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Adaptation of *Candida albicans* to environmental pH induces cell wall remodelling and enhances innate immune recognition

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Abstract

*Candida albicans* is able to proliferate in environments that vary dramatically in ambient pH, a trait required for colonising niches such as the stomach, vaginal mucosal and the GI tract. Here we show that growth in acidic environments involves cell wall remodelling which results in enhanced chitin and β-glucan exposure at the cell wall periphery. Unmasking of the underlying immuno-stimulatory β-glucan in acidic environments enhanced innate immune recognition of *C. albicans* by macrophages and neutrophils, and induced a stronger proinflammatory cytokine response, driven through the C-type lectin-like receptor, Dectin-1. This enhanced inflammatory response resulted in significant recruitment of neutrophils in an intraperitoneal model of infection, a hallmark of symptomatic vaginal colonisation. Enhanced chitin exposure resulted from reduced expression of the cell wall chitinase Cht2, via a Bcr1-Rim101 dependent signalling cascade, while increased β-glucan exposure was regulated via a non-canonical signalling pathway. We propose that this “unmasking” of the cell wall may induce non-protective hyper activation of the immune system during growth in acidic niches, and may attribute to symptomatic vaginal infection.

Author Summary

To be able to colonise a host or cause infection, microbes must be able to adapt and respond to changes in their environment. One of the most important environmental signals for opportunistic pathogens is ambient pH, with changes in external pH resulting in phenotypic, metabolic and physical changes in fungi and bacteria (i.e. *E. coli, Salmonella, Aspergillus, Candida*). *Candida albicans* is an opportunistic fungal pathogen of humans that can colonise and infect niches of varying pH including the acidic mucosa of the vagina. Here we show that growth in an acidic environment results in structural modification of the fungal cell wall, a dynamic organelle key to immune recognition. These cell wall perturbations resulted in enhanced immune
recognition of the fungal pathogen, a strong proinflammatory immune response and enhanced recruitment of neutrophils. Therefore, colonisation of acidic mucosa may result in the unmasking of cell wall components that trigger hyper-activation of the innate immune response and could contribute to immunopathology associated with vaginal candidiasis.
Introduction

The opportunistic fungal pathogen *Candida albicans* is a commensal in up to 80% of the population, and can cause superficial mucosal infections in healthy individuals (1, 2) and invasive disease in immune suppressed patients (3, 4). Mucosal infections increase population morbidity and are expensive to treat, while disseminated disease is associated with high mortality rates (5, 6).

One attribute of *C. albicans* that has made it such a successful opportunistic pathogen is its ability to adapt and proliferate in a broad range of host environments. One of the most important environmental conditions that fluctuate between different niches is ambient pH. *C. albicans* is able to grow in media ranging from pH2 to pH10, and *C. albicans* has been isolated from a range of anatomical sites that vary dramatically in ambient pH including the stomach (pH2) (7), vagina (pH4-5) (8) and the oral mucosa (pH6) (9), suggesting that adaptation to environmental pH is key to the pathogenicity of *C. albicans*. Adaptation of *C. albicans* to acidic environments regulates key biological processes including morphogenesis (10), white-to-opaque switching, and mating (11). However, the impact environmental adaptation has on the structure and composition of the fungal cell wall, the first point of contact between the fungus and host, is not well defined.

The fungal cell wall is a complex, multi-layered structure of mannoproteins, β-glucans and chitin that provides rigidity and shape and protects the fungus from the environment (12). These protein and carbohydrate motifs are immunogenic and play important roles in innate immune recognition (12). Environmental adaptation influences the cell wall proteome and impacts on the structure of the glycans that decorate the cell wall proteins (13). For instance, growth in blood or lactate media decreases the structural complexity of the outer mannan fibrils (14-16), while antifungal drug treatment influences β-glucan exposure (17, 18). Structural changes
in the cell wall, as a result of mutation in key glycolytic cell wall assembly enzymes, confirm that modulation of the cell wall has profound implications for innate immune recognition (19). Therefore, understanding how environmental adaptation impacts on cell wall biogenesis, and the consequence this cell wall remodelling has on innate immune recognition is an important, but understudied, area of fungal biology. Here, we investigate how adaptation to acidic environments that mimic the pH of the vaginal mucosa impact on the structure and composition of the C. albicans cell wall and deduce how this cell wall remodelling affects the innate immune recognition of the pathogen.

Results

Growth in acidic environments alters the ultrastructure of the fungal cell wall.

The impact of adaptation to environmental pH on the ultrastructure of the cell wall was investigated via transmission electron microscopy. The cell wall of mid-log phase yeast cells grown in YPD buffered at pH2, pH4 pH6, pH8 and standard YPD were imaged. Under all tested conditions, the cell wall maintained two distinct layers: an inner layer comprised of glucan and chitin, and an outer, fibrillar layer of mannoproteins. However, adaptation to acidic conditions resulted in a significant loss of structural organisation in the outer cell wall, which appeared to be less fibrilar (Figure 1a). Quantification of the thickness of the outer cell wall layer confirmed that adaptation to acidic environments significantly reduced mannan fibril length (pH2 23.23 ± 3.58 nm (p = 0.0001), pH4 39.49 ± 4.61 nm (p = 0.0495), compared to pH6 60.22 ± 10.73 nm; Figure 1b). We hypothesized that this loss of fibrilar structure might be due to changes in the underlying architecture of the cell wall. Therefore, we investigated whether pH similarly influences chitin structure.
Acidic environments induce reorganisation of the fungal cell wall through modulation of chitin.

Although chitin only forms a small fraction of the fungal cell wall (3-5% by dry weight), a major compensatory mechanism of *C. albicans* to cell wall stress is to up-regulate chitin synthesis to provide increased cell wall integrity (20). To identify whether the observed disorganisation of the outer cell wall layer was a result of increased chitin incorporation, the cell wall of acidic adapted cells was stained with Calcofluor White (CFW). Quantification of CFW fluorescence indicated that the chitin content of the cell wall was only elevated at pH2 (Figure 2a), a result that was confirmed by HPLC (Table 1), suggesting that adaptation to environments of very low pH requires chitin synthesis. However, staining the cell wall with wheat germ agglutinin (WGA), a lectin that binds surface exposed chitin, indicated that even adaptation to pH4 required reorganisation of cell wall chitin, with chitin becoming increasingly exposed during adaptation to acidic pH (Figure 2b). Microscopy confirmed that cells adapted to acidic pH showed significant de-cloaking of chitin around the cell periphery, with intense WGA staining occurring at bud scars at pH2 (Figure 2c). To deduce whether this de-cloaking phenomenon is an active process, we analysed the de-cloaking of chitin in response to pH in dead cells. De-cloaking of cell wall chitin only occurred in live cells (Supplemental Figure 1a), confirming that the observed increase in surface exposure of chitin was not a physical effect of the pH simply degrading the cell wall. Thus, adaptation to acidic environments is a two-stage process with moderate acidic environments (pH4) causing active de-cloaking of chitin, and strong acidic environments (pH2) inducing *de novo* chitin synthesis and further de-cloaking of chitin.

Neither Mkc1, Hog1, or Crz1 regulate chitin exposure in response to moderate pH stress.
Chitin synthesis is regulated by the cell wall salvage, calcium/calcineurin and high osmolarity glycerol (HOG) signalling pathways (21). To assess whether these pathways are involved in chitin reorganisation, activation of these signalling cascades in response to environmental pH was determined. Hog1 was not activated during exponential growth in environments of differing pH (Figure 3a). In agreement with this, deletion of HOG1 did not impact on the surface exposure of chitin during adaptation to pH4 environments (Figure 3b). Although the cell wall salvage pathway was activated in response to acidic environments (Figure 3c), deletion of MKC1 and RLM1 did not prevent the cell wall remodelling that results in de-cloaking of the underlying chitin (Figure 3b, d). Deletion of CRZ1, the transcription factor downstream of the calcium/calcineurin pathway, also had no impact on chitin exposure in acid adapted *C. albicans* cells (Figure 3d). Therefore, these pathways are not involved in the reorganisation of chitin during adaptation to pH4 environments.

Rim101 and Bcr1 coordinate correct localisation of chitin in the cell wall through regulation of Cht2.

