Replicative Aging Remodels the Cell Wall and Is Associated with Increased Intracellular Trafficking in Human Pathogenic Yeasts

Vanessa K. A. Silva, Somanon Bhattacharya, Natalia Kronbauer Oliveira, Anne G. Savitt, Daniel Zamith-Miranda, Joshua D. Nosanchuk, Bettina C. Fries

Division of Infectious Diseases, Department of Medicine, Stony Brook University, Stony Brook, New York, USA
Department of Microbiology and Immunology, Renaissance School of Medicine, Stony Brook University, Stony Brook, New York, USA
Department of Microbiology and Immunology, Albert Einstein College of Medicine, Bronx, New York, USA
Division of Infectious Diseases, Department of Medicine, Albert Einstein College of Medicine, Bronx, New York, USA
Veterans Administration Medical Center, Northport, New York, USA

Vanessa K. A. Silva and Somanon Bhattacharya share first authorship. Both have generated the majority of the data. Vanessa is in first position because she led the investigations and wrote the majority of the manuscript.

ABSTRACT Replicative aging is an underexplored field of research in medical mycology. Cryptococcus neoformans (Cn) and Candida glabrata (Cg) are dreaded fungal pathogens that cause fatal invasive infections. The fungal cell wall is essential for yeast viability and pathogenesis. In this study, we provide data characterizing age-associated modifications to the cell wall of Cn and Cg. Here, we report that old yeast cells upregulate genes of cell wall biosynthesis, leading to cell wall reorganization and increased levels of all major components, including glucan, chitin, and its derivatives, as well as mannan. This results in a significant thickening of the cell wall in aged cells. Old-generation yeast cells exhibited drastic ultrastructural changes, including the presence of abundant vesicle-like particles in the cytoplasm, and enlarged vacuoles with altered pH homeostasis. Our findings suggest that the cell wall modifications could be enabled by augmented intracellular trafficking. This work furthers our understanding of the cell phenotype that emerges during aging. It highlights differences in these two fungal pathogens and elucidates mechanisms that explain the enhanced resistance of old cells to antifungals and phagocytic attacks.

IMPORTANCE Cryptococcus neoformans and Candida glabrata are two opportunistic human fungal pathogens that cause life-threatening diseases. During infection, both microorganisms have the ability to persist for long periods, and treatment failure can occur even if standard testing identifies the yeasts to be sensitive to antifungals. Replicative life span is a trait that is measured by the number of divisions a cell undergoes before death. Aging in fungi is associated with enhanced tolerance to antifungals and resistance to phagocytosis, and characterization of old cells may help identify novel antifungal targets. The cell wall remains an attractive target for new therapies because it is essential for fungi and is not present in humans. This study shows that the organization of the fungal cell wall changes remarkably during aging and becomes thicker and is associated with increased intracellular trafficking as well as the alteration of vacuole morphology and pH homeostasis.

KEYWORDS Candida glabrata, Cryptococcus neoformans, cell wall, replicative lifespan, vesicle trafficking

Fungal pathogens are an emerging threat to global health due to the rise of immunosuppressed patients (1). Their prevalence may even be affected by the evolving...
climate change (2, 3). Among the human fungal pathogens, Candida and Cryptococcus account for most of the invasive fungal infections (4). Cryptococcus neoformans (Cn), a member of the phylum Basidiomycota, is the leading agent of fungal meningitis and affects mostly patients living with advanced HIV disease (5). Candida glabrata (Cg), a member of the phylum Ascomycota, is the second most common fungal pathogen that causes bloodstream infections in North America and is associated with rapidly acquired antifungal resistance (6, 7). Currently, there are no vaccines for fungal diseases, and antifungal therapeutic options are still limited (8, 9).

The fungal cell wall is an ideal target for antifungal chemotherapies since it is essential for fungal viability and virulence but absent in human cells (10). The fungal cell wall is composed of a complex matrix of glucose, proteins, lipids, and pigments (11). The cell wall of Cn is mainly composed of polymers of glucose (α- and β-glucans), N-acetylglucosamine (GlcNAc) (chitin and chitooligomers), glucosamine (chitosan), and glycoproteins (mannoproteins) (12). Similarly, the cell wall of Cg is mainly composed of α- and β-glucans, mannoproteins, and chitin (10).

