Amino Acid Substitutions at the Major Insertion Loop of *Candida albicans* Sterol 14alpha-Demethylase Are Involved in Fluconazole Resistance

Nidia Alvarez-Rueda1*, Audrey Fleury1, Florent Morio1,2, Fabrice Pagniez3, Louis Gastinel3, Patrice Le Pape1,2*

1 Département de Parasitologie et de Mycologie Médicale, Université de Nantes, Nantes Atlantique Universités, EA1155 – IICiMed, Faculté de Pharmacie de Nantes, Nantes, France, 2 Laboratoire de Parasitologie-Mycologie, CHU de Nantes, Nantes, France, 3 Laboratoire de Pharmacologie des Immunosuppresseurs en Transplantation, INSERM UMR 850, Université de Limoges, Limoges, France

**Abstract**

**Background:** In the fungal pathogen *Candida albicans*, amino acid substitutions of 14alpha-demethylase (*CaErg11p*, *CaCYP51*) are associated with azole antifungals resistance. This is an area of research which is very dynamic, since the stakes concern the screening of new antifungals which circumvent resistance. The impact of amino acid substitutions on azole interaction has been postulated by homology modeling in comparison to the crystal structure of *Mycobacterium tuberculosis* (MT-CYP51). Modeling of amino acid residues situated between positions 428 to 459 remains difficult to explain to date, because they are in a major insertion loop specifically present in fungal species.

**Methodology/Principal Finding:** Fluconazole resistance of clinical isolates displaying Y447H and V456I novel *CaErg11p* substitutions confirmed in vivo in a murine model of disseminated candidiasis. Y447H and V456I implication into fluconazole resistance was then studied by site-directed mutagenesis of wild-type *CaErg11p* and by heterogeneously expression into the *Pichia pastoris* model. CLSI modified tests showed that Y447H and V456I are responsible for an 8-fold increase in fluconazole MICs of *P. pastoris* mutants compared to the wild-type controls. Moreover, mutants showed a sustained capacity for producing ergosterol, even in the presence of fluconazole. Based on these biological results, we are the first to propose a hybrid homology structure-function model of *Ca-CYP51* using 3 different homology modeling programs. The variable position of the protein insertion loop, using different liganded or non-liganded templates of recently solved CYP51 structures, suggests its inherent flexibility. Mapping of recognized azole-resistant substitutions indicated that the flexibility of this region is probably enhanced by the relatively high glycine content of the consensus.

**Conclusions/Significance:** The results highlight the potential role of the insertion loop in azole resistance in the human pathogen *C. albicans*. This new data should be taken into consideration for future studies aimed at designing new antifungal agents, which circumvent azole resistance.

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**Introduction**

*Candida albicans* is an opportunistic fungal pathogen that causes severe blood and disseminated infections (BSIs). The incidence of these infections has markedly increased over the past decade, due to the increase of immunocompromised and neutropenic patients after organ transplantation, cancer therapy or AIDS [1]. In these populations, *C. albicans* is associated with a high mortality rate and the obvious economic consequences [2,3].

Although some antifungal drugs are available (see review [4,5]), fluconazole (FLC) is still considered the drug of choice to treat most *Candida* infections [6–8]. However, long-term exposure to FLC, as well as FLC underdosage during the empirical regimens, leads to an increased resistance phenomena [9–16]. Thus, the understanding of the molecular mechanisms underlying azole resistance in *C. albicans* is necessary in order to discover new antifungal agents that circumvent drug resistance [17,18] and could guide the choice of the appropriate antifungal treatment at the onset of infection.

Among the molecular mechanisms involved in the resistance of *C. albicans* to azole antifungals we can distinguish the over-expression of the gene encoding efflux pumps CDR1, CDR2 and MDR1 [19–22] and/or upregulation or amino acid substitutions of lanosterol 14alpha-demethylase (*CaErg11p*, *CaCYP51*) [11,12,23–26]. *CaErg11p* participates in ergosterol biosynthesis, an essential requirement for yeast viability [27]. Mutations in the *CaErg11* gene can result in amino acid substitutions in the *CaErg11p* which lead to the decreased affinity of FLC for its target and can lead to toxic lanosterol accumulation [28].

Among the many *CaErg11p* substitutions described to date in *in vitro* azole-resistant clinical isolates of *C. albicans* (See review [29]), only a few...
of them have been implicated in resistance, using site-directed mutagenesis [12,30–32]. These mutations are clustered into three hot spots, ranging from amino acids 105 to 165, 266 to 287 and 405 to 488 [13].

In a previous study, the CaErg11 gene sequencing of 73 C. albicans clinical isolates displaying various levels ofazole susceptibility, allowed us to identify two novel amino acid substitutions, Y447H and V456I, in two clinical isolates (CAAL61 and CAAL37 respectively) displaying a high-level of FLC resistance (64 \mu g/ml) [29]. Although V456I substitution has been previously mentioned in the literature by Sanglard D and coll. [33], its involvement inazole resistance has not been investigated so far. Both amino acid residues located in the third hot spot (amino acid residues 405 to 488) are conserved in various yeast and filamentous fungal species [29,34]. Whereas, substitutions at these positions could be involved in azole resistance. However, the interaction of amino acid residues between positions 428 to 459 with FLC is still difficult to explain because of a major insertion loop specifically present in fungal species [35]. To the best of our knowledge, molecular information of this region is lacking, due to the absence of amino acid sequence homology between CaErg11p and bacterial- or trypanosomatid- CYP51 molecular models [34,36–40]. Taken together, the involvement of both Y447H and V456I substitutions in FLC resistance remains difficult to solve using the current homology models of CaCYP51.

