOH 9.5:0.5 v/v),  $R_f = 0.23$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.05 (s, 1H, NH-thiourea), 10.00 (s, 1H, NH-thiourea), 8.22 (t, J = 5.9 Hz, 1H, NH-CH<sub>2</sub>), 8.20 (s, 1H, triaz), 7.84 (s, 1H, triaz), 7.70 (d, J = 8.9 Hz, 2H, Ar), 7.55 (dd, J = 8.8, 21.2 Hz, 4H, Ar, 7.43 (d, J = 7.1 Hz, 2H, Ar), 7.39 (d, J= 9.0 Hz, 2H, Ar), 7.29 (m, 2H, Ar), 6.05 (s, 1H, OH, ex), 5.74  $(dd, J = 14.3, 20.3 \text{ Hz}, 2H, CH_2\text{-triaz}), 3.86 (dd, J = 6.7, 14.0 \text{ Hz},$ 

1H,  $C\underline{H_2}$ –NH), 3.63 (dd, J = 5.2, 14.1 Hz, 1H,  $C\underline{H_2}$ –NH). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  179.97 (C, C=S), 167.33 (C, C=O), 150.96 (CH, triaz), 145.38 (CH, triaz), 142.70 (C, Ar), 142.45 (C, Ar), 141.48 (C, Ar), 138.75 (C, C-Cl), 132.14 (C, C-Cl), 129.85 (C, Ar), 129.84 (2× CH, Ar), 128.23 (2× CH, Ar), 128.17 (2× CH, Ar), 126.24 (2× CH, Ar), 125.74 (2× CH, Ar), 122.74 (2× CH, Ar), 76.48 (C, C-OH), 57.40 (CH<sub>2</sub>-triaz), 48.35 (CH<sub>2</sub>-NH). HPLC (method B): 99%, RT = 4.78 min. HRMS (ESI, m/z). Calcd for  $C_{25}H_{22}Cl_2N_6O_2S$  [M + H]<sup>+</sup>, 541.0980; found, 541.0971.

4-(3-(4-Chlorophenyl)thioureido)-N-(2-(2,4-dichlorophenyl)-2hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (11b,  $R^1 = Cl$ ,  $R^2 = Cl$ , X = S). Prepared from 4-amino-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1*H*-1,2,4-triazol-1-yl)propyl) benzamide (10b) (0.17 g, 0.41 mmol) and 4-chlorophenyl isothiocyanate (0.1 g, 0.62 mmol). Purified by gradient column chromatography eluting with 3.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.13 g (56%). M.p.: 138-140 °C. TLC (CH2Cl2-MeOH 9.5: 0.5 v/v),  $R_{\rm f} = 0.32$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.06 (s, 1H, NHthiourea), 10.03 (s, 1H, NH-thiourea), 8.47 (t, J = 5.9 Hz, 1H, NH- $CH_2$ ), 8.34 (s, 1H, triaz), 7.74 (s, 1H, triaz), 7.73 (d, J = 8.8 Hz, 2H, Ar), 7.57 (m, 4H, Ar), 7.51 (d, J = 8.9 Hz, 2H, Ar), 7.39 (d, J = 8.9Hz, 2H, Ar), 7.29 (dd, J = 2.3, 8.7 Hz, 1H, Ar), 6.46 (s, 1H, OH-ex),  $5.05 \text{ (d, } J = 14.4 \text{ Hz, } 1H, \text{ CH}_2\text{-triaz}), 4.68 \text{ (d, } J = 14.4 \text{ Hz, } 1H, \text{ CH}_2\text{-}$ triaz), 4.02 (dd, J = 5.7, 14.2 Hz, 1H, CH<sub>2</sub>-NH), 3.97 (dd, J = 6.5, 14.2 Hz, 1H, CH<sub>2</sub>-NH).  $^{13}$ C NMR (DMSO-d<sub>6</sub>):  $\delta$  179.90 (C, C=S), 168.02 (C, C=O), 151.01 (CH, triaz), 145.59 (CH, triaz), 142.89 (C, Ar), 138.76 (C, Ar), 138.19 (C, C-Cl), 133.26 (C, C-Cl), 132.17 (C, C-Cl), 132.04 (C, C-Cl), 131.54 (CH, Ar), 130.41 (CH, Ar), 128.81 (2× CH, Ar), 128.32 (2× CH, Ar), 127.27 (CH, Ar), 125.72 (2× CH, Ar), 122.70 (2× CH, Ar), 76.99 (C, C-OH), 54.18 (CH<sub>2</sub>triaz), 46.05 (CH<sub>2</sub>-NH). Anal. calcd for C<sub>25</sub>H<sub>21</sub>Cl<sub>3</sub>N<sub>6</sub>O<sub>2</sub>S (575.89): C 52.13%, H 3.68%, N 14.59%. Found: C 52.29%, H 3.82%, N 14.71%.

