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# Exploiting interkingdom interactions for the development of small molecule inhibitors of *Candida albicans* biofilm formation

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14 **Running Head:** Hydroxy Alkylquinolone signals target *Candida* biofilm.

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### 19 Abstract

A rapid decline in the development of new antimicrobial therapeutics has coincided with the emergence of new and more aggressive multidrug resistant pathogens. Pathogens are protected from antibiotic activity by their ability to enter an aggregative biofilm state. Therefore, disrupting this process in pathogens is a key strategy for the development of next generation antimicrobials. Here we present a suite of compounds, based on the Pseudomonas aeruginosa 2-heptyl-4(1H)-quinolone (HHQ) core quinolone interkingdom signal structure, that exhibit non-cytotoxic antibiofilm activity towards the fungal pathogen Candida albicans. In addition to providing new insights into what is a clinically important bacterial-fungal interaction, the capacity to modularize the functionality of the quinolone signals is an important advance in harnessing the therapeutic potential of signaling molecules in general. This provides a platform for the development of potent next generation small molecular therapeutics targeting clinically relevant fungal pathogens. 

### 41 Introduction

With the ever increasing emergence of antibiotic resistant pathogens and the lack of new 42 antibiotics coming to market, we are entering a 'post-antibiotic era' (1-3). This realization has 43 underpinned a global initiative to identify new and innovative approaches to infection 44 management. As such, targeting virulence as a potential strategy for developing new anti-45 microbial drugs has been the focus of several research initiatives (4-11). In principle, suppressing 46 virulence behavior and locking pathogens in a vegetative non-biofilm forming lifestyle renders 47 them less infective and more susceptible to conventional antibiotics (4, 12). While some success 48 has been achieved against bacterial pathogens (6, 10, 13-19), less focus has been placed on 49 50 fungal infections which nevertheless continue to cause serious complications and mortality in 51 patients (8, 20-22). Indeed, despite the medical and economic damage caused by fungal biofilms, there remains an urgent and largely unmet need for the identification of compounds able to 52 specifically and selectively target and inhibit this mode of growth in clinically relevant fungal 53 pathogens (23). 54

The predominant nosocomial fungal pathogens, which include *Candida* spp., Aspergillus spp., 55 and Fusarium spp., are difficult to diagnose and cause high morbidity and mortality, even 56 following antifungal therapy (21). Candida albicans causes a variety of complications ranging 57 from mucosal disease to deep seated mycoses, particularly in immunocompromised individuals 58 59 (21, 24). Along with other fungal and yeast pathogens, C. albicans are known to form structured communities known as biofilms on medical devices either pre- or post-implantation leading to 60 recurring infections and in some cases death (25, 26). Once established in the biofilm phase, C. 61 albicans presents a significant clinical problem with current treatment options severely limited 62

by the intrinsic tolerance of fungal biofilms to antimycotics (20, 27, 28). Recent combination 63 therapies incorporating antibacterial and antifungal agents have provided some success (29). 64 However, as with all antibiotic based strategies, reports of resistance continue to emerge (27), 65 66 and biofilms themselves are considered a breeding ground for the emergence of antibiotic resistant strains, effectively hastening the onset of the perfect storm whereby the rapid decline in 67 68 new antibiotic production has been met by an equally rapid increase in multidrug resistant organisms (1). Thus the need to consider new anti-infective strategies that do not target essential 69 processes in the target organism. While blocking biofilms in these organisms' remains a major 70 71 clinical challenge (26, 30), exploiting our increased understanding of microbial signaling networks to control virulence and biofilm behavior is one innovative approach with significant 72 potential. 73

Many sites of infection are colonized by communities of mixed fungal and bacterial organisms, 74 and several layers of communication impact significantly on the dynamics and flux of these 75 76 populations (31, 32). For example, C. albicans is known to co-exist with Pseudomonas aeruginosa in the cystic fibrosis (CF) lung, and interkingdom communication between the two 77 organisms has previously been reported (16, 33). The Pseudomonas Quinolone Signal (PQS), 2-78 79 heptyl-3-hydroxy-4-quinolone, and its biological precursor 2-heptyl-4-quinolone (HHQ) are important virulence factors produced by *P. aeruginosa*. Structurally, POS and HHQ differ by the 80 presence of a hydrogen at C3 in HHQ and a hydroxyl group in PQS, giving rise to the increased 81 82 interest in modulating this position to assign biological function to the structure of these molecules (34-37). Previously, we have shown that HHQ, but not PQS, suppresses biofilm 83 formation of C. albicans (10). In response, C. albicans produces farnesol which has been shown 84

to modulate PQS production in *P. aeruginosa* (33). As both PQS and HHQ promote virulence
and pathogenicity of *P. aeruginosa* (38, 39), their utility as an anti-*Candida* treatment falls short
of being a viable anti-fungal treatment. However, the amenability of these small molecules to
chemical modification provides an opportunity to develop compounds with specificity of
function.