Site-directed mutagenesis of the CaErg11 gene and its heterologous expression in the yeast Saccharomyces cerevisiae is the most commonly employed model to investigate the potential involvement of CaErg11p point substitutions on azole resistance [31,41,42]. More recently, Pichia pastoris has emerged as a practical expression system and is considered a viable alternative to bacterial and S. cerevisiae expression models [43–45]. Pichia pastoris has also been suggested as an efficient method for mutant library creation, based on the selection of single copy plasmid integrated transformants on low antibiotic concentrations [46]. Furthermore, the P. pastoris expression model has some advantages such as: (i) the repression of endogenous genes with low production levels of native proteins, (ii) the genetic stability of the P. pastoris transformants by gene integration, (iii) the capacity for eukaryotic post-translational modifications such as glycosylation and (iv) the possibility of a high production of mutated recombinant proteins.

In the present study, we introduced single mutations into the CaErg11 gene by site-directed mutagenesis. We cloned the CaErg11 gene and mutants into P. pastoris to study the contribution of Y447H and V456I amino acid substitutions in FLC resistance. We also investigated the effect of these Erg11p substitutions on ergosterol biosynthesis. Based on the recently published CYP51 ortholog sequence alignments and structures, we first proposed a hybrid model of CaCYP51 in order to localize amino acid substitutions in the major insertion loop and to explain their possible implication on azole resistance.

**Results**

**Azole susceptibility of clinical isolates**

Minimal inhibitory concentrations (MICs) of the CAAL37 and CAAL61 clinical isolates against FLC, ITC and VRC were determined in vitro (Table 1). The two isolates were resistant for FLC (MIC = 64 \mu g/ml). The ITR and VRC MICs were changed in these isolates, although not reaching clinical breakpoints (CAAL37: 0.125 and 0.06 \mu g/ml respectively; CAAL61: 0.25 and 0.25 \mu g/ml respectively). FLC resistance was also confirmed in vivo in a murine model of disseminated candidiasis. The median survival of mice groups receiving C. albicans isolates CAAL37, CAAL61 and CAAL97 in the absence of drug treatment, were respectively 2, 3 and 4 days. Virulence of C. albicans was not significantly different between these drug-free groups (p = 0.102 between groups). Whereas 5 mg/kg of FLC had a significant survival effect on mice infected by the susceptible CAAL97 isolate (median survival of 13 days after FLC treatment), both CAAL37 and CAAL61 clinical isolates showed a drastically reduced FLC susceptibility in vivo (median survival of 3 days). The Kaplan-Meier survival plot clearly revealed a significant mortality disadvantage of CAAL37 and CAAL61 isolates after treatment with FLC compared to the CAAL97 isolate (p = 0.02 vs CAAL97) (Figure S1).

**Construction of P. pastoris transformants expressing mutated and wild-type CaErg11p**

The general strategy of mutagenesis into P. pastoris is shown in Figure S2 and Figure S3. A single colony of each P. pastoris mutants, wild-type and mock controls was used for expression studies in shake flasks. After one day of cell growth on BMGY medium at 30°C, 250 rpm, the BMGY medium was then switched to BMMY medium containing 0.5% methanol. The yeasts were fed with methanol every 24 h for a period of 96 h. The transformed strain, with the parental pPIC3.5K plasmid, was used as a control.

SDS-PAGE shows that heterologous expression of the CaErg11p protein was successfully expressed into P. pastoris. After Coomassie gel staining, a band of around 56 kDa was detected in 0.5% methanol-induced clones in comparison to the mock control (Figure 1A). The overall levels of specific intracellular CaErg11p protein and wild-type control are likely to be relevant after Immunoblot with a polyclonal rabbit anti-Candida Erg11p analysis compared to the mock control (Figure 1B). No band was detected in the induced strain transformed with the parental plasmid (Mock control). With respect to protein dosage, CaErg11p expression seems to be lower in Y447H and V456I mutants than wild-type control under the same experimental conditions.

**Azole susceptibility of P. pastoris CaErg11p transformants**

The susceptibility of P. pastoris transformants expressing the wild-type and mutant CaErg11p proteins to azole compounds was tested by a microdilution reference method with some modifications. The K143R substitution was used as resistance-positive control according to the site-directed mutagenesis studies showing that this substitution was responsible of a 64-fold increase of FLC MIC. P. pastoris transformants containing single mutations (Y447H, V456I or K143R) presented no difference in their MIC to FLC following growth on a non-inducing medium, BMGY (data not shown). CaErg11p was then induced in BMMY during 72 h. The
susceptibility to azole compounds after induction was measured by modified-reference CLSI test with some modifications. Under these conditions, methanol was added to 96-well plates every 24 h. These experiments showed that growth and induction into a BMMY medium led to a significant increase of FLC MIC (Figure 2A). The MIC value for P. pastoris transformants containing the Y447H and V456I substitutions was 8 μg/ml for FLC, similar to K143R positive-control mutant. Compared to the wild-type and the parental plasmid transformants, these values represented an 8-fold increase in FLC resistance. The MICs to VRC and ITR following growth and induction into a BMMY medium of P. pastoris CaErg11p transformants containing single substitutions were similar, indicating that Y447H and V456I mutations were specific for the FLC binding to CaErg11p. The susceptibility of CaErg11p P. pastoris transformants to FLC was also qualitatively measured by spotting serial dilutions of each clone on BMMY medium containing various FLC concentrations (Figure 2B). Under these conditions, 0.5% methanol was added during CaErg11p pre-induction in BMMY and only at the beginning of spot tests. The wild-type and mock P. pastoris transformants were hypersensitive to FLC at 2 μg/ml, whereas the growth of Y447H, V456I and K143R P. pastoris mutants remained unchanged. After 72 h, P. pastoris mutants showed a better capacity to growth at high yeast dilution than controls when they were treated with 4 and 8 μg/ml FLC (c, d).