N-(2-(4-Chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1yl)propyl)-4-(3-(4-chlorophenyl)ureido)benzamide (12a,  $R^1 = Cl$ ,  $R^2 = H$ , X = O). Prepared from 4-amino-N-(2-(4-chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10a) (0.16 g, 0.43 mmol) and 4-chlorophenyl isocyanate (0.1 g, 0.64 mmol). Purified by gradient column chromatography eluting with 6% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.14 g (63%). M.p.: 226-228 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f = 0.35$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  8.99 (s, 1H, NHurea), 8.90 (s, 1H, NH-urea), 8.25 (s, 1H, triaz), 8.22 (t, J = 5.7Hz, 1H, NH-CH<sub>2</sub>), 7.83 (s, 1H, triaz), 7.54 (dd, J = 3.2, 8.9 Hz, 2H, Ar), 7.48 (m, 4H, Ar), 7.43 (d, J = 8.8 Hz, 2H, Ar), 7.31 (m, 4H, Ar), 6.16 (s, 1H, OH-ex), 4.60 (dd, J = 14.3, 21.5 Hz, 2H,  $CH_2$ -triaz), 3.85 (dd, J = 6.7, 14.1 Hz, 1H,  $CH_2$ -NH), 3.63 (dd, J= 5.2, 14.0 Hz, 1H, CH<sub>2</sub>-NH).  $^{13}$ C NMR (DMSO-d<sub>6</sub>):  $\delta$  167.51 (C, C=O), 152.63 (C, C=O), 151.00 (CH, triaz), 145.45 (CH, triaz), 143.01 (C, Ar), 142.50 (C, Ar), 141.52 (C, Ar), 138.88 (C, C-Cl), 132.12 (C, C-Cl), 129.10 (C, Ar), 128.80 (2× CH, Ar), 128.38 (2× CH, Ar), 128.09 (2× CH, Ar), 120.34 (2× CH, Ar), 117.66 (2× CH, Ar), 76.34 (C, C-OH), 57.14 (CH<sub>2</sub>-triaz), 48.24 (CH<sub>2</sub>-NH). Anal. calcd. for  $C_{25}H_{22}Cl_2N_6O_3$ , (525.39): C 57.15%, H 4.22%, N 15.99%. Found: C 57.53%, H 4.28%, N 15.94%.