Chitin can be remodelled in the cell wall through the actions of four chitinases (Cht1-4) (22). Therefore, the role of these chitinase enzymes in the de-cloaking of chitin in response to environmental pH was determined. Deletion of CHT2 resulted in enhanced chitin exposure at pH6 (Figure 4a) compared to parental control strains. To determine whether the expression of CHT2 is regulated by ambient pH, semi-quantitative RT-PCR was performed. CHT2 expression was repressed in acidic environments compared to pH6 or pH8 environments (Figure 4b). Large scale transcriptional profiling suggests that expression of CHT2 is largely dependent on the transcription factor Bcr1 (23). To confirm this, the expression of CHT2 in a *bcr1* mutant strain was determined. CHT2 expression was significantly reduced in the *bcr1* mutant.
mutant at all pH conditions tested (Figure 4c), confirming a dominant role for Bcr1 in
the expression of CHT2.

To investigate the mechanism by which CHT2 is repressed during adaptation to
acidic environments, we focused on the Rim101 transcription factor. Rim101 has
previously been shown to regulate the expression of the cell wall modifying enzymes
Phr1 and Phr2 in response to environmental pH (24), has been implicated in cell wall
reorganisation (25), and is an activator of gene expression in alkaline environments
(26). The expression of CHT2 was repressed in the rim101 mutant in environmen-
t above pH6 (Figure 4c), with CHT2 expression levels at pH6 or pH8 being
comparable to pH4, suggesting that Rim101 is a pH-dependent activator of CHT2
expression. Alkaline induced expression of CHT2 was dependent on Bcr1, as CHT2
levels remain low under all pH conditions in the bcr1 mutant (Figure 4c).

To determine how the deregulation of CHT2 expression affects chitin exposure, the
bcr1 and rim101 mutants were grown in YPD buffered at pH4 and pH6, and stained
with WGA. Deletion of BCR1 resulted in increased chitin exposure at pH6 (p =
0.0076) and pH4 (p = 0.0491), but maintained some pH dependency, while deletion
of RIM101 resulted in constitutively high chitin exposure (Figure 4d). To investigate
whether expression of Rim101 at pH4 would affect the phenotype, we quantified
chitin exposure in a strain expressing constitutively active Rim101 (CARP1-1). FACS
analysis confirmed that expression of active Rim101 in acidic environments reduced
chitin exposure (Figure 4e), but was still insufficient to completely mask the chitin.
These results confirm that Rim101 plays an important role in the regulation of cell
wall genes pivotal to correct chitin incorporation.

The mRNA of CHT2 is a target of She3, which transports mRNAs to the cell wall
(27). Therefore, we determined whether this complex is also required for correct
incorporation of chitin into the cell wall. Deletion of SHE3 resulted in enhanced de- 197 cloaking of chitin in environments above pH4 (Figure 4f), similar to the cht2 and bcr1 198 mutants. Taken together, these results suggest that incorporation of chitin into the 199 inner cell wall at pH6 is regulated by Rim101 and Bcr1 and requires She3 dependent 200 delivery of Cht2 to the cell wall, while growth in an acidic environment inactivates this 201 pathway leading to reduced CHT2 expression and enhanced exposure of chitin at the 202 cell surface.

203

**Low pH promotes unmasking of the underlying β-glucan.**

β-glucan is a major component of the cell wall, which is highly immunostimulatory 206 and consequently is normally masked by a dense layer of glycosylated proteins. As a 207 result, fungal cells have a degree of resistance to extracellular glucanases, which 208 break down the cell wall and cause cell lysis. Therefore, the sensitivity of *C. albicans* 209 to recombinant β-glucanase can be used as an indirect measure of β-glucan exposure (28). Growth of *C. albicans* in acidic environments enhanced β-glucanase 210 sensitivity (Figure 5ai and ii) compared to pH6 or pH8 grown cells, suggesting that β- 211 glucan is more accessible in cells grown in media buffered to pH4. In agreement with 212 this, immunofluorescent staining of the cell wall with a monoclonal β1,3-glucan 213 antibody revealed enhanced staining around the cell wall periphery (Figure 5b). 214 Quantification of the immunofluorescent staining by FACS revealed that acid adapted 215 cells had almost 4-fold more surface exposed β-glucan (Figure 5c) than cells grown 216 in YPD media, suggesting that adaptation to acidic environments induce unmasking 217 of β-glucan. To rule out the possibility that the pH simply degrades the cell wall, 218 exposing the underlying β-glucan, dead cells were exposed to YPD at pH2, 4 and 6 219 for 4 h and surface exposure quantified by FACS. Only live cells unmasked their β- 220 glucan in response to acidic pH (Supplemental Figure 1b), confirming that, like the 221 chitin exposure, unmasking of β-glucan is an active process.
To determine whether β-glucan synthesis was required for this unmasking effect, total glucan levels were quantified by staining the cells with Aniline Blue. Binding of Aniline Blue to the fungal cell wall was consistent across all pH conditions, suggesting that total β-glucan levels remained constant (Figure 5d). Furthermore, HPLC analysis confirmed that the amount of glucose in the cell wall was not significantly different in cells grown in media buffered to different pHs (Table 1). Therefore, the increase in β-glucan exposure occurs as a result of cell wall remodelling, and not enhanced β-glucan synthesis.

To deduce whether de-cloaking of the cell wall was a general adaptation response of *C. albicans*, pH-dependent cell wall reorganisation of a series of clinical isolates from different sites and types of infection was examined. All clinical isolates de-cloaked their cell wall in response to acidic environments (Figure 5e), confirming that this phenomenon is not restricted to laboratory-evolved strains.

To investigate whether the form of acid used to pH the media affects unmasking, YPD media was buffered using lactic acid an organic acid produced by Lactobacilli during colonisation of the vaginal mucosa. Growth in YPD buffered to pH4 with lactic acid still induced unmasking of the cell wall, similar to reducing pH with HCl (Figure 6a). Therefore, physiologically relevant acids drive unmasking of the cell wall in a pH-dependent manner. In order to determine whether environments that more closely mimic the environment of the female reproductive tract induce cell wall remodelling, *C. albicans* was grown in Vaginal Simulation Media (VSM) buffered to pH4, pH6 or pH7. *C. albicans* unmasked significantly more β-glucan (p = 0.0022) when the VSM was buffered to pH4 than either pH6 or pH7 (Figure 6b), suggesting that the environment within the female reproductive tract has the potential to induce exposure of β-glucan. Because infection sites are know to contain a mixture of yeast and hyphal cells, *C. albicans* hyphae were generated in acidic media in elevated...
concentrations of carbon dioxide, a potent inducer of morphogenesis (29) and in the presence of vaginal epithelial cells. Both yeast and hyphal cells showed increased chitin and β-glucan exposure compared to their respective controls (Figure 6c, d). Therefore, both yeast and hyphal cells undergo cell wall remodelling during adaptation to acidic environments, which results in the surface exposure of β-glucan.

As adaptation to acidic pH resulted in surface exposure of both chitin and glucan we investigated whether these carbohydrates became exposed at the same or different points in the cell wall. Dual immunofluorescent imaging using TRITC-WGA and FITC-glucan, confirmed that while chitin exposure occurred mainly at bud scars at pH4 with increase exposure occurring in the lateral cell wall at pH2, glucan exposure was continuously localised to punctate patches around the cell periphery (Figure 7). As the exposed carbohydrates were not co-localised we hypothesised that they are not dependent on each other, and may be regulated via different processes.

One potential hypothesis to explain the unmasking response to acidic pH is that the chemical cleavage of the outer phosphomannan could reduce the mannan complexity and result in a more porous outer mannan shield, permitting access to the underlying cell wall layers. To test this hypothesis, carbohydrate exposure after adaptation to acidic pH in the mnn4Δ mutant, which is devoid of phosphomannan (30), was examined. The mnn4Δ mutant de-cloaked its cell wall similarly to the parental control strain (Supplementary Figure 2), suggesting that this phenomenon is not a result of the loss of the phosphomannan and likely involves significant active cell wall reorganisation.

*Candida tropicalis* also unMASKS β-glucAN in response to acidic pH.