FLC inhibition of ergosterol biosynthesis in P. pastoris CaErg11p transformants

The effect of single substitutions on fluconazole resistance was determined by measuring the catalytic activity of P. pastoris transformants CaErg11p through the analysis of yeast sterol composition by gas chromatography-mass spectrometry. The activity was expressed as the ratio of ergosterol biosynthesis compared to the toxic lanosterol accumulation (E/L). As expected, under untreated conditions i.e without FLC (Figure 3A), sterol composition of wild-type and mock P. pastoris transformants, showed ergosterol as the dominant sterol (median E/L ratio of 64 for each control). No significant difference was observed for K143R, Y447H and V456I P. pastoris mutants (median E/L ratio of 35, 48 and 47, respectively). Under un-treated conditions a significant difference in E/L ratio was observed between P. pastoris mutants and wild-type control. The controls and mutants also showed small quantities of the intermediates of the ergosterol biosynthesis pathway, such as zymosterol, fecosterol and episterol.

The wild-type, mock and all CaErg11p P. pastoris transformants were then treated with FLC at two concentrations around their FLC MICs. Wild-type and mock P. pastoris transformants showed a marked reduction of sterol 14α-demethylase activity after treatment with FLC (E/L ratio = 1) which concords with their susceptibility to FLC. In contrast, the control mutant K143R maintained enzymatic activity despite FLC treatment with 4 and 8 μg/ml (median E/L ratio of 9.5 and 5.4, respectively). Interestingly, demethylated activity of Y447H and V456I mutants was significantly higher than wild-type control after treatment with 4 μg/ml of FLC as reflected in the E/L ratio of 11.5 and 11 respectively (Figure 3B). When they were treated with 8 μg/ml FLC, the E/L ratio were also significantly higher than wild-type control (6.4 for Y447H mutant and 9.3 for V456I) (Figure 3C).

CaCYP51 Insertion loop model and azole-resistant substitution mapping

CaCYP51 (P10613) was modeled using 3 different homology modeling programs (Swiss-model, ModWeb and YASARA). All CaCYP51 structure models, obtained with the different programs, present the same overall fold typical of CYP51 proteins. The best model obtained by ModWeb and YASARA used a human CYP11A, respectively ketoconazole (3ld6) and econazole (3jus) liganded versions presenting 45% sequence identity (residues 49–525). All the models differ essentially in the position in space of some connecting loops between alpha-helices of the all alpha-helicial domain of CYP51 and in the position of the large insertion loop characteristic of fungi phyla CYP51 family members. ModWeb and YASARA modeled the sequence D262E-TAAA-KANSVS8N6SDEVDYGFPGKVKVSPPSS9 of CaCYP51 protein as a long coil pointing out from the protein core (Figure 4A and 4B). A small sheet, consisting of two small beta-strands, is located at the tip of the loop in the YASARA model. The last good Swiss-model represented this loop as a long 2 beta-strands hairpin-like structure with the DYGF motif at the top (Figure 4C). This loop interacts with the alphaB helix and the loop containing the GGGHR (alpha-K') motif.

Interestingly, one of the two best ModWeb models using the CYP450 from zebrafish (3b98) represents this insertion with two small beta-stranded beta-sheet structures that exactly superpose CaCYP51 Y447 residue over the Y409 residue of the zebrafish version in close proximity to the catalytically important G448GGRHR motif (Figures 4D and S4). Figure 4 focuses the azole-resistant amino acid positions mapped on the hybrid models. More than 140 amino acid substitutions of CaCYP51 were reported in literature (reviewed recently by Morio F. and coll. [29]. Using a criterion of reduced susceptibility isolates and azole-resistance which have been experimentally proven, we localized Y132F, Y132H, N136Y, K143R, G307S, S405F, Y447H, G448E, G448V, G450E and V456I. Three of these positions involved the substitution of a glycine in a bulkier amino acid, possibly
demonstrating the importance of a reduced local flexibility for azole-resistance phenotype due to conformational restriction.

**Discussion**

The understanding of the molecular mechanisms underlying the resistance to azole in *C. albicans* is essential, both in providing guidance in the selection of the appropriate antifungal agent at the onset of infection, and in the discovery of new antifungals which circumvent azole resistance. Intense research in recent years has highlighted azole resistance as a complex molecular process in which multiple mechanisms are involved, including overexpression of the efflux pumps genes [12,20–22] and upregulation or substitution of the 14alpha-demethylase target enzyme CaErg11p [11,23,24]. Several of these mechanisms are frequently combined in a single clinical azole-resistant isolate. Sterol 14alpha-demethylase Erg11p encoded by the *Erg11* gene participates in ergosterol biosynthesis, a major sterol of the fungal cell membrane that is required for yeast viability, membrane fluidity and permeability [27]. From a medical perspective, Erg11p is the primary target ofazole antifungal drugs. The reduction ofazole affinity for Erg11p caused by amino acid substitutions is one of the most frequent mechanisms involved in azole resistance in *C. albicans* [28]. In a previous study, we identified two novel CaErg11p substitutions, Y447H and V456I possibly involved in azole resistance from two FLC-resistant (MIC = 64 μg/ml) clinical isolates (CAAL61 and CAAL37, respectively). These isolates presented other CaErg11p substitutions, G307S for CAAL61, and E266D, G464S, V488I for CAAL37 [29]. G307S has been previously identified in resistant clinical isolates [12]. G464S substitution has been confirmed to be the cause of FLC resistance in *C. albicans* due to reduced affinity for CaErg11p using site-directed mutagenesis [30]. G307S is often associated with other confirmed azole-resistant substitutions such as G450E and G464S.

