Candida albicans Mrv8, is involved in epithelial damage and biofilm formation

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One sentence summary: Mrv8 is involved in mycelial maturation, epithelial damage and biofilm-associated caspofungin tolerance.

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ABSTRACT

Candida albicans is the most common human fungal pathogen that can cause superficial and deep-seated infections in susceptible individuals. Despite its medical importance, the vast majority of C. albicans genes remain of unknown function. Here, we report a role for the lineage-specific gene, MRV8, in host pathogen interactions, mycelial microcolony maturation and biofilm formation. In silico analysis indicated that MRV8 encodes a four-pass transmembrane protein unique to the closely related pathogens C. albicans and Candida dubliniensis. Deletion of MRV8 did not affect C. albicans adherence to, or initial invasion into human oral epithelia, but inhibited mycelial development and strongly reduced epithelial damage. mrv8Δ/Δ cells exhibited a media-dependent defect in biofilm formation and mutant biofilm metabolic activity was enhanced by cyclosporin A. mrv8Δ/Δ biofilms were more tolerant to treatment with caspofungin, but not to fluconazole or amphotericin B. Co-stimulation with calcium chloride and calcofluor white rescued biofilm growth in the presence of caspofungin, and this rescue-effect was Mrv8-dependent. Together, our data demonstrate an important role for a lineage-specific gene (MRV8) in C. albicans biofilm formation, drug tolerance and host–pathogen interactions.

Keywords: Candida albicans; biofilm; Marvel family; caspofungin; host-pathogen interaction; calcium homeostasis

INTRODUCTION

Candida albicans is normally a commensal member of the human microbiota (mycobiota) and can cause disease when an imbalance in host immunity or a disturbance of the normal microbiota composition allows the transition to a pathogenic lifestyle (Gow and Hube 2012). The pathogenicity of this fungus is mediated by a number of factors including adhesion to host cells, invasion and damage, which is predominantly caused by secretion of the cytolytic toxin, candidalysin (Moyes et al. 2016). Biofilms also play an important role in pathogenicity as they can serve as reservoirs, driving recurrent candidaemia.

Biofilms are surface-associated communities of cells encased within an extracellular matrix. Biofilm development emerges in
several phases and can be divided into distinct stages. In early stages, yeast cells adhere to the substratum forming the biofilm basal layer, followed by co-aggregation of cells and colonization. The intermediate stage is characterized by the formation of hyphae and pseudohyphae and these filamentous cells continue to develop to the mature stage when the biofilm exhibits a three-dimensional structure composed of all three morphological cell types embedded in extracellular matrix material. Finally, disperser cells can be released from the biofilm to reintegrate the process, or serve as an infection reservoir (For reviews, see: Mathe and Van Dijck 2013; Desai, Mitchell and Andes 2014; Ramage, Robertson and Williams 2014; Nobile and Johnson 2015; Gulati and Nobile 2016; Soll and Daniels 2016).

Candida albicans biofilms can grow on host tissues, such as epithelia and on abiotic medical and dental devices. The organization of the biofilm structure protects its inhabitants from attack by host immunity and biofilms can exhibit extremely high resistance to antifungal drugs. Indeed, C. albicans biofilms can exhibit 30 to 200 times higher antifungal drug resistance than planktonic cells (Hawser and Douglas 1995). Biofilm cells can be isolated and cast, or serve as an infection reservoir (For reviews, see: Mathe and Van Dijck 2013; Desai, Mitchell and Andes 2014; Ramage, Robertson and Williams 2014; Nobile and Johnson 2015; Gulati and Nobile 2016; Soll and Daniels 2016).