The transcriptional data and microscopic imaging described in this study have implicated 90 91 components of the cell wall as key factors in the response of C. albicans to P. aeruginosa alkylhydroxyquinolone (AHO) signaling. Furthermore, the biological activity of each class of 92 analogue in bacterial, fungal and host systems provides a new insight into the possible 93 94 interkingdom role of AHQs, particularly in a clinical setting such as the CF lung where all three systems co-exist. From a translational perspective, lead HHQ analogues were identified with four 95 key features: (1) potent anti-biofilm activity towards C. albicans, (2) selective non-cytotoxicity 96 towards mammalian cell lines, (3) non-agonistic and (4) potentially antagonistic to the virulent 97 pathogen P. aeruginosa. Several analogues retained the significant potency of the parent HHQ 98 99 molecule against C. albicans biofilm formation, whilst simultaneously becoming inactive in P. 100 *aeruginosa* quorum sensing. This suggests that these molecules have the potential to be further optimized for use as anti-Candida infectives without the concomitant limitation of P. aeruginosa 101 102 virulence augmentation.

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104 Materials and Methods

105 *C. albicans* stock maintenance and culturing conditions.

*C. albicans* strain SC5314 was sub-cultured from 15% (v/v) glycerol stocks at -80°C onto Yeast
Peptone Dextrose (YPD) medium [1% (w/v) yeast extract, 2% (w/v) peptone and 2% (w/v)
dextrose] and incubated at 30°C overnight.

### 109 *P. aeruginosa* stock maintenance and culturing conditions.

110 *P. aeruginosa* strains, PAO1 and  $pqsA^{-}$  mutant, containing the chromosomally inserted  $pqsA^{-}$ lacZ promoter fusion on plasmid pUC18-mini-Tn7, were maintained on Luria Bertani (LB) agar 111 plates, supplemented with carbenicillin (200 µg/ml) and X-gal (40 µg/ml), and incubated at 37°C 112 113 overnight. Single colonies were inoculated into LB broth (20 ml), supplemented with carbenicillin (200 µg/ml), and incubated at 37°C, shaking at 180 rpm overnight. For subsequent 114 experiments, the  $OD_{600nm}$  was recorded and a starting  $OD_{600nm}$  of 0.02 was inoculated into fresh 115 LB broth, supplemented with carbenicillin (200  $\mu$ g/ml) and incubated at 37°C, shaking at 180 116 117 rpm.

### 118 Structural modification of HHQ

The synthesis of HHQ, PQS (40, 41) and other HHQ-based analogues (36, 37) was carried out
via previously described methods. Novel compounds and compounds which required modified
syntheses are described *vide infra* and in the supporting information (Supplemental Data).

### 122 Thin Layer Chromatographic (TLC) analysis.

Silica TLC plates, activated by soaking in 5% (w/v)  $K_2HPO_4$  for 30 min were placed in an oven at 100°C for 1 hr (42). Analogues (5 µl, 10 mM) were spotted approximately 1 cm from the bottom. The spots were dried and the plate placed in a mobile phase comprising 95:5 dichloromethane:methanol. The plate was viewed under UV light when the mobile phase had run5 cm below the top of the plate.

### 128 Biofilm formation, quantification, and visualization.

C. albicans biofilm formation was carried out in 96 well plates, as previously described (43). 129 130 Seeding densities for all subsequent experiments (n=3) were  $OD_{600nm}$  0.05. Biofilm formation was measured as previously described using a semi-quantitative tetrazolium salt, 2,3-bis-(2-131 methoxy-4-nitro-5-sulfophenyl)-2H tetrazolium-5-carboxanilide inner salt (XTT) reduction assay 132 133 (44). Experiments were repeated at least three times, with at least eight technical replicates. Visualization of biofilm formation was performed on glass coverslips in 6-well plates using 134 Confocal Scanning Laser Microscopy (CSLM). All images were captured using the Zeiss HBO-135 100 microscope illuminating system, processed using the Zen AIM application imaging program 136 and converted to JPGs using Axiovision 40 Ver. 4.6.3.0. A minimum of three independent 137 biological repetitions were carried out. 138

### 139 Viable colony biofilm assay

*C. albicans* biofilms, supplemented with analogues and parent compounds, were grown in 6-well plates and incubated overnight at 37°C. Briefly, *C. albicans* Yeast Nitrogen Base (YNB) cultures were measured at OD<sub>600nm</sub>, diluted to 0.05 in YNB-NP supplemented with analogues, plated onto 6-well plates and incubated for 1 hr at 37°C. Media was removed, wells were washed twice with sterile PBS and supplemented with fresh YNB-NP with analogues. Plates were incubated overnight at 37°C after which media was removed and wells washed with sterile PBS. For serial dilutions, biofilms were cell-scraped into 1 ml PBS, vortexed, and serially diluted into sterile

PBS. Serial dilutions were plated (100 µl) onto YPD agar and incubating overnight at 37°C.
Colonies were counted and recorded the next day.

### 149 *C. albicans* growth curves

Overnight *C. albicans* cultures grown in YNB were diluted to 0.05 in YNB supplemented with
analogues. Cultures (200 µl) were added to each well of a 100 well plate and grown for a 24 hr
period on a Bioscreen C spectrophotometer (Growth Curves USA).