Glucans and chitin are the most important structural components of fungal cell walls (13). Chitin can be degraded to chitooligomers, chitin-derived structures composed of 3 to 20 residues of GlcNAc (14). Chitooligomers are found around nascent buds and participate in connecting the cryptococcal cell wall to the capsule (14, 15). Also, chitin can be enzymatically deacetylated to chitosan (16). Most fungal pathogens expose chitin, but Cn is one of the few species that can evade the host immune system by replacing the chitin in its cell walls with chitosan (17). Mannose residues by N- or O-linkages can be found in association with proteins and are highly immunogenic (11).

The fungal cell wall is a dynamic structure that regulates cellular morphology (18) and remodels its components in response to environmental stresses, morphogenesis, cellular growth, and cell division (19). Aging is the consequence of cell division and a conserved natural process among eukaryotic cells. Replicative aging is the result of asymmetric cell divisions and is an important contributor to pathogenesis (20). During replicative aging, the fungal cell wall becomes thicker, which is associated with increased resistance to antifungals and phagocytic uptake (21, 22). However, the changes in cell wall composition and architecture during aging have not been elucidated. In this study, we investigate cell wall remodeling during aging in Cn and Cg. We report the increased expression of essential genes related to fungal cell wall biosynthesis. These transcriptome changes concur with our observation of enhanced levels of significant cell wall components (chitin, chitooligomers, and glucans) and enhanced cell wall thickness. Cell wall remodeling during aging is accompanied by formation of intracellular vesicles and multivesicular bodies, which suggests intensification of vesicular trafficking. In addition, our data indicate modifications in the vacuole morphology and pH, underscoring the importance of this organelle in the dynamics of progressive aging. Our data provide insights into the complex aging-related changes in fungal pathogens.

RESULTS

Regulation of cell wall biosynthesis genes is altered in older-generation cells.

Old yeast cells previously biotin-labeled and conjugated to streptavidin were separated when they reached the desired generation using a magnetic field, while nonlabeled young cells were washed off from the system. After isolation, we first performed a focused transcriptome analysis on known essential cell wall biosynthesis genes. Of the eight chitin synthases in Cn and four in Cg, CHS3 generates the majority of chitin (16, 23). CHS3 was upregulated in old Cn (3.3-fold change, \( P < 0.0001 \)) (Fig. 1A) but not in old Cg (Fig. 1B). Likewise, we observed upregulations of CHI22, an endochitinase involved in chitooligomer formation (24) in older Cn (2.38-fold change, \( P \leq 0.0001 \)) (Fig. 1A) and CTS1, an endochitinase (2.66-fold change, \( P \approx 0.0001 \)), in older Cg (Fig. 1B).

In Cn, chitin can be converted to chitosan by four putative chitin deacetylases (CDAs) (17, 25). We found that CDA1 transcription was downregulated (0.45-fold...
Old yeast cells presented increased levels of cell wall components. Flow cytometry was used to quantify the relative levels of cell wall carbohydrates in both young and old Cn and Cg (Fig. S1). Mean fluorescence intensity (MFI) levels confirmed increased chitin levels in old Cn (P < 0.0001; Fig. 4A) and Cg (P = 0.014; Fig. 4B) compared to young cells. In agreement with microscopic findings, both older Cn and Cg exhibited significantly more chitooligomer content (P < 0.0001 and P = 0.0008, change, P ≤ 0.0001), whereas both CDA3 and CDA4 were upregulated (2.27-fold change and 2.08-fold change, respectively, P ≤ 0.001) in old Cn (Fig. 1A). In Candida species, chitosan is present only in the chlamydospore cell wall and is generated by two genes (CDA1 and CDA2) (26), of which CDA2 was upregulated (>2-fold change, P ≤ 0.0001) in old Cg (Fig. 1B). This is different from the Cn, where CDA2 was not up-regulated in the old cells (Fig. 1A).

Transcription of AGS1, which encodes an α-1,3-glucan synthase, was upregulated in old Cn (9-fold change, P = 0.0001) (Fig. 1A). In contrast, transcription of α-1,4-glucan synthase (GLC3) of Cg was not significantly altered Fig. 1B). Genes relevant for β-glucan synthesis, FKS1 and SKN1, did not significantly change in old Cn (Fig. 1A) but were downregulated in old Cg, genes FKS2 (0.27-fold change, P ≤ 0.05) and SKN1 (0.46-fold change, P ≤ 0.0001). FKS3 was increased (2.48-fold change, P ≤ 0.0001), whereas KRE2 and KRES remained not markedly altered Fig. 1B).

