Clearing the Fungal FoG: Perseverance, a property distinct from resistance, is associated with clinical persistence

Alexander Rosenberg¹*, Iuliana V. Ene²*, Alon M. Dahan¹, Ella Shtifman Segal¹, Arnaldo L. Colombo³, Richard J. Bennett², Judith Berman¹**

1. School of Molecular Cell Biology and Biotechnology, Tel Aviv University, Ramat Aviv, Israel
2. Department of Molecular Microbiology and Immunology, Brown University, Providence, Rhode Island, USA
3. Department of Medicine, Federal University of São Paulo, São Paulo, Brazil

*Equal contributions
**Corresponding Author: judithberman11@gmail.com, +972-52-584-2031
Abstract

Drug resistance, defined by the minimal inhibitory concentration (MIC), often does not predict whether fungal infections will respond to drug treatment in the clinic. Here we define and quantify an antifungal response, termed ‘perseverance’, that correlates with clinical outcomes in patients treated with fluconazole, the most widely used antifungal drug. Perseverance is defined by the ability of fungal cells to grow at drug concentrations above the MIC, and is measured either as the ‘fraction of growth’ (FoG) in drug disk diffusion assays, or as the degree of ‘supra-MIC growth’ (SMG) in broth microdilution assays. Perseverance is distinct from resistance and is related to the proportion of cells that form colonies at supra-MIC drug concentrations. Higher perseverance correlates with shorter lag times at high drug concentrations, highlighting a critical distinction from ‘tolerance’ in bacterial populations. Importantly, several adjuvant drugs, including inhibitors of Hsp90 and calcineurin, eliminate perseverance without altering the MIC. We also find that perseverance and resistance are sensitive to mutations in different pathways, underscoring the distinct nature of the two phenomena. Analysis of isolates recovered from clinically persistent or non-persistent infections in immunocompetent patients reveals that persistent isolates show higher levels of perseverance than isolates cleared by antifungals. Thus, perseverance is an intrinsic property of fungal isolates that correlates with the success or failure of treatment in the clinic, and may provide a useful parameter for predicting clinical persistence and choosing appropriate antifungal therapies.
Introduction

Antimicrobial drug resistance, the ability to grow well at high drug concentrations irrespective of treatment duration, involves several distinct mechanisms including alteration in drug targets, drug efflux or drug inactivation. Of note, infections generally follow the “90/60” rule for predicting therapeutic outcomes based on in vitro susceptibility testing: approximately 90% of susceptible isolates and 60% of resistant isolates respond to therapy\(^1\text{-}\!^5\). This is important as it implies that drug resistance, as defined by in vitro assays, is often not sufficient to explain clinical outcomes, and that effective treatment is likely influenced by patient attributes as well as by additional characteristics of the infecting isolates.

Two microbial variables that have been established to affect drug responses in bacteria are “tolerance” and “persistence”. Importantly, both tolerant and persistent isolates require similar levels of antibiotic drugs for killing as susceptible isolates. However, the duration of drug treatment necessary to achieve killing of tolerant or persistent strains is much longer than that necessary for susceptible strains. Mechanistically, both tolerance and persistence involve the ability of cells to survive and/or grow during transient exposure to high drug concentrations.

Toxin/antitoxin-modules are responsible for promoting tolerance and persistence in some cases, and isolates with more toxin/antitoxin modules therefore tend to exhibit higher levels of persistence\(^6\). A major distinction between the two phenomena is the size of the subpopulations involved: tolerance is observed in 10-90% of cells whereas persistence occurs in <1% of cells (reviewed in\(^7\)). Both tolerance and persistence typically involve slower growth or a longer lag phase, which effectively reduces metabolism and toxicity of the drug in the exposed cells. This is reflected in the minimum duration for killing (MDK), a quantitative parameter which is useful for studying cidal, but not static, drug responses\(^7\text{-}\!^9\).

The term “tolerance” has been discussed for fungal isolates, and is loosely defined as survival above intrinsic inhibitory concentration levels. Currently, the most widely used antifungal drugs are azoles, which interact with sterols in the fungal membrane. Azoles are
primarily fungistatic in *Candida albicans*, one of the most prevalent human fungal pathogens.

With respect to fungistatic drugs, tolerance is likely synonymous with “trailing growth”, defined as the residual growth of cells at high antifungal concentrations in liquid cultures, and which has been detected in a subset of clinical isolates. Tolerance has been proposed to promote the appearance of ‘persister cells’ that can cause recurrent infections, although this has not been demonstrated. Strains that showed evidence of trailing growth did not have a clear difference in virulence in a systemic model of candidiasis or in short term responses to antifungal treatments in HIV patients. It is currently unclear whether trailing growth impacts the clinical outcome of infections by *Candida* species, yet 25-60% of isolates show some evidence of trailing growth. However, conventional susceptibility assays have been optimized to avoid detection of trailing growth by determining the MIC at 18-24 h, at which time the trailing effect is minimal or absent. Nonetheless, we posit that both tolerance and trailing growth yield increased numbers of cells dividing for long periods of time in the presence of antifungals, and can potentially alter the ability to adapt in the patient, a property termed “clinical persistence”.

*C. albicans* is a diploid species that lacks meiosis and thus relies on mitotic processes and a parasexual cycle for genetic variation. Each individual largely carries their own strain, often inherited from their mother, with isolates from different patients exhibiting broad divergence illustrated by thousands of unique SNPs. There are currently only four classes of approved antifungal drugs, and combination therapy using antifungals together with adjuvant drugs is considered a promising route to extend the effectiveness of current antifungals.

Resistance to fluconazole, the most widely used fungistatic drug, is an increasing problem in the clinic and is generally driven by increased drug efflux and/or changes in ergosterol biosynthesis, the target ofazole drugs. A broad range of adjuvant drugs have been reported to enhance the activity of fluconazole, with inhibitors of Hsp90 and calcineurin the most prominent among them. Targeting Hsp90, a highly abundant chaperone, or its client protein calcineurin, has
been proposed as a “powerful, broadly effective therapeutic strategy for fungal infectious disease”\(^{54}\).

In the clinic, persistence refers to a subset of infections that do not respond to drug treatment despite the infecting isolate being drug susceptible when evaluated in vitro. Host immune status may play a key role in determining the persistence or recurrence of fungal infections\(^{59-61}\). However, in the absence of host immune deficiencies or drug resistant isolates, it is unclear to what extent the intrinsic properties of fungal isolates impact clinical persistence.

Strain diversity is an important parameter that is often overlooked in laboratory studies of antifungal responses. There are multiple major and minor clades of *C. albicans*\(^{35-37}\) and some exhibit clade-specific levels of susceptibility to specific antifungals\(^{38}\). Clinical persistence of *C. albicans* has been observed in many patients, with particularly high incidence rates in neonates\(^{62-64}\). The inability to clear infection is associated with increased mortality; in one study the case-fatality rate was 54\% among infections with persistent candidemia and only 3\% among those with non-persistent candidemia\(^{65}\). The mechanisms underlying *C. albicans* clinical persistence have yet to be defined and it is not understood how fungal infections can persist in the bloodstream despite having MIC levels that indicate they should be effectively cleared by antifungal treatment.

We set out to characterize drug-response variables that could influence clinical persistence of fungal infections and be generalizable across all *C. albicans* clades. This work utilized an image analysis pipeline, *diskImageR*, for quantitative examination of disk diffusion assays (DDAs)\(^{66}\). This approach examines both the radius (RAD) of the zone of inhibition (ZOI), a parameter that correlates with the MIC, as well as the fraction of growth (FoG) within the ZOI, a parameter that correlates with ‘trailing growth’ or ‘tolerance’\(^{10,15,67-70}\). We emphasize that FoG is an independent variable that does not correlate with MIC\(^{66}\) or with *C. albicans* clade distinctions\(^{23}\). We characterized FoG in disk diffusion assays and its equivalent, supra-MIC growth (SMG), in broth microdilution assays (BMDAs), and demonstrated that the two
parameters correlate with one another and that they provide a measure of what we term ‘perseverance’. Perseverance correlates with the proportion of cells in the population that can grow at high drug concentrations. Strains with high perseverance show a shorter lag phase following drug exposure, in contrast to what is observed for bacterial tolerance to cidal drugs. Surprisingly, inhibitors of Hsp90 and of calcineurin eliminated perseverance completely but did not affect the MIC in any of the >50 isolates tested. Importantly, we found that perseverance was significantly higher in clinically-persistent isolates of *C. albicans* than in those readily cleared by fluconazole treatment, thereby providing support for the idea that perseverance is a parameter with clinical implications.

**Results**

**Slow, supra-MIC growth is distinct from resistance**

*C. albicans* isolates display high genotypic and phenotypic variability, including distinct responses to antifungal drugs. We examined the responses of isolates from different genetic backgrounds and types of infections to some of the most frequently used antifungals drugs using *diskImageR*, a quantitative disk diffusion assay analysis tool. *diskImageR* can quantify two parameters: the radius of the zone of inhibition (ZOI) at 20% inhibition (RAD20) as an indicator of susceptibility/resistance; and the fraction of growth within the ZOI (FoG20), calculated as the area under the curve relative to the maximum possible area (Fig. 1A). A screen of 219 clinical isolates (Supplementary Table 1), revealed that FoG levels on fluconazole (FLC) range from 0.10 to 0.85 and do not correlate with RAD20 levels (Fig. 1B,C), indicating that RAD and FoG measure two independent responses to this drug.