### 153 RNA isolation and qRT-PCR transcriptional analysis.

Overnight C. albicans cultures were diluted to 0.05 at OD<sub>600nm</sub> in either YNB or YNB-NP 154 155 (Difco). YNB cultures were supplemented with methanol whereas YNB-NP cultures were 156 supplemented with either 100  $\mu$ M HHQ or the methanol volume equivalent. Cultures were grown at 37°C with agitation (180 rpm) for 6 hr after which they were centrifuged at 4000 rpm, 157 supernatants discarded and pellets frozen at -20°C until processing. RNA was isolated using the 158 MasterPure Yeast RNA purification kit (Cambio Ltd, Cambridge UK) according to 159 manufacturer's specifications, and was quantified using a ND-1000 Spectrophotometer 160 161 (NanoDrop Technologies, USA). Genomic DNA was enzymatically removed using Turbo DNAfree DNase (Ambion), and samples were confirmed DNA free by PCR. RNA was converted to 162 cDNA using random primers and AMV reverse transcriptase (Promega) according to 163 164 manufacturer's instructions. qRT-PCR was carried out using the Universal ProbeLibrary (UPL) system (Roche) according to manufacturer's specifications, and samples were normalized to C. 165 166 albicans actin transcript expression (ACT1). A full list of primers and UPL probes used in this study is detailed in Supplemental Table 1. 167

P. aeruginosa strains were cultured as described above for 24 hr, with the addition of analogues 169  $(100 \mu M)$ , and pyocyanin was extracted as described previously (45). Cultures were centrifuged 170 171 at 4000 rpm for 10 minutes and the cell free supernatant (5 ml) removed. Chloroform (3 ml) was added, and mixed by vortex. After centrifugation at 4000 rpm for 5 min, the lower aqueous phase 172 was transferred to 0.2 M HCl (2 ml). Samples were mixed by vortex and centrifuged at 4000 rpm 173 174 for 5 min to separate the phases. An aliquot of the top phase (1 ml) was removed and spectrophotometrically analyzed at OD<sub>570nm</sub>. Phenazine production was calculated using the 175 following formula:  $OD_{570nm} \ge 2 \ge 17.072$  and the units expressed in  $\mu$ g/ml. 176

### 177 **Promoter fusion based expression analysis**.

Promoter fusion analyses were performed in a 96-well format, with  $\beta$ -galactosidase activity 178 measured as described previously (46). Briefly, overnight cultures of wild-type PAO1 pgsA-lacZ 179 (pLP0996) and mutant strain PAO1 pgsA<sup>-</sup> pgsA-LacZ were diluted to OD<sub>600nm</sub>=0.02 in LB. 180 Analogues at 100 µM final concentration were added, mixed, aliquoted into 96 well plates and 181 incubated overnight at 37°C with shaking. The next day, OD<sub>600nm</sub> values were recorded in a plate 182 reader. Aliquots of cells (0.02 ml) were permeabilized [100 mM dibasic sodium phosphate 183 184 (Na<sub>2</sub>HPO<sub>4</sub>), 20 mM KCl, 2 mM MgSO<sub>4</sub>, 0.8 mg/mL CTAB (hexadecyltrimethylammonium bromide), 0.4 mg/mL sodium deoxycholate, 5.4 µL/mL beta-mercaptoethanol] and added to 185 substrate solution [60 mM Na<sub>2</sub>HPO<sub>4</sub>, 40 mM NaH<sub>2</sub>PO<sub>4</sub>, 1 mg/mL o-nitrophenyl-β-D-186 Galactoside (ONPG), 2.7 μL/mL β-mercaptoethanol]. The kinetics of color development was 187 monitored and the reactions were stopped using 1M NaCO<sub>3</sub>. OD<sub>420nm</sub> were recorded as above. 188

189 Miller units were calculated using the following equation;  $1000 \times [OD_{420nm}/(OD_{600nm}) \times 0.02 \text{ ml}]$ 190 x reaction time (min)].

### 191 Cytotoxicity Assay.

Lactate dehydrogenase (LDH) release from a panel of mammalian cells was assayed as a 192 measure of cytotoxicity using an LDH colorimetric kit (Roche) according to manufacturers' 193 instructions (36). Briefly, IB3-1 lung epithelial cells, A549 human lung adenocarcinoma 194 epithelial cells, DU-145 human prostate cancer cells and HeLa cervical cancer cells were seeded 195 196 onto 96 well plates and treated with methanol (control) and analogues. Following 16 hr incubation at 37°C and 5% CO<sub>2</sub>, supernatants were removed and added to catalyst reaction 197 mixture in a fresh plate and further incubated at 37°C and 5% CO<sub>2</sub> for 30 min to allow for color 198 development. After this period, the plate was analyzed on an ELISA plate reader at  $OD_{490nm}$ . 199 Cytotoxicity was expressed as a percentage of cells treated with 0.1% (v/v) Triton (100% 200 cytotoxicity). 201

### 202 Statistical analysis.

All graphs were compiled using GraphPad Prism (version 5.01) unless otherwise stated. All data were analyzed using built-in GraphPad Prism (version 5.01) functions as specified. The level of significance was set at p = 0.05 (\*) and post-hoc comparisons between groups were performed using the Bonferroni multiple-comparison test.

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### 212 Results

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# 214 Key modifications of the quinolone framework retain anti-biofilm activity towards *C*. 215 *albicans*

216 The HHQ molecule has previously been shown to suppress biofilm formation in C. albicans at concentrations from  $10 - 100 \,\mu\text{M}$  (2.47 - 24.7  $\mu\text{g/ml}$ ) independent of any effects on the growth 217 of planktonic cells (10). Previous structure function analysis of the activity of the quinolone 218 framework had implicated the C-3 position as a key component of interspecies anti-biofilm 219 activity (36). We undertook further modification of the HHO parent molecule with the aim of 220 developing viable anti-biofilm compounds to target C. albicans. These compounds were 221 incorporated into a larger collection of alkylquinolone analogues, systematically modified at 222 223 different positions on the molecule and classified on the basis of their substitutions relative to the parent framework HHQ (Table 1). 224