Yeast cell wall architecture is reshaped during aging. Next, we assessed the cell wall main components by fluorescence microscopy (Fig. 2). In old Cn, chitin staining (by calcofluor white [CFW]) was enhanced surrounding the cell surface compared to in younger cells (Fig. 2A). In old Cg, chitin staining was enhanced but remained localized to bud scars in the cell wall (Fig. 2B). A similar pattern was noted with the binding of wheat germ lectin (WGA), which binds to chitooligomers. Again, it was abundant throughout the cell wall of most old Cn cells. In young Cn, chitooligomers are located in the area of the emerging developing buds (Fig. 2C). The latter pattern was similar to chitooligomer staining in young Cg. WGA staining of old Cg is also bound only to the numerous persistent bud scars (Fig. 2D), similar to the calcofluor binding to chitin. The deacetylated form of chitin, chitosan, binds specifically to the anionic dye eosin Y (25). Chitosan is uniformly enhanced and distributed through old Cn cell walls (Fig. 2E). In contrast, chitosan was not detected in Cg cell walls (data not shown).

The Cn polysaccharide capsule prevents binding of the monoclonal antibodies (MAb) MOPC-104E (α-glucan) and Fc-dectin (β-glucan); hence, like others, we were not able to assess α- and β-glucan levels (27, 28). Both young and old Cg exhibited comparable β-glucan staining of the cell surface (Fig. 2F). Finally, we found that mannoproteins were enriched in the region next to the developing buds in both young and old Cn. However, old Cn presented slightly greater intensity than young cells (Fig. 2G). Furthermore, mannoproteins of old cells also displayed a dotted staining pattern and colocalized in the capsule (Fig. 2H), which was not observed in young cells (Fig. 3A). In Cg, continuous staining of mannoproteins on the outer surface of the cell wall of young and old Cg cells was comparable (Fig. 2H).
Although chitoooligomers can be quantified by labeling with WGA, CFW penetrates the cell surface more efficiently, and it also binds to other chitin derivatives (28). As expected, chitosan levels were also significantly higher in old Cn compared to young cells ($P = 0.0058$; Fig. 4E).

Regarding the mannoprotein content, old Cn and Cg presented higher levels respectively; Fig. 4C and D). Although chitoooligomers can be quantified by labeling with WGA, CFW penetrates the cell surface more efficiently, and it also binds to other chitin derivatives (28). As expected, chitosan levels were also significantly higher in old Cn compared to young cells ($P = 0.0058$; Fig. 4E).
compared to younger cells (P = 0.0019 and P = 0.0015, respectively; Fig. 4G and H). β-β-glucan content in the Cn cell wall could only be quantified using a biochemical assay. These data indicated old Cn contained increased levels in comparison to young cells (P = 0.0062; Fig. 3C). Also, a higher level of β-glucan content was found in the cell wall of old Cg compared to that in young cells (P = 0.0025; Fig. 4F), which could not be documented by microscopy.

Old yeast cells developed thicker and robust cell walls, giant vacuoles, and increased multivesicular body-like structures. As expected, transmission electron microscopy (TEM) analysis showed that older Cn had a significantly thicker cell wall compared to that of younger cells (235.9 versus 123.7 nm; P < 0.0001). Specifically, we found in young Cn that the inner cell wall layer measured on average 66.76 nm, and the outer layer measured 91.29 nm. In old cells, the average thickness of the inner layer was 103.4 nm and grew to 138.3 nm (P < 0.0001). These results suggest that both cell wall layers enlarged equally during replicative aging. Old Cg also showed notable growth in the total thickness of the cell wall compared to young cells (132.5 versus 93.11 nm; P < 0.0001). Here, the inner cell wall layer was thicker than the outer. Specifically, in old Cg the inner layer measured 103.5 nm versus 78.21 nm in young cell (P = 0.0007). Conversely, the outer cell layer of young Cg was thicker than that of old Cg (34.75 nm versus 25.39 nm; P = 0.0017) (Fig. 5B).

Next, we tested if replicative aging affected cell wall strength. Cell wall integrity is vital
Old *C. neoformans* (Cn) and *C. glabrata* (Cg) presented increased cell wall content levels. (A to H) By flow cytometry, analysis of levels in young (RC2 Y and BG2 Y) and old cells of Cn (RC2 O) and Cg (BG2 O) of total chitin (A and B), exposed chitooligomers (C and D), chitosan (E), β-1,3-glucan (F), and mannoproteins (G and H). Unstained cells were sorted as controls to determined positive labeling. For each group, a total of 10,000 events were gated, and levels of chitin (blue; calcofluor white [CFW]), chitooligomers (orange; WGA-TRITC), chitosan (green; eosin Y [EY]), β-glucan (light blue; (Continued on next page)
for maintaining cell wall functionality and structure (29). We assessed the sensitivity of old cells to cell wall stressors, including Congo red (inhibits assembly of cell wall polymers such as chitin), caffeine (interferes with cell wall-related signal transduction pathways), and sodium dodecyl sulfate (disrupts plasma membrane) (28, 30). These data showed aging had no impact on resistance of the cell wall in old Cn and Cg (Fig. S2).