We selected a set of seven isolates to represent the breadth of diversity among clinical isolates for further characterization of FoG and the growth parameters associated with it (Fig. 1C, colored data points). We note that for drug-resistant strains such as T101 (MIC = 64, Supplementary Fig. 2A), the very small ZOI makes FoG measurements less accurate. For a
given isolate, both FoG and RAD levels were reproducible when evaluating cells taken from different colonies or when cells were taken from inside or outside the ZOI and re-tested for their drug responses (Fig. 1D). These observations indicate that FoG, the level of growth inside the ZOI, is a stable and heritable property of a given \textit{C. albicans} strain. Moreover, cells growing within the ZOI are not drug resistant and yield progeny indistinguishable from the rest of the cells in the population.

FoG and RAD levels exhibited different dynamics in the disk diffusion assays. While RAD levels were constant after 24 h, FoG was often undetectable at 24 h and only became fully apparent at 48 h (Fig. 1E). Thus, the detection of RAD is generally time-independent while the detection of FoG is largely time-dependent. By definition, drug susceptibility is a concentration-dependent parameter and, as expected, when drug concentration in the disk was increased, the ZOI size (and thus RAD) also increased (Fig. 1F, Supplementary Table 1). In contrast, FoG levels were similar irrespective of the drug concentration in the disk, indicating that this drug response is concentration-independent. Taken together, these results highlight the distinctive nature of the drug response measured as FoG, which is more reminiscent of the clinically ignored trailing effect, and is different from classical MIC measurements.

A range of FoG levels also was detectable with different antifungal drugs including other azoles (posaconazole and ketoconazole), 5-fluorocytosine and, albeit to a lesser degree, echinocandins (micafungin and caspofungin) and polyenes (amphotericin B) (Supplementary Fig. 1A). Overall, FoG levels were generally lower in response to drugs considered to be fungicidal (e.g., echinocandins and polyenes) rather than fungistatic (e.g., azoles, 5-fluorocytosine). We also measured drug responses in different fungal species including \textit{C. tropicalis}, \textit{Candida krusei}, \textit{Candida glabrata}, \textit{Candida parapsilosis} and \textit{S. cerevisiae}, resulting in a range of RAD and FoG levels to each of the antifungals tested (Supplementary Fig. 1B). \textit{C. parapsilosis} exhibited the highest susceptibility to azoles (highest RAD) with intermediate FoG levels. In response to azoles, \textit{C. glabrata} exhibited the highest FoG levels (but intermediate
Figure 1

A. Diagram showing the relationship between RAD and FoG. The graph on the right shows the range of FoG in Callicans isolates.

B. Scatter plot showing the correlation between RAD and FoG.

D. Illustration of parental, Inside ZOI, and Outside ZOI with their respective RAD and FoG values.

E. Graphs showing pixel intensity over time (24h and 48h).

F. Graphs showing RAD and FoG for different FLC concentrations (25 µg, 50 µg, 300 µg).
Figure 1. Measuring drug responses of *C. albicans* clinical isolates in disk diffusion assays (DDAs).

(a) Illustration of *diskImageR* analysis, which measures pixel intensity corresponding to cell density, along 72 radii every 5°. The average radius (RAD) threshold is the distance in mm corresponding to the point where 20%, 50% or 80% growth reduction occurs (light, medium, dark blue dots). The Fraction of Growth inside the ZOI (FoG) is the area under the curve (red hatched area) at the RAD threshold level, divided by the maximum area proscribed by the RAD threshold (max). (b) Range of FoG levels in 219 *C. albicans* clinical isolates (from Dr. R. Ben-Ami). Red, blue and green lines estimate high, medium and low levels of FoG respectively. Unless otherwise specified, DDAs were performed using a single 25 µg fluconazole disk analyzed after 48h at 30°C. (c) Comparison of FoG<sub>20</sub> and RAD<sub>20</sub> for the 219 isolates (*P* < 0.0064, n=219). (d) RAD<sub>20</sub> and FoG<sub>20</sub> for a parental strain and progeny colonies picked from inside and outside the ZOI. Left panel, DDAs for strain SC5314, and corresponding RAD (middle, black outlined dots) (One way ANOVA, *P* < 0.99, n groups = 3, *F* < 0.0032) and FoG (right, dots without outlines) (One way ANOVA, *P* < 0.97, n groups = 3, *F* < 0.029) levels for diverse *C. albicans* isolates (corresponding to indicated colors)

(e) Effect of incubation time on RAD and FoG values. Left panel, illustration of *diskImageR* analysis at 24 h and 48 h; corresponding RAD (middle panel) (Paired t-test, *P* < 0.094, t=2, df=6, n pairs-7) and FoG (right panel) levels (Paired t-test, *P* < 0.0071, t=4, df=6, n pairs-7) for strains as in (d)

(f) RAD is concentration-dependent and FoG is concentration independent as measured with disks containing increasing concentrations of fluconazole (25, 50 and 300 µg fluconazole) for strain SC5314 (left panel); corresponding RAD (middle panel) and FoG (right panel) levels for strains as in (d).
RAD), followed by *C. tropicalis* and *C. krusei*. We posit that the high FoG levels of *C. glabrata* could contribute to the high incidence of azole resistance and to the intrinsic low susceptibility of the species to this class of antifungals\textsuperscript{71-73}. Most *S. cerevisiae* strains displayed low FoG levels in response to azoles, but had intermediate FoG levels on echinocandins and polyenes. Clearly, FoG is a property observed in a range of yeast species and is not specific to fluconazole.

Broth microdilution assays (BMDAs) are considered the clinical assay of choice for quantifying drug susceptibility. We therefore measured MIC\textsubscript{50} values, the lowest drug concentration that inhibits 50% of growth, in the seven *C. albicans* isolates chosen above. Trailing growth is frequently observed in BMDAs as increased growth at 48 h relative to 24 h, but this phenomenon is not usually quantified or otherwise used in clinical assays. We quantified supra-MIC growth (SMG) in BMDAs as the average growth per well (at 48 h) above the MIC\textsubscript{50} (at 24 h) normalized to growth levels in the absence of drug (Fig. 2A). Using this parameter, most isolates exhibited growth at drug concentrations above the MIC, while the MIC levels did not change between 24 and 48 h. Thus, SMG provides a parameter that quantifies the trailing growth effect and is independent of resistance/MIC. Furthermore, SMG, like FoG, is dependent upon longer assay times and is more readily detectable at 48 h rather than 24 h (Fig. 2B,C). Similar to FoG, cells taken from wells above or below the MIC\textsubscript{50} were indistinguishable from parental cells when re-analyzed for MIC and SMG levels (Supplementary Fig. 2A).

Analysis of the relationship between parameters measured in BMDAs and DDAs revealed that SMG and FoG correlate well ($R^2 = 0.82$, $P < 0.01$, n – 12, Supplementary Fig. 2B), whereas there was no relationship between SMG and MIC ($R^2 = 0.25$, $P < 0.01$, n – 28 Supplementary Fig. 2B). \textit{This indicates that FoG and SMG reflect similar features of growth at supra-MIC antifungal concentrations, and that this response is independent from drug resistance.}

Of note, BMDAs cannot detect the individual sub-populations that are detectable on agar
Figure 2. Measuring drug responses of *C. albicans* clinical isolates in liquid broth assays. (a) Illustration of MIC and Supra-MIC Growth (SMG) calculations. MIC values were calculated at 24 h as the fluconazole concentration at which 50% of the growth in the absence of fluconazole is inhibited. Supra-MIC Growth (SMG) was calculated as the average growth per well above the MIC divided by the level of growth in the well without drug. Fluconazole was used in two-fold dilutions (0, 0.125, 0.25, 0.5, 1, 2, 4, 8, 16, 32, 64, and 128 μg/ml). (b) Heatmaps illustrating OD$_{600}$ levels for concentrations above the MIC (yellow bar) in cyan and beyond the MIC in yellow for the same seven *C. albicans* isolates from Fig. 1. Maps show OD$_{600}$ at 24 and 48 h. (c) Corresponding MIC (t-test, P=1, t=0, df=6, n pairs=7) and SMG (t-test, P=0.0356 ,t=2.7, df=6, n pairs=7 ) levels for these strains measured at 24 and 48 h.
medium and thus are more prone to jackpot events (overrepresentation of rare mutants with a selective growth advantage).

**Growth characteristics of FoG and SMG**

Different environmental cues that influence trailing growth include temperature, pH, carbon source and the type of surface cells are growing on\(^{25,26,67}\). Similarly, we found that FoG and SMG levels were influenced by different growth conditions. For example, FoG and SMG levels were reduced significantly at pH 4.5 relative to pH 7, albeit to different degrees in different strains (Supplementary Fig. 3A,B). Most clinical MIC assays are performed at 37°C, while laboratory assays are often performed at lower temperatures (30°C) to avoid confounding morphologies such as filamentous growth. Temperatures above 30°C had little effect on RAD and MIC levels but had a significant impact on both FoG and SMG, and this effect was dependent on strain background (Supplementary Fig. 3C,D). *C. albicans* isolates were very sensitive to the shift from 30°C to 37°C, but did not display significant differences in either FoG or SMG between 37°C and 39°C/41°C (Supplementary Fig. 3C,D), suggesting that the shift from normal body temperature to febrile temperature does not necessarily alter drug responses. In addition, DDAs performed using rich medium (YPD) often gave slightly different FoG and RAD levels than those using casitone medium (Supplementary Fig. 3E). Nonetheless, the relative differences between the degree of FoG in different *C. albicans* strains was generally maintained between the two media (Supplementary Fig. 3E).