4-(3-(4-Chlorophenyl)ureido)-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide  $(12b, R^1 = Cl, R^2 = Cl, X = O)$ . Prepared from 4-amino-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10b) (0.16 g, 0.43 mmol) and 4-chlorophenyl isocyanate (0.1 g, 0.64 mmol). Purified by gradient column chromatography eluting with 6% MeOH in CH2Cl2. Product obtained as a white solid, yield 0.11 g (52%). M.p.: 216-218 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f = 0.37$ . <sup>1</sup>H NMR (DMSO- $d_6$ ):  $\delta$  9.06 (s, 1H, NH-urea), 8.97 (s, 1H, NH-urea), 8.42 (t, J = 6.0 Hz, 1H, NH-CH<sub>2</sub>), 8.34 (s, 1H, triaz), 7.72 (d, J = 11.0 Hz, 2H, Ar), 7.70 (s, 1H, triaz), 7.57 (dd, J = 8.6, 11.0 Hz, 2H, Ar), 7.49 (m, 4H, Ar), 7.33 (d, J = 8.9 Hz, 2H, Ar), 7.28 (dd, J = 2.2, 8.6 Hz, 1H, Ar), 6.53 (s, 1H, OH, ex), 5.05 (d, J = 14.3 Hz, 1H, CH<sub>2</sub>-triaz), 4.67 (d, J = 14.3 Hz, 1H, CH<sub>2</sub>-triaz), 4.02 (dd, J = 5.5, 14.1 Hz, 1H, CH<sub>2</sub>-NH), 3.95 (dd, J = 6.3, 14.1 Hz, 1H, CH<sub>2</sub>-NH). <sup>13</sup>C NMR (DMSO- $d_6$ ):  $\delta$  168.21 (C, C=O), 152.66 (C, C=O), 151.00 (CH, triaz), 145.45 (CH, triaz), 143.18 (C, Ar), 138.90 (C, Ar), 138.25 (C, C-Cl), 133.25 (C, C-Cl), 132.02 (C, C-Cl), 131.55 (CH, Ar), 131.24 (CH, Ar), 130.04 (CH, Ar), 129.10 (2× CH, Ar), 128.93 (CH, Ar), 127.27 (CH, Ar), 120.35 (2× CH, Ar), 117.64 (2× CH, Ar), 77.03 (C, C-OH), 54.19 (CH<sub>2</sub>-triaz), 46.12 (CH<sub>2</sub>-NH). Anal. calcd. For C<sub>25</sub>H<sub>21</sub>-Cl<sub>3</sub>N<sub>6</sub>O<sub>3</sub> (559.83): C 53.64%, H 3.78%, N 15.00%. Found: C 53.84%, H 3.68%, N 14.63%.

4-Chloro-N-(4-((2-(4-chlorophenyl)-2-hydroxy-3-(1H-1,2,4triazol-1-yl)propyl) carbamoyl) phenyl)benzamide (13a,  $R^1 = Cl$ ,  $R^2$ = H, Y = C(O)). Prepared from 4-amino-N-(2-(4-chlorophenyl)-2hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10a) (0.15 g, 0.42 mmol) and 4-chlorobenzoyl chloride (0.1 g, 0.64 mmol). Purified by gradient column chromatography eluting with 6% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.16 g (80%). M.p.: 222-224 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f = 0.33$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.50 (s, 1H, NHamide), 8.28 (t, J = 5.7 Hz, 1H, NH-CH<sub>2</sub>), 8.26 (s, 1H, triaz), 7.99 (d, J = 8.8 Hz, 2H, Ar), 7.84 (s, 1H, triaz), 7.83 (dd, J =3.8, 8.9 Hz, 2H, Ar), 7.76 (d, J = 8.9 Hz, 2H, Ar), 7.62 (d, J =8.8 Hz, 2H, Ar), 7.44 (d, J = 8.8 Hz, 2H, Ar), 7.30 (m, 2H, Ar), 6.14 (s, 1H, OH-ex), 4.59 (dd, J = 14.4, 21.9 Hz, 2H, CH<sub>2</sub>triaz), 3.86 (dd, J = 6.8, 14.1 Hz, 1H, CH<sub>2</sub>-NH), 3.64 (dd, J =5.3, 14.1 Hz, 1H, CH<sub>2</sub>-NH).  $^{13}$ C NMR (DMSO-d<sub>6</sub>):  $\delta$  167.38 (C, C=O), 165.13 (C, C=O), 151.00 (CH, triaz), 145.45 (CH, triaz), 142.26 (C, Ar), 141.49 (C, Ar), 137.10 (C, C-Cl), 133.76 (C, Ar), 132.14 (C, C-Cl), 130.17 (2× CH, Ar), 129.34 (C, Ar), 128.97 (2× CH, Ar), 128.53 (2× CH, Ar), 128.09 (2× CH, Ar), 126.25 (2× CH, Ar), 119.91 (2× CH, Ar), 76.33 (C, C-OH), 57.10 (CH<sub>2</sub>-triaz), 48.24 (CH<sub>2</sub>-NH). Anal. calcd. for  $C_{25}H_{21}Cl_2N_5O_3$ (510.38): C 58.83%, H 4.15%, N 13.72%. Found: C 58.78%, H 4.07%, N 13.68%.