The ultramicroscopic images showed that old cells of both yeasts exhibited thicker cell walls and an accumulation of vesicle-like structures in the cytosol. (A and B) Quantification of the thickness of the inner (squares), outer (triangles), and total (inner + outer) (circles) layers of young (blue) and old (red) C. neoformans (Cn) (A) and C. glabrata (Cg) (B). Data represent the analysis of 30 individual cells for each group by imageJ/Fiji software. An unpaired t test with Welch’s correction was used to compare the pairs (young and old) of each group, and error bars represent the standard deviation (***, P ≤ 0.0001; ***, P ≤ 0.001; **, P ≤ 0.01; ###, P ≤ 0.0001; ##, P ≤ 0.01). The asterisk symbol (*) represents differences between the groups young and old, whereas the pound symbol (#) represents differences within the groups young or old. (A to J) Representative electron micrographs showing the ultrastructural changes of young and old cell walls of Cn (RC2) (C to F) and Cg (BG2) (G to J). White arrows point to vesicle-like particles in the cytoplasm of old yeast cells.

**FIG 4** Legend (Continued)

Fc-dectin1/DyLight 405), and mannan (red; concanavalin A, Texas Red conjugated) were represented by the mean fluorescence intensity (MFI). The following lasers were used: violet, 405 nm (for CFW and DyLight 405); YelGreen, 561 nm (for WGA-TRITC and CoA-TR); blue, 488 nm (for EY). All experiments were done in biological triplicates, and statistical analyses were performed by t test (****, P ≤ 0.0001; *** P ≤ 0.001; ** P ≤ 0.01; * P ≤ 0.05).
expanded vacuoles with amorphous lumen content, occupying a significant fraction of the total cell (Fig. 5E and I). Interestingly, the cytoplasm and cell wall (Fig. S3) of old cells contained a large number of electron-dense particles (Fig. 5E and I), suggestive of vesicles derived from the conventional secretion pathway. The vacuoles in old 
\textit{C. neoformans}\ (Fig. 6D) and 
\textit{C. glabrata}\ (Fig. 6H) contained vesicles resembling multivesicular bodies (MVBs). MVB-like structures accumulated in 67.31\% of old 
\textit{C. neoformans}, whereas they were found in only 24.53\% of young cells. In a similar fashion in old 
\textit{C. glabrata}, MVB-like structures were present in 33.96\% of old cells and only in 4\% in young cells (Table 1).

**Old \textit{C. neoformans} accumulated intracellular GXM and upregulated secretion gene \textit{SEC14}**. 
\textit{C. neoformans} extracellular vesicles (EVs) contain glucuronoxylomannan (GXM) (31). Analysis of ultrathin sections of both young and old 
\textit{C. neoformans} labeled with MAb 18B7 for GXM (32) revealed a significant increase in GXM detection in the intracellular space in old compared to young 
\textit{C. neoformans} (Fig. 7A to E). Immunogold-labeled antibodies detected GXM in the vacuole (Fig. 7D) and also bound to vesicle-like structures in the cell wall (Fig. 7E) of old yeast. We also tested transcription of two genes essential for secretion, \textit{SEC14} and \textit{SAV1} (33–35). \textit{SEC14} transcription was upregulated (3.09-fold change), whereas \textit{SAV1} transcription was not significantly altered in old 
\textit{C. neoformans} (Table 2).

| Ordinary | Oversized, MVB-like |
|----------|---------------------|
| Young    | ![Young Ordinary](image1.png) | ![Young Oversized](image2.png) |
| Old      | ![Old Ordinary](image3.png)   | ![Old Oversized](image4.png)   |
| \textit{C. neoformans} | ![C. neoformans](image5.png) |

| Young    | ![Young Ordinary](image6.png) | ![Young Oversized](image7.png) |
| Old      | ![Old Ordinary](image8.png)   | ![Old Oversized](image9.png)   |
| \textit{C. glabrata} | ![C. glabrata](image10.png) |