**Growth properties of cells withstanding supra-MIC drug concentration**

To characterize the growth properties of cells underlying the FoG/SMG response, we measured the growth of individual colonies inside the ZOI. All strains exhibited nearly 100% viability in the presence of drug relative to growth in the absence of drug despite having a broad range of FoG/SMG levels (Fig. 3A). This result is consistent with the fungistatic nature of fluconazole and illustrates that neither FoG nor SMG is a measure of cell viability.
We next measured the proportion of the population that formed colonies at supra-MIC drug concentrations. The number of colonies that grew on 10 µg/ml fluconazole (a concentration 10-20-fold above the MIC of the average strain) ranged between 25% and 98% and correlated well with FoG levels on DDAs (Fig. 3B, R² = 0.91, P < 0.01, n=11). This indicates that FoG is due to having a relatively large subpopulation of cells that is able to grow, albeit slowly, at supra-MIC drug concentrations.

We turned to time-lapse microscopy to follow the dynamics of early cell growth in the presence of 10 µg/ml fluconazole. As expected from previous studies, exponentially growing cells plated onto supra-MIC drug concentrations did not immediately stop growing in the presence of fluconazole. Rather, the majority of cells continued dividing for ~4-6 h, and then divided more slowly to different degrees (Fig. 3C and Supplementary videos 1-3). Isolates with low FoG (e.g., P87) generally formed fewer growing microcolonies over a 24 h period than those with high FoG (e.g., SC5314, Fig. 3C) (similar numbers of cells were plated in each well). Thus, analysis of early cell growth, before colonies were detectable on plates, reflected the two trends seen with assays that measure visible colony growth: higher FoG isolates had shorter time of colony appearance (Fig. 3E) and greater number of cells growing on supra-MIC drug concentrations (Figs. 3B).

We next measured the dynamics of colony growth in the presence of supra-MIC drug concentrations (10 µg/ml fluconazole) using ScanLag, an assay that measures growth parameters with a flatbed scanner and colony size image analysis (Fig. 3D). ScanLag records the initial time of colony appearance (ToA, a proxy for lag phase length), the area occupied by light pixels (a proxy for colony size), and the change in this area over time (a measure of growth rate). Studies of tolerance in bacteria with ScanLag revealed that tolerant strains have longer ToA (lag times), a property thought to facilitate their survival. We adapted ScanLag for use with C. albicans and applied it to a series of isolates spanning a range of FoG/SMG levels (Fig. 3E and Supplementary Fig. 4). In the absence of drug, ToA values differed across the set of
isolates, indicating that ToA is a strain-specific parameter which reflects the genetic diversity of the isolates used. As expected, ToA was longer in the presence of supra-MIC drug concentrations (10 µg/ml fluconazole) than in medium without drug, except for truly resistant strains (Fig. 3E). High FoG isolates formed visible colonies ~10 h faster than low FoG isolates (Fig. 3E). This suggests that high FoG strains can overcome the inhibitory pressures of the antifungal more efficiently than low FoG strains. Of note, ToA was the only ScanLag parameter found to be associated with FoG; neither colony growth rate nor final colony size correlated with FoG/SMG levels (Fig. 3F and Supplementary Fig. 5A,B).

We performed a parallel analysis examining perseverance in two additional lineages that included strains with differences in growth rates (Supplementary Table 1). Thus, we tested two faster growing derivatives of the slow-growing isolate P37005 that were obtained via passaging\textsuperscript{77}, as well as heterozygous and homozygous null mutants of the Clb4 mitotic cyclin that result in slower growth\textsuperscript{78}. Each of these strains were compared on DDAs using fluconazole. We found no correlation between FoG and doubling time or between FoG and lag duration for either of the two lineages (Supplementary Fig. 5B). Thus, the two parameters that correlate with higher FoG values are (1) the proportion of cells that form colonies in the presence of drug (Fig. 3B), and (2) the more rapid appearance of colonies in the presence of drug, presumably reflecting shorter lag periods (Fig. 3E).

Taken together, these results indicate that two aspects of growth dynamics contribute to differences between low and high FoG isolates: high FoG strains have larger subpopulations that grow to form visible colonies at supra-MIC drug concentrations, and high FoG strains also form colonies more rapidly than strains with low FoG levels. This contrasts with tolerant bacterial cells with longer lag or quiescent periods that enable survival during bactericidal drug exposure. The distinctive growth properties of perseverant cells measured here as FoG/SMG in cells exposed to fungistatic drugs, and those measured for bacteria tolerant to cidal drugs indicate that mechanisms mediating drug responses are different in the two kingdoms. For
Figure 3

A. Drug disk diffusion assay

B. Replica plating

C. FoG

D. FLC (10 µg/ml)

E.cell clones that continue dividing

F. cell clones that have stopped dividing

G. FoG

H. cell clones that have stopped dividing

I. FoG

J. no drug

K. FLC (10 µg/ml)

L. no drug

M. FLC (10 µg/ml)
Figure 3. Growth analysis of cells growing within the ZOI (above MIC concentration). (a) Cells that grow within the ZOI are viable, as seen by replica plating of disk diffusion assays grown on casitone without fluconazole and incubated at 30°C for 48 h. (b) The degree of FoG for a given strain correlates with the proportion of colonies that are able to grow on 10 µg/ml fluconazole relative to growth on plates without fluconazole (n=11, P<0.001). (c) Microcolony analysis at supra-MIC concentrations of fluconazole (10 µg/ml). Black triangles indicate cells that continued dividing and black circles indicate cells that slowed or stopped dividing, respectively, over 24 h. Time lapse videos of these cells are available as Supplementary videos 1-3. FoG correlates with faster time of colony appearance. (d) Schematic of ScanLag analysis that measures time of colony appearance (ToA), colony growth rate and colony size using desktop scanners. (e) Time of Appearance of colonies on medium without drug (blue) or with 10 µg/ml fluconazole (red) for resistant isolate T101, and isolates with different FoG levels. ~500 cells were plated per plates with or without 10 µg/ml fluconazole and scanned every 30 min for 96 h. Histograms show the number of colonies that appeared (y-axis) at each time point (x-axis). Data for additional isolates is presented in Supplementary Figure 4. (F) Correlation between FoG values and the difference (Δ) in the time of appearance of colonies in the presence vs absence of fluconazole (ΔToA = ToA with 10 µg/ml fluconazole – ToA without fluconazole). (n=7, P=0.0027)
example, tolerant bacterial cells do not grow within the ZOI of a DDA unless the drug disk is
replaced with additional fresh medium, while perseverant cells, by definition, appear within the
ZOI in DDAs even in the presence of drug. It is also likely that bacterial tolerance and fungal
perseverance reflect different strategies of growth in cidal vs. static drugs. To highlight this
distinction we chose a new term, *perseverance*, to describe the ability of subpopulations of
fungal cells to grow in the presence of supra-MIC drug concentrations.

**Clinical persistence is associated with higher levels of perseverance**

Fluconazole is often the first line of therapy for treating systemic *C. albicans* infections and is also the major antifungal used in third world countries. However, treatment with fluconazole often fails despite isolates being susceptible to the drug when tested in vitro. We obtained sets of clinical isolates that were either efficiently cleared by fluconazole treatment (non-persistent isolates) or that persisted in the host despite extended fluconazole therapy.

Clinical persistence is defined here as two or more blood cultures positive for *C. albicans* on one or more days apart despite antifungal therapy. Non-persistent isolates (n = 7) were cleared from the bloodstream soon after fluconazole treatment was initiated. In contrast, persistent infections in 12 patients yielded serial isolates both prior to and throughout the course of treatment, with 3 to 9 isolates per patient (Fig. 4A and Supplementary Table 5). Clinical persistence in these patients occurred despite clinical MIC assays having established that all isolates were fluconazole susceptible, with MIC levels less than 1 µg/ml.

Each patient had a different clinical trajectory, all of them having central venous catheters that, in most cases, were removed upon detection of fungal infection. Importantly, none of the patients from which these isolates were collected were neutropenic or otherwise immunocompromised. All patients were treated with fluconazole as the first drug of choice and, in some cases, antifungal therapy was continued with either caspofungin or amphotericin B.
the 19 patients, 7 patients died during the 30 day follow up, 6 of which were unable to clear the fungal infection, although a causality between infection and death could not be determined.

We examined the drug responses of persistent and non-persistent isolates using DDAs and BMDAs as described above, and included three of the seven strains utilized above as additional controls (termed ‘susceptible’ isolates). As expected, persistent and non-persistent isolates displayed similar RAD and MIC levels of 0.25 to 1 µg/ml (Fig. 4B and Supplementary Fig. 6A-B), levels well below those of resistant strains (clinical MIC breakpoint = 4 µg/ml). The sole exception to this was the final isolate from series 3 (S03), which became fluconazole-resistant (MIC > 128 µg/ml) (R, Fig. 4C). For this series, the penultimate isolate, which had not acquired resistance (MIC = 0.5 µg/ml), was used in subsequent analyses. Strikingly, both FoG and SMG levels differed significantly between the two groups of clinical isolates (Fig. 4B,C):

non-persistent isolates had average FoG values of 0.38 while persistent isolates had average FoG values almost 2-fold higher (0.59). For most of the series, FoG levels did not change significantly as the infection progressed, despite ongoing fluconazole therapy (Supplementary Fig. 6A). Similarly, SMG levels were an average of 1.4-fold higher in persistent than non-persistent isolates and this distinction in SMG levels between the two groups was clearly evident when SMG was measured at 48 h (Fig. 4C). By contrast, MIC levels did not change when analyzed at 24 h or 48 h (Supplementary Fig. 6B), reinforcing the idea that perseverance is time-dependent whereas susceptibility/resistance is not. SMG and FoG levels again correlated for both persistent and non-persistent isolates (Supplementary Fig. 6C), indicating that the two parameters measure features of the same phenomenon.