The suite of analogues was first tested to establish their potency as anti-biofilm compounds 225 against C. albicans using an optimized XTT assay, a commonly used quantitative method to 226 assess Candida biofilm mass and growth (47). As previously described (10), HHQ significantly 227 suppressed biofilm formation when compared to untreated and methanol treated cells, whereas 228 PQS appeared to induce biofilm formation (Figure 1). When all analogues were similarly 229 screened by XTT assay, several had similar anti-biofilm activity to HHQ [1 and 2 (class I; 230 modified at C-3)], 3, 4, 6, 7, 9 (class II; modified anthranilate ring), and 12 (class III; modified 231 232 alkyl chain) (Figure 2a). These analogues were diverse members of classes I, II, and III

suggesting that several components of the HHO framework contribute to anti-biofilm activity of 233 the parent compound. A number of substitutions led to intermediate anti-biofilm activity, 234 including 5, 8, 10, 11, (class II; modified anthranilate ring) and 15 (class V; modified 235 236 anthranilate ring and alkyl chain length), while some analogues had completely lost the ability to suppress C. albicans biofilm formation e.g. 13 (class IV; modified C-4), and the class V 237 238 compounds 14, 16, and 17 (Figure 2a). While modification of the C-3 position to produce PQS led to loss of anti-biofilm activity (Figure 1), incorporation of an  $-N-NH_2$  moiety (2) at the 3-239 position or substitution of C-3 with NH (1), did not affect the ability to suppress C. albicans 240 241 biofilm formation (Figure 2a). Addition of Cl at the C6 and C8 positions of the anthranilate ring (6 and 7) also did not lead to loss of anti-biofilm activity. In contrast, the introduction of 242 considerable steric bulk with the addition of an *n*-hexyl alkyl chain at C6 of the anthranilate ring 243 (5), or elaboration of the aromatic group as with the naphthyl compound (8), resulted in 244 compounds with significantly less potent anti-biofilm activity relative to HHQ. These data 245 246 suggest an exquisite level of specificity for the interaction between HHQ and the C. albicans 247 biofilm intracellular machinery. Modification of the C2 alkyl chain from *n*-heptyl (HHQ) to *n*nonyl C9 (12) did not affect anti-biofilm activity, while parallel modification of the anthranilate 248 ring resulted in a complete loss, as with the Class V compounds 16 or 17. Modifying the C-4 249 position (C=O to C=S) (13), the quinolone thiol exhibited an increase in XTT activity (P<0.05) 250 relative to controls (Figure 2a), comparable to the increase observed in the presence of PQS 251 252 (Figure 1). Previously, we have shown that HHQ elicits a dose-dependent reduction in C. albicans biofilm formation (10). In order to determine if this also applied to the analogues that 253 retain anti-biofilm activity, dose response analysis of selected compounds 1, 2, 4, 6, and 12 254

representing classes I, II, and IV, was undertaken. This revealed compound specific responses, 255 with 10 µM of compounds 2, 4, and 12 being sufficient to elicit a statistically significant 256 reduction in biofilm formation (Figure 2b). All five compounds reduced biofilm formation when 257 258 applied at 50  $\mu$ M and 100  $\mu$ M. To further confirm the anti-biofilm activity of the lead compounds, viable colony counts were performed on selected analogues using the maximum 100 259 260 µM compound dose. This confirmed the outputs from the XTT assays; all analogues, along with HHQ, significantly reduced viable biofilm cells in comparison to the control (Figure S1a). 261 Importantly, the anti-biofilm activity was found to be independent of planktonic growth, which 262 263 was unaffected in the presence of selected compounds (Figure S1b).

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### 265 Microscopic staining reveals structural changes in *C. albicans* biofilms

The formation of biofilms in bacteria, yeast and fungi is a highly ordered process involving 266 267 multicellular behavior and has been defined in several stages (22). Confocal microscopy combined with intracellular staining was used to assess structural integrity and cellular 268 morphology of *C. albicans* incubated on coverslips. Biofilms were individually stained with each 269 270 of the dyes and multiple fields of view were visualized to accurately represent the effect of the analogues. The biofilm observed for methanol and untreated controls displayed all the 271 characteristics of a typical C. albicans biofilm and were classified as wild-type (Figure 1). 272 Calcofluor, Concanavalin A and FUN-1 staining revealed a uniform distribution of 273 274 chitin/cellulose and cell wall mannosyl/glucosyl residues indicative of viable wild-type 275 morphology (Figure 1). Those analogues identified by XTT assay as causing impaired biofilm

formation (1, 2, 3, 4, 6, 7, 9, 12, and 15) exhibited markedly disrupted structures when grown on 276 coverslips and were classified as atypical morphologies (Figure 3). Cells treated with class I 277 analogues were found to be largely compromised in their biofilm forming capabilities and were 278 279 classified as morphologically atypical. Biofilms produced with both 1 and 2 were significantly distorted, displaying a spindle-like phenotype. Hyphae were short in length, and predominantly 280 281 displayed yeast cell types rather than hyphal structures. Other structure disrupting analogues were from Class II, III and V, suggesting that specific modifications on the anthranilate ring and 282 alkyl chain variation do not significantly affect the anti-biofilm activity compared to the parent 283 284 compound (Figure 3).