**Fig 6** Alterations of vacuolar morphologies in young and old 
\textit{C. neoformans}\ (\textit{Cn}) and 
\textit{C. glabrata}\ (\textit{Cg}). Representative transmission electron microscopy (TEM) images of young and old cells of \textit{C. neoformans} in the upper panel (A to D) and \textit{C. glabrata} in the lower panel (E to H). Boxed areas illustrating vacuoles (ordinary or oversized, with or without intravacuolar vesicles) (left panels), were magnified in the right panels. Scale bars represent 500 nm.
TABLE 1 Distribution of multivesicular body (MVB)-like structures in young and old cells of C. neoformans (Cn) and C. glabrata (Cg)*

| Strain       | Vacuole                  | Young (%) (n) | Old (%) (n) |
|--------------|--------------------------|---------------|-------------|
| Cn (RC2)     | Ordinary                 | 64.15 (34)    | 32.69 (17)  |
|              | Oversized, MVB-like      | 24.53 (13)    | 67.31 (35)  |
|              | Other                    | 11.32 (6)     | 0           |
| Cg (BG2)     | Ordinary                 | 80 (40)       | 49.05 (26)  |
|              | Oversized, MVB-like      | 4 (1)         | 33.96 (18)  |
|              | Other                    | 16 (8)        | 16.99 (9)   |

*Transmission electron microscopy (TEM) images of at least 50 cells from each group were analyzed according to the size of the vacuole and the presence of MVB-like structures.

**The number, size, and pH of vacuoles are affected in old yeast cells.** In fungi, the vacuole participates in secretory pathways, Ca\(^{2+}\) storage, and pH regulation (36, 37). Because reduced vacuolar acidity has been reported in aged Saccharomyces cerevisiae (38), we further investigated modifications in vacuole morphology and pH in the old cells. First, we stained the vacuolar membranes with FM 4-64 dye (Fig. 8A and D). The average vacuole area was significantly larger in old compared to young cells, corresponding to an increase of 3.46-fold for Cn (0.7481 \(\mu\)m\(^2\) versus 0.2162 \(\mu\)m\(^2\), \(P < 0.0001\)) and 4-fold in Cg (0.04 nm\(^2\) versus 0.01 nm\(^2\), \(P < 0.0001\)). The surface area of old cells was 1.82-fold increased in Cn (41.38 \(\mu\)m\(^2\) versus 22.68 \(\mu\)m\(^2\), \(P < 0.0001\)) and 1.63-fold in Cg (12.93 nm\(^2\) versus 7.89 nm\(^2\), \(P < 0.0001\)) (Fig. S4). Calculations of the ratio of vacuole and cell size confirmed disproportional vacuolar growth in old Cn (\(P = 0.0155\); Fig. 8B) and Cg (\(P < 0.0001\); Fig. 8F). The average number of vacuoles in young cells was higher in Cn (5.9 versus 7.8 for old and young cells, respectively, \(P = 0.0231\); Fig. 8C) and Cg (3.9 versus 6.4 for old and young, respectively, \(P < 0.001\); Fig. 8F). We hypothesize that the enlargement of vacuoles in aging cells could be the result of the fusion of multiple vacuoles.

Next, the vacuole pH of young and old cells was assessed with quinacrine staining, a fluorescent probe that increases under acidic conditions (39). In Cn, higher pH was observed in the vacuoles of old compared to young cells (2.98 \(\times\) \(10^6\) MFI versus 3.85 \(\times\) \(10^6\) MFI, \(P = 0.0155\)) (Fig. 8G). Consistent with this finding, ALL2 expression, which is linked to the homeostasis of intracellular pH in Cn (40), trended up, and CMR1, which promotes calcium homeostasis (41), was significantly upregulated (4.84-fold increase, \(P < 0.005\)) (Table 2). In contrast, old Cg cells exhibited lower pH in vacuoles compared to young cells (3.6 \(\times\) \(10^6\) MFI versus 2.6 \(\times\) \(10^6\) MFI, \(P = 0.0007\)) (Fig. 8H).

A simplified illustration model summarizes the phenotypic characteristics observed in this study during replicative aging in Cn and Cg in Fig. 9.

**DISCUSSION**

The cell wall remains an attractive target for developing fungal vaccines (9) and antifungal drugs (10) because it is essential for fungal virulence (11). This study demonstrates that critical genes for cell wall synthesis are upregulated during replicative aging of Cn and Cg. The development of a thicker cell wall in old cells is probably facilitated by increased intracellular trafficking, associated with the formation of oversized vacuoles that contain MVB-like structures. Additionally, our results indicate that the morphology and pH of vacuoles are affected in old yeast cells. The observed reorganization of the cell wall partially explains the enhanced resistance to phagocytosis and antifungals, which has been observed in Cn, Cg, Candida albicans and Candida auris (21, 22, 42, 43).