Taken together these results reveal that persistent isolates display significantly higher levels of growth at high drug concentrations compared to isolates that are clinically cleared by fluconazole treatment. In fact, many of the persistent isolates had FoG levels > 0.5, a level at which more than half of the cells in the population are able to grow, albeit more slowly, in the presence of antifungals (Fig. 3B). This is important because it indicates that perseverance is an
Figure 4.

(a) Clinical history of 7 non-persistent isolates and 12 series of clinically persistent isolates showing the age, sex and associated conditions of the patients from whom they were isolated. ‘X’ symbols indicate the time when isolates were collected, shaded areas indicate the duration of antifungal therapy received (fluconazole, pink; caspofungin, yellow; amphotericin B, violet), red lines mark death events over a 30-day follow-up. (b) FoG and RAD levels for drug-susceptible (S, n = 4) isolates SC5314, GC75, P78042 and P87, resistant (R, n = 1) isolate P60002, non-persistent isolates (NP, n = 7) and the first patient isolate from each of the series of clinically persistent isolates.
strains (P, n = 12). Asterisks indicate significant differences between persistent and non-persistent isolates (unpaired t-test, P = 0.67 for RAD, P < 0.001 for FoG). (c) Broth microdilution assays showing MIC and SMG levels at 24 and 48 h for the susceptible control strains (S, n = 4), the non-persistent isolates as in (b) (NP, n = 7) and for both the initial and final isolates for each of the 12 clinically persistent series (S01-12). The final isolate in S03 became fluconazole resistant (R), therefore the penultimate isolate in this series was included as well. Asterisks indicate significant differences between persistent and non-persistent isolates (unpaired t-test, NP vs P-first P = 0.061 and NP vs P-last P = 0.053 for SMG at 24 h, NP vs P-first P < 0.001 and NP vs P-last P < 0.001 for SMG at 48 h).
intrinsic property of clinical isolates distinct from drug resistance. Furthermore, perseverance contributes to the ability of an antifungal to clear an infection and thereby directly impacts clinical outcome.

Drug combinations primarily affect perseverance rather than resistance

Combination therapy is currently being explored as a strategy to extend the lifespan of available antifungal drugs. We tested several adjuvant drugs reported to have additive or synergistic effects with fluconazole for their effects on FoG/SMG and RAD/MIC. These include aureobasidin A, an inhibitor of sphingolipid biosynthesis, rapamycin, an inhibitor of the mTOR signaling pathway, fluoxetine, a serotonin inhibitor reported to inhibit efflux pump activities, and fluphenazine, an antipsychotic drug that stimulates ABC transporter efflux pumps and indirectly inhibits calcineurin via calmodulin (Fig. 5A). We found that none of these adjuvant drugs caused a significant change in susceptibility to fluconazole (i.e., there was no change in RAD), but all of them significantly reduced perseverance (FoG) to a baseline level. Adjuvant treatment led to clearing of FoG for all 7 strains tested, indicating that this is a common characteristic of diverse C. albicans isolates.

Inhibitors of Hsp90 (geldanamycin and radicicol), as well as inhibitors of calcineurin (cyclosporine A and FK506), yielded similar results in drug response assays. Thus, each of these compounds had no significant effect on RAD/MIC but significantly reduced both FoG and SMG to baseline levels (Fig. 5B and Supplementary Fig. 7A). Again, this was consistent across all isolates tested, including a series of clinical isolates from a single HIV patient that had acquired increased MIC levels over time (Supplementary Fig. 7B,C). Similarly, for both the persistent and non-persistent clinical isolates FoG levels were reduced to baseline by the addition of adjuvant drugs including geldanamycin, rapamycin, FK506, fluphenazine and doxycycline (Supplementary Fig. 7D). As a consequence, FoG levels of persistent strains became indistinguishable from those of non-persistent strains. In contrast, and as seen with
Figure 5. Adjuvant drugs significantly reduce fluconazole perseverance but not resistance and render fluconazole cidal rather than static. (a) Disk diffusion assays performed with 25 µg fluconazole disks on casitone plates supplemented with adjuvant drugs 20 µg/ml fluoxetine, 5 ng/ml aureobasidin A, 0.5 ng/ml rapamycin or 10 µg/ml fluphenazine for strain SC5314 (left), and RAD (middle panels) (1-way ANOVA, n groups=5, P=0.9998) and FoG (right panels) (1-way ANOVA, n groups=5, P<0.0001) levels for strains as in Figure 1. (b) Disk diffusion assays as in (a) performed with casitone plates supplemented with Hsp90 inhibitors (0.5 µg/ml geldanamycin or 0.5 µg/ml radicicol) or calcineurin inhibitors (0.5 µg/ml FK506 or 0.4 µg/ml cyclosporine A) for strain SC5314 (left), and RAD (middle panels) (1-way ANOVA, n groups=5, P=0.9994) and FoG (right panels) (1-way ANOVA, n groups=5, P=0.0004) levels. (c) Effect of drug adjuvants and inhibitors on the viability of cells growing inside the ZOI. Fluconazole disk diffusion assays from (a) and (b) were replica plated (after removal of the drug disk) onto casitone plates (without fluconazole or the adjuvant drugs/inhibitors) and incubated at 30°C for 48 h.
other isolates, RAD levels remained unchanged in response to most of these adjuvants (Supplementary Fig. 7D).

We conclude that exposure to inhibitors of either Hsp90 or calcineurin reduced growth within the ZOI (FoG) as well as growth above MIC levels in BMDAs (SMG). However, none of the inhibitors induced a significant change in MIC or RAD when measured at 24 h as recommended. Of note, geldanamycin (Supplementary Fig. 7E), as well as all the other adjuvant drugs, significantly reduced cell viability within the ZOI (Fig. 5C).

**Genetic pathways affecting perseverance and/or resistance**

We next explored whether genetic pathways that are known to contribute to drug responses affect perseverance. Toward this goal, we collected a series of mutant isolates, most of them constructed in various backgrounds of laboratory strain SC5314 (Supplementary Fig. 8A). As expected, most of the parental isolates exhibited similar RAD/MIC levels (Supplementary Table 2 and Supplementary Fig. 8B). Surprisingly, however, perseverance levels were significantly reduced in some of the parental control strains. Specifically, FoG levels dropped by ~40-50% in CAI-4 and many of its derivatives (Supplementary Table 2, Supplementary Fig. 8B and Supplementary Text 1). The inadvertent deletion of *IRO1* (because it is adjacent to *URA3*) is likely responsible for decreased perseverance levels in CAI-4 and many of its derivatives, as deletion of *IRO1* alone reduced FoG by ~42.5%. This implies that *IRO1*, which encodes a transcription factor involved in iron utilization, promotes higher FoG levels in *C. albicans* strains. Despite the effect of strain background differences, relative changes in FoG and RAD could be measured by comparing each mutant strain to the corresponding parental strain and, in some cases, to strains in which the deleted gene had been reintroduced (Supplementary Fig. 8B).

Given that calcineurin inhibitors reduced perseverance, we first tested null mutants lacking genes that encode the calcineurin subunits Cna1 and Cnb1, along with
Independently constructed mutants lacking Crz1\(^91\), the major transcription factor activated by calcineurin in response to drug stress. Null mutants in CNA1, CNB1 and CRZ1 constructed in two different parental strains (SC5314 and SN192) had significantly reduced perseverance (50-60%) yet displayed no significant change in RAD/MIC (Supplementary Table 3 and Fig. 6A). These results indicate that calcineurin promotes perseverance via the Crz1 transcription factor.

In contrast, deletion of genes encoding regulators of calcineurin, including RCN1 and RCN2, had little effect on RAD or FoG when compared to isogenic parental strains (Supplementary Table 3 and Fig. 6A). Consistent with this, RAD and FoG levels in an rcn1\(^{\Delta \Delta}/crz1\(^{\Delta \Delta}\) mutant were similar to those of the crz1\(^{\Delta \Delta}\) single mutants (Supplementary Table 3 and Fig. 6A). However, these data do not exclude the possibility that RCN1 and RCN2 could have functionally redundant roles in drug responses.

VPS21, a gene reported to affect ‘tolerance’ but not susceptibility or resistance to azoles was analyzed because estimates from published BMDA data indicated that vps1\(^{\Delta \Delta}\) null mutants do not affect MIC at 24 h yet exhibit very high SMG at 48 h\(^{10}\). VPS21 encodes a Rab GTPase that regulates endosomal trafficking through the late endosome. As expected, vps21\(^{\Delta \Delta}\) null mutants did not alter RAD but displayed high FoG levels in DDAs relative to the parental strain (Supplementary Table 3 and Fig. 6A).