4-Chloro-N-(4-((2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl) carbamoyl)phenyl)benzamide (13b,  $R^1 = Cl$ ,  $R^2 = Cl$ , Y = C(O)). Prepared from 4-amino-N-(2-(2,4-dichlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10b) (0.16 g, 0.39 mmol) and 4-chlorobenzoyl chloride (0.16 g, 0.6 mmol). Purified by gradient column chromatography eluting with 6% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.16 g (76%). M.p.: 190–192 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>–MeOH 9.5: 0.5 v/v),  $R_f = 0.32$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.51 (s, 1H, NH-amide), 8.49 (t, J = 0.32).

5.9 Hz, 1H, NH–CH<sub>2</sub>), 8.35 (s, 1H, triaz), 7.99 (d, J = 8.6 Hz, 2H, Ar), 7.84 (d, J = 8.8 Hz, 2H, Ar), 7.79 (d, J = 8.8 Hz, 2H, Ar), 7.74 (s, 1H, triaz), 7.62 (d, J = 8.6 Hz, 2H, Ar), 7.57 (dd, J = 8.6, 13.4 Hz, 2H, Ar), 7.29 (dd, J = 2.2, 8.6 Hz, 1H, Ar), 6.50 (s, 1H, OH, ex), 5.07 (d, J = 14.3 Hz, 1H, CH<sub>2</sub>-triaz), 4.68 (d, J = 14.3 Hz, 1H, CH<sub>2</sub>-triaz), 4.04 (dd, J = 5.5, 14.0 Hz, 1H, CH<sub>2</sub>-NH), 3.97 (dd, J = 6.5, 14.0 Hz, 1H, CH<sub>2</sub>-NH). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  168.02 (C, C=O), 165.14 (C, C=O), 151.00 (CH, triaz), 145.58 (CH, triaz), 142.38 (C, Ar), 138.21 (C, Ar), 137.11 (C, C-Cl), 133.75 (C, C-Cl), 133.26 (C, C-Cl), 132.04 (C, Ar), 131.55 (CH, Ar), 130.41 (CH, Ar), 130.17 (2× CH, Ar), 129.06 (C, Ar), 128.98 (2× CH, Ar), 128.65 (2× CH, Ar), 127.27 (CH, Ar), 119.89 (2× CH, Ar), 76.99 (C, C-OH), 54.16 (CH<sub>2</sub>-triaz), 46.68(CH<sub>2</sub>-NH). Anal. calcd. for C<sub>25</sub>H<sub>20</sub>Cl<sub>3</sub>N<sub>5</sub>O<sub>3</sub> (544.82): C 54.39%, H 3.65%, N 12.68%. Found: C 54.05%, H 3.80%, N 12.31%.