Higher chitin levels result from budding and are implicated in increasing antifungal resistance, albeit not in all fungal species (44–46). For instance, chitin content is associated with caspofungin resistance in Cn (45), but not in Cg (44). Specifically, increasing chitin levels were noted throughout the life span in both yeasts, similar to Saccharomyces cerevisiae (47). Staining patterns for chitooligomers mimicked the chitin staining pattern with more diffuse staining throughout the cell wall of old Cn and are strictly bud scar-associated in Cg. In Cn, binding of chitooligomer-specific antibodies increases fungicidal efficacy.
of amphotericin B (48). As chitooligomers are fungi-specific, this novel therapeutic approach could be explored to treat the enhanced antifungal tolerance of old Cn and possibly Cg.

In Cryptococcus species, chitosan is a critical virulence factor, and its levels can exceed those of chitin (16, 49, 50). It is produced from chitin by deacetylating enzymes (25), and CDA3 transcription was upregulated in old Cn. Cda3 activity was more important during infection by Cryptococcus gattii (49); however, Cda1 (51) and Cda2 were the main
regulators for Cn pathogenesis under host conditions (52). Chitosan inhibits protective Th1-type adaptive immune responses (53), maintains cell integrity, and allows melanization, thereby enhancing antifungal resistance (54–56). Our data suggest that higher chitosan content in the cell wall promotes enhanced melanization of old Cn cells during replicative aging (21). A common architecture for chitosan-containing elements in the fungal cell wall from ascomycetes to basidiomycetes has been proposed based on nuclear magnetic resonance (NMR) studies (57). Although Cg produces a melanin-like pigment (58), it is unknown if it accumulates during replicative aging. Chitosan was not detected in any Cg cells, similar to other Candida species, where expression is limited to the chlamydocyst (26).

Higher β-1,3-D-glucan levels were confirmed in old Cn by biochemical assays and in old Cg by fluorescent staining. Two of the three FKS genes involved in the synthesis were found to be upregulated in old Cg (42). However, FKS1 levels in old Cn were not significantly increased. Transcript and protein levels do not always correlate, and unstable mRNA can be translated more efficiently (59). In Cn lacking posttranscriptional gene regulator PUF4, the FKS1 mRNA and Fks1p levels did not correspond (45). Unfortunately, we were unable to quantify α-glucan with the available methods, but the α-glucan synthase AGS was upregulated in old Cn, indicating that this glucan is likely also increased. Nevertheless, it is still unclear whether the quantification of the cell wall content may be influenced by the enlargement of old cells. Further analysis, such as high-performance liquid chromatography, would be required to resolve this issue (60, 61). However, the low yield of old cells isolated from yeast culture is still currently a limitation for performing this technique.

Fungal pathogens can evade host immune recognition by masking β-glucans at the cell wall surface (62, 63). The exposure of β-glucans is reduced by Cn’s capsule (60) and by mannan in the outer cell wall layer in Candida species (44, 64). Mannan is present in Cn polysaccharide capsule (65, 66); thus, we propose that the observed accumulation of mannan could be due to augmented vesicle export. Alternatively, an altered permeability in the polysaccharide capsule in old cells (67) may efficiently expose the mannan epitopes. Altered mannoprotein content may contribute to the poor recognition of old yeast by host phagocytes (21, 22), which is suggested by data from C. gattii (68) and C. albicans (69).

Comparisons of TEM images of Cn and Cg identified differential growth of their inner and outer cell wall layer. The outer cell wall layer was thicker in old Cn, although the inner cell wall layer dominated in Cg. Chitinous structures and chitosan are evenly distributed across the cell wall of Cn but must be flexible enough to permit even expansion of the cell wall. We hypothesize that the heterogeneous composition of the inner cell wall in Cg results from chitin breaks in the bud scars, similar to S. cerevisiae (47), and causes the wavy cell wall. It is conceivable that the mannan layer in old Cg is more compactly arranged than in young cells. Reduced fibril length on the cell surface has been correlated with increased exposure on β-1,3-glucan (44). Alternatively, the outer layer of aged Cg can be more fragile and could have partially detached during ultrastructural processing. The increase in the inner cell wall layer thickness avoids a hyperosmotic shock and favors fungal cell survival (70). Most importantly, analogous to S. cerevisiae (47), old and young cells exhibit comparable tolerance to common cell wall stressors. Furthermore, comparable doubling times of 10th-generation (10GEN)

| Gene ID  | Gene name | Young (fold-change) | Old (fold-change) |
|---------|-----------|---------------------|------------------|
| CNAG_02817 | SAV1 | 1.034 | 0.517 |
| CNAG_03153 | SEC14 | 1.065 | 3.099* |
| CNM_02200 | ALL2 | 1.041 | 1.800 |
| CNAG_01704 | CMRI | 1.121 | 4.846* |