To determine whether unmasking of β-glucan is specific to *C. albicans*, the impact of acidic environments on non-albicans Candida species and on *Saccharomyces*
cerevisiae was assessed. In stark contrast to *C. albicans*, growth in acidic conditions reduced the sensitivity of *S. cerevisiae* to β1,3-glucanase, suggesting that in response to acidic environments *S. cerevisiae* masks its β-glucan (Figure 8), which has been previously reported (28). Likewise, *Candida parapsilosis* also displayed a mild, statistically insignificant, decreased sensitivity to β1,3-glucanase when grown under acidic conditions (Figure 8). On the other hand, *Candida glabrata* and *Candida dublioniensis* did not show any pH-dependent modulation of β-glucan exposure (Figure 8). Unexpectedly, *Candida krusei* unmasked its β-glucan in response to alkaline environments (Figure 8), while *Candida tropicalis* unmasked its β-glucan in response to low pH (Figure 8). Therefore, of the isolates examined only *C. albicans* and *C. tropicalis* unmask β-glucan in response to low pH.

Unmasking of β-glucan involves a non-canonical cell wall remodelling pathway.

Investigation of the major signal transduction pathways known to be involved in cell wall remodelling (i.e. Mkc1, Hog1, Crz1) confirmed that these pathways are not required for pH-dependent β-glucan unmasking (Supplementary Figure 3a,b). Due to the involvement of Rim101 and Bcr1 in chitin de-cloaking, we hypothesised that these transcription factors may also be involved in β-glucan unmasking. However, *rim101Δ* and *bcr1Δ* mutants still unmasked β-glucan in a pH-dependent manner (Supplementary Figure 3b, c). As Rim101 does not undergo C-terminal processing at acidic pH, unmasking of β-glucan in a strain constitutively expressing active Rim101 (CARP1-1) was also assessed. The CARP1-1 strain still displayed pH-dependent unmasking of β-glucan (Supplementary Figure 3c), suggesting that the Rim101 signalling cascade is not required for pH-dependent β-glucan unmasking. Therefore, while Rim101/Bcr1 regulate chitin exposure, a non-canonical signalling pathway regulates pH-dependent β-glucan unmasking.
Acid adapted C. albicans cells are readily phagocytosed and induce a strong innate immune response.

The cell wall is the first point of contact between the fungus and the host's immune system and thus the cell wall plays a major role in innate immune recognition of fungi. Therefore, the role of the pH-dependent cell wall de-cloaking in regulating innate immune responses was investigated. C. albicans cells grown in acidic media (pH2 and pH4) were more readily phagocytosed by macrophages and neutrophils than C. albicans cells grown in standard YPD media (Figure 9a,b).

To deduce whether adaptation of C. albicans to low pH results in a heightened pro-inflammatory innate immune response, the cytokine response of peripheral blood monocytes (PBMCs) exposed to C. albicans adapted to different environmental pH conditions was examined. C. albicans adapted to acidic environments elicited a much stronger proinflammatory cytokine response from PBMCs than C. albicans cells grown in more alkaline conditions (Figure 9c-f). Specifically, C. albicans cells grown in YPD buffered to pH4 induced higher secretion of TNFα, IFN-γ, IL-6 and IL-1β compared to cells grown in YPD buffered to pH6 or pH8. Therefore, de-cloaking of the fungal cell wall in response to acidic pH promotes innate immune recognition of C. albicans.

The rim101Δ and bcr1Δ mutants displayed levels of chitin exposure at pH6 similar to those observed at pH4. As chitin has been shown to supress the innate immune system (31, 32), we tested whether the increased chitin in these mutants affected the innate immune response. Phagocytosis rates in both mutants were reduced compared to the parental control strain (Supplemental Figure 4a,b). However, this did not impact on the cytokine response (Supplemental Figure 4c). Therefore, the altered cell wall structure in these mutants affects phagocytosis, but is not sufficient to affect pro-inflammatory cytokine responses.
Enhanced phagocytosis is mediated by recognition of β-glucan by Dectin-1.

β-glucan is recognised by the C-type lectin-like receptor Dectin-1 (33). As the monoclonal anti-β1,3-glucan specific antibody confirmed that growth in acidic conditions resulted in β-glucan unmasking, the accessibility of this exposed β-glucan to Dectin-1 was assessed. Immunofluorescent staining of acid grown cells using Fc-Dectin-1 confirmed that Dectin-1 more readily bound to cells grown in acidic conditions (Figure 10a). To ascertain whether this enhanced binding of Dectin-1 to the surface of acid grown cells resulted in the increased phagocytosis rate, Dectin-1 was expressed on the surface of fibroblasts. Fibroblasts expressing Dectin-1 more readily bound *C. albicans* cells grown in acidic conditions than YPD or pH8 grown cells (Figure 10b). Enhanced adhesion was not observed on fibroblasts which did not express Dectin-1, and could be blocked using glucan phosphate (Figure 10c), confirming that the enhanced adhesion is due to a specific interaction between Dectin-1 and surface exposed β-glucan, and not due to differences in electrochemical properties of the cell wall.

To further confirm the role of Dectin-1, the Dectin-1 receptor was blocked with glucan phosphate. Blocking of Dectin-1 in the J774.1A macrophage cell line, resulted in a pH-dependent decrease in phagocytosis (Figure 10d), but did not affect the association of *C. albicans* with the macrophage, with acid adapted cells still displaying enhanced association (Figure 10e). Therefore, Dectin-1 is required for the enhancement of phagocytosis, but other pattern recognition receptors are responsible for the initial attachment of acid-adapted *C. albicans* to the surface of innate immune cells.

Adaptation to acidic pH results in increase immune cell recruitment *in vivo*. 
Assessment of the innate immune response to acid adapted C. albicans cells confirmed that the increased exposure of glucan results in a heightened proinflammatory immune response. To investigate the in vivo significance of this discovery, live C. albicans cells adapted to different pH conditions were injected into the peritoneal cavity of mice, and the recruitment of innate immune cells was determined after 4 hours. C. albicans cells adapted to acidic environments recruited more CD45+ lymphocytes than C. albicans cells adapted to pH6 (Figure 1a, p=0.007), including significantly more neutrophils than C. albicans cells adapted to pH6 media (Figure 1b, p=0.010). However, analysis of the overall cell population confirmed that although the total number of recruited immune cells was increased, there was no significant difference between the percentages of each cell type recruited (Figure 1c, p=0.540). Therefore, C. albicans cells adapted to acidic environments initiate a stronger proinflammatory innate immune response, and recruit significantly more immune cells to the site of infection.

Discussion

C. albicans has a remarkable ability to respond and adapt to a multitude of environmental signals. Here, we demonstrate that adaptation to low pH results in significant remodelling of the C. albicans cell wall. The most striking effects include the de-cloaking of the underlying chitin and β-glucan. Unmasking of these underlying carbohydrate epitopes was an active process, and was independent of phosphomannan loss, suggesting that chemical cleavage of this outer cell wall component did not sufficiently deplete the outer mannan armour to reveal internal pathogen associated molecular patterns (PAMPs).

Chitin is remodelled in the cell wall through the actions of chitinases. C. albicans expresses four chitinases (22), however only deletion of CHT2 prevented the pH-
dependent de-cloaking of chitin. Cht2 is mainly expressed in yeast cells, and is GPI-anchored to the cell wall (34). pH-dependent de-cloaking of cell wall chitin also was dependent on the Rim101 and Bcr1 transcription factors. Rim101 is C-terminally processed in response to alkaline environments through the RIM101 signalling cascade, and activates the expression of alkaline induced genes, while repressing the transcription of acid induced genes (26). Bcr1 is essential for biofilm formation and controls the expression of many cell wall associated genes including HWP1, ALS1, ALS3 and HYR1 (35). The expression of CHT2 was dependent on Bcr1, and showed Rim101 pH-dependent expression in environments above pH6. This alkaline dependent increase in expression was lost in the bcr1Δ mutant, suggesting that Bcr1 is essential for CHT2 expression, as indicated in large scale transcriptomic analyses (36). However, Bcr1 also positively regulates RIM8 (37), suggesting that inactivation of BCR1 would also decrease processing of Rim101, linking the two pathways (Figure 12a).