MATERIALS AND METHODS

Strain construction

For generation of a strain containing the open reading frame and 376 base pairs upstream and 282 base pairs downstream sequences were amplified from SCS314 C. albicans genomic DNA using the Phusion High Fidelity DNA Polymerase Kit (Finnzymes). The PCR product was purified with the QIAquick PCR Purification kit (Qiagen) and then cloned into the Pcr2.TOPO vector (Invitrogen). The plasmid was digested with SalI and MluI, following gel extraction with the QIAquick Gel Extraction kit (Qiagen). The plasmid CIP10 was also digested with SalI and MluI and the restriction enzymes were heat inactivated by incubation at 65°C/20 min. Next, the linearized plasmid was dephosphorylated with calf intestinal alkaline phosphatase (New England BioLabs) and gel extracted using the QIAquick Gel Extraction kit (Qiagen). The MRV8 insert and CIP10 plasmid were ligated at 25°C/30 min using the Quick Ligation Kit (New England BioLabs). Five μl of the ligation product was used to transform Escherichia coli DH5α and the positive clones were selected on LB agar [1% bacto-tryptone, 0.5% yeast extract, 1% NaCl, 2% agar] plates supplemented with 50 μg ml⁻¹ ampicillin. The plasmid carrying the insert was re-isolated using the miniprep (Peqlab) and midiprep (Qiagen) kit and controlled by digestion with Stul, Ncol, and Mfet. The plasmid was digested with Stul prior to the transformation into the uridine auxotrophic C. albicans strain mrv8Δ/Δura-. Positive clones were selected on SD agar [2% glucose, 0.17% yeast nitrogen base, 0.5% ammonium sulfate, 2% agar] plates without amino acids and correct integration was verified by PCR using primers RPF1/URA3-F2 and MRV8-INT-F/RPF1-R1. The strains and primers used in this study are listed in Table 1 and Table 2, respectively.

Biofilm development

Biofilms were grown as described previously (Nobile and Mitchell 2005; Junqueira et al. 2011). The strains were cultivated in YPD [1% yeast extract, 2% bacto-peptone, 2% D-glucose] at 30°C/24 h. The cultures were adjusted to OD600 at 0.5 in Spider medium [1% nutrient broth, 1% mannitol, 0.4% K2HPO4, pH 7.2] and 250 μl was added to the wells of a 96-well flat-bottom microwell plate (TPP) followed by incubation at 37°C for 1.5 h with shaking at 75 rpm (Quimia). After washing the wells, fresh medium was added and the plate was incubated at 37°C/48 h at 75 rpm. The medium was changed every 24 h.

The protocol for XTT colorimetric assay was described by Jin et al. (2003). Briefly, the biofilms were washed twice and treated with 197.5 μl of PBS, 50 μl of XTT (1 mg ml⁻¹) (Sigma-Aldrich) and 2.5 μl of menadione (0.4 mM) (Sigma-Aldrich) in the dark at 37°C/3 h. Next, 100 μl of the supernatant was transferred to a new well and the colorimetric change was measured at 492 nm using a microtiter plate reader (Thermo Plate, TP Reader NM).

Scanning electron microscopy (SEM)

The biofilms were grown on gamma radiation-sterilized and 24 h fetal bovine serum (FBS) (Sigma-Aldrich) pre-treated polystyrene discs measuring 8 mm in diameter and 2 mm in thickness in 24-well plates (TPP) for 60 h at 37°C with agitation. The biofilm method was performed as described above with media and PBS adjusted to 2 ml. First, the biofilms were photographed for visual appearance analysis and, then, fixed in 2.5% glutaraldehyde for 1 h and dehydrated by alcohol solutions with increasing concentrations of ethanol (10%, 25%, 50%, 75% and 90% for 20 min, and 100% for 1 h). The discs were dried at 37°C/24 h (Costa et al. 2011). The discs were then transferred to aluminum stubs and covered with gold for 120 s at 40 milliamps (BAL-TEC 50D 050 Sputter Coater) and imaged with Jeol JSM5600.

Stress susceptibility

The spotting assay was carried out according to the method of Wilson et al. (2014) on SD or YPD agar containing 1.5, 2 and 2.5 M NaCl (Roth), 1.5, 2, 2.5 M Sorbitol (Roth), 2 mM H2O2, 450 μg ml⁻¹ Congo red (Sigma-Aldrich), 800 μg ml⁻¹ calcofluor white (CFW) (Sigma-Aldrich), 3% ethanol, 0.1%–0.6% SDS (Sigma-Aldrich) and incubated at 37°C for up to 7 days. For UV stress, plates were exposed to 5 mJ of UV-C light using a UV-crosslinker at 254 nm.
Table 1. C. albicans strains used in this study.

| Strain       | Genotype                                                                 | Reference       |
|--------------|---------------------------------------------------------------------------|-----------------|
| BWP17 + Clp30| ura3::Δ/Δimm434/ura3::Δ/Δimm434 arg4::hisG/arg4::hisG                     | Mayer et al. (2012) |
| mrv8Δ/Δ      | orf19.3908Δ::ARG4/orf19.3908Δ::HIS1 + Clp10 (URA3)                       | Wilson et al. (2014) |
| mrv8Δ/Δ::MRV8| orf19.3908Δ::ARG4/orf19.3908Δ::HIS1 + Clp10 (ORF19.3908, URA3)           | This study      |