In the first part of this study, we aimed to investigate the potential in vitro contribution of Y447H and V456I CaErg11p substitutions on FLC resistance. We used site-directed mutagenesis methodology of the wild-type CaErg11 gene and heterologous expression into *P. pastoris* transformants, in order to produce *P. pastoris* transformants expressing CaErg11p mutant proteins. We clearly demonstrated that both Y447H and V456I substitutions were responsible for an 8-fold increase in FLC MICs determined by a broth microdilution method when expressed into *P. pastoris* transformants compared to the wild-type parental plasmid controls using CLSI tests. That resistance level was equivalent to the K143R positive control mutant. Next, we examined the activity of CaErg11p containing single substitutions,
by measuring the sterol composition of *P. Pastoris*
transformants before and after FLC treatment. Under un-treated conditions, we observed a significant difference between wild-type, mock controls and mutated *P. pastoris* transformants. This suggests that punctual mutations (K143R, Y447H and V456I) possibly have an effect on CaErg11p expression. This would be particularly relevant regarding the lower CaErg11p expression by westernblot compared to controls. In spite of the lowest ergosterol formation by mutant clones, these punctual mutations represent an significant azole resistance advantage when they were treated by FLC compared to wild-type and mock controls. They showed a sustained capacity for producing ergosterol even in the presence of FLC. Seen together, these results clearly highlight the fact that Y447H and V456I are involved in fluconazole resistance confirming previous hypotheses with clinical isolates CAAL61 and CAAL37 [29].

Although, *Saccharomyces cerevisiae* is the most commonly employed model for studying the role of *C. albicans* Erg11p substitutions on azole resistance [31,41,42], we used in this study, the *P. pastoris* expression system as an alternative to the expression of mutated CaErg11p. Importantly, our results using *P. pastoris* displaying mutated CaErg11p with a modified broth microdilution method and sterol biosynthesis experiments, concord with the site-directed mutagenesis studies into *S. cerevisiae*. Moreover, *P. pastoris* has been proposed as an adapted expression system for mutagenesis studies [43–45]. Several elements supported these experiments: (i) the use of methanol inducible AOX1 promoter allows weak expression of native proteins; (ii) the selection of single copy-gene transformants avoids the over-expression of mutated proteins; (iii) *P. pastoris* leads to the production of post-translational modifications of mutated proteins such as glycosylation. Fourthly, in the area of directed evolution of CaErg11p by site-directed mutagenesis, the choice of an appropriate expression plasmid system is important in order to express and measure enzyme activity in their natural microsomal environment. Finally, *P. pastoris* allows the production and the

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**Figure 3. CaErg11p activity of *P. pastoris* transformants in the presence of FLC.** *P. pastoris* CaErg11p methanol-induced transformants were treated with FLC in BMMY medium for 24 h at 30°C. Non-saponifiable lipids (sterols) were extracted as described in Material and Methods. Sterol identification was done in reference to the relative retention times and mass spectra previously reported [54,55]. Activity results were expressed as the ratio of ergosterol biosynthesis compared to the lanosterol accumulation (E/L). (A): Untreated *P. pastoris* clones, (B): 4 µg/ml FLC and (C): 8 µg/ml FLC (n = 4).

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purifying high levels of recombinant proteins for crystallographic and structural analyses by using extracellular expression plasmid systems.

Because all the fungal CYP51 that have been characterized were membrane bound microsomal proteins, their crystal structures have been difficult to resolve to date. The impact of CaCYP51 substitutions on interaction with azole antifungals was explained after 3D homology modelling using the amino acid sequence and the crystal structure of *Mycobacterium tuberculosis* as a reference [36–38]. Using this reference, the involvement of Y447H, V456I and the other amino acid substitutions located at the fungi-specific insertion sequence is difficult to question. Here, we proposed, for the first time ever, a novel hybrid model (YASARA) keeping the general overall core structure of the known CYP51 ortholog structures as described so far. This model helped us to map experimentally proved azole-resistant mutants (see Figure 4). The insertion sequence specifically found in fungal CYP51 orthologs was modeled as a long coil pointing out from the surface of the core protein. Because of the absence of homologous fungal CYP51 crystal structures, the position and interaction of this loop with the secondary structure core protein is difficult to model. In each of the 30 models obtained with YASARA, as well as the two models obtained by ModWeb and the swissmodel, the variable position of the insertion suggests its inherent flexibility. The insertion is localized between K' alpha-helix and the long L alpha-helix following the conserved catalytic C470 residue. Moreover, the insertion containing 39 possible positions in the recently published complete alignment of CYP51 family members [34] varies greatly in length. Depending on the fungal species, the insertion varies from 37 amino acid residues (*Pichia kudriavzevii*) to 27 amino acids (*Schizosaccharomyces pombe*). *C. albicans* CYP51 having a 31 amino acid insertion. The half-N-terminal part of the insertion is variable in size and in sequence, unlike its half-C-terminal part which contains a short portion rich in acidic residues followed by an almost invariable sequence with the motif DYG[FY]Gx[VI][ST]KG. This motif is not shared with any protein, although it has a few similarities to a short sequence found in bacteria GTPase protein EngB.

Using the hybrid model, substitution Y447H is found to be solvent-accessible and is located in the middle of the invariable sequence portion of the insertion loop and could possibly modify the CaCYP51 local structure by disturbing the hydrogen binding

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**Figure 4. Localization of the major long insertion sequence of CaErg11p and azole-resistance substitutions mapping on CaErg11p structures obtained by homology modeling.** (A): ModWeb model with human CYP51 ketoconazole liganded (3ld6). (B): YASARA model with human CYP51 liganded with econazole (3jus) with protoporphyrin IX and econazole represented as a stick respectively in orange and yellow. (C): SwissModel with *M. tuberculosis* 4-phenylimidazole liganded structure as template (1e9x). (D): ModWeb model zebrafish prostacyclin synthase CYP450 Ba1 free (3b98). All the models are aligned, based on their secondary structure with the N-terminal part of the beta-strand rich domain left and the alpha-helices rich domain right. The secondary structure of the models are color-coded according to their type, red = beta-stands, blue = alpha-helices, green = turn, cyan = coil, magenta = the long fungi specific amino acid sequence insertion. Amino acid positions that have been proven to be responsible for azole-resistance are indicated with the amino acid name of the WT CaCYP51 with yellow for substrate binding site hotspots and magenta in the insertion fungi specific sequence. N- and C-terminal are labeled by single letters N and C.