One hypothesis for the de-cloaking of chitin is that during growth in neutral or alkaline environments Rim101 and Bcr1 activate the expression of CHT2. She3 then delivers CHT2 mRNA to the cell wall, where it is translated, and incorporated into the cell wall via its GPI anchor. Cht2 is predicted to possess endo-chitinase activity. Therefore, once in the cell wall, Cht2 would act internally on the chitin microfibrils, decreasing their length, and permitting correct incorporation in the inner layer of the cell wall. However, during growth in acidic conditions, Rim101 is inactive, resulting in lower transcriptional levels of CHT2 and other cell wall regulatory enzymes. In this case, there will be less Cht2 to act on the chitin microfibrils, which may increase the length of these microfibrils, hindering their embedment into the inner layer of the cell wall and resulting in increased exposure of chitin at the cell surface (Figure 12b).
More striking than the de-cloaking of chitin was the unmasking of the highly pro-inflammatory PAMP, β-glucan. Dissection of key pathways known to be involved in the regulation of cell wall biosynthesis and pH sensing suggested that these conventional pathways are not responsible for regulating the change in distribution of β-glucan. In *S. cerevisiae*, adaptation to acidic environments has opposing effects inducing masking of β-glucan, which is mediated by the Hog1 signal transduction pathway (28). Deletion of *HOG1* had no impact on β-glucan unmasking in *C. albicans*, suggesting that significant transcriptional rewiring has occurred during evolution between *S. cerevisiae* and *C. albicans*.

*C. albicans* is isolated from 95% of vulvovaginal candidiasis (VVC) cases, with the remaining 5% of infections being mainly caused by *C. glabrata* (38). During colonisation of the vaginal mucosa *C. albicans* is exposed to many environmental conditions including low mucosal pH (vaginal pH = 3.5-4.5). Unlike bacterial vaginosis, which is associated with alkalisation of the vaginal mucosa, during VVC mucosal pH remains low (39). Therefore, during both colonisation and infection of the vaginal mucosa, *C. albicans* is exposed to acidic conditions.

The pH-dependent unmasking of the cell wall results in enhanced recruitment of innate immune cells *in vivo*, which correlated with enhanced production of proinflammatory cytokines. Our *in vitro* assays confirm that this enhanced immune recognition was mediated via the C-type lectin like receptor Dectin-1, a dominant receptor in fungal innate immunity known to recognise β-glucan (40).

Symptomatic vaginal colonisation by *C. albicans* results in excessive neutrophil migration, with vaginal secretions displaying enhanced chemotactic potential and increased mucosal damage (41). Neutrophil depletion reduces inflammation and symptoms associated with VVC (42), suggesting that neutrophil recruitment in VVC is
non-protective, in contrast to oral candidiasis where neutrophil recruitment and
activation of Th17 responses are protective (43).

Growth of *C. albicans* in VSM (a media which more closely resembled vaginal fluid)
confirmed that the unmasking of β-glucan is not media specific, but is pH dependent,
with unmasking only occurring at acidic pH. This raises questions regarding the link
between β-glucan unmasking and VVC, as mouse VVC models, which elicit similar
immunopathologies to human samples, have a neutral, not an acidic, vaginal pH
(44). Although the mouse models can replicate some conditions experienced during
VVC, these still do not provide a complete model of VVC. For example, mice are not
naturally colonised by *C. albicans* in the vaginal tract, and are normally immune
suppressed with oestrogen to maintain colonisation and the high inoculum used to
establish the infection is unlikely to reflect fungal burdens during the initiation of VVC
(44). The cells used for intravaginal inoculation will also be immunogenically different
to those acquired from the GI tract or skin, which is the primary source of vaginal
seeding in human VVC. Finally, the neutral pH of the mouse vaginal mucosa would
favour rapid hyphal development of *C. albicans*, which will activate the
inflammasome pathway more rapidly resulting in increased epithelial activation and
secretion of proinflammatory cytokines (45). Therefore, although the mouse model
parallels the symptoms of human VVC we still do know whether the observed
immunopathology arises from the same underlying mechanism. For example, it has
recently been reported that neutrophils also promote unmasking of β-glucan (46),
which would also increase inflammation. Therefore, although we have animal models
for VVC, these do not currently directly reflect human infection. It is conceivable that
in human VVC, a gradual increase in fungal burden in combination with pH-
dependent unmasking of the immunostimulatory β-glucan in both yeast and hyphal
cells, as a response to environmental changes within the vaginal niche, might initiate
a earlier, stronger proinflammatory cytokine response, resulting in increased
neutrophil recruitment causing symptomatic vaginal colonisation.

Materials and Methods

Ethics

All animal work was carried out by competent researchers under UK Home Office
project licence PPL 70/9027 (awarded to Dr Donna MacCallum), which was reviewed
and approved by the University of Aberdeen Animal Welfare and Ethical Review
Body (AWERB) and the UK Home Office. Animal experiments adhered to the UK
Animals (Scientific Procedures) Act 1986 (ASPA) and European Directive
2010/63/EU on the protection of animals used for scientific purposes. All animal
experiments were designed with the 3Rs in mind and were reported using the
ARRIVE guidelines. The Institutional Review Board of the School of Biosciences at
the University of Birmingham approved the protocol for blood collection, and isolation
of PBMCs and neutrophils from healthy volunteers. Blood donations were
anonymous, and all volunteers provided written informed consent for samples to be
included in this study.

Strains, media and growth conditions

Unless stated otherwise, all media and consumables were purchased from Sigma-
Aldrich UK. Yeast strains were maintained on YPD agar (1% yeast extract, 1% bacto-
peptone, 2% glucose and 2% agar). For liquid cultures, yeasts were cultured in YPD
(1% yeast extract, 1% bacto-peptone, 2% glucose). Buffered YPD was made by
supplementing YPD with 3.57% HEPES and adjusting the pH accordingly. VSM
contained 58 mM NaCl, 18 mM KOH, 2 mM Ca(OH)₂, 1.75 mM glycerol, 6.7 mM
urea, 33 mM glucose, and 0.67% yeast nitrogen base (YNB) and buffered to pH4,
pH6 or pH7 with 22 mM lactic acid and 17 mM acetic acid according to (47). Strains used in the study are listed in Supplementary Table 1.

**High pressure freeze substitution transmission electron microscopy**

The ultrastructure of the cell wall of wild type (NGY152) cells grown at different pH was visualised by TEM. Exponentially growing cells at 37°C were frozen under high pressure in liquid nitrogen and embedded in resin as described previously (48). Ultrathin sections (70 nm) were cut sections and mounted on 400 mesh copper grids and stained in 4.5% uranyl acetate in 1% acetic acid for 45 min and Reynolds lead citrate for 7 min. Images were acquired using a Jeol 1230 at 80 kV accelerating voltage fitted with a Gatan 791 multiscan camera. Five cells were selected at random for each condition and multiple images taken around the cell periphery. The thickness of the inner and outer cell wall layers was measured from at least 30 different points of each cell using ImageJ. Data were analysed by two-way ANOVA followed by post-hoc Tukey’s multiple comparisons test at 95% confidence.

**Immunofluorescent staining of cell wall components**

*C. albicans* was grown overnight in appropriately buffered YPD at 37°C, 200 rpm. Cells were sub-cultured in fresh media and grown to exponential phase at 37°C, 200 rpm. Cells were harvested by centrifugation at 3500 rpm for 3 min, fixed on ice for 30 min in 4% PFA in PBS and washed three times in PBS. To kill *C. albicans* to test whether cell viability is required for cell wall unmasking, overnight cultures of SC5314 were either fixed with 4% PFA for 45 mins, heat killed at 65 °C for 2 h, treated with 1 KJ UV light, or treated with 100 mM thimerosal for 45 min. Following killing, *C. albicans* cells were washed in PBS and incubated in YPD buffered at pH2, pH4 and pH6 for 4 h. To generate hyphal cells in acidic environments, 2 x10⁵ yeast cells were inoculated into DMEM buffered at pH4 of pH6 and incubated under static conditions in 5% CO₂ either with or without 2 x10⁵ A431 vaginal epithelial cells (Sigma) in 24
well plates for 4 h. For controls (yeast cells), *C. albicans* were incubated in YPD buffered at pH4 or pH6, and DMEM buffered at pH4 and pH6 at 37°C, 150 rpm, in a 24 well plate. Quantification of WGA staining in the presence of vaginal epithelial cells could not be performed due to non-specific binding of the lectin to the epithelial cells.