Table 2. Primers used in this study.

| Primer       | Sequence                                                                 | Reference       |
|--------------|---------------------------------------------------------------------------|-----------------|
| MRV8-Rec-F   | GTCGACAGTAATCTATACATTACACAACC                                               | This study      |
| MRV8-Rec-R   | ACGGCTATACATCAGGGATGTTGC                                                    | This study      |
| MRV8-Int-F1  | GTTGGGCTATTGGCGTTATTTTG                                                    | This study      |
| RPF-F1       | GAGCAGTGTTACACACACACATCTTG                                                  | Wilson et al. (2014) |
| RPF1-R1      | CGCCAAAGAGTTCITATTATAC                                                     | This study      |
| URA3-F2      | GAGATTGGATTAGTAGATGAAAAGTGATGG                                              | Gola et al. (2003) |
| HIS-F2       | GGAGTTGGATTAGAGGGTGCAACC                                                   | Gola et al. (2003) |
| HIS-R2       | CAAGCAATGGAAGAGGCTGTGCAACC                                                  | Gola et al. (2003) |
| ARG4-F2      | GATATGTTGGCTACTGATTAC                                                      | Martin et al. (2011) |
| ARG4-R2      | AATGGAATCTGTTGGCAACC                                                      | Gola et al. (2003) |

(Bio-Link, Vilber-Lourmat) and incubated at 37°C for up to 7 days. Plates were incubated at 42°C for 4–6 days to assess growth under thermal stress. Heat shock was performed by incubating serial 10-fold dilutions (range 10^6–10^1) at 55°C for 10, 15, 20 and 30 min, followed by spotting onto YPD agar plates and incubation at 37°C for up to 7 days. Each experiment was performed at least twice.

Growth assay

For growth assays under nutrient limitation, agar containing 0.67% YNB plus ammonium sulfate without amino acids (Difco) was supplemented with 2% glucose, 2% potassium acetate, 2% glycerol, 2% galactose, 2% maltose or 2% olive oil as carbon source. For alternative nitrogen sources, agar containing yeast carbon base medium (Difco) was supplemented with 0.5% BSA (Sigma-Aldrich), lysine (Sigma-Aldrich), histidine (Sigma-Aldrich), glycine (Sigma-Aldrich), cysteine (Sigma-Aldrich) or methionine (Sigma-Aldrich) at a concentration of 1 nM. Spot dilution assays (range 10^6–10^1) were prepared and plates were incubated at 37°C/3-7 days depending on the carbon and nitrogen source. All experiments were repeated at least twice.

Filamentation assays

For the filamentation assays, the cells grown overnight in YPD were washed twice in PBS and the cell density was determined using a hemocytometer. Filamentation on solid media was performed in the following media and incubation times: Spider agar at 30 or 37°C/7 days; agar supplemented with 10% FBS at 30 or 37°C/7 days; YPS agar [1% yeast extract, 2% bacter-peptone, 2% D(+)-saccharose, 2% agar] seeded by embedding at 25°C/5 days (Chaput et al. 1983). The colonies were analyzed visually and the experiments were performed on two different occasions.

For filamentation analysis in liquid media, 5 × 10^4 fungal cells were added per well in a 12-well cell culture plate containing 10% FBS, YPD, YPD plus 10% FBS, Spider medium, Lee’s medium at pH 6.5, RPMI-1640 medium (Invitrogen) or Dulbecco Modified Eagle Medium (DMEM) and the plates incubated at 37°C/3 h, with 5% CO2 for RPMI and DMEM. The percentage of filament forms was determined by microscopy observation and the length of the filaments was directly measured using LAS software (Leica Application Suite). The experiments were performed in duplicate and repeated twice.

Induction of filamentation was carried out on oral epithelial cell monolayers (TR146) (Nicholls et al. 2009). These cells were infected with 10^5 C. albicans cells and incubated at 37°C/3-6 h with 5% CO2 atmosphere. The hyphal cells were then differentially stained according to the invasion assay protocol described below and the length of the hyphae measured using the LAS software (Leica Application Suite). The experiment was performed in triplicate on two different occasions.