Research Article

N-(2-(4-Chlorophenyl)-2-hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)-4-((4-chlorophenyl) sulfonamido)benzamide (14a R<sup>1</sup> = Cl, R<sup>2</sup> = H, Y = S(O<sub>2</sub>)). Prepared from 4-amino-N-(2-(4-chlorophenyl)-2-hydroxy-3-(1*H*-1,2,4-triazol-1-yl)propyl)benzamide (10a) (0.30 g, 0.82 mmoL) and 4-chlorobenzenesulfonyl chloride (0.26 g, 1.23 mmol). Purified by gradient column chromatography eluting with 3.5% MeOH in CH<sub>2</sub>Cl<sub>2</sub>. Product obtained as a white solid, yield 0.35 g (77%). M.p. 236-238 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5: 0.5 v/v),  $R_f = 0.45$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.73 (br.s, 1H, NHSO<sub>2</sub>), 8.23 (s, 1H, triazole), 8.21 (t, J = 6.0 Hz, 1H, NH), 7.81 (s, 1H, triazole), 7.79 (d, J = 8.8 Hz, 2H, Ar), 7.63 (m, 4H, Ar), 7.41 (d, J = 8.7 Hz, 2H, Ar), 7.28 (m, 2H, Ar), 7.16 (m, 2H, Ar), 6.04 (s, 1H, OH), 4.56 (dd, J = 14.4, 25.0 Hz, 2H, CH<sub>2</sub>-triazole), 3.79 (dd, J =6.6, 14.0 Hz, 1H, NHCHaHb), 3.59 (dd, J = 5.3, 14.0 Hz, 1H, NHCHaHb). <sup>13</sup>C NMR (DMSO-d<sub>6</sub>):  $\delta$  167.19 (C, C=O), 151.00 (CH, triazole), 145.44 (CH, triazole), 141.45 (C, Ar), 140.85 (C, Ar), 138.65 (C, Ar), 138.48 (C, C-Cl), 132.15 (C, C-Cl), 130.03 (2× CH, Ar), 129.76 (C, Ar), 129.15 (2× CH, Ar), 129.04 (2× CH, Ar), 128.35 (2× CH, Ar), 128.06 (2× CH, Ar), 119.09 (2× CH, Ar), 76.25 (C, C-OH), 57.04 (CH<sub>2</sub>-triazole), 48.18 (CH<sub>2</sub>-NH). HPLC (method B): 100%, RT = 4.77 min. HRMS (ESI), m/z. Calcd for  $C_{24}H_{21}Cl_2N_5O_4S$  $[M + H]^+$ , 546.0769; found, 546.0769.

4-((4-Chlorophenyl)sulfonamido)-N-(2-(2,4-dichlorophenyl)-2hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (14b,  $R^1 = Cl$ ,  $R^2 =$ Cl,  $Y = S(O_2)$ ). Prepared from 4-amino-N-(2-(2,4-dichlorophenyl)-2hydroxy-3-(1H-1,2,4-triazol-1-yl)propyl)benzamide (10b) (0.17 g, 0.42 mmoL) and 4-chlorobenzenesulfonyl chloride (0.13 g, 0.63 mmoL). Purified by gradient column chromatography eluting with 3.5% MeOH in CH2Cl2. Product obtained as a white solid, yield 0.2 g (83%). M.p. 228-230 °C. TLC (CH<sub>2</sub>Cl<sub>2</sub>-MeOH 9.5:0.5 v/v),  $R_f = 0.46$ . <sup>1</sup>H NMR (DMSO-d<sub>6</sub>):  $\delta$  10.76 (br.s, 1H, NHSO<sub>2</sub>), 8.40 (t, J = 6.0 Hz, 1H, NH), 8.32 (s, 1H, triazole), 7.79 (d, J = 8.8Hz, 2H, Ar), 7.72 (s, 1H, triazole), 7.60 (dd, J = 4.8, 8.9 Hz, 4H, Ar), 7.54 (dd, J = 3.2, 5.5 Hz, 2H, Ar), 7.27 (dd, J = 2.2, 8.7 Hz, 1H, Ar), 713 (d, J = 8.8 Hz, 2H, Ar), 6.36 (s, 1H, OH), 5.04 (d, J = 14.4Hz, 1H, CHaHb-triazole), 4.63 (d, J = 14.4 Hz, 1H, CHaHbtriazole), 4.01 (dd, J = 5.7, 14.1 Hz, 1H, CHaHbNH), 3.89 (dd, J = 6.3, 14.0 Hz, 1H, CHaHbNH).  $^{13}$ C NMR (DMSO-d<sub>6</sub>):  $\delta$  167.78 (C, C=O), 151.00 (CH, triazole), 145.56 (CH, triazole), 141.01 (C, Ar), 138.65 (C, Ar), 138.48 (C, Ar), 138.15 (C, C-Cl), 133.26 (C, C-Cl), 132.01 (C, C-Cl), 131.50 (CH, Ar), 130.41 (CH, Ar), 130.03 (2× CH,

Ar), 129.44 (C, Ar), 129.26 (2× CH, Ar), 129.08 (2× CH, Ar), 127.27 (CH, Ar), 119.02 (2× CH, Ar), 76.91 (C, C–OH), 45.09 (CH<sub>2</sub>-triazole), 45.96 (CH<sub>2</sub>–NH). HPLC (method B): 99.7%, RT = 4.76 min. HRMS (ESI), m/z. calcd for  $C_{24}H_{20}Cl_3N_5O_4S$  [M + H]<sup>+</sup>, 580.038; found, 580.0304.