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network. Moreover, we could predict a possible pH influence of the mutant, because of the possible environmental dependence protonation of the H447 nitrogen atom of the azole ring. Moreover, the modeling of the CaCYP51 insertion sequence with the structurally similar sequence F_{190}L{NADTRTEKDDFFK\_}GARVKY_{199}PSVP_{113} found in zebrafish CYP450 structure (3b9k) position, put the critical Y447 CaCYP51 in a position superposed over that of the Y409 zebrafish and in direct proximity to the catalytic important GGRHRRC motif. In contrast, the V456 position is located in the invariable sequence of the insertion loop. This position is mutated to isoleucine in almost 50% of the reported sequences in fungi phyla and could be described as a polymorphic position.

We note, with interest, that several other substitutions such as G448E, G448R, G448V, F449S, F449V and F449Y, previously associated withazole resistance, have been reported in this insertion region [15,23,24,28,47]. Moreover, protein sequence alignment of CaCYP51 with other fungal pathogens reveals that Y447 corresponds to Y431 in \textit{Aspergillus fumigatus}, a residue (Y431C) involved in pan-azole resistance in this fungal species [49]. Looking at these models we could make a hypothesis of a loop-secondary structure substrate- or an azole-dependent conformation transition that could give better substrate availability or enzyme stability. Moreover, the flexibility of the region is probably enhanced by the relatively high glycine content occurring in the consensus sequence. Interestingly, the Y447H mutant is characterized by FLC resistance but remains highly susceptible to voriconazole which has the best \(K_d\) value for the wild-type CaCYP51 protein [49]. As previously hypothesized for H71T [49], FLC resistance in the Y447H mutant could be attributed to reduced affinity for its target.

The fact that amino acids at the insertion loop are conserved in several fungal species, probably indicates their implication in substrate-azole interaction. Clearer insights into the resistance mechanism of these mutants could be obtained by \textit{in vitro} characterization of enzyme activities using purified wild-type and mutant proteins, recording circular dichroism and the \(1^H\)-NMR spectra of purified proteins in the presence of substrates/inhibitors in order to look for possible changes in secondary structure composition and by the elucidation of the \textit{C. albicans} CYP51 crystallographic structure.

The construction of a heterologous expression system of CaErg11p mutants and a molecular model of clinical relevant amino acid substitutions highlight the potential role of the CaErg11p insertion loop in azole resistance. The rationale of preliminary knowledge of the resistance mechanisms before the design of new azole antifungals candidates, proposed by this comprehensive investigation, should be taken into consideration for future studies aimed at designing new azole antifungals against the human pathogen \textit{C. albicans}.

Materials and Methods

Cells and vectors

\textit{Escherichia coli} TOP10F\textsuperscript{\textcopyright} (Invitrogen, Inc.) competent cells were used for the transformation and propagation of recombinant plasmids. The \textit{pBluescript} (SK\textsuperscript{-}) (Stratagene GmbH) plasmid was used for \textit{CaErg11} gene cloning and site-directed mutagenesis. The \textit{pPIC3.5K} plasmid (Invitrogen) was used for sub-cloning and for the intracellular expression of \textit{CaErg11} protein mutants into the \textit{P. pastoris} strain (KM71, \textit{mut1}, \textit{arg4}, \textit{his4}). The \textit{mut1} designation «methanol utilization slows» phenotype was due to the loss of \textit{AOX1} gene. The gene coding for \textit{CaErg11p} was integrated behind the \textit{AOX1} promoter.

Reagents

Restriction enzymes BamHI, NotI, SacI, EcoRI, T4 ligase, calf intestinal phosphatase (CIP) and DNA molecular weight markers were purchased from New England Biolabs Inc. Phusion high fidelity DNA polymerase and buffers came from Finnzymes. GoTag Flexi DNA polymerase was from Promega. Agarose was purchased from invitrogen.

The Nuclease plasmid preparation kit and the Nucleospin Extract II DNA fragment extraction kit from agarose gel were purchased from Macherey-Nagel GmbH. Acid washed beads, genetin G418, ampicillin, kanamycin antibiotics, BCA assay reagents, methanol, n-hexane and glycerol and all oligonucleotide primers were obtained from Sigma Aldrich Co. Molecular weight marker for SDS-PAGE, electrophoretic transfer cell for western blotting and SDS-PAGE equipment were purchased from Biorad. A polyclonal anti-\textit{Candida} Erg11p was kindly provided by Dr. Diane Kelly from the Institute of Life Science, School of Medicine, Swansea University, Swansea, UK. Anti-rabbit HRP conjugate was purchased from Cell Signaling Technology Inc. Super signal West Pico substrate solution was purchased from Thermo Scientific. FLC was purchased from Sigma, itraconazole (ITR) from Janssen-Cilag, and voriconazole (VRC) from Sigma Aldrich Co.

\textbf{In vitro and in vivo FLC susceptibility of \textit{C. albicans} clinical isolates}

Antifungal susceptibility to FLC, ITR and VRC was determined \textit{in vitro} for \textit{CAAL37} and \textit{CAAL61} clinical isolates using the broth microdilution reference method, as recommended by the Clinical and Laboratory Standards Institute (CLSI) document M27-A2 [50]. MIC, that is the lowest drug concentration that resulted in 50% growth inhibition relative to the growth in the control well, was visually determined after 48 h of incubation at 35°C.