For induction of hyphal micro-colonies, approximately 50 fungal cells were added to oral epithelial cell monolayers (TR146) grown according to the invasion assay protocol described below. The plates were incubated at 37°C/24 h in the presence of 5% CO2. Next, the cells were fixed with 4% paraformaldehyde and stained with calcofluor white solution. Control micro-colonies were grown on plastic and fixed with 4% paraformaldehyde prior to observation. The dimensions of at least 50 micro-colonies per strain were determined using an inverse microscope (Leica DMIL) and the LAS software (Leica Application Suite). The tests were performed on three different occasions.

Around 100 Candida cells were plated on a layer of Spider agar or 10% FBS containing 1.5 or 4% agar and subsequently overlaid with a second layer of medium for invasive growth analysis. Plates were incubated at 30 and 37°C/7 days. The colonies were examined and photographed. The experiment was performed in duplicate on two different occasions.

Antifungal tests on biofilm

Susceptibility of biofilms to antifungal drugs and growth in the presence of different substances were verified using the method described by Pierce et al. (2008). Briefly, overnight YPD cultures were washed and resuspended in pre-warmed RPMI-1640 buffered with MOPS (165 mM, pH 7.0) and adjusted to a concentration of 1 × 10^6 cells ml^-1. One hundred microliters of the inocula were added to the well of a 96-well plate and incubated...
at 37°C/24 h statically. Afterwards, 200 μl of the antifungal solutions in RPMI was added to the washed biofilms and the plates were incubated for additional 24 h at 37°C. The biofilms were exposed to the chemicals over the developing time, 48 h. After the treatments, the biofilm metabolic activity was determined by XTT assay as described above. To determine the quantity of dispersed cells, the plate was shaken in an orbital shaker for 5 min (Solab). Aliquots of serial dilutions from the supernatant were plated onto Sabouraud dextrose agar plates and incubated at 37°C/48 h to determine colony forming units.

The antifungal drugs fluconazole, amphotericin B and caspofungin were used ranging from 2-1024 mg l⁻¹, 0.03125-16 mg l⁻¹, and 0.0625-32 mg l⁻¹, respectively. The biofilms were treated with the chemicals 75 μg ml⁻¹ FKS06, 20 μM nikkomycin Z, 0.025 M CaCl₂, 12.5 μg ml⁻¹ CFW and 75 μg ml⁻¹ Cyclosporin A, all of which were purchased from Sigma-Aldrich and the stock solution prepared in distilled water, except for Cyclosporin A and amphotericin B that were diluted in DMSO (Sigma-Aldrich).

**Oral epithelial cells: adherence, invasion and damage**

The human buccal carcinoma epithelial cell lineage TR-146 (Cancer Research Technology, London) (Rupniak et al. 1985) was used in the following methods and cultivated as described by Wilson et al. (2014).

Invasion rates of the strains were determined and analyzed as previously described by Park et al. (2005) and Mayer et al. (2012).

For adherence assay, the counting of adhered cells was performed using the slides prepared for the invasion assay at the time point of 3 h. The number of adhered cells was determined by counting 50 high power fields of 150 x 150 μm size on three different occasions.

The method to examine epithelial cell damage infected with Candida cells for 24 h has previously been described in detail (Mayer et al. 2012).

**Statistical analysis**

Differences for the in vitro tests and host-pathogen interaction experiments among the strains were analyzed by analysis of variance (ANOVA) and the Tukey test. P values < 0.05 were considered to be statistically significant. GraphPad Prism version 6.00 program was used for the statistical analysis.

**RESULTS**

**Mrv8 is required for C. albicans mycelial micro-colony development and epithelial damage**

As part of a larger attempt to characterise unknown function genes in C. albicans, we had previously generated a mrv8Δ/Δ mutant and found it to exhibit defective epithelial damage (Wilson et al. 2014).

We therefore generated a mrv8Δ/Δ/+Mrv8 complemented strain and analysed the impact of Mrv8 deletion on Candida-epithelial interactions in greater detail.