To stain for surface exposed chitin, 2 x10⁶ cells were incubated with 100 μg/ml FITC conjugated WGA (Molecular Probes, Life Technologies) for 30 min. To stain for total chitin, fixed cells were stained with 3.5 μg/ml CFW for 5 min. To stain for total glucan, PFA fixed cells were incubated with Aniline Blue fluorochrome (Bioscience supplies) for 15 min. To stain for surface exposed β1,3-glucan, cells were blocked with 2% BSA in PBS, for 30 min and then incubated with a monoclonal anti-β1,3-glucan antibody (Bioscience Supplies, Australia) diluted 1:800 in PBS, 2% BSA on ice for 30 min. Cells were washed three times with PBS, and incubated with 1:200 diluted anti-mouse IgG, conjugated to FITC (Invitrogen) on ice for 30 min. Alternatively, PFA fixed *C. albicans* cells were stained with 3 μg/ml Fc-Dectin-1 (a kind gift from Prof G. Brown, University of Aberdeen) (49), and goat anti-human IgG Fc, conjugated to Alexa Flour 488 (Invitrogen). To stain for total mannan, 2 x10⁶ cells were incubated with 100 μg/ml ConA conjugated to TRITC (Molecular Probes, Life Technologies) for 30 min and washed with PBS. Cells were imaged using a Nikon Eclipse TE 2000U, Plan Apo 60x/1.40 NA oil DIC objective magnification using the appropriate filter set, or analysed on an Attune FACS machine (50 mW Blue/Violet standard configuration), with 10,000 events observed. CFW and Aniline Blue fluorescence intensities were quantified using the 405 nm laser on the Attune in combination with 603/48 and 650DPL filters, FITC labelled cells were quantified using the 488 nm laser in combination with 530/30 and 555DLP filters, and TRITC fluorescence was quantified using the 488 nm laser in combination with 574/26 and 650DLP filters. The MFI was corrected for background fluorescence and expressed as a ratio compared to YPD.
grown cells. FACS data were analysed by Kruskal-Wallis test followed with a post-hoc Dunn’s multiple comparisons test at 95% confidence.

**β1,3-glucanase sensitivity assay.**

Yeast cells were inoculated in YPD and grown overnight at 37°C, 200 rpm. Cells were sub-cultured into YPD buffered at the appropriate pH to an OD₆₀₀ of 0.1 and grown until mid exponential phase at 37°C, 200 rpm. Cells were harvested, washed once in sterile water and resuspended in fresh assay buffer (40 mM 2-mercaptoethanol, 50 mM Tris-HCl pH 7) at an OD₆₀₀ of 1.0 and 190 μl added to triplicate wells of a 96-welled plate. β1,3-glucanase (Sigma-Aldrich, UK) was resuspended in sterile ultra pure water to (2 U/ml) and 10 μl (0.02 units) added to each well. The OD₆₀₀ was recorded every 2 min as a measure of cell lysis, and data expressed as a percentage of the OD₆₀₀ at the initial time point. The rate of cell lysis was determined from the first 100 min and data were analysed by Kruskal-Wallis test followed by a post-hoc Dunn’s multiple comparisons test at 95% confidence.

**HPLC analysis of the carbohydrate component of the cell wall**

NGY152 was grown to exponential phase in YPD media, or YPD media buffered at the appropriate pH at 37°C and the carbohydrates from the cell wall extracted as described previously (48). Lyophilised cell wall material (3 mg) was acid hydrolysed with trifluoroacetic acid for 3 h at 100°C, washed, resuspended to 10 mg/ml and diluted 1:2 for HPLC analysis. Data were analysed by two-way ANOVA followed by post-hoc Tukey’s multiple comparison test at 95% confidence.

**Activation of cell wall remodelling pathways**

To deduce whether Hog1 was activated during prolonged exposure to different environmental pH, a *C. albicans* strain expressing a GFP tagged version of Hog1 was utilised (hAHGL). GFP-Hog1 was grown to exponential phase in YPD at the
appropriate pH, fixed and immediately imaged for GFP-Hog1 localisation. The percentage of cells positive for nuclear localisation (and hence activation), of GFP-Hog1 was quantified in ImageJ from 200 cells per condition per repeat. As a positive control for Hog1 activation, cells growing exponentially in YPD were exposed to 1 M NaCl for 30 min prior to fixation and imaging.

To deduce whether the cell wall salvage pathway was activated upon adaption to environmental pH, NGY152 was grown in appropriately buffered YPD media overnight, diluted 1:100 into fresh media at the appropriate pH and grown at 37°C, 200 rpm until exponential phase. As a positive control for Mkc1 activation, 0.032 μg/ml caspofungin was added to exponentially growing YPD cells for 30 min. Cells were harvested by centrifugation, and immediately snap frozen in liquid nitrogen. Pellets were defrosted in 500 μl RE buffer (50 mM HEPES pH 7.5, 150 mM NaCl, 5 mM EDTA, 1% Triton X-100) supplemented with Roche complete proteinase inhibitor cocktail, washed and resuspended in 500 μl RE buffer. Cells were lysed using a bead beater (6 x 2 cycles 6000 rpm) with 5 min between each cycle. Cell lysis was confirmed by microscopy, and lysates cleared by centrifugation. Total protein concentration was determine by Bradford assay in comparison to a BSA standard curve and 15 μg of total protein was separated by SDS-PAGE on a 4-12% NuPAGE Bis-Tris gel. Proteins were transferred on PVDF membrane, which was blocked with 5% BSA in PBST. Activated (phosphorylated) Mkc1 and Cek1 was detected using an anti-phospho-p44/p42 rabbit monoclonal antibody (Cell Signalling technologies) diluted 1:2000 in 5% BSA, PBST. Protein-antibody complexes were detected using an anti-rabbit IgG-HRP antibody (Invitrogen), diluted 1:5000 in 5% BSA, PBST. Membranes were washed in PBST and signal detected using enhanced chemiluminescence (ECL) kit (Bio-Rad) as per the manufacturer’s recommendations.

**Phagocytosis assay**
J774.1A macrophages (Sigma-Aldrich, UK) were maintained in DMEM media supplemented with 10% FBS, 100 mM L-glutamine and 100 mM penicillin/streptomycin at 37°C, 5% CO₂. 1 x10⁵ J774.1A macrophages were seeded onto 13 mm diameter glass coverslips in 24-well plates and allowed to attach for 24 h. Immediately prior to phagocytosis assays, J774.1A macrophages were serum starved in serum free DMEM media for 1 h with 1.5 µg/ml PMA. Yeast cells were inoculated in YPD and grown overnight at 37°C, 200 rpm. Cells were sub-cultured into YPD buffered at the appropriate pH to an OD₆₀₀ of 0.1 and grown until mid-exponential phase at 37°C, 200 rpm. Cells were harvested, washed three times in sterile, endotoxin free PBS (Sigma-Aldrich, UK) and resuspended in PBS to 1 x10⁷ cells/ml. PMA containing media was aspirated from the macrophages and replaced with fresh serum free DMEM media, to which 5 x10⁵ Candida cells were added (MOI = 5). Cells were co-incubated for 1 h, non-phagocytosed Candida cells were removed by repeated washing with sterile PBS and cells fixed with 4% PFA for 15 min. To block Dectin-1 recognition, cells were incubated with 50 µg/ml glucan phosphate in serum-free media for 1 h prior to addition of C. albicans. Media was replaced with fresh serum free media containing 5 x10⁵ C. albicans cells and 50 µg/ml glucan phosphate (Kind gift from Prof. D. Williams, East Tennessee State University), and cells co-incubated for 1 h. To distinguish between attached and phagocytosed yeasts, coverslips were stained for 30 min with 50 μg/ml ConA conjugated to TRITC (Molecular Probes, Life Technologies), washed three times with PBS and imaged using a Nikon TE2000. At least six images were taken per sample with approximately 100 macrophages/image. Phagocytosis was scored in ImageJ. Candida cells stained with ConA-TRITC were considered attached to the exterior of the macrophage while, non-stained yeasts were considered as internalised and phagocytosed. Data were analysed by Kruskal-Wallis test followed with a post-hoc Dunn's multiple comparisons test at 95% confidence.
**Fibroblast association assay**

To assess attachment of *C. albicans* to fibroblasts (NIH3T3) or fibroblasts expressing human Dectin-1 (NIH3T3-Dectin-1, kind gift from Prof G. Brown, University of Aberdeen) (50), 1 x10^5 fibroblasts were seeded on glass coverslips and allowed to attach for 24 h at in DMEM supplemented with 10% FBS, 100 mM L-glutamine and 100 mM penicillin/streptomycin 37°C, 5% CO_2. Immediately prior to the association assay, fibroblasts were serum starved in serum free DMEM media for 1 h. *C. albicans* adapted to different pH conditions were added to the fibroblasts at an MOI = 5. Cells were co-incubated for 1 h, non-attached *Candida* cells were removed by repeated washing with sterile PBS and cells fixed with 4% PFA for 15 min. To block Dectin-1 recognition, cells were incubated with 50 µg/ml glucan phosphate in serum-free media for 1 h prior to addition of *C. albicans*. Media was replaced with fresh serum free media containing 5 x10^5 *C. albicans* cells and 50 µg/ml glucan phosphate, and cells co-incubated for 1 h. Attachment was scored in ImageJ. Data were analysed by Kruskal-Wallis test followed with a post-hoc Dunn’s multiple comparisons test at 95% confidence.