The \textit{in vivo} antifungal activity of FLC for the \textit{CAAL37} and \textit{CAAL61} clinical isolates was evaluated in a mouse disseminated candidiasis infection model, according to a previous study [51] with the following modifications. \textit{CAAL97} isolate was used as a FLC susceptible control. Briefly, four-week-old female Swiss mice purchased from Elevages Janvier, were housed at the Experimental Therapy Unit (Faculty of Medicine, Nantes, France). Immunosuppression in mice was induced by subcutaneous administration of prednisolone (30 mg/kg; Sigma Aldrich Co) one day before infection. Disseminated candidiasis was induced by intravenous infection with \(5 \times 10^5\) \textit{C. albicans} cells (\textit{CAAL37}, \textit{CAAL61} and \textit{CAAL97} clinical isolates) in 0.1 ml suspension. Fluconazole (5 mg/kg) was then administered orally once a day, for 5 days, starting 1 h after infection. Virulence control groups were inoculated with \textit{C. albicans} isolates and were treated with PBS as described above. Survival was monitored every day, for two weeks after infection. The survival rate was compared to the control group by using the logrank test and a \(p\) value of less than 0.05 was considered significant.

C. albicans 14 α-Demethylase and Azole Resistance

Culture medium for bacteria and yeast

Luria broth medium supplemented with 100 µg/ml ampicillin was used for our bacterial culture. 10X YNB: 13.4% yeast nitrogen base with ammonium sulfate without amino acids. YPD medium: 1% yeast extract, 2% peptone, 2% dextrose. YPD-geneticin plates: the same as YPD plus 2% agar and variable amounts of geneticin. Minimal glycerol medium without histidine (MGY): 1.34% YNB, 1% glycerol, 4 \(\times 10^{-5}\) \% biotin. Regeneration dextrose medium without histidine (RD): 1 M sorbitol, 2% dextrose, 1.34% YNB, 4 \(\times 10^{-5}\) \% biotin, 0.005% amino acids. RDB plates: the same as RD with 2% agarose. Buffered glycerol-
complex medium (BMGY): 1% yeast extract, 2% peptone, 100 mM potassium phosphate, pH 6.0, 1.34% YNB, 1% glycerol. Buffered-methanol complex medium (BMMY): the same as BMGY with 0.5% methanol replacing glycerol. Buffering bufer: 50 mM sodium phosphate, pH 7.4, 1 mM PMSF, 1 mM EDTA, 5% glycerol.

Isolation and cloning of CaErg11 gene into pBluescript(SK-)

The CaErg11 gene was isolated from a wild-type C. albicans strain isolated from the Mycobank of the Parasitology and Medical Mycology Department, Nantes, FR. The ORFBamHI and ORFNotI primers containing a Kozak consensus sequence, were designed to amplify a 1745 bp cDNA fragment based on the reported sequence from the GeneBank nucleotide sequence database accession number X132961.1. PCR conditions were as follows: initial denaturation at 98°C for 5 min, followed by 35 cycles each at 98°C for 1 min, 58°C for 1 min and 72°C for 1 min. Finally one cycle was performed at 72°C for 10 min.

The CaErg11 gene fragment and pBluescript(SK-) were prepared for restriction endonuclease digestion with BamHI and NotI. These primers facilitated the sub-cloning of the coding sequence into pPIC3.5K. The plasmid (10 µg) and cDNA fragment (1 µg) were mixed separately with 10 U/µg BamHI, 20 U/µg NotI, 5 µl 10X BSA, 5 µl 10X NEB3 buffer to a final volume of 50 µl. After incubation for 1 hour at 37°C, the digests were purified on 1% agarose gel by using the NucleoSpin Extract II kit (Macherey-Nagel). For cloning, 20 ng of digested pBluescript(SK-) and 60 ng of pBluescript(SK-) mutant coding region was sequenced using internal primers [52,53].

Site-directed mutagenesis

Site-directed mutagenesis was carried out using a QuikChange mutagenesis kit (Stratagene). The cloned CaErg11p gene (into pBlueScript(SK-)) was used as the starting material for constructing all the mutants. Mutagenic primer sequences for studying Y447H and V456I substitutions are presented in Table 2. The K143R substitution was selected as resistance-positive control according to the site-directed mutagenesis studies published by Chau et al., 2004 [15]. This substitution was responsible of a 64-fold increase of FLC substitution was selected as resistance-positive control according to site-directed mutagenesis studies published by Chau et al., 2004 [15]. This substitution was responsible of a 64-fold increase of FLC

Construction of recombinant Pichia pastoris transformants

The general strategy of mutagenesis into P. pastoris is shown in Figure S2. The CaErg11 gene was isolated from pBluescript(SK-) by using BamHI and NotI enzymes and purified after agarose gel electrophoresis with a NucleoSpin Extract II kit. Purified fragment was subcloned into the P. pastoris intracelluar expression plasmid pPIC3.5K. The recombinant clones were confirmed through PCR analysis with ORFBamHI and ORFNotI primers and sequencing (Figure S3.A).

The recombinant pPIC3.5K-CaErg11 was used to transform P. pastoris KM71 strain by electroporation. Briefly, 10 µg of recombinant plasmids were linearized by 10 U/µg SacI leading to targeting plasmid for the P. pastoris chromosome at the AOX1 locus. Linearized plasmid were purified from agarose and mixed with 80 µl competent P. pastoris cells. The mixture was transferred into an ice-cold 0.2 cm cuvette and an electric shock was given at 2 Kvolts for integration into the P. pastoris genome. Then, 1 M sorbitol was immediately added. The 0.6 ml of mixture was poured onto the top of RDB plates and incubated at 30°C until colonies appeared. The positive transformants that produced histidine were screened for the ability to grow on YPD-geneticin plates ranging from 0.25 mg/ml to 2 mg/ml geneticin. The yield of plasmid transformation was about 3000 colonies per transformation. Thirty colonies of pPIC3.5K-CaERG11-K143R, 27 colonies of pPIC3.5K-CaERG11-V447H, 17 colonies of pPIC3.5K-CaERG11-Y447H and 20 colonies of pPIC3.5K-CaERG11-wild-type were gathered from the YPD-geneticin plates at the 0.5 and 0.75 mg/ml concentrations, in order to test single-copy colonies. The His"Mut" (methanid utilization slow) transformants were picked onto RDB plates and resuspended in YPD containing 15% glycerol and stored at −80°C.