During the early stages of infection (3 h), the mrv8Δ/Δ mutant adhered to and invaded epithelia and formed hyphae at rates almost identical to that of the wild type (Fig. 1A–C). However, at later phases mrv8Δ/Δ caused significantly less epithelial damage than the wild type, and complementation with a single copy of MRV8 restored damage to near wild type levels (Fig. 1D). As this damage defect was unlikely the result of impaired hypha formation, adhesion or invasion during the early stages of the interaction (Fig. 1A–C), we performed morphological analysis of late phase C. albicans-epithelial interactions. We infected epithelia with very low numbers of C. albicans yeast cells (around 50 per well), and assessed the morphology and dimensions of resultant micro-colonies formed by the different strains. Figure 1E shows that, by 24 h post-infection, deletion of MRV8 resulted in significantly smaller C. albicans micro-colonies on oral epithelia. To test whether this defect was specific for growth on epithelium, we repeated the experiment in cell culture medium on plastic surfaces in the absence of host cells. Again, the mrv8Δ/Δ formed smaller micro-colonies than the wild type and complemented strains (Fig. 1E). These data indicate that Mrv8 is dispensable for the initial filamentation, adhesion and invasion events associated with C. albicans-epithelial infection, but that the mutant’s attenuated damage capacity is likely due to defective mycelial-related growth.

**Candida albicans MARVEL family member, Mrv8, is involved in biofilm growth despite showing no other in vitro defects**

Having confirmed its role in epithelial damage, we next sought to functionally characterise Mrv8. The mrv8Δ/Δ mutant was subject to exhaustive functional characterisation tests, revealing no altered phenotype compared to the wild type. mrv8Δ/Δ was able to grow at wild type rates on yeast growth media (YNB agar) in the presence of selected sole carbon sources, such as glucose, potassium acetate, glycerol, galactose, maltose or olive oil and on YCB agar in the presence of BSA and the amino acids lysine, histidine, glycine, cysteine, or methionine as sole nitrogen sources. This indicates that Mrv8 is dispensable for yeast phase growth. The mrv8Δ/Δ mutant formed filamentous colonies comparable to the wild type on Spider medium and serum-containing agar plates, as well as when embedded in YPS-, Spider- or serum-agar. At the single cell level, the mutant also formed hyphae of similar lengths in liquid hypha-inducing media on a plastic surface, including 10% FBS, YPFD plus 10% FBS, Spider medium, Lee’s medium at pH 6.5, RPMI-1640 and DMEM with CO₂. Deletion of MRV8 also had no obvious impact on the ability of C. albicans to grow in the presence of thermal stress, the osmotic stressors NaCl and sorbitol, oxidative stress (H₂O₂), the cell wall stressors calcifluor white and Congo Red, and the cell membrane stressors ethanol and SDS or following exposure to UV radiation (data not shown).

When tested for biofilm formation, mrv8Δ/Δ exhibited lower metabolic activity in Spider medium (Fig. 2A), but not in RPMI (below). Upon visual inspection, mrv8Δ/Δ biofilms appeared aberrant and patchy biofilm compared to the wild type and revertant strains (Fig. 2B). Scanning electron microscopy images revealed a mature wild type biofilm coating the substrate surface and consisting of yeast, pseudohyphae and hyphal cells with evidence of extracellular matrix. In contrast, the mrv8Δ/Δ biofilm was sparse and consisted mainly of yeast cells, with little evidence of filamentation. Complementation with a single copy of MRV8 resulted in an intermediate phenotype: the biofilm covered most of the underlying substrate and all three morphotypes were present (Fig. 2B). Therefore, Mrv8 plays a role in biofilm development in Spider medium.
Figure 1. Mrv8 is required for mycelial maturation and damage of oral epithelial cells. Host-pathogen interaction tests were performed on TR146 oral epithelial cell monolayers. (A), Candida albicans cells adhered to host cells after 3 h of interaction were counted from 50 fields in three independent experiments. (B), The percentage of invasive hyphae was determined by dividing the number of invaded cells by the total number of cells on epithelia. The results represent the mean ± SD of three independent experiments performed in triplicate. (C), The length of hyphae grown on epithelia for 3 and 6 h was measured using LAS software on two separate occasions carried out in triplicate. (D), Damage capacity was determined for TR146 monolayers infected with C. albicans cells for 24 h by measuring lactate dehydrogenase (LDH) levels in the supernatant. Results are the mean ± SD of at least three independent experiments performed in triplicate. Difference statistically significant for multiple comparisons, Tukey’s test \( P < 0.001 \). (E), At least 50 microcolonies grown in DMEM medium for 24 h on epithelial cells or on plastic were measured and the results expressed as the mean ± SD of at least three independent experiments. On epithelia, fungi were stained with calcofluor white. Representative pictures of micro-colonies formed by the wild type, mutant and revertant strains, respectively, scale bars: 250 μm on TR146 and 50 μm on plastic. Difference statistically significant for multiple comparisons on TR146 or on plastic, Tukey’s test \( P < 0.0001 \).
of caspofungin, with drug tolerance increasing at higher concentrations. Interestingly, at several drug concentrations (Fig. 3C), biofilms of mrv8Δ/Δ exhibited increased caspofungin tolerance. We also tested the sensitivity of planktonic cells to these three drugs and observed no differences amongst the strains (data not shown).