*Statistical difference was considered when the fold-change was greater than 2-fold.
FIG 8 Vacuole morphology and pH are affected in yeast old cells. (A and D) Images depicting vacuolar morphology (white arrows) stained with FM4-64 in young (Y) and old (O) cells of *C. neoformans* (Cn) (A) and *C. glabrata* (Cg) (D). DIC, differential interference contrast. (B and E) The ratio of vacuolar/cell area was significantly increased in old (red) Cn (B) and Cg (E) compared to younger cells (blue). An unpaired t test with Welch’s correction was used to generate the P-values (*, P = 0.0245; ****, P < 0.0001). (C and F) The average number of vacuoles per yeast cell was significantly lower in old (red) Cn (C) and Cg (F) compared to the respective young cells (blue). An unpaired t test with Welch’s correction was used (**, P = 0.0098; ****, P < 0.0001). For all analyses, 25 cells of each group were measured using ImageJ/Fiji software. Measurement of quinacrine levels with a spectrophotometer showed a significant decrease in fluorescence in aged Cn compared to young cells (G). This was confirmed with microscopy. In contrast, significantly increased fluorescence was measured in old cells of Cg compared to the young cells (H). This was also confirmed with microscopy. The assay was done in biological triplicate and an unpaired t test with Welch’s correction was used to generate the P values (*, P = 0.0155; ***, P = 0.0007).
old Cn as well as 14GEN old Cg cells (22, 71) indicate that old cells are equally fit in the host environment.

Another important discovery is the development of oversized vacuoles with multiple vesicle-like structures in the cytoplasm of old yeast cells. Enlarged vacuoles have been described in old as well as nutrient-deprived S. cerevisiae cells (37, 38, 72). Vacuoles play a role in membrane trafficking (73) and protein sorting (37). Secretion mechanisms can also be influenced by autophagic processes (74). The autophagosome can merge first with MVBs and then with the cell membrane to deliver heterogeneous cargo outside the cell (75). In Cn, MVBs participate in laccase trafficking (76), a cell wall-associated virulence factor (77); in addition, GXM and melanin are exported through EVs (31, 78). The cloning of a “mother cell enriching mutant” in Cn and Cg, similar to the one developed in S. cerevisiae (85), would permit prolonged growth of old cells, with isolation and analysis of their EV content.

The enlarged vacuole in old Cn and Cg cells is most likely the result of the fusion of smaller vacuoles. Vacuolar fusion in S. cerevisiae is necessary and sufficient for life span extension (86). Vacular acidification is also a critical determinant of replicative life span and declines continuously with age, thereby limiting the replicative life span of a cell (87, 88). In C. albicans, vacuolar pH is a regulator of fission and fusion of vacuoles (89). We demonstrated loss of vacuolar acidity in old compared to young Cn similar to S. cerevisiae (38) but different from old Cg, where the vacuole pH was more acidic. Different factors, including glucose starvation, can also affect vacuolar pH (90). It is possible that Cg is experiencing more carbon starvation and has not advanced in its life span as much as Cn. Differences in pH regulation during aging and carbon starvation have been described in different Cn strains (91). Cn has a unique protein (All2p) involved in pH homeostasis (92), which accumulates during aging in low-glucose conditions (40). More extensive analysis will be necessary to better understand the mechanisms involved in age-associated regulation of intracellular pH in these yeast cells, including the use of other pH probes that consistently colocalize within the vacuoles.
It is noteworthy that these studies were conducted on a single strain for each pathogen, and therefore, further work including several strains of different genotypes is required.

In summary, these data provide insights regarding the dynamics of cell wall rearrangement during aging of Cn and Cg. The data highlight the divergence between these pathogens in bud scar formation throughout the life span and uncover differences such as cell wall composition, especially concerning mannan and chitosan. This work opens new avenues of investigation, such as the impact of cell wall reorganization in old yeast cells on the host innate response and sensitivity to cell wall-targeting antifungals. Further work is also needed to elucidate the implications of EV secretion during aging, including cargo and “cross talk” between young and old pathogenic yeast cells and explore the association among aging, intracellular pH, and mitochondrial function.