Analysis of P. pastoris recombinant transformants

The geneticin-resistant colonies were also grown in YPD for 24 h, then the genomic DNA was purified. PCR amplification of the CaErg11 gene was carried out with 5’ AOX1 and 3’ AOX primers. Another PCR analysis was also done using 5’ AOX1 and a CaErg11 internal R2 primer. The PCR product of recombinant clones was a band at 1956 bp corresponding to the CaErg11 gene (1745 bp) and a fragment of pPIC3.5K plasmid (214 bp) [Figure S3,B]. A band at 714 bp corresponding to a piece of AOX1 promoter (30 bp) and 664 bp of the 5’ extremity of the CaErg11 gene indicates that all P. pastoris transformants were recombinated [Figure S3,C]. Recombinant P. pastoris with the parental plasmid was used as PCR control.

Sequencing analysis

DNA sequencing was performed using an Applied Biosystems 3730 sequencer using internal primers [53]. Nucleotide sequences were assembled using Seqscape navigator software (Applied Biosystems). For each mutant, the entire CaErg11 open reading frame sequence

Table 2. Oligonucleotide primers used in this study.

| Primer | DNA sequence |
|--------|--------------|
| ORFBamHI | 5’-CCGATCACATTTGTGTGAATAGTG-3’ |
| ORFNotI | 5’-GATCGCCGCTTAAAAACATACAGTTTCTTTTT-3’ |
| K143Rfor | 5’-GGACAAAGAAGTCTTGAATTTGCG-3’ |
| K143Rrev | 5’-GAAATTTAGACGACTTCATGTTCC-3’ |
| Y447Hfor | 5’-GATGAGATGATGACGTGGTTG-3’ |
| Y447Hrev | 5’-CCGAAACCGTTAGCAGCCTTTTAC-3’ |
| V456Ior | 5’-CTTCTGAAGGAGCTTTTCGATAC-3’ |
| V456Irev | 5’-GATGAAAGAATCCTTTGAGAAA-3’ |
| CaCYP51R2 | 5’-AATATGTTGCGAATAATGAAACG-3’ |
| 5’AOX1 | 5’-GACTGTCTCAATTGCAAAGCG-3’ |
| 3’AOX1 | 5’-GCAAATGCGTCTGACATC-3’ |

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was compared to a previously described CaErg11 sequence (accession number X13296). The Erg11 sequences of the strains displaying new amino acid substitutions have been submitted to GenBank database under accession numbers EU385935 (CAAL37 isolate) and EU385936 (CAAL61 isolate). An exhaustive list of amino acid substitutions of each clinical isolate used in this study are presented in Table 1. The alignment data indicates that Y447H substitution was homonymous and V456I heteronymous.

Small scale intracellular expression of recombinant CaErg11p proteins

P. pastoris cells transformed with wild-type, mock, K143R, Y447H and V456I mutants of the CaErg11p protein were analyzed for protein expression. Briefly, single colony transformants from RDB plates were inoculated into 1 ml of BMGY medium and incubated, whilst being agitated, at 30°C to reach an absorbance of 2 at 600 nm (18 h), then this 1 ml was inoculated into 10 ml of BMMY medium at 30°C. Incubation was prolonged for 96 h with the addition of methanol (0.5% v/v final concentration) every 24 h. Cells were collected every day by centrifugation at 3000 g for 10 min and analyzed for the expression level, protein activity and absorbance at 600 nm.

Western blot analysis

For evaluation of the intracellular accumulation of proteins, cell pellets were resuspended in 100 µl ice-cold lysis buffer. Cells were mashed with 100 m of glass beads (0.2 mm) by doing several 30 s vortex applications and a cooling cycle. Cell-mashed slurry was centrifuged at 800 g for 5 min at 4°C. The collected supernatants (soluble cytosolic proteins), were centrifuged at 13000 g for 10 min at 4°C. The protein content of clear supernatant was quantified using BCA assay. Forty µg protein per well were loaded into 10% SDS-PAGE gels. Electrophoresis was performed according to the Laemmli method. After electrophoresis, the gels were stained with Coomassie brilliant blue R-250 and destained with a mixture of acetic acid-methanol-electrophoresis, the gels were stained with Coomassie brillant was performed according to the Laemmli method. After electrophoresis, the gels were stained with Coomassie brilliant blue R-250 and destained with a mixture of acetic acid-methanol-water (10:25:65 v/v/v). Proteins were also transferred to PVDF membranes and incubated with a 1:100 dilution of polyclonal anti-Candida Erg11p produced in rabbit overnight at 4°C and a 1:2000 dilution of goat-anti rabbit antibody coupled to HRP 1 h at room temperature. The signals were visualized using the super signal west pico staining reagent.

Aazole susceptibility tests for P. pastoris transformants

The susceptibility of P. pastoris transformants expressing the wild-type and mutant CaErg11p proteins to azole compounds was tested using two methods. First, a broth microdilution reference method, as recommended by the Clinical and Laboratory Standards Institute (CLSI) document M27-A2 [50]. In some experiments RPM11640 medium, which is usually used in this method, was replaced by BMMY liquid medium. A final concentration of 0.5% methanol at was added every 24 h to each well in order to guarantee protein induction. The MIC was defined as the antifungal concentration giving a 50% or less reduction in the optical density at 450 nm compared to the OD of the corresponding drug-free incubation medium. Secondly, a qualitative test was carried out by spotting serial dilutions of induced yeast onto agar plates containing BMMY medium. The azole compounds were diluted at different concentrations into BMMY plates. The P. pastoris transformants were induced at 30°C for 72 h in BMMY liquid medium and diluted to 2 x 10⁹ cells per ml in 0.9% NaCl. Five microliters of this suspension and 5 µl of serial dilutions of each yeast culture were spotted onto each type of plate. The plates were incubated for 72 h and 96 h at 30°C.