Caspofungin tolerance can be induced by a Ca$^{2+}$/calcineurin-mediated cell wall salvage pathway which causes an increase in chitin content (Munro et al. 2007; Walker, Gow and Munro 2013; Rueda, Cuenca-Estrella and Zaragoza 2014). As MRV8 encodes a predicted plasma membrane protein, we reasoned that the observed differences in caspofungin tolerance in the mrv8Δ/Δ mutant may be due to altered Ca$^{2+}$/calcineurin signaling and/or chitin synthesis. In order to test this hypothesis, biofilms were grown for 24 h in the presence of sub-inhibitory concentrations of the calcineurin inhibitors, FK506 and cyclosporin A (CsA), the chitin synthesis inhibitor, Nikkomycin Z or CaCl$_2$/CFW (a combination treatment which increases chitin synthesis and caspofungin tolerance) (Walker et al. 2008), then treated further with 0.5 mg l$^{-1}$ caspofungin and the chemicals for a further 24 h and biofilm integrity assessed by measuring XTT activity.

Cyclosporin A treatment alone enhanced mrv8Δ/Δ biofilm formation compared to the wild type, whilst the other treatments had no effect (Fig. 3D). Pre-treatment of biofilms with cyclosporin A did not affect caspofungin sensitivity whilst both FK506 and Nikkomycin Z increased its antifungal effect. Interestingly, treatment of biofilms with the chitin-stimulatory combination of Ca$^{2+}$/CFW increased caspofungin tolerance in wild type, but not in mrv8Δ/Δ biofilms. This is unlikely to be the result of defective chitin synthesis because both chitin and calcium levels were high in mrv8Δ/Δ yeast cells following the co-stimulation (Fig. S1, Supporting Information). Indeed, Ca$^{2+}$/CFW treatment significantly (P = 0.0285) enhanced wild type biofilm tolerance against caspofungin by 5.3-fold but had virtually no effect on mrv8Δ/Δ biofilms (1.2-fold) and the mrv8Δ/Δ + MRV8 complemented strain exhibited an intermediate (2.3-fold) increase in biofilm activity, which was not significant.

We also determined biofilm dispersal under these different treatment conditions (Fig. 3E). Overall, the level of dispersal reflected biofilm activity as determined by XTT assay (Fig. 3D vs. E). However, whilst caspofungin-treated mrv8Δ/Δ biofilms exhibited increased metabolic activity (Fig. 3D), they released fewer disperser cells but this was not statistically significant (Fig. 3E). Together, these data suggested that mrv8Δ/Δ biofilms may be associated with altered calcium/calcineurin signaling or chitin synthesis in response to stimulation.

**DISCUSSION**

*Candida albicans* is a member of the CTG-Ser Clade of fungi—a group which contains pathogens, non-pathogens, and species with lower virulence than *C. albicans*. Because of this, our group has postulated that lineage-unique genes may contribute to the high virulence potential of *C. albicans*. In a previous study, we deleted a series of lineage unique genes and found several of them to be important for host cell damage. There, we observed a reduction in epithelial and endothelial damage by a mrv8Δ/Δ mutant (Wilson et al. 2014).

To examine the mechanism underlying this defect, the mrv8Δ/Δ and mrv8Δ/Δ + MRV8 strains were evaluated for adhesion, invasion and filamentation in association with epithelia. Complementation of mrv8Δ/Δ with a single copy of MRV8 fully restored epithelial damage, demonstrating that this defect was...
Figure 3. The effect of calcineurin-inhibition and chitin-inhibition and stimulation on Mrv8-dependent biofilm caspofungin tolerance. Biofilms were formed for 24 h and then treated with the antifungal drugs fluconazole (mg l⁻¹) (A), amphotericin B (mg l⁻¹) (B), or caspofungin (mg l⁻¹) (CSF) (C) for an additional 24 h. The susceptibility of the biofilms to the drugs was then evaluated by measuring the metabolic activity with XTT-assay. The results are expressed as the percentage of growth in relation to the control (100%). The experiments were performed in triplicate on three different occasions. Alternatively, biofilms were allowed to form for 24 h in the presence of 0.75% DMSO, 75 μgm l⁻¹ FK506, 75 μgm l⁻¹ cyclosporin A, 20 μM Nikkomycin Z, or 0.025 M CaCl₂/12.5 μgm l⁻¹ CFW before treatment with 0.5 mg l⁻¹ caspofungin and the chemicals for a further 24 h. Biofilm metabolic activity (XTT assay [D]) or dispersal (CFU (Log10)/well [E]) was then determined. Results are the mean ± SD of three independent experiments performed in triplicate. Difference statistically significant for multiple comparisons, Tukey’s test *P < 0.05.