**Neutrophil isolation**

Peripheral whole blood (18 ml) was taken from healthy volunteers and immediately laid under dual Percoll (GE Healthcare) density gradients of 1.098 and 1.079. Gradients were centrifuged at 150 x g for 8 min, followed by 1200 x g for 10 min. The neutrophil layer was removed to a fresh tube containing 3 volumes of red blood cell lysis buffer (0.83% NH_4Cl, 0.1% KHCO_3, 0.004% Na_2EDTA.2H_2O and 0.25% BSA) and gently agitated for 3 min to lyse contaminating red blood cells. Neutrophils were centrifuged at 400 x g for 6 min, and the resulting pellet washed twice in sterile endotoxin free PBS. Neutrophils were resuspended in serum free RPMI supplemented with 100 mM L-glutamine at a cell concentration of 1 x10^5 cells/ml.
Neutrophil phagocytosis assay

Two hundred microlitres of neutrophils (0.2 x 10^5 cells) were co-incubated with 1 x 10^5 C. albicans cells (MOI = 5) grown to mid-exponential phase in buffered YPD for 1 h. Non-phagocytosed C. albicans cells were removed by repeated washing with sterile PBS and cells fixed with 4% PFA for 15 min. Cells were washed once with PBS to remove PFA and cells were wet mounted and immediately imaged in duplicate. At least 300 neutrophils were counted for each condition per experiment. Data were analysed by one-way ANOVA followed by post-hoc Tukey test at 95% confidence.

Stimulation of peripheral blood mononuclear cells

PBMCs were isolated as described above, except the monocyte layer was extracted from the Percoll gradient. NGY152 was grown to exponential phase in YPD buffered at the appropriate pH, washed in PBS and resuspended to the desired cell concentration. Yeast cells were fixed with 4% PFA for 30 min washed three times with PBS and 0.5 x 10^5 C. albicans cells were added to 2.5 x 10^5 PBMCs (MOI = 0.5) in a final volume of 200 μl. Samples were incubated at 37°C, 5% CO_2 for 24 h. Samples were centrifuged and supernatant transferred into a fresh 96-welled plate and stored at -20°C. Extracellular cytokines were measured using commercially available ELISA kits (R&D) according to the manufacturer’s recommendations. Data were analysed by two-way ANOVA, followed by post-hoc test at 95% confidence level.

Neutrophil recruitment assay

C. albicans NGY152 yeast cells were inoculated in YPD and grown overnight at 37°C, 200 rpm. Cells were sub-cultured into YPD buffered at the appropriate pH to an OD_{600} of 0.1 and grown until mid-exponential phase at 37°C, 200 rpm. Cells were harvested, washed three times in sterile, endotoxin free saline and resuspended in
saline to $1 \times 10^6$ cells/ml for subsequent inoculation into mice. Cell counts were verified by plating for CFUs.

BALB/c female mice ($n=5$/group, 8-12 weeks old, Envigo, UK) were injected intraperitoneally with the prepared inocula at $10^7$ C. albicans in 100 μL sterile saline. Group size was determined from previous experiments as the minimum number of mice needed to detect statistical significance ($p<0.05$) with 90% power. Mice were randomly assigned to groups by an investigator not involved in the analysis and the fungal inocula were randomly allocated to groups. Mice were housed in Individually Ventilated Cages (IVCs) and were provided with food and water ad libitum. Inocula were delivered in an unblinded fashion. After 4 h, mice were sacrificed by IV euthatal injection. Immune infiltrates were collected by peritoneal lavage (51). Cells were stained to discriminate live/dead cells using a UV fixable live/dead dye (Thermo) and then fixed in 2% PFA. Cells were stained for CD45, CD11b, and Ly-6B.2 (7/4 clone) to differentiate lymphocytes, and F4/80 and Ly6G to differentiate neutrophils and macrophages. Using flow cytometry, 10,000 cells were analysed for each mouse. Cells were analysed on a BD Fortessa flow cytometer, with automatic compensation protocols. Statistical analyses were performed using Graph Pad Prism (v 7). Significance was determined using Welch’s t-test for unpaired data. Bars represent 95% CI. Variance between the groups for total cell counts was statistically different (F test to compare variances, $p=0.0354$). All animal experiments were performed under UK Home Office project license PPL 70/902760/4135 granted to DMM in accordance with Home Office ethical guidelines. Work was performed by DMM and ERB.

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We would like to acknowledge Professor Gordon Brown, University of Aberdeen for providing Fc-Dectin-1 and Dectin-1 expressing fibroblast cell lines, Professor David Williams, East Tennessee State University, for providing glucan phosphate, the Candida community for strains and reagents, the Microscopy and Histology Core facility in the Institute of Medical Sciences, University of Aberdeen where the high pressure freezing was performed, staff at the University of Aberdeen Medical Research Facility, Raif Yuecel and Linda Duncan in the Iain Fraser Cytometry Centre (Aberdeen University) for their expert help with the cytometry experiments, and all members of the host and pathogen interaction (HAPI) group at the University of Birmingham for fruitful discussions.
References

1. Sobel JD. Vagi


Figure 1. Adaptation to environmental pH alters the ultrastructure of the fungal cell wall. a) Electron micrographs showing the ultrastructure of the cell walls of wild type C. albicans (NGY152) grown to mid-log phase in YPD buffered to the appropriate pH. Scale bar represents 100 nm. b) Quantification of the thickness of the inner cell wall layer and mannoprotein fibril length. Data represent the mean ± SEM from 5 individual cells. Each cell was measured at 30 different points around the cell periphery (* p < 0.05, ** p < 0.01, *** p < 0.001).

Figure 2. Adaptation to acidic environments promotes surface exposure of chitin. a) Wild type cells (NYG152) were grown in YPD buffered at the appropriate pH and stained with CFW. The CFW fluorescence was quantified by FACS. The fold increase in CFW staining was determined from the background subtracted MFI values from FACS analysis, and normalised to YPD grown cells. b) Fold increase in FITC fluorescence of FITC-WGA stained wild type cells (NYG152) grown at different pH, relative to growth in YPD as quantified by FACS analysis. All data represent the mean ± SEM from three independent experiments. c) Microscopy of WGA stained cells. Arrowheads indicate chitin exposure. Scale bar = 10 μm.

Figure 3. Mkc1, Hog1 and Crz1 are not required for de-cloaking of chitin in response to acidic pH. a) A C. albicans strain expressing Hog1-GFP was grown in YPD buffered at the appropriate pH for 4 h and the localisation (cytoplasmic vs. nuclear) of Hog1 scored in 200 cells per condition. As a positive control of Hog1 activation, cells grown in YPD were incubated with 1 M NaCl for 30 min prior to imaging. The data represent the mean ± SEM from three independent experiments. b) Increase in WGA and CFW staining in kinase mutants grown at pH4 relative to YPD as quantified by FACS analysis. Data represent the mean ± SEM from three independent experiments. DAY286 is the parental control strain of the two kinase mutants c) Wild type C. albicans (NGY152) was grown to mid-log phase in YPD.
buffered at the appropriate pH, total protein extracted and activation of Mkc1 and Cek1 assessed via western blot using the MAPK P42/p44 antibody which cross reacts with both kinases. Arrow indicates Mkc1, while arrowhead indicates Cek1.  