We also tested a strain lacking CAS5 because of its reported role in fluconazole responses. CAS5 encodes a transcription factor originally identified in C. albicans for its role in caspofungin survival\(^{92}\) and was shown to enhance the cidality of fluconazole, especially when analyzed at 48 h rather than 24 h after drug exposure\(^{93}\). Indeed, the cas5\(^{\Delta \Delta}\) mutant reduced FoG by 90% and increased RAD by 25% (Supplementary Table 3 and Fig. 6A), indicating that Cas5 affects perseverance dramatically and resistance more modestly.

We next analyzed the role of genes encoding transcription factors known to have roles in fluconazole resistance. These include Upc2, a transcription factor that regulates sterol uptake and ergosterol biosynthetic genes (ergosterol is the target of azoles\(^{94}\)), Tac1 and Mrr1, which
Figure 6. Effects of different pathways on perseverance and/or susceptibility to fluconazole. Disk diffusion assays performed using a series of mutants carrying deletions in (a) genes encoding calcineurin subunits Cna1 and Cna2, two separately constructed isolates lacking the calcineurin responsive transcription factor Crz1, a vacuolar trafficking protein Vps21, the Cas5 transcription factor up-regulated by fluconazole or (b) mutants lacking transcription factors known to regulate ergosterol gene expression (Upc2) or efflux pump expression (Tac1, Mrr1). Results for both crz1 isolates were similar and only one is illustrated. These mutants as well as mutants lacking calcineurin regulators Rcn1 or Rcn2, and an rcn1 crz1 double mutant were analyzed by diskImageR, and RAD and FoG levels, relative to the isogenic parental strain backgrounds are illustrated in the middle and lower rows, respectively. Pictures show representative images of two biological replicates (n = 2).
regulate the expression of ABC and MFS efflux genes, respectively. Like the cas5\Delta\Delta null mutant, upc2\Delta\Delta mutants affected both RAD (increased by 25\%) and FoG (decreased by ~70\%).

In contrast, tac1 and mrr1 mutants had no significant effects on RAD or FoG (Supplementary Table 3 and Fig. 6B). This is consistent with the idea that there are multiple efflux pump genes and transcriptional regulators that control them, and that any one of them is likely to be redundant with efflux activity of the others sufficient for wild-type activity\textsuperscript{95-97}.

Together, these results show that different genetic pathways, such as calcineurin activity via Crz1, or that involving Iro1, contribute to increased perseverance levels while Vps21 and vacuolar membrane trafficking/homeostasis contribute to reduced perseverance levels. Other Transcription factors such as Cas5 or Upc2, affect both resistance and perseverance, either by independently regulating pathways in both processes or by additional mechanisms that remain to be discovered.

**Discussion**

Currently, decisions about therapeutic strategies to treat Candida infections are often based upon patient status, the infecting species and the level of resistance (MIC) to antifungal drugs. Here, we quantified and characterized a new drug response parameter termed perseverance that appears to be relevant for the persistence of C. albicans isolates in the clinic.

Perseverance, measured as FoG or SMG, quantifies features previously described qualitatively as ‘trailing growth\textsuperscript{10,11,25,26,67,69,98,99}, ‘tolerance\textsuperscript{12-16} or ‘Hsp90-dependent resistance\textsuperscript{40,57,100-102}.

Previous studies have suggested no strong correlation between these supra-MIC growth phenomena and virulence in either mouse models\textsuperscript{17,18} or infection outcomes in the human host\textsuperscript{20}. However, most of these studies analyzed immediate responses rather than clinical persistence or recurrence over longer time frames.

We show that perseverance is concentration-independent, is correlated with the size of the subpopulation that grows at drug concentrations above the MIC, and is inversely correlated
with the length of lag phase for these colonies. The mechanisms that determine the size of the subpopulation that can grow at supra-MIC concentrations remain to be determined. Importantly, we measure SMG and FoG relative to the MIC, utilizing standard assays measured on consecutive days, to provide information about both resistance and perseverance. We show that perseverance is influenced by pH, temperature, and nutritional inputs, as previously reported for trailing growth and tolerance$^{26,66,67,70}$. The term “perseverance” was chosen as it describes the process rather than the assay, and it avoids confusion with tolerance mechanisms that promote survival in cidal drugs via longer lag phases$^{7,9}$. While current descriptions of tolerance in bacteria have analyzed survival in cidal drugs, it remains possible that a growth strategy akin to fungal perseverance may be found in bacteria that divide in supra-MIC concentrations of bacteriostatic drugs.

Perseverance is dependent upon pathways that do not affect the MIC, and is sensitive to adjuvant drugs that not only reduce perseverance but also increase the cidality of fluconazole, which is generally fungistatic. This ability to ‘clear the FoG’ suggests that the adjuvant drugs must inactivate stress response pathways that are essential for perseverance and viability in the drug. The idea that Hsp90 and calcineurin affect drug responses in fungi is certainly not new, and the idea that they can affect the cidality of antifungals has also been reported$^{15,53,69,93,103-105}$. However, this study clearly shows that these proteins, and the pathways they regulate, are important for a response to drug that is distinct from resistance and that specifically affects cell survival at supra-MIC antifungal concentrations.

Multiple pathways influence perseverance, some of which we identify in this study. Vps21, a protein involved in membrane trafficking, clearly impacts perseverance via its ability to increase cytoplasmic calcium stores$^{10,106}$, which, in turn, is likely to increase calcineurin activity. However, not all adjuvant drugs may act via the Hsp90/calcineurin pathway. For example, it remains to be determined how the TOR pathway (inhibited by rapamycin), sphingolipid synthesis (inhibited by AbA), as well as the antidepressant fluoxetine and the antipsychotic
fluphenazine affect perseverance (Supplementary Tables 2 and 3). Interestingly, Pkc1 inhibitors phenocopy inhibitors of Hsp90 or calcineurin and render fluconazole cidal\textsuperscript{101}, suggesting that perseverance is likely to be mediated, at least in part, via the Pkc1 pathway. Furthermore, the effect of Pkc1 inhibitors on the clinical isolate series used in Supplementary Figure 5 was indistinguishable from the effect of geldanamycin on these isolates\textsuperscript{101}, thus we posit that Pkc1 is critical for perseverance but not for \textit{bona fide} resistance. However, since genetic experiments found that \textit{PKC1} is not required for calcineurin-dependent drug responses\textsuperscript{101}, the relationship between Pkc1 and perseverance is likely to require further study.

It is also unclear how \textit{IRO1}, a gene that was inadvertently deleted in a commonly used \textit{C. albicans} lineage, contributes to perseverance. One possibility is that it relates to the requirement for iron in the function of Erg11, which is a cytochrome p450 enzyme and the direct target of fluconazole. Of note, doxycycline reduces tolerance by chelating limited iron stores in the medium\textsuperscript{15}, and iron deprivation also regulates the expression of \textit{HSP90} and \textit{CRZ1} via calcineurin signaling\textsuperscript{107}. Furthermore, the effect of doxycycline on iron availability is important to consider when studying azole responses using doxycycline to activate or repress the Tet promoter in engineered strains\textsuperscript{108,109}.

Similarly, we do not yet understand how transcription factors \textit{CAS5} and \textit{UPC2} affect both resistance and perseverance. One option is that they may regulate multiple independent pathways, some of which influence resistance and others that affect perseverance. A recent study supports this idea, given that \textit{cas5\Delta/\Delta} strains cause altered promoter occupancy of RNA polymerase II at over 600 genes\textsuperscript{110}. Alternatively, genes affecting resistance may also reduce the degree to which an antifungal drug causes cellular stress and thus may alter the requirement for the stress response mechanisms that drive perseverance.

Of more direct significance, this analysis of a well-matched set of clinically persistent and non-persistent isolates supports the hypothesis that strains with higher FoG/SMG levels may relate to the failure of antifungal drugs to clear an infection despite an isolate being drug
susceptible. The prolonged presence of isolates in the host could also contribute to the emergence of resistance by providing cells with a greater chance of acquiring mutations that render them drug resistant, a process that occurred in one of the patients with a persistent isolate. Measuring the perseverance level of an infecting isolate could facilitate useful clinical studies to determine the degree to which perseverance may predict which isolates are more likely to give rise to persistent and/or recurrent infections. In addition, we posit that quantifying perseverance may enable clinicians to determine when adding an adjuvant drug to increase the cidality of fluconazole would be most effective. This may be especially relevant when antifungals other than fluconazole are less accessible, for example in developing countries.

Methods

Strains. All strains (Supplementary Table 1) were streaked onto YPD plates and grown for 48 h at 30°C. A single colony from each strain was arbitrarily selected and frozen in 15% glycerol and stored at -80°C for all assays. For persistent and nonpersistent strains obtained from Brazilian patients, all ethical regulations were observed and the study was approved by the Ethical Committee of the Federal University of São Paulo (January, 2016, NO 9348151215).

Strain construction. To disrupt the two alleles of the RCN2 gene we amplified (using primers BP1440 and BP1441, Supplementary Table 4) the flanking sequence of C7_01700W ORF using BJB-T 2 (HIS1) and BJB-T 61 (ARG4) plasmids as a template (Supplementary Table 4). Strain YJB-T65 was transformed with the PCR product to generate heterozygote mutant YJB-T2214 (rcn2::HIS1) and subsequently homozygote mutant YJB-T2227 (rcn2::HIS1 / rcn2::ARG4). The disruptions were verified using primers BP1444 and BP1445.