Some analogues, including those that exhibited intermediate activity in the XTT assay, did not 285 alter the biofilm structure, with 5, 11, 13, 14 and 16 all placed into the wild-type morphology 286 287 group. Biofilms formed in the presence of 13 showed hyper-production of short hyphae, creating a dense mycelial network (Figure S2). The remaining analogues from Class II; 8 and 10 (Figure 288 289 S1) caused significant biofilm disruption with fragmented hyphae, stunted vegetative growth and considerably large cell debris fields. Cells incubated with the Class V molecule 17 induced a 290 severely compromised phenotype (Figure S2) where debris fields comprising yeast cells and 291 blastospores characterized the structural phenotype. 292

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294 Enhanced gene transcript expression of *HWP1*, *ECE1*, *ALS3*, *IHD1* and the 295 uncharacterized open reading frame *ORF* 19.2457 provides a molecular mechanism for 296 alkyl quinolone activity towards *C. albicans* 

In addition to providing new insights into the interkingdom relationship between these important 297 pathogens, there has recently been a strong emphasis placed on ligand receptor interactions and 298 the need to provide molecular mechanisms for the action of any potential therapeutic compound 299 300 (48). We previously implicated *TUP1* in the HHQ-mediated suppression of biofilm formation in C. albicans suggesting a role for the cell-wall in this interaction (10, 16). More recently, several 301 302 reports have shown changes in expression of cell-wall associated genes linked to biofilm formation in this organism (16, 20, 28, 49-51). These included a cohort of eight genes that are 303 proposed to constitute the core filamentous response network; namely ALS3, ECE1, HGT2, 304 305 HWP1, IHD1, RBT1, DCK1, and the gene of unknown function open reading frame orf19.2457 (51). Therefore, transcript expression of a cohort of genes implicated in cell wall biogenesis, 306 hyphal development, biofilm formation and other related functions that were previously shown to 307 308 be upregulated during the morphological transition from yeast to filamentous growth was investigated (Table S1) (16, 51). The housekeeping gene ACT1 was chosen for normalization 309 310 based on previous biofilm studies (52). We observed that several transcripts were hyperexpressed in a HHQ dependent manner; specifically, HWP1, ECE1, ALS3, IDH1 and the as yet 311 uncharacterized open reading frame (ORF) 19.2457 (Figure 4). The remaining transcripts 312 313 (CPH1, EFB1, ESS1, RBT1, TUP1, BCR1, DCK1, and HGT2) yielded expression patterns 314 similar to control cells (Figure S3). It was perhaps somewhat surprising that, while treatment of C. albicans with P. aeruginosa supernatants has previously been shown to downregulate 315 316 expression of the RBT1, RBT5 and RBT8 genes (16), expression of RBT1 was unaltered in the presence of HHO (Figure S3). Taken together, these data suggest that HHO induces a specific 317 subset of cell wall proteins in C. albicans. Further work is needed to identify the upstream 318

components of this response, although *in silico* screening of *C. albicans* genome sequences has
ruled out the presence of an obvious PQS receptor (unpublished data).

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### 322 Lead compounds display reduced cytotoxic activity towards specific mammalian cell lines

323 Evaluating the cytotoxicity of synthetic compounds is crucial in the context of developing targeted and highly optimized molecular therapeutics that are benign to human cellular 324 physiology and ideal for use in a clinical environment. In previous work, we showed that 325 326 analogue 1 was significantly less cytotoxic than HHQ, with an 80% reduction in LDH release relative to the parent compound (36). Therefore, the suite of analogues was tested for in vitro 327 cytotoxicity towards IB3-1 airway epithelial cells. Class I analogues exhibited reduced 328 cytotoxicity to IB3-1 cells with 2 displaying approximately 34% toxicity (Figure 5a). Several 329 class II analogues (4, 6, and 9) exhibited reduced cytotoxicity relative to IB3-1 cells treated with 330 HHQ, with 7 not reaching statistical significance. The class III analogue 12 was comparable to 331 HHQ. Of the analogues that did not retain anti-biofilm activity, 5, 8, 10 and 11 exhibited variable 332 333 cytotoxicity to IB3-1 cells whereas 13 exhibited considerably reduced cytotoxicity to IB3-1 cells 334 (Figure S4). Finally, 16 exhibited very low levels of cytotoxicity, while 17 was reduced relative to HHQ treated cells. Compound 15 was the most toxic killing approximately 91% of all cells 335 (Figure 5a). 336

In order to achieve a more comprehensive understanding of the selective toxicity of the lead compounds, several additional cell lines were tested (**Figure 5b**). LDH release assays were performed in A549, DU145, and HeLa cell lines in the presence of 100  $\mu$ M of the lead compounds revealed distinct cytotoxicity profiles, with **1** and **9** consistently proving the least cytotoxic of the compounds tested. Compounds **4** and **6** exhibited reduced cytotoxicity in DU145 cells (although not statistically significant) but were comparable to HHQ in both the A549 and HeLa cell lines, while compound **2** exhibited increased cytotoxicity relative to HHQ in DU145 cells (**Table 2**). These data suggest that cell-specific cytotoxicity analysis will need to be performed prior to the introduction of these compounds in an applied setting.

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### 347 HHQ analogues display a spectrum of agonist activity towards *P. aeruginosa* virulence.