Figure 4. Masking of chitin in the cell wall is regulated by Rim101 and Bcr1 and requires Cht2.  
a) Respective strains were grown in YPD buffered at pH4 or pH6 to mid-log phase, cells were fixed and stained with FITC-WGA. Fluorescence was quantified by FACS.  
b) SC5314 was grown to mid-log phase in YPD buffered at pH4, pH6 and pH8, snap frozen and total RNA extracted. Expression of CHT2 was determined by semi quantitative RT-PCR using 50 ng of total RNA. Expression levels were normalized to ACT1.  
c) Relative expression of CHT2 in rim101Δ and bcr1Δ mutants exponentially growth in YPD buffered at pH4, pH6 and pH8 and determined by semi quantitative RT-PCR. Expression levels were normalized to ACT1.  
d) FACS analysis of WGA staining in rim101Δ and bcr1Δ mutants.  
e) FACS analysis of WGA staining in CARP1-1 which expresses constitutively active Rim101  
f) FACS analysis of WGA staining in she3Δ mutants. Data represent the mean ± SEM from three independent experiments. (** p < 0.001, *** p < 0.001, ** p < 0.01, * p < 0.05).  

Figure 5. Acidic environments unmask β-glucan in C. albicans.  
a) i) Wild type C. albicans (NGY152) was grown to mid-log phase in YPD buffered at the appropriate pH, and incubated with recombinant β1,3-glucanase. The decrease in OD₆₀₀ represents cell lysis as the β1,3-glucanase digests the cell wall and is expressed as a percentage of the starting OD₆₀₀.  
ii) The initial rate of cell lysis as calculated from the first 100 min. Data represent the mean ± SEM from four independent repeats  

b)
Immunofluorescent imaging of β-glucan exposure of exponentially growing NGY152 cells using a anti-β1,3-glucan monoclonal antibody. Scale bar = 10 μm. c) Quantification of β-glucan exposure by FACS counting 10,000 events per repeat. Fold increased is relative to unbuffered YPD. Data represent the mean ± SEM from six independent experiments. d) Quantification of Aniline Blue staining of exponentially growing NGY152 cells by FACS analysis counting 10,000 events per repeat. Fold increased is relative to unbuffered YPD. Data represent the mean ± SEM from three independent experiments. e) β-glucan exposure of clinical C. albicans isolates grown to mid-log phase in YPD buffered to pH4 relative to YPD. Data represent the mean ± SEM from three independent experiments (* p < 0.05).

Figure 6. pH-dependent cell wall remodeling occurs in response to physiologically relevant acids and is not restricted to yeast cells. a) C. albicans cells (NGY152) were grown in YPD, or YPD buffered to pH4 by the addition of HCl or lactic acid to mid-log phase, fixed with 4% PFA and stained for β-glucan and chitin exposure. Fluorescence intensity was quantified by FACS. Data represent the mean ± SEM from three independent repeats (**** p < 0.0001). b) C. albicans cells (NGY152) were grown in vaginal secretion medium (VSM) buffered at pH4, pH6 or pH7 for 4 h, fixed and stained for β-glucan. Fluorescence intensity was quantified by FACS. Data represent the mean ± SEM from six independent repeats (**** p < 0.01). c) C. albicans cells (NGY152) were grow in 24-well plates in either YPD buffered at pH4 and pH6, DMEM buffered at pH4 and pH6 at 37C, 150 rpm, or grown in DMEM buffered at pH4 or pH6 with 5%CO2 in the presence or absence of A431 vaginal epithelial cells for 4 h. Cells were fixed in 4% PFA and stained for β-glucan and d) chitin exposure. Results are fold increase relative to pH6. Data represent the mean ± SEM from three independent repeats.
Figure 7. Exposure of β-glucan and chitin do not co-localise. Wild type *Candida albicans* cells (NGY152) were grown the mid-log phase in YPD buffered at pH2, pH4 and pH6, washed, fixed with 4% PFA and stained with CFW, TRITC-labeled WGA, and Fc-Dectin-1 (FITC). White arrowheads indicate patches of β-glucan exposure. Scale bar represents 10 μm.

Figure 8. Unmasking of β-glucan in response to environmental pH is specific to *C. albicans* and *C tropicalis*. a) Exponentially growing cells in YPD, YPD buffered to pH4 and YPD buffered to pH8 were incubated with recombinant β1,3-glucanase. The decrease in OD$_{600}$ represents cell lysis as the β1,3-glucanase digests the cell wall and is expressed as a % of the starting OD$_{600}$. Data represent the mean ± SEM from four independent experiments. b) Initial rate of cell lysis was calculated from the first 100 min after the addition of β-glucanase. Data represent the mean ± SEM from four independent repeats (* p < 0.05, ** p < 0.01).

Figure 9. Adaptation to acidic environments increases immune recognition. *C. albicans* was grown in YPD at the appropriate pH to mid-log phase, co-incubated with a) J774.1A macrophages b) neutrophils at an MOI = 5 for 1 h and the rate of phagocytosis determined. Data represent the mean ± SEM from four independent repeats. PBMCs were incubated with PFA fixed mid-log phase wild type *C. albicans* cells (NGY152), at an MOI of 0.5 for 24 h. Cytokine secretion was quantified by ELISA, c) TNFα, d) IL-6, e) IL-1β, f) IFNγ. Data represent the mean ± SEM from six donors (* p < 0.05, ** p < 0.01, *** p < 0.001, **** p < 0.0001).

Figure 10. Enhanced immune recognition of acidic adapted cells is mediated via Dectin-1. a) Fc-Dectin-1 binding to wild type *C. albicans* cells grown to mid-log phase in YPD buffered to the appropriate pH, as quantified by FACS counting 10,000 events per repeat. Fold increased is relative to unbuffered YPD. b) *C. albicans* was
grown in YPD at the appropriate pH to mid-log phase, co-incubated with fibroblasts for 1 h, fixed and the association index (number of fungal cells either attached and phagocytosed/100 macrophages) determined. c) Association of acidic adapted C. albicans cells to fibroblasts expressing Dectin-1 in the presence of glucan phosphate. d) J774.1A macrophages were pre-incubated with glucan phosphate and infected with C. albicans at an MOI of 5 and the phagocytosis index (number of fungal cells phagocytosed/100 macrophages) determined after 1 h. e) J774.1A macrophages were pre-incubated with glucan phosphate and infected with C. albicans at an MOI of 5 and the association index determined after 1 h. Data represent the mean ± SEM from three independent repeats (* p > 0.05. *** p > 0.001).

Figure 11. C. albicans cells adapted to acidic environments recruit more innate immune cells in vivo. a) Total number of CD45+, CD11b+ and Ly-6B.2+ (7/4 clone) cells (including monocytes and neutrophils) recruited to the peritoneal cavity following 4 h exposure to C. albicans incubated in pH6 or pH4 YPD (p=0.007). b) Total number of neutrophils (further identified using F4/80) recruited to the peritoneal cavity (p=0.010) c) Percentage of neutrophils in the total population of recruited innate immune cells (p=0.540).

Figure 12. Regulation of cell wall remodeling during adaptation to environmental pH. a) In environments above pH5.5, the Rim101 pathway is activated resulting in C-terminal processing of Rim101 and enhance expression of CHT2. The She3 complex then transports CHT2 mRNA to the cell wall, where it is translated and attached via its GPI anchor. At pH4 Rim101 is not activated, resulting in reduced expression of Cht2. b) (i) When present in the cell wall (i.e. above pH5.5), Cht2 hydrolyses the growing chitin polymer into shorter fragments which may hydrogen bound and form short chitin microfibrils that are embedded deep within the
cell wall. (ii) When there is reduced amounts of Cht2 in the cell wall (i.e. environments below pH4), the growing chitin polymer in the cleaved less efficiently, resulting in the incorporation of longer chitin polymers that are more exposed on the outer surface of the cell wall.

Supplementary Figure 1. De-cloaking of the fungal cell wall in response to environmental ph is an active process. C. albicans cells were grown overnight in YPD. Cells were killed by fixing with 4% PFA, heat killing at 65°C for 2 hrs, treatment with 1 Kj UV light, or 100 mM thimerosal for 45 mins. Cells were washed and incubated in YPD buffered at pH2, 4 and 6 for 4 hrs. Cells were stained for a) chitin and b) β-glucan exposure. Data represent the mean and SEM from three biological repeats (* p < 0.05, ** p < 0.01, **** p < 0.001).

Supplementary Figure 2. Release of the outer phosphomannan does not result in unmasking of inner cell wall components. C. albicans strains were grown to mid-log phase in YPD and YPD buffered at pH4, fixed with 4% PFA and carbohydrate exposure quantified by immunofluorescence. Fluorescence was quantified by FACS analysis of 10,000 events per strain, per condition, per repeat and is expressed as the fold-increase at pH4 relative to YPD. Data represent the mean ± SEM from three independent repeats.