Broth micro dilution assays. Minimal inhibitory concentration (MIC) for each strain was measured using CLSI M27-A2 guidelines with minor modifications as follows. Strains were
streaked from frozen culture onto YPD agar and incubated for 24 h at 30°C. Colonies were suspended in 1 ml PBS and diluted to 10^3 cells/ml in a 96 well plate with casitone containing a gradient of two-fold dilutions per step of fluconazole, with the first well contain no drug. For persistent and non-persistent clinical isolates, cells were grown overnight in YPD at 30°C and diluted to 10^4 cells/ml in YPD containing a gradient of two-fold dilutions per step of fluconazole. MIC_{50} levels were determined after 24 h or 48 h by taking an optical density reading (OD_{600}) after manual agitation of all wells on a Tecan plate reader (Infinite F200 PRO, Tecan, Switzerland). MIC_{50} levels (shown as yellow lines on BMDA heatmaps) were determined as the point at which the OD_{600} had been reduced by ≥50% compared to the no-drug wells.

**Disk diffusion assays.** The CLSI M44-A2 guidelines for antifungal disk diffusion susceptibility testing were followed with slight modifications. In brief, strains were streaked from frozen culture onto YPD agar and incubated for 24 h at 30°C. Colonies were suspended in 1 ml PBS and diluted to 1 x 10^6 cells/ml. 2 x 10^5 cells were spread onto 15 ml casitone (9 g/l Bacto casitone, 5 g/l yeast extract, 15 g/l Bacto agar (BD Sparks, MD USA), 11.5 g/l sodium citrate dehydrate and 2% glucose (Sigma-Aldrich, St. Louis, MO USA). For persistent and non-persistent clinical isolates, cells were grown overnight in YPD and 10^5 cells were spread onto 15 ml YPD plates. To facilitate comparisons between casitone and YPD disk diffusion assays, a subset of control strains with diverse FoG levels were included in both types of assays (SC5314, GC75, P87). For disk assays with drug adjuvants, the following concentrations of drugs were used: 0.5 µg/ml geldanamycin, 0.5 µg/ml radicicol, 0.5 µg/ml FK506, 0.4 µg/ml cyclosporine A, 20 µg/ml fluoxetine, 5 ng/ml aureobasidin A, 0.5 ng/ml rapamycin, 10 µg/ml fluphenazine and 50 µg/ml doxycycline. A single 25 µg fluconazole disk (6 mm, Oxoid, UK or Liofilchem, Italy) was placed in the center of each plate, plates were then incubated at 30°C for 48 h, and each plate was photographed individually. Analysis of the disk diffusion assay was done using the *diskImageR* pipeline and the R script is available at
ScanLag assay. The ScanLag assay was adapted from\textsuperscript{75} with minor modifications. Strains were streaked from frozen culture onto YPD agar and incubated for 24 h at 30°C. Colonies were suspended in 1 ml PBS, diluted to $10^4$ cell/ml and $2 \times 10^3$ cells (200 μl) were spread onto 15 ml casitone plates with or without 10 μg/ml fluconazole (DMSO, Ofer Chemical Laboratory, Israel). Plates were placed on the scanners at 30°C and scanned every 30 min for 72 h. Analysis of the images was done in MATLAB using the "ScanLag" script\textsuperscript{75} that was adjusted to yeast cells by changing the identification of the size of the colony to a minimum of 20 pixels.

Viability assays. Replica plating was performed from disk diffusion plates that were incubated at 30°C for 48 h. Master plates were inverted and pressed firmly on a sterile cotton velveteen stamp and then transferred to new casitone plates containing no drug. Replica plates were incubated at 30°C for 48 h, and then each plate was photographed individually.

Growth assays. To determine growth parameters (doubling time and lag phase duration), \textit{C. albicans} cells were seeded at a concentration of $2 \times 10^5$ cells/ml in 96-well plates containing YPD media. Plates were grown for 48 h at 30°C with continuous shaking in a Tecan plate reader and the optical density reading ($\text{OD}_{600}$) of all wells was recorded every 15 min. Measurements of doubling time and lag phase duration were obtained from 3 biological replicates, each performed in duplicate.

Microcolony assays. Microcolony assays were adapted from\textsuperscript{74} with minor modifications. Colonies were suspended in 3 ml casitone media and grown overnight in 30°C. Cultures were
diluted 1:30 in casitone and incubated at 30°C until cultures reached logarithmic phase (~3-4 h) and diluted to 10^4 cells/ml in casitone.

Microscopy. Glass-bottomed 24-well plates (De-GROOT, Israel) were coated with 1 ml of a 200 µg/ml Concanvalin A solution (Type VI, Sigma-Aldrich, St. Louis, MO USA) for 3-4 h. Wells were washed twice with 1 ml ddH20 and 10^4 cells/ml in a 2 ml volume were added to wells with or without 10 µg/ml fluconazole. Plates were sealed with an optically clear film (Axygen breathing sealing film, Corning, Israel) and spun at 1200 rpm for 1 min. Images where captured in 100 fields per well for 48 h in 1 h intervals using a Nikon Ti Eclipse microscope equipped with a full-stage environmental chamber (set to 30°C) with a Nikon Plan Apo 10x (0.45 numerical aperture) objective using Nikon Elements AR software. Focus coordinates and the positions over the plate surface was assigned using homemade R- and C-based scripts (adapted from by Dr. Naomi Ziv). Automated Image analysis, including image processing and microscope control routines were written in Matlab, R, C, and shell scripts were adapted from 74.

Statistical analyses. All experiments represent the average of three or more biological repeats, with two technical repeats of each. Error bars represent the standard deviation. Statistics were performed using Microsoft Excel 2016 (Microsoft Corporation) and Prism 6 (GraphPad). The statistical tests performed were two-tailed Student’s t tests, one way ANOVAs for comparisons between groups, and Tukey’s multiple comparison tests. R squared tests were used for linear regressions. Significance was assigned for P values smaller than 0.05, asterisks denote P values as follows: ***, P < 0.001; **, P < 0.01; *, P < 0.05.

Data availability. The authors declare that all data supporting the findings of the study are available in this article and its Supplementary Information files, or from the corresponding author upon request.
Acknowledgements

We thank members of the Berman and Bennett labs for stimulating discussions throughout the work. G. Palmer, M. Lohse, O. Homann, A. Johnson, R. Ben-Ami, T. White, D. Soll, D. McCallum, F. Odds, D. Sanglard, L. Cowen, P.T. Magee, B. Cormack, P. Magwene, M. Kupiec, W. Fonzi, S. Noble, B.M. Vincent, S. Lindquist, and A. Colombo for providing strains and Naomi Ziv, Naama Droor and Stephanie Diezmann for helpful comments on the manuscript.

This work was funded by European Research Council Advanced Award 340087 (RAPLODAPT) to JB, by Fundação de Amparo a Pesquisa do Estado de São Paulo (FAPESP), Brazil to ALC, by National Institutes of Health grant AI081704 to RJB and by a Brazil Initiative Collaborative Grant from the Watson Institute to RJB and IVE.

Author contributions

IVE, RJB and JB designed the study; AR, IVE and AD collected the data; AR and IVE analyzed the data, ESS constructed mutant isolates, ALC provided clinical isolates and clinical input to the paper; AR, IVE, RJB and JB wrote the manuscript.
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Clearing the Fungal FoG: Perseverance, a property distinct from resistance, is associated with clinical persistence

Alexander Rosenberg**, Iuliana V. Ene**, Alon M. Dahan¹, Ella Shtifman Segal¹,
Arnaldo L. Colombo³, Richard J. Bennett², Judith Berman¹***

Supplementary Material
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Supplementary text (describing parental strains in Supplementary Fig. 8)

Lab strain CAI-4 differs from SC5314 in that both copies of \textit{URA3} and the adjacent gene, \textit{IRO1}, were sequentially deleted\textsuperscript{1,2}. In addition, the strain had undergone several homozygosis events\textsuperscript{3}. Transformation of CAI-4 with \textit{URA3} (Supplementary Table 2, CAI-4+\textit{URA3}) did not restore FoG to SC5314 levels, but introduction of a single copy of both \textit{URA3} and \textit{IRO1} into derivative SN76 was sufficient to restore FoG to SC5314 levels (Supplementary Fig. 8 and Supplementary Table 2, SN95). This suggests that deletion of \textit{IRO1} is responsible for a difference in perseverance levels in many derivatives of SC5314 and that different LOH events did not affect FoG levels, although we note that the hemizygosity acquired in BWP17\textsuperscript{4} may have contributed to the exceptionally low perseverance levels in this specific strain (Supplementary Fig. 8 and Supplementary Table 2). Indeed, deletion of \textit{IRO1} alone in strain SN152 (\textit{MTLa/MTL}\textsubscript{Δ}) reduced FoG to levels similar to those seen in derivatives of CAI-4 (Supplementary Fig. 8A,B). Thus, deletion of \textit{IRO1} in most lab strains is sufficient to explain the reduced FoG in these strains.