#### CaCYP51 susceptibility testing

Antifungal MIC values were obtained according to the CLSI microdilution method for yeasts.  $^{28}$  *C. albicans* strains CA14 and SC5314 were tested in triplicate. Cultures were diluted to  $2.5 \times 10^3$  cells per mL in RPMI 1640 (Sigma), buffered with 0.165 M MOPS, pH 7.0. Antifungal compounds were dissolved in DMSO and added at a final concentration of 1% v/v DMSO. Plates were incubated at 37 °C and read at 48 h.

#### CaCYP51 IC<sub>50</sub>

C. albicans CYP51 was overexpressed in E. coli and purified and CaCYP51 reconstitutions assays performed.  $^{10,37}$  The final reaction volume of 500  $\mu$ L, contained; 1  $\mu$ M C. albicans CYP51, 2  $\mu$ M Homo sapiens cytochrome P450 reductase (CPR) (UniProtKB accession number P16435), 60  $\mu$ M lanosterol, 50  $\mu$ M dilaurylphosphatidylcholine, 4% w/v 2-hydroypropylcyclodextrin, 0.4 mg mL $^{-1}$  isocitrate dehydrogenase, 25 mM trisodium isocitrate, 50 mM NaCl, 5 mM MgCl $_2$ , and 40 mM MOPS (morpholinepropanesulfonic acid, pH 7.2). Antifungal compounds were added in 2.5  $\mu$ L DMSO. The reaction was started by the addition of 4 mM NADPH-tetrasodium salt. Samples were shaken for 15 min at 37 °C and sterol metabolites extracted with ethyl acetate and derivatised with BSTFA.

#### Binding affinity $(K_d)$

The ligand–CaCYP51 complex was determined by non-linear regression (Levenberg–Marquardt algorithm) using a rearrangement of the Morrison equation for tight ligand binding,  $^{38}$  while weak ligand binding was calculated by the Michaelis–Menten equation. Curve fitting for novel azole saturation curves was performed using the computer program ProFit 6.1.12 (QuantumSoft, Zurich, Switzerland) for Mac OSX.  $K_d$  values were determined for each of the three replicate titrations per azole compound and then mean  $K_d$  values and standard deviations calculated.

#### Preparation of S. cerevisiae strain Y2760

The *HsCYP51* ORF (NP\_000777.1) was codon harmonized (ch) using DNA 2.0 for expression in *S. cerevisiae* and synthesized by ATUM (Newark, Ca, USA). The 3' end of the ORF was extended to encode EQKI of the ScErg11 C-terminus, followed by a GGR linker and a 6xHis tag. Strain Y2514 was prepared by transforming host strain Y1857 with a transformation cassette including the above construct together with the *ScPGK* transcription terminator and *URA3* selection marker located downstream, bordered by upstream and downstream parts of *ScPDR5* gene identical to sequences in ScErg11 over-

expressing strain Y941. Strain Y2514 overexpressed HsCYP51ch-6xHis and URA3 from the ScPDR5 locus.

Strain Y2757 (Y2514, PDR15::HsCPRch-6xHis-LoxP-HIS1-LoxP) was prepared by co-expressing the cognate reductase HsCRP in strain Y2514. Codon harmonisation and synthesis of human NADPH-cytochrome P450 reductase isoform 2 (HsCPRch, NP\_001369584.1) ORF was performed by ATUM. The gene was 3' extended with a sequence encoding GGR-6xHIS. A transformation cassette was prepared by recombinant PCR to include the HsCPRch ORF with 657 nucleotides the ScPDR5 promoter upstream, and the ScPGK terminator followed by the LoxP flanked ScHIS1 selection marker downstream. ScPDR15 specific arms bordering the construct enabled the cassette to be integrated between the nucleotide 441 of the 5' UTR and nucleotide 1 of the 3' UTR in PDR15.