due to MRV8 deletion. Adhesion, invasion and initial hypha formation on epithelia was virtually identical for all strains, indicating that early morphogenic events are independent of Mrv8 function. However, at later phases of mycelial development on epithelia, the mrv8Δ/Δ mutant exhibited significantly smaller micro-colonies. This was not an epithelial-specific phenomenon, as mrv8Δ/Δ micro-colonies were also reduced in size in the absence of host cells. We propose that Mrv8 plays a role in later stages of hyphal development, and the resultant defects in hyphal maturation and ramification may account for the epithelial damage defect of the mrv8Δ/Δ mutant. These defects in hyphal maturation may also contribute to the altered biofilm phenotypes observed in this study. The precise mechanism by which Mrv8 contributes to hyphal maturation is currently unknown. Because Mrv8 is a predicted transmembrane protein, it is possible that it acts as a sensor, relaying an extracellular cue to the intracellular signal transduction pathways and transcriptional machinery governing morphogenesis.

Deletion of MRV8 did not affect biofilm sensitivity to fluconazole or amphotericin B. Interestingly, however, mrv8Δ/Δ exhibited increased tolerance to inhibitory concentrations of caspofungin (Fig. 3C).

Caspofungin targets β-1,3-glucan via inhibition of the cognate synthase Fks1 and hot-spot mutations in FKS1 have been associated with caspofungin resistance (Walker et al. 2008; Walker, Gow and Munro 2013). Increased caspofungin tolerance can also be induced by elevations in cell wall chitin (Walker et al. 2008). This cell wall salvage pathway is induced by increased CHS1, CHS3, CHS2, and CHS8 expression, elevated chitin synthase activity and increased chitin content in the cell wall in response to caspofungin (Walker et al. 2008; Rueda, Cuenca-Estrella and Zaragoza 2014). These events can be mediated through the protein kinase C (PKC), high osmolarity glycerol response (HOG), and Ca²⁺-calcineurin signaling pathways (Walker et al. 2008; Walker, Gow and Munro 2013). Calcium stimulation activates the calmodulin-calcineurin pathway and
results in Cnr1-dependent induction of CHS2 and CHS8 (Munro et al. 2007). We hypothesized that altered calcineurin signaling may be involved in the caspofungin tolerance phenotype of mrv8Δ/Δ mutant biofilms.

In line with this, in the absence of caspofungin, cyclosporin A treatment actually enhanced biofilm metabolic activity for the mrv8Δ/Δ mutant suggesting that calcineurin signaling may indeed be altered in this strain.

Exposure to calcium and CFW is known to protect C. albicans cells against caspofungin by increasing chitin synthesis (Munro et al. 2007; Walker et al. 2008; Walker, Gow and Munro 2013; Rueda, Cuenca-Estrella and Zaragoza 2014). We also observed this protective effect with wild type, but not mrv8Δ/Δ biofilms. Therefore, in the absence of MRV8, CaCl2/CFW does not protect C. albicans from caspofungin.

As MARVEL proteins are implicated in plasma membrane organization and Mrv8 itself contains four predicted transmembrane helices (Douglas, Wang and Konopka 2013), it is possible that this protein functions at the cell surface, sensing the calcium/CFW signal and possibly signaling through the calcineurin pathway.

In summary, we determined the association of a specific MARVEL family member with pathogenicity and biofilm drug tolerance. MRV8 was involved in hyphal maturation on both abiotic and epithelial surfaces, damage of epithelial cells, and biofilm-associated caspofungin tolerance, possibly via interactions with calcium- and calcineurin signaling pathways.

SUPPLEMENTARY DATA
Supplementary data are available at FEMSYR online.

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Conflicts of Interest. None declared.

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