Accordingly, all analysis of mutant isolates are presented relative to the isogenic parental strain from which they were derived.
Summary of lineage changes in lab strains derived from SC5314:

**SC5314**

**CAF2-1** (derived from SC5314)
- LOH to: allele A @ Chr7L near CEN

**CAI4** (derived from CAF-2)
- ura3Δ/Δ  ired1Δ/Δ
- LOH to: allele A @ Chr7L near CEN;
- LOH to: allele B @ Chr3R near CEN;
- Trisomy of: Chr1 &/or 2

**RM10** (derived from CAI-4)
- his1Δ/+ ura3Δ/+ ired1Δ/Δ
- LOH to: allele A @ Chr2L;
- LOH to: allele A @ Chr7L near CEN;
- LOH to: allele B @ Chr3R near CEN;
- Trisomy of: Chr2

**RM100#13** (derived from RM10)
- his1Δ/+ ura3Δ/+ ired1Δ/Δ
- Monosomy of Chr5 terminal 30 kb allele A
- LOH to: allele A @ Chr2L
- LOH to: allele A @ Chr7L near CEN
- LOH to: allele B @ Chr3R near CEN

**RM1000#6** (derived from RM100)
- his1Δ/Δ ura3Δ/Δ ired1Δ/Δ
- Monosomy of Chr5 terminal 30 kb allele A
- LOH to: allele B @ Chr3R near CEN
- LOH to: allele A @ Chr2L
- LOH to: allele A @ Chr7L near CEN

**BWP17** (derived from RM1000#6)
his1Δ/Δ ura3Δ/Δ iro1Δ/Δ arg4Δ/Δ

LOH to: allele A @ Chr2L

LOH to: allele B @ Chr3R near CEN

LOH to: allele A @ Chr4 near right telomere

LOH to: allele A @ Chr7L near CEN

Monosomy of Chr5 terminal 30 kb allele A

SN76 or RM1000#2 (derived from RM100)

his1Δ/Δ ura3Δ/Δ iro1Δ/Δ

Homozygosis of Chr5 terminal 30 kb to disomic allele A

LOH to: allele B @ Chr3R near CEN

LOH to: allele A @ Chr2L

LOH to: allele A @ Chr7L near CEN

SN95 (derived from SN76)

MTLa/α his1Δ/Δ ura3Δ/+ iro1Δ/+ arg4Δ/Δ

Homozygosis of Chr5 terminal 30 kb to disomic allele A

LOH to: allele B @ Chr3R near CEN

LOH to: allele A @ Chr2L

LOH to: allele A @ Chr7L near CEN

SN152* (derived from SN95)

MTLa/Δ leu2Δ/+ his1Δ/+ ura3Δ/+ iro1Δ/+ arg4Δ/+ 

Homozygosis of Chr5 terminal 30 kb to disomic allele A

LOH to: allele B @ Chr3R near CEN

LOH to: allele A @ Chr2L

LOH to: allele A @ Chr7L near CEN

SN152 iro1Δ* (derived from SN152)

MTLa/Δ leu2Δ/+ his1Δ/+ ura3Δ/+ iro1Δ/Δ arg4Δ/+ 

Homozygosis of Chr5 terminal 30 kb to disomic allele A

LOH to: allele B @ Chr3R near CEN
LOH to: allele A @ Chr2L

LOH to: allele A @ Chr7L near CEN

*Not sequenced to detect additional possible LOH events
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Supplementary Figure 1. Perseverance is detected for different drugs and fungal species. (a) Disk diffusion assays measure RAD and FoG levels for a set of 19 *C. albicans* isolates in response to diverse antifungal drugs containing fluconazole (25 µg), ketoconazole (10 µg), poseconazole (10 µg), 5-fluorocytosine (25 µg), micafungin (5 µg), caspofungin (5 µg), or amphotericin B (50 µg). (b) Disk diffusion assays performed with 25 µg fluconazole disks detect RAD and FoG levels for different pathogenic yeast species including *C. tropicalis*, *C. krusei*, *C. glabrata*, *C. parapsilosis* and *S. cerevisiae*. Note that FoG levels in azoles were particularly low for *S. cerevisiae*. 
Supplementary Figure 2. Measuring drug responses of C. albicans isolates using broth microdilution assays (BMDAs). (a) Comparison of MIC and SMG for parental isolates compared with progeny from wells with drug concentration below (cyan heatmaps) or above (yellow heatmaps) the MIC₅₀ (yellow lines) are very similar. MIC₅₀ (top right) and SMG (lower right) levels are similar for parental and progeny cells. (b) Correlation analyses for log₂(MIC₅₀) vs SMG (n-28, P=0.0064), SMG vs FoG₂₀ (n-12, P<0.0001), log₂(MIC₅₀) vs RAD₂₀ (n-28, P<0.0001) for the series of isolates analyzed in Figure 1.
Supplementary Figure 3

A

pH 7

pH 4.5

RAD<sub>20</sub>  18  17

FoG<sub>20</sub>  0.68  0.46

B

GC75  SC5314  P78  L26  AM2  P87  T101

pH 7  pH 4.5

FLC (µg/ml) 0 → 128

SMG

Log<sub>2</sub>(MIC<sub>20</sub>)

SMG

Log<sub>2</sub>(MIC<sub>20</sub>)

pH 7  pH 4.5

growth (OD<sub>600</sub>)

below MIC

above MIC

0  1.6

C

30°C  39°C

RAD<sub>20</sub>  16  24.5

FoG<sub>20</sub>  0.74  0.32
Supplementary Figure 3. Impact of growth conditions on FoG and SMG levels.

Disk diffusion assays (a,c,e) and broth microdilution assays (b,d) measure effect of different pH (a,b) (RAD (Paired t-test, t=2.2, df=6 n pairs=7, P = 0.0656), FoG (Paired t-test, t=3.4, df=6 n pairs=7, P = 0.0138), MIC (Paired t-test, t=1.33, df=6 n pairs=7, P = 0.2380),SMG (Paired t-test, t=2.3, df=6 n pairs=7, P = 0.0693) )
temperature (c,d) (RAD (One way ANOVA, n groups=3, P = 0.6382), FoG (One way ANOVA, n groups=3, P = 0.0302), MIC (One way ANOVA, n groups=4, P = 0.9646),SMG (One way ANOVA, n groups=4, P = 0.7366) )
and nutrient media conditions (e) (RAD (Paired t-test, t=0.75, df=6 n pairs=7, P = 0.4797)
FoG (Paired t-test, t=1.7, df=6 n pairs=7, P = 0.1491)) on RAD/MIC and FoG/SMG levels. Disk diffusion assays performed with strain SC5314 (left panels), RAD levels (middle panels) and FoG levels (right panels) calculated with for range of C. albicans strains as in Figure 1. Broth microdilution assays were analyzed as in Figure 2 for the same set of strains in medium at pH 7 and pH 4.5 (b) and at 30, 37, 39 and 41°C (d).
Supplementary Figure 4. Time of initial colony appearance (ToA) for strains with a range of FoG levels. ToA, determined using ScanLag, on a range of isolates with different FoG levels, as in Figure 4. Histograms show the time of new colony appearance and ΔToA indicates the difference in the time of appearance of colonies with vs without drug (ΔToA = ToA with 10 µg/ml fluconazole – ToA without fluconazole).
Supplementary Figure 5. Relationships between growth parameters and perseverance. (a) Correlation analyses for FoG and colony size (left) or FoG and growth rate (right) on casitone plates with 10 µg/ml fluconazole (n = 9). (b) Correlation analyses between FoG and doubling time or FoG and lag phase duration for related C. albicans isolates with differences in growth rates. Red data points indicate isolates from the P37055 lineage (P37 parental strain, and P37-PA,B passaged isolates), blue data points indicate isolates with different numbers of the mitotic cyclin CLB4 gene (BWP17 clb +/-, clb +/-, clb -- A,B). Data points represent the average of 3 biological replicates (n = 3, P = 0.075 for FoG vs Doubling time, P = 0.16 for FoG vs Lag time).
Supplementary Figure 6. FoG and SMG levels differ between persistent and non-persistent clinical isolates. (a) Fluconazole disk diffusion assays performed on all isolates from each patient series with a clinically persistent infection (Figure 5A) were analyzed using diskImageR for RAD (left) and FoG (right). The final isolate in the S03 series became resistant during therapy and is shown in a black square (R). (b) 24 and 48 h MIC levels from BMDAs for drug-susceptible isolates (SC5314, GC75, P78042 and P87), non-persistent isolates (NP1-7), and the first and the last isolate in each of the 12 persistent series (S01-12). The final isolate in S03 became fluconazole resistant (R), therefore the penultimate isolate in this series was included. (c) Correlation analysis between FoG and SMG levels for susceptible control strains (dark blue), persistent (red) and non-persistent isolates (light blue) (n = 35, P < 0.0001).
Supplementary Figure 7