Taken together, compounds 1, 4, 6, and 9 pass both the first and second criteria described above, 348 i.e. they retain anti-biofilm activity towards C. albicans while exhibiting reduced selective 349 cytotoxicity towards specific host cell lines. However, both HHO and POS are co-inducers of the 350 virulence associated LysR-Type Transcriptional Regulator; PqsR (41). The structural moieties 351 that underpin the interaction between HHQ/PQS and PqsR remain to be fully characterized, 352 although recent studies have reported diverse classes of PqsR antagonist (53-55), and implicated 353 the hydrophobic pocket situated within the PqsR protein (56). Therefore, in order to assess 354 whether the lead compounds could elicit a virulence response from *P. aeruginosa*, phenazine 355 356 production and *pqsA* promoter activity (57) were monitored in a *pqsA* mutant where the capacity to produce native HHQ and PQS had been lost. 357

Both HHQ and PQS restored phenazine production in the *pqsA*- strain (**Figure 6A**). In contrast, the majority of analogues did not restore phenazine production in this strain, with the notable exception of compound **9**. Several analogues from different classes did partially restore phenazine production in the mutant background, including 10, 12, and 17 (Figure 6 and Figure
S5). None of the analogues interfered with phenazine production in the wild-type PAO1 strain,
suggesting that they are ineffective as PQS antagonists (Figure S5).

Similarly, while some degree of PqsR agonist activity was observed in the presence of 364 compounds 6, 9, 10, 12, 13, and 17, only HHQ and PQS significantly induced promoter activity. 365 All other analogues did not influence promoter activity in this system (Figure 6B and Figure 366 367 **S5**). Somewhat surprisingly, antagonistic activity towards pqsA promoter activity was not observed, with almost all analogues failing to significantly suppress *pqsA* promoter activity in 368 the wild-type strain (Figure S5). The relative ineffectiveness of these analogues as PQS 369 370 antagonists may in part be due to hydroxylation of HHQ analogues (H at C3) to PQS analogues (OH at C3), thus establishing the non-antagonistic behavior explained by a recent report by Lu 371 and colleagues, where the action of PqsH rendered anti-PQS compounds ineffective through 372 373 bioconversion (55).

374

### 375 **Discussion**

Current antimicrobial therapies tend to be non-pathogen-specific and there is evidence to suggest that the availability of relatively non-toxic broad-spectrum therapies has contributed to the emergence of resistance among both targeted and non-targeted microbes (58, 59). Consequently, there is an urgent need to innovate new options for the targeted prevention of microbial infection while avoiding the inevitable emergence of resistance that is the hallmark of broad spectrum antibiotic therapies (59, 60). Increasingly, industry, academia and regulatory bodies have become interested in single-pathogen therapies to treat highly resistant or totally resistant bacterial pathogens, rightly viewed as an area of high unmet need (61-63). Exploiting interkingdom communication networks, and the mode of action of the chemical messages or signals employed therein, offers us a powerful platform from which to deliver on this.

386 Previously we had shown that the HHQ interkingdom signal molecule from *P. aeruginosa* could suppress biofilm formation in C. albicans at concentrations ranging from 10 - 100 µM (2.47 -387 388 24.7 µg/ml) (10). This suppression occurred independent of any growth limitation in planktonic cells, and morphogenesis on spider media was also found to be unaffected (10). The design and 389 subsequent analysis of a suite of analogues based on the core HHQ quinolone framework has led 390 391 to the identification of several lead compounds that retain anti-biofilm activity towards C. albicans but exhibit significantly reduced cytotoxicity towards IB3-1 epithelial cells when 392 compared with the parent HHQ molecule. The selective cytotoxicity of the lead compounds 393 together with the dose dependent anti-biofilm effects will be key considerations in determining 394 the cell line specific therapeutic index of lead analogues as part of the ongoing development of 395 396 these compounds. Furthermore, unlike HHQ, these lead compounds are now inactive towards the 397 P. aeruginosa PqsR quorum sensing system, a critical requirement for their potential future development as anti-biofilm therapeutics. In addition, the ability to generate hydrochloride salts 398 399 of the compounds ((36) and data not shown) suggests that solubility of future therapeutics based on these scaffolds will not be a bottleneck. Several strategies have been proposed for the 400 implementation of anti-biofilm compounds as clinical therapeutics to target C. albicans biofilm 401 402 infections (64). As the HHQ analogues possess anti-biofilm, but not anti-Candida activity, they would disrupt the formation of biofilms but not likely remove the planktonic cells that remain at 403 the site of infection. Therefore, combination with conventional anti-fungal compounds would be 404

required for effective clearance. Alternatively, where the potency of the anti-biofilm activity can
be synthetically enhanced through further derivatization, clearance by the immune system might
also be realistic.

408 The molecular mechanisms through which AHQs and the lead compounds identified in this study disrupt the formation of biofilms by C. albicans remains to be fully elucidated. Previously we 409 have shown that HHQ does not affect adhesion, but rather impacts directly on the subsequent 410 developmental stages in a TUP1-dependent manner (10). In this study we have shown that the 411 expression of several cell-wall associated genes is increased in response to HHQ during the 412 413 switch to hyphal growth. These genes have previously been implicated in the formation of C. albicans biofilms and have been shown to exhibit increased levels of expression during the 414 hyphal transition (50, 51, 65). Therefore, anti-biofilm compounds might be expected to suppress 415 416 this induction rather than enhance it. However, five of the target genes tested exhibited an increase in expression relative to control cells under inducing conditions. This may be a 417 418 reflection of the previous observation that HHQ interferes with the later stages of biofilm 419 development (10). Alternatively, this hyper-expression phenotype may affect the capacity of the cell to engineer a community based biofilm. Future studies will focus on elucidating the 420 421 pathways through which C. albicans perceives and responds to challenge with HHQ with the aim of identifying potential therapeutic targets. 422

Further work using defined *in vivo* models of biofilm and infection will be required to progress the development and evaluation of these small molecules as anti-biofilm compounds. Models are now available for the investigation of infections involving medical devices such as vascular catheters, dentures, urinary catheters, and subcutaneous implants, as well as mucosal biofilm infections (66). The ongoing development of cell-based or animal models to study *in vivo*infections (66-69), whether as single pathogen or co-culture systems (70), has provided a wellequipped tool-kit for the pre-clinical assessment of these AHQ-based compounds.