Supplementary Figure 3. Unmasking of β-glucan in response to environmental pH is not mediated via conventional cell wall or pH sensing pathways. a) β-glucan unmasking in kinase mutants grown to mid-log phase in YPD buffered to pH4 as quantified by FACS analysis of immunofluorescent staining and repressed as fold change relative to YPD. Data represent the mean ± SEM from three independent experiments. b) β-glucan unmasking in C. albicans transcription factor mutants grown to mid-log phase in YPD buffered to pH4 as quantified by FACS analysis of
immunofluorescent staining and repressed as fold change relative to YPD. Data represent the mean ± SEM from three independent experiments. c) β-glucan unmasking in Rim101 pathway mutants grown to mid-log phase in YPD buffered to pH4 as quantified by FACS analysis of immunofluorescent staining and repressed as fold change relative to YPD. Data represent the mean ± SEM from three independent experiments.

Supplemental Figure 4 The rim101Δ and bcr1Δ mutants display reduced phagocytosis. C. albicans strains were grown in YPD at the appropriate pH to mid-log phase, co-incubated with J774.1A macrophages at an MOI = 5 for 1 h and the a) phagocytosis index and b) association index determined. Data represent the mean ± SEM from three independent repeats. c) PBMCs were incubated with PFA fixed mid-log phase cells at an MOI of 0.5 for 24 h and TNFα secretion quantified by ELISA. Data represent the mean ± SEM from three donors in triplicate (* p < 0.05).

Table 1 Relative dry weight proportions (%) of glucan and chitin in the cell wall quantified by HPLC

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Supplemental Table 1. Strains used in this study.

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<td><em>C. dublinensis</em></td>
<td>Typed strain</td>
<td>(57)</td>
</tr>
<tr>
<td>AM16/0701</td>
<td><em>C. krusei</em></td>
<td>Clinical isolate</td>
<td>From D. MacCallum, University of Aberdeen</td>
</tr>
<tr>
<td>CLIB214</td>
<td><em>C. paraosilosis</em></td>
<td>Typed strain</td>
<td>(58)</td>
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<tr>
<td>ATCC 2001</td>
<td><em>C. glabrata</em></td>
<td>Typed strain</td>
<td>ATCC</td>
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<tr>
<td>rim101Δ</td>
<td><em>C. albicans</em></td>
<td>arg4Δ/arg4Δ leu2Δ/leu2Δ his1Δ/his1Δ URA3/ura3Δ:: λimm434 IRO1/iro1Δ:: λimm434 rim101::LEU/rim101::HIS1</td>
<td>(59)</td>
</tr>
<tr>
<td>Gene</td>
<td>Organism</td>
<td>Genetic Modifications</td>
<td>References</td>
</tr>
<tr>
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<td>---------------------------------------------------------------------------------------</td>
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</tbody>
</table>
| mkc1Δ  | C. albicans | λimm434/ura3Δ::λimm434  
\(\text{his}1\Delta \text{hisG/}\text{his}1\Delta::\text{hisG} \text{arg}4\Delta::\text{hisG-ARG}4-\text{URA3}/\text{arg}4\Delta::\text{hisG} \text{mkc}1::\text{URA3/mkc}1::\text{ARG}3\) | (60)       |
| rlm1Δ  | C. albicans | arg4Δ/arg4Δ leu2Δ/leu2Δ  
\(\text{his}1\Delta/\text{his}1\Delta \text{URA3/ura3Δ::λimm434} \text{IRO1/iro1Δ::λimm434} \text{rlm1::LEU/rlm1::HIS}1\) | (59)       |
| bcr1Δ  | C. albicans | arg4Δ/arg4Δ leu2Δ/leu2Δ  
\(\text{his}1\Delta/\text{his}1\Delta \text{URA3/ura3Δ::λimm434} \text{IRO1/iro1Δ::λimm434} \text{bcr1::LEU/bcr1::HIS}1\) | (59)       |
| crz1Δ  | C. albicans | arg4Δ/arg4Δ leu2Δ/leu2Δ  
\(\text{his}1\Delta/\text{his}1\Delta \text{URA3/ura3Δ::λimm434} \text{IRO1/iro1Δ::λimm434} \text{crz1::LEU/crz1::HIS}1\) | (59)       |
| efg1Δ  | C. albicans | arg4Δ/arg4Δ leu2Δ/leu2Δ  
\(\text{his}1\Delta/\text{his}1\Delta \text{URA3/ura3Δ::λimm434} \text{IRO1/iro1Δ::λimm434} \text{efg1::LEU/efg1::HIS}1\) | (59)       |
| czf1Δ  | C. albicans | arg4Δ/arg4Δ leu2Δ/leu2Δ  
\(\text{his}1\Delta/\text{his}1\Delta \text{URA3/ura3Δ::λimm434} \text{IRO1/iro1Δ::λimm434} \text{czf1::LEU/czf1::HIS}1\) | (59)       |
| cht1Δ  | C. albicans | λimm434/ura3Δ::λimm434  
\(\text{hisG/his}1\Delta::\text{hisG} \text{arg}4\Delta::\text{hisG-ARG}4-\text{URA3}/\text{arg}4\Delta::\text{hisG}\) | (61)       |
<table>
<thead>
<tr>
<th>Strain</th>
<th>Species</th>
<th>Transformation</th>
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<tr>
<td>cht2Δ</td>
<td>C. albicans</td>
<td>λimm434/ura3Δ::λimm434, hisG/his1Δ::hisG arg4Δ::hisG-URA3/arg4Δ::hisG, cht2::URA3/cht2::ARG3</td>
<td>(61)</td>
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<td>cht3Δ</td>
<td>C. albicans</td>
<td>λimm434/ura3Δ::λimm434, hisG/his1Δ::hisG arg4Δ::hisG-URA3/arg4Δ::hisG, cht3::URA3/cht3::ARG3</td>
<td>(61)</td>
</tr>
<tr>
<td>CARP1-1</td>
<td>C. albicans</td>
<td>prr1Δ::hisG/prr1Δ::hisG-URA3-hisG ura3Δ: λimm434/ura3Δ:: λimm434, RIM101&lt;sup&gt;1697A&lt;/sup&gt;</td>
<td>(62)</td>
</tr>
<tr>
<td>hog1Δ</td>
<td>C. albicans</td>
<td>λimm434/ura3Δ::λimm434, hisG/his1Δ::hisG arg4Δ::hisG-URA3/arg4Δ::hisG, hog1::URA3/hog1::ARG3</td>
<td>(51)</td>
</tr>
<tr>
<td>hAHGI</td>
<td>C. albicans</td>
<td>λimm434/ura3Δ::λimm434, hog1::hisG/hog1::hisG ACT1p-HOG1-GFP::leu2/LEU2</td>
<td>(63)</td>
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<table>
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<tr>
<th>Isolate</th>
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<th>Institution</th>
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<tbody>
<tr>
<td>302</td>
<td>C. albicans</td>
<td>Clinical isolate from urine</td>
<td>National Institute for Health</td>
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<tr>
<td>300</td>
<td>C. albicans</td>
<td>Clinical isolate from blood</td>
<td>Research</td>
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<td>C. albicans</td>
<td>Clinical isolate from sputum</td>
<td>Surgery</td>
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<td>316</td>
<td>C. albicans</td>
<td>Clinical isolate from burn wound</td>
<td>Reconstruction and Microbiology</td>
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<td>317</td>
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<td>Clinical isolate from wound drain</td>
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<tr>
<td>320</td>
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<td>Clinical isolate from wound</td>
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<tr>
<td>Reference</td>
<td>Organism</td>
<td>Strain Characteristics</td>
<td>Year</td>
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<td>-----------</td>
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<td>CDH14</td>
<td><em>C. albicans</em></td>
<td>ura3Δ::λimm434/ura3Δ::λimm434, mnn4Δ::hisG/mnn4Δ::hisG, RP10::[MNN4-URA3-RP10]</td>
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<tr>
<td>CDH15</td>
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<td>ura3Δ::λimm434/ura3Δ::λimm434, mnn4Δ::hisG/mnn4Δ::hisG, RP10::URA3</td>
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<tr>
<td>she3Δ</td>
<td><em>C. albicans</em></td>
<td>she3Δ/she3Δ; ura3Δ/ura3Δ</td>
<td>27</td>
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</table>