A

SC5314
P78
L26
AM2
P87
T101

GCD75

No drug
Geldanamycin
Radicicol
FK506
Cyclosporine A

FLC (µg/ml) 0 → 128

B

TW01
TW17

No drug
Geldanamycin
Radicicol
FK506
Cyclosporine A

time

C

RAD73

No drug
Geldanamycin
Radicicol
Cyclosporine A
FK506

D

RAD73

casitone

No drug
Geldanamycin
Radicicol
FK506

Drug in media

E

FoG20

DDA 25 µg FLC
replica plate no drug

DDA 25 µg FLC 0.5 µg/ml geld
replica plate no drug
**Supplementary Figure 7.** Fluconazole adjuvants significantly reduce perseverance. (a,b) Broth-micro dilution assays on casitone media supplemented with 0.5 µg/ml geldanamycin, 0.5 µg/ml radicicol, 0.5 µg/ml FK506 or 0.4 µg/ml cyclosporine A for 7 *C. albicans* isolates (a) and a clinical isolate series isolated from a single HIV patient (b). MIC\(_{50}\) values (yellow bars) were calculated as the fluconazole concentration resulting in 50% growth inhibition relative to wells with no fluconazole. Cyan heatmaps indicate growth below MIC, while yellow heatmaps indicate growth above MIC levels. Graphs show MIC (1-way ANOVA, n groups=5 , P< 1) and SMG (1-way ANOVA, n groups=5 , P < 0.0001) levels for the 16 isolates in the HIV series. (c) RAD and FoG levels from fluconazole disk diffusion assays performed on casitone media supplemented with Hsp90 inhibitors (0.5 µg/ml geldanamycin or 0.5 µg/ml radicicol) or calcineurin inhibitors (0.5 µg/ml FK506 or 0.4 µg/ml cyclosporine A) on the series of 16 clinical isolates from a single HIV patient and analyzed using *diskImageR*. RAD (1-way ANOVA, n groups=5 , P<0.9999) and FoG (1-way ANOVA, n groups=5 , P < 0.0001) levels. (d) Fluconazole disk diffusion assays for a subset of non-persistent (NP, n = 4, cyan dots) and persistent (P, n = 10, red dots) isolates from Figure 5 measured at 48 h on casitone media supplemented with adjuvant drugs (doxycycline 50 µg/ml, 0.5 µg/ml FK506, 10 µg/ml fluphenazine, 0.5 µg/ml geldanamycin or 0.5 ng/ml rapamycin) (for RAD P < 0.001 for no drug vs DOX and no drug vs FK506, p > 0.05 for the other drugs; for FoG P < 0.01 for all drug comparisons). (e) Effect of geldanamycin on viability of cells growing in the ZOI. Fluconazole disk diffusion assays on casitone media (top two rows, as in Figure 3) or casitone medium supplemented with 0.5 µg/ml geldanamycin (bottom two rows), were replica-plated onto casitone plates without drug and incubated at 30°C for 48 h.
Supplementary Figure 8. Impact of IRO1 on FoG levels in a standard laboratory lineage. (a) Cartoon of the SC5314 lineage and its derivatives CAF2-1, CAI4, RM10, RM1000, BWP17, SN76, SN95, and SN152 strain containing one or no IRO1 alleles present (stars indicate location of deleted genes on the respective chromosomes). (b) Fluconazole disk diffusion assays performed using a series of laboratory strains in which the IRO1 gene was inadvertent deleted or reintegrated. +/- signs indicate presence or absence of IRO1 (+/+, both IRO1 alleles present, +/-, one IRO1 allele present, -/- no IRO1 alleles present). RAD and FoG levels are shown relative to the most ancestral isolate in the lineage (SC5314). Pictures show fluconazole disk assays performed on an iro1 null mutant and corresponding parental strain. Histograms show RAD (middle panel) and FoG (right panel) for strains SC5314, CAF2-1, CAI4, RM1000, BWP17, SN76, SN95, and 2 versions of the SN152 strain containing one or no IRO1 alleles present (n = 2).
## Supplementary Table 2

| Strain          | Casitone RAD<sub>20</sub> | Casitone FoG<sub>20</sub> | URA3 ura<sup>3</sup>+/+ | URA3 ura<sup>3</sup>+- | IRO1 iro<sup>+</sup>+/- | IRO1 iro<sup>+</sup>-/- | FoG responsive to Ca<sup>2+</sup> |
|-----------------|---------------------------|---------------------------|--------------------------|--------------------------|--------------------------|--------------------------|----------------------------------|
| SC5314          | 1.00                      | 1.00                      | ura<sup>3</sup>+/-     | ura<sup>3</sup>+/-     | iro<sup>+</sup>+/-     | iro<sup>+</sup>+-        | yes                             |
| CAF2-1          | 1.00                      | 1.04                      | ura<sup>3</sup>+-      | ura<sup>3</sup>+-      | iro<sup>+</sup>+-     | iro<sup>+</sup>+-        | yes                             |
| CAI4            | 0.95                      | 0.54                      | ura<sup>3</sup>+-      | ura<sup>3</sup>-/-     | iro<sup>+</sup>+-     | iro<sup>+</sup>-/-      | no                              |
| CAI4 + URA3     | 0.95                      | 0.54                      | ura<sup>3</sup>+-      | ura<sup>3</sup>-/-     | iro<sup>+</sup>-/-     | iro<sup>+</sup>+-        | yes                             |
| RM1000 #6       | 0.85                      | 0.67                      | ura<sup>3</sup>-/-     | ura<sup>3</sup>-/-     | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |
| BWP17           | 0.85                      | 0.39                      | ura<sup>3</sup>+-      | ura<sup>3</sup>-/-     | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |
| RM1000 #2       | 0.90                      | 0.70                      | ura<sup>3</sup>-/-     | ura<sup>3</sup>-/-     | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |
| SN76            | 0.85                      | 0.67                      | ura<sup>3</sup>+-      | ura<sup>3</sup>-/-     | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |
| SN95            | 1.05                      | 1.00                      | ura<sup>3</sup>+-      | ura<sup>3</sup>+-      | iro<sup>+</sup>+-     | iro<sup>+</sup>-/-      | no                              |
| SN152*          | 1.13                      | 0.94                      | ura<sup>3</sup>+-      | ura<sup>3</sup>+-      | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |
| SN152* iro1Δ    | 1.26                      | 0.57                      | ura<sup>3</sup>+-      | ura<sup>3</sup>+-      | iro<sup>+</sup>-/-     | iro<sup>+</sup>-/-      | yes                             |

**Supplementary Table 2.** RAD and FoG levels of SC5314-derived laboratory strain lineage.
## Supplementary Table 3

| Function/pathway tested | Relevant wild-type | Mutant          | RAD<sub>20</sub> | FoG<sub>20</sub> |
|--------------------------|--------------------|-----------------|------------------|-----------------|
| **calcinerin pathway**   | SC5314             | SC5314          | 1.00             | 1.00            |
|                          |                    | *cna1ΔΔ*        | 1.00             | 0.44            |
|                          |                    | *cnb1ΔΔ*        | 1.00             | 0.48            |
|                          |                    | *crz1ΔΔ*        | 1.00             | 0.50            |
|                          | SN152              | *crz1ΔΔ*        | 1.10             | 0.39            |
| **calcineurin regulators** | SC5314             | *rcn1ΔΔ*        | 1.00             | 0.46            |
|                          |                    | *crz1ΔΔ*        | 1.05             | 0.94            |
|                          |                    | *rcn2ΔΔ*        | 1.05             | 0.91            |
|                          | SN95               | *rcn1ΔΔ*        | 1.05             | 0.94            |
| **Calcium channels**     | SC5314             | *mid1ΔΔ*        | 1.10             | 0.39            |
|                          |                    | *cch1ΔΔ*        | 1.00             | 1.06            |
| **vacuole/intracellular calcium concentration** | CAI-4             | **CAI4**        | 1.00             | 1.00            |
|                          |                    | *vps21ΔΔ*       | 1.00             | 2.34            |
| **cidality of fluconazole** | SN152             | **SN152**       | 1.00             | 1.00            |
|                          |                    | *cas5ΔΔ*        | 1.25             | 0.13            |
| **regulator of ergosterol biosynthesis genes** | SN152             | *upc2ΔΔ*        | 1.25             | 0.29            |
|                          |                    | *tac1ΔΔ*        | 1.05             | 0.95            |
|                          |                    | *mrr1ΔΔ*        | 1.10             | 0.89            |
| **regulators of efflux pump genes** | SN152             | **G5**          | 1.00             | 1.00            |
|                          |                    | *mdr1ΔΔ*        | 1.14             | 0.87            |
|                          |                    | **CAI4**        | 1.00             | 1.62            |
|                          |                    | **MDR1 OE**     | 1.00             | 0.92            |

**Supplementary Table 3.** RAD and FoG levels of null mutants of interest.
Supplementary Table 4

| Primer | Sequence                                                                 |
|--------|--------------------------------------------------------------------------|
| BP1440 | 5' CCACCTCTTTTCATTATTTCCCAATCAAATTATTCTTTTTCTTACTCTTTA TTTGCGttttccagtcacgcgtt 3' |
| BP1441 | 5' GTGATTGTAGGGGACGATACAAAATCTTGTTTTTTTCTTATTAAGGAGA CCAGGCtgtaattgtaggag 3' |
| BP1444 | 5' gtccttcgcacatctt 3'                                                   |
| BP1445 | 5' ccacgagacacgacatcataat 3'                                           |

| Name   | Plasmid       | Reference                  |
|--------|---------------|----------------------------|
| BJB-T2 | pGEM-His1     | Wilson et al., 1999        |
| BJB-T61| pSN69-Arg4    | Noble and Johnson, 2005    |

**Supplementary Table 4.** Plasmids and primers used in this study.
Supplementary Videos 1-3. Time-lapse videos of cells from strains with similar MIC and different FoG levels. Cells were plated in liquid casitone medium containing 10µg/ml fluconazole, which is ~ 20-fold above their MIC levels. (1) Strain P87-low FoG (FoG-0.29, RAD-18), (2) Strain AM2- medium FoG (FoG-0.41, RAD-17) (3) SC5314- high FoG (FoG-0.68, RAD-18).