430

### 431 Conclusions

432 In this study, we have functionalized the important microbial signaling molecules HHQ and PQS in order to exploit their interkingdom role to control biofilm formation in C. albicans. In addition 433 to deciphering further insights into the molecular mechanism through which these chemical 434 435 messages elicit a biofilm suppressive response from C. albicans, the bioactivity of several lead compounds has provided a viable platform for the development of next generation therapeutics. 436 Crucially, some of these compounds are non-toxic to mammalian cells and have been rendered 437 incapable of activating *P. aeruginosa* virulence systems, thus highlighting their potential utility 438 as an effective therapy combatting human infection. 439

440

### 441 Author Contributions

FJR, GPM and FOG conceived and designed the investigation. FJR, JPP, LG, and DW
performed the biological experimentation, while RC, RS, and EOM conducted the chemical
synthesis. FJR, JPP and FOG wrote the manuscript and all authors read and edited the final draft.

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## **Table 1. Compound Data**

Compound	Structure	Yield [%] <sup>a</sup>	MW	Rf <sup>b</sup>	Class <sup>c</sup>
1	NH NH NH NH NH	76	244.3	0.319	Ι
2*	Nr NH <sub>2</sub>	3	259.3	0.907	I
3	MeO MEO MEO	34	273.4	0.252	П
4		31	257.4	0.286	II
5		35	327.5	0.504	Π
6	CI C	21	277.8	0.403	Π
7		19	277.8	0.630	Π
8	O N H H G	46	293.4	0.294	II

	o ci o				
9*	$CI \xrightarrow{O}_{H} \xrightarrow{V}_{H} \xrightarrow{V}_{6} \xrightarrow{V}_{6} \xrightarrow{CI}_{H} \xrightarrow{O}_{H} \xrightarrow{V}_{6}$	6	277.8	0.361	Π
10	Meo H H	33	273.4	0.261	Π
11	O We H OMe H H G	12	273.4	0.504	Π
12	C N N N N N N N N N N N N N N N N N N N	23	271.4	0.395	III
13*	SH N N H 6	48	259.4	1	IV
14		16	355.6	0.538	V
15	MeO HeO H H H H	28	301.4	0.286	V
16	C H H H	51	321.5	0.462	V
17	F H H H	16	289.4	0.504	V

- <sup>a</sup> % yields are isolated yields over all steps
- <sup>b</sup> TLC on silica plates with Dichloromethane:MeOH (95:5) mobile phase
- <sup>c</sup> Class I modified C-3; Class II modified anthranilate ring; Class III modified alkyl chain;
- 648 Class IV modified C-4; Class V modified anthranilate ring and alkyl chain.
- 649 \* New compounds synthesized in this study

IB3-1 ** *	A549 ***	DU145 **	HeLa ****
		**	****
*			
	**	*	*
*	*	*	*
**	****	****	****
*	****	**	****
**	***	*	****
*	**	*	**
**	***	**	****
	* **	* **** ** *** ** **	* *** ** ** *** * ** *** * * ** *

# **Table 2. Selective Toxicity Index of Lead Compounds**

**Figure 1.** *C. albicans* **biofilms are altered in the presence of HHQ**. Filamentous *C. albicans* biofilms grown in the presence of PQS and HHQ (100  $\mu$ M) were assessed structurally by confocal microscopy and metabolically using the XTT biofilm assay. Data (means  $\pm$  SEM) are representative of three independent biological experiments and are presented relative to the untreated control. Two-tailed paired student's t-test was performed by comparison of *C. albicans* in the presence of HHQ and PQS with *C. albicans* treated with methanol or ethanol (\*, p-value  $\leq$ 0.05).

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Figure 2. Decoration of HHQ exhibits variable biofilm activity against C. albicans. (a) A 677 panel of HHO derivatized analogues incubated with filamentous C. albicans and screened for 678 biofilm formation using the metabolic XTT biofilm assay. Data is presented as OD<sub>492nm</sub> 679 spectrophotometric output normalized to the untreated control, and is representative of at least 680 three independent biological replicates, with error bars representing SEM. (b) Dose-dependent 681 XTT analysis of selected anti-biofilm compounds applied at 10, 50, and 100 µM. Data is the 682 average of at least two independent biological replicates, each constituting eight technical 683 684 replicates. Statistical analysis of both datasets was performed by one-way ANOVA with Bonferroni corrective testing, and is presented relative to the MeOH control; \*  $p \le 0.05$ , \*\*  $p \le$ 685 0.01 and \*\*\*  $p \le 0.001$ . 686

Figure 3. Microscopic analysis reveals altered biofilm structures. Analogues that lead to reduced *C. albicans* biofilm formation in the XTT assay (1, 2, 3, 4, 6, 7, 9, 12, and 15) exhibit compromised biofilm structures. Filamentous *C. albicans* biofilm in the presence of analogues (100  $\mu$ M) was stained for chitin and cellulose (calcofluor; blue), lectins which binds to sugars, glycolipids and glycoproteins (concanavalin A; green) and live-dead cells (FUN-1; red).