Strain Y2758 (Y2757 AHIS1) was made using the following steps. Strain Y2757 was transformed with the pSH69 pGAL1-cre plasmid (Euroscarf, Germany) via selection on hygromycin (75 µg mL<sup>-1</sup>) containing YPD plates. A single hygromycin positive colony was inoculated in 1% yeast extract, 2% peptone and 2% of galactose broth and incubated for 16 hours at 30 °C with shaking. The diluted culture was then plated on YPD agar and replicaplated on YPD + hygromycin and SD-HIS dropout agar plates. Deletion of the HIS selection marker downstream of HsCPRch-6xHis in the PDR15 locus was confirmed in double His and hygromycin negative isolates by PCR and DNA sequence analysis.

The endogenous ScERG11 ORF in strain Y2758 was removed by transforming with a LoxP- ScHIS1 containing cassette with arms complementing from +387 nucleotides in the 5'UTR and -60 nucleotides in the 3'UTR of the ScERG11 locus, resulting in strain Y2760 (ADAA PDR5::HsCYP51ch-6×HIS-URA3; PDR15::  $HsCPRch6 \times HIS-LoxP$ ,  $\Delta ERG11-LoxP-HIS1-LoxP$ ).

#### Antifungal susceptibility testing of recombinant S. cerevisiae strains expressing CYP51

Minimum inhibitory concentration (MIC) were determined according to recommendations outlined in the Clinical and Laboratory Standards Institute (CLSI) document M27-Ed4E<sup>39</sup> with modifications. Sabouraud dextrose (SD) medium (pH 6.8) was used instead of RPMI-1640 for inoculum and growth of the S. cerevisiae strains and cultures were incubated at 30 °C. Assays were performed in round-bottomed 96-well plates (Corning, NY, US) and MICs were measured as the lowest concentrations of each antifungal agent that resulted in 80% inhibition of growth as compared with a drug-free, positive control. Stock solutions of each agent were prepared in DMSO and the final concentration of DMSO was 1% (v/v) in SD. The final concentration of fluconazole ranged from 0.25 to 128 µg mL<sup>-1</sup>. The final concentration of all other compounds ranged from 0.03-16 µg mL<sup>-1</sup>. Three biological repeats were performed.

#### Computational studies

CaCYP51 protein-ligand complexes of all final compounds (R- and S-enantiomers) were obtained with Molecular Operating Environment (MOE) software<sup>33</sup> using the crystal structure of wild-type CaCYP51 (PDB 5FSA).<sup>34</sup> Minimisations were performed with the MMFF94 force field (ligands) and partial charges were automatically calculated. The charge of the haem iron at physiological pH was set to 3<sup>+</sup> (geometry d2sp3) through the atom manager in MOE. Results were refined using the MMFF94 force field and scored by applying the London  $\Delta G$  scoring function. The output database dock file was created with different poses for each ligand and arranged according to the final S-score function, which is the score of the last stage that was not set to zero. Molecular dynamics (MD) simulations were performed on CaCYP51ligand complexes, which were optimised with the protein preparation wizard in Maestro (Schrödinger release 2020-1), 35 by assigning bond orders, adding hydrogens, and correcting incorrect bond types. A default quick relaxation protocol was used to minimise the MD systems with the Desmond programme.<sup>36</sup> The orthorhombic water box allowed for a 10 Å buffer region between protein atoms and box sides. Overlapping water molecules were deleted, and the systems were neutralised with Na<sup>+</sup> ions and a salt concentration of 0.15 M. Force-field parameter assignment used the OPLS 2005 force field, for a 200 ns molecular dynamic run in the NPT ensemble (T = 300 K) at a constant pressure of 1 bar. Energy and trajectory atomic coordinate data were recorded at 1.2 ns intervals.

## Data availability

The data supporting this article have been included as part of the ESI.†

#### Author contributions

MA and FAB performed the chemical synthesis and ET performed computational studies supervised by CS. AGW performed the binding affinity studies supervised by DEK and SLK. JEP performed the MIC and IC50 assays and BCM and MVK contributed to the S. cerevisiae model assays. All authors contributed to manuscript preparation.

#### Conflicts of interest

There are no conflicts to declare.

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