FLC effect on ergosterol biosynthesis by P. pastoris CaErg11p transformants

After P. pastoris CaErg11p induction for 72 h at 30°C in BMMY medium, yeast cells were harvested and treated with various concentrations of FLC in BMMY medium for 24 h at 30°C. The treated cells were then harvested and resuspended in 3 ml of 60% (wt/v) KOH and saponified by heating at 80°C for 2 h. Non saponifiable lipids (sterols) were extracted from the saponified mixture, twice, with 2 ml of n-hexane pooled, and dried under nitrogen. The sterols were suspended in 100 µl of bis(trimethylsilyl) trichloride for 30 min for silylation. The silylated sterols were analyzed by gas chromatography-mass spectrometry (Agilent Technologies). The sterol identification was done in reference to the relative retention times and mass spectra previously reported [54,55] (n = 4).

CaCYP51 homology modeling

CaCYP51 (P10613) was modeled using 3 different homology modeling programs. The Swiss-model repository already contains a CaCYP51 model using Mycobacterium tuberculosis 4-phenylimidazole liganded structure as a template (1le9x). 2 homology models were retrieved from the server version of modeler (ModWeb) with human CYP51 ketocamazole liganded (3bo6) and zebrafish prostacyclin synthase CYP450 8a1 free (3989) as templates. Finally, the homology modeling module of the YASARA program (www.yasara.org) [56] using the latest CaCYP51 crystallographic structures that have been resolved, was used. YASARA homology modeling uses 6 different crystallographic structures as templates from different phyla, M. tuberculosis liganded (2w9b and 2w0a), human econazole liganded (2ps2) and T. brucei liganded (3g5q and free (3g5q), C. elegans liganded (2w2x) selected automatically by PSI-BLAST optimized sequence alignments with the target sequence. After secondary structure prediction, loop construction and amino acid rotamers selection, molecular dynamic steps were performed with the YASARA default force field parameters and the 6 energetically lowest structures per template were retained after stereochemistry validation. A total of 30 models were then scored, sorted by Z-score and a unique hybrid model consisting of the best part of the 30 models was obtained. This hybrid model was submitted to a final molecular dynamics round and its quality evaluated and retained if its pertinence had improved. The effects of F447H and V456I mutations at a molecular level were then further analyzed using the Foldx plugin within the program YASARA.

Ethics Statement

This study was carried out in strict accordance with the recommendations of the Directive 86/609/EEC on the protection of animals used for experimentals and other scientific purposes. The protocol was approved by the Committee on Ethics of Animal Experiments of the Experimental Therapy Unit (UTE) of the Faculty of Medicine, University of Nantes, France (C-44015). The mice were anesthetized by inhalation of an isoflurane-air mix (2%, 1 l/min) before any manipulation and all efforts were made to minimize suffering.

Supporting Information

Figure S1 FLC susceptibility of clinical isolates. The In vivo antifungal activity of FLC against the clinical isolates was evaluated in a mouse disseminated candidiasis infection model. Briefly,
four-week-old female Swiss mice were immunosuppressed by subcutaneous administration of prednisolone (30 mg/kg) one day before infection. Disseminated candidiasis was induced by intravenous infection with 5 x 10^7 C. albicans cells. After 1 h after infection, mice were treated either with PBS or FLC (5 mg/kg) orally once a day, for 5 days. Survival was monitored every day, for two weeks after infection. The survival rate was compared to the control group by using the logrank test and a p value of less than 0.05 was considered significant. CAAL37, CAAL37+ FLC, CAAL61, CAAL61+ FLC, CAAL97, and CAAL97+ FLC were determined by PCR amplification of the CaErg11 gene with 5’ AOX1 and 3’ AOX primers. (C): PCR amplification of the CaErg11 gene with 5’ AOX1 and a CaErg11p internal R2 primer. Lane 1: CaErg11-K143R, lane 2: CaErg11-Y447H, lane 3: CaErg11-V456I, lane 4: CaErg11-wild-type and lane 5: PCR of pPIC3.5K transformed P. pastoris. (EPS)

**Figure S3** Construction of recombinant Pichia pastoris transformants. (A): Integration of CaErg11 mutant genes into mutagenesis pBSSK(-) plasmid confirmed by PCR using ORF BamHI and ORF NcoI primers. Lane 1: pBSSK-CaErg11-K143R, lane 2: pBSSK-CaErg11-Y447H, lane 3: pBSSK-CaErg11-V456I, lane 4: pBSSK-CaErg11-wild-type and lane 5: PCR of pBSSK(-) mock control. (B): The genomic DNA from representative P. pastoris transformants was purified. PCR amplification of the CaErg11 gene with 5’ AOX1 and 3’ AOX primers. (C): PCR amplification of the CaErg11 gene with 5’ AOX1 and a CaErg11p internal R2 primer. Lane 1: CaErg11-K143R, lane 2: CaErg11-Y447H, lane 3: CaErg11-V456I, lane 4: CaErg11-wild-type and lane 5: PCR of pPIC3.5K transformed P. pastoris. (Eps)

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**Author Contributions**

Conceived and designed the experiments: NA-R PLP. Performed the experiments: NA-R AF FM. Analyzed the data: NA-R AF FM FP LG. Contributed reagents/materials/analysis tools: PLP. Wrote the paper: NA-R PLP.

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