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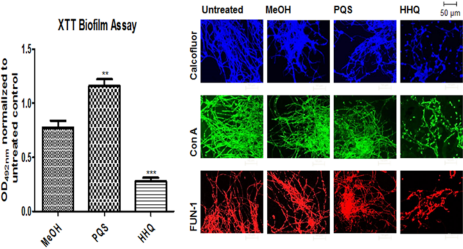
**Figure 4. Hyphal pathway genes are hyper-expressed in response to HHQ.** Transcript expression analysis (Real Time RT-PCR) of a panel of biofilm genes was assessed in *C. albicans* grown in YNB-NP (filamentous inducing media) in 100  $\mu$ M HHQ for 6 hours at 37°C. All data was normalized to a housekeeper gene (*ACT1*). Error bars represent SD of three independent biological replicates. Two-tailed paired student's t-test was performed by comparison of HHQ treated cells with methanol control in YNB-NP inducing medium (\*, p-value  $\leq 0.05$ ).

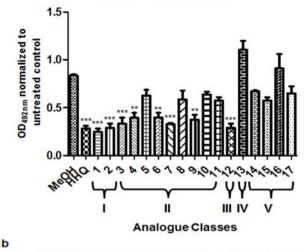
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Figure 5. Cytotoxicity towards specific mammalian cell lines is reduced in lead compounds. (a) Cytotoxicity, measured as a percentage of total lactate dehydrogenase (LDH) released from IB3-1 cells treated with 0.1% Triton X-100 (100% cytotoxicity), was significantly reduced in the presence of several lead compounds. Data (means  $\pm$  SEM) are representative of three independent biological experiments. (b) Selected lead compounds were tested against A549, DU145, and HeLa cell lines. Data represents four independent biological replicates and all datapoints are normalized to Triton X-100 as above. A one-way ANOVA was performed with Bonferroni corrective testing on all datasets and comparison relative to MeOH control is presented; \*  $p \le 0.05$ , and \*\*\*  $p \le 0.001$ .

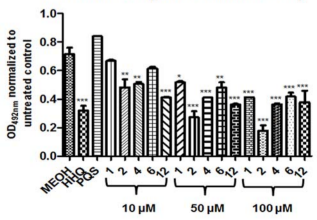
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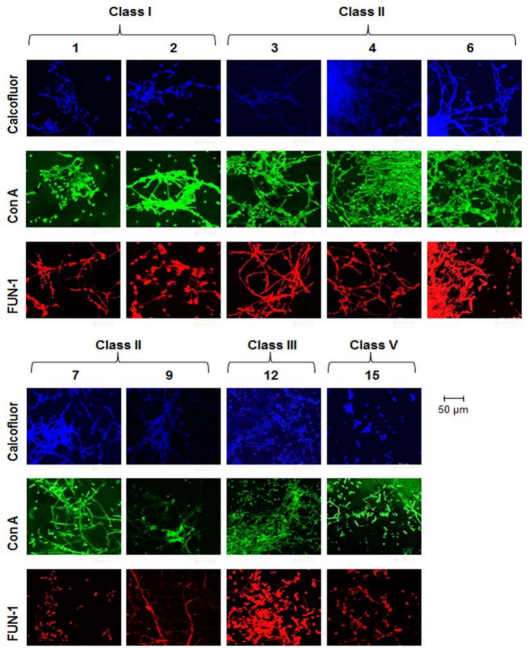
Figure 6. Influence of HHQ analogues on PQS-dependent virulence phenotypes in *P*. *aeruginosa*. (a) Phenazine production and (b) pqsA-lacZ promoter activity quantified in a PAO1 *pqsA* mutant in the presence of HHQ, PQS and lead compounds. Data is presented as mean +/-SEM and is representative of at least three independent biological replicates. A one-way ANOVA was performed with Bonferroni corrective testing and statistical significance relative to the MeOH control is presented; \*\* p  $\leq$  0.01 and \*\*\* p  $\leq$  0.001.

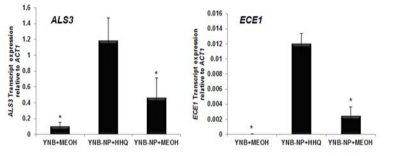


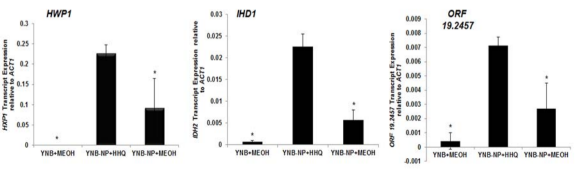


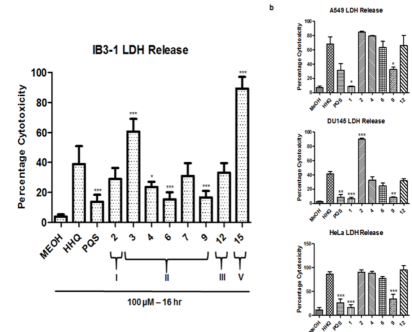




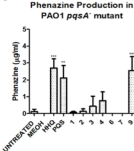








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