MATERIALS AND METHODS

Strains and media. The fungal strains Cn RC2 (ATCC 24067 variant) and Cg BG2 were kept on yeast extract-peptone-dextrose (YPD) agar (Difco BD) plates. Synthetic medium (SM) (43) was used for cultivation of yeast cells.

Isolation of young and old Cn and Cg cells. Yeast cells (10⁶) from an overnight culture were washed with phosphate-buffered saline (PBS) and labeled with 8 mg/mL sulfo-succinimidyl-6-(biotin-amido) hexanoate (Sulfo-NHS-LC-LC-Biotin, 21338; Thermo Fisher Scientific) with phosphate-buffered saline (PBS) and labeled with 8 mg/mL sulfosuccinimidyl-6-(biotin-amido) hexanoate (Sulfo-NHS-LC-LC-Biotin, 21338; Thermo Fisher Scientific) for 30 min at room temperature (RT). Subsequently, cells were washed with PBS and grown for 5 to 7 doubling times. Then, 100 µL of magnetic streptavidin microbeads (130-048-101; Miltenyi Biotec) were added to every 10⁶ cells/mL, following incubation for 15 min at 4°C. Biotin-labeled yeasts were isolated using autoMACS magnetic columns (130-021-101; Miltenyi Biotec) and the autoMACS Pro separator (Miltenyi Biotec). The biotin-streptavidin-labeled older cells were passed through a magnet where they get stuck, while the younger unlabeled population flowed through. The labeled cells were then recovered after the magnetic field was removed. The positive cell fraction was grown again in SM to the desired generation (10GEN Cn, 14GEN Cg) and isolated as outlined above. Young cells washed off from the magnetic columns were kept as controls. The purity of fractions was verified by microscopy. In this work, “old” Cn are 10 generations old, and “old” Cg are 14 generations old.

RNA isolation and quantitative reverse transcriptase PCR (RT-qPCR). RNA isolation and RT-qPCR were done as described (22). Briefly, the RNAeasy kit (Qiagen) was used with on-column DNase digestion following the manufacturer’s instructions. The concentration of total RNA was measured using a NanoDrop instrument (Eppendorf). The quantity of RNA was adjusted to 250 ng and converted to cDNA using the Verso cDNA synthesis kit (Thermo Scientific). cDNA was diluted and used in a mix that contained Power SYBR green mastermix (Applied Biosystems), together with oligo(dT) primers (Table S1). The experiment was performed using the high-performance real-time PCR (Roche) and LightCycler 480 systems (Roche). Gene expression was computed using ∆∆CT. ACT1 was used to normalize the gene expression, and the gene expression in young cells was used as the reference. Values above 2-fold were significant. Threshold cycle (Ct) values of ACT1 in old and young cells were similar to one another, signifying that ACT1 gene expression does not vary between the young and old cells. Each experiment was repeated at least twice on two separate days.

Cell wall staining. Young and old cells (10⁶) were fixed in 4% paraformaldehyde (PFA) for 15 min at RT, washed with PBS, and stained with calcofluor white (CFW; cell wall staining of chitin; 5 µg/mL for 10 min at 37°C), concanavalin A, Texas Red conjugate (ConA-Texas Red; cell wall staining of mannoproteins; 50 µg/mL for 45 min at 37°C), eosin Y (EY, cell wall staining of chitosan; 300 µg/mL for 10 min at 37°C (94), or tetramethylrhodamine-labeled wheat germ agglutinin (TRITC-WGA; cell wall staining of chitooligosomers; 5 µg/mL for 30 min at 37°C) (14).

To stain β-glucan and GXM, fungal cells were blocked with 1% bovine serum albumin (BSA) for 1 h at 37°C and incubated, respectively, with the primary antibody Fc-dectin 1 (AdipoGen; 4G-408-0138) (5 µg/mL for 40 min at 4°C) (28) or MaB 18B7 (10 µg/mL; 1 h at 37°C) (32). After washing with PBS, cells were incubated with secondary antibody DyLight 405 goat anti-human IgG + IgM (5 µg/mL; 30 min at 4°C) or with an anti-murine IgG DyLight 488 (10 µg/mL; 30 min at 37°C) (Jackson Immunoresearch).

Prior to analysis using a BD LSRFortessa flow cytometer, cells were washed and resuspended in PBS. The following lasers were used: blue, 488 nm (for EY); YelGreen, 561 nm (for ConA-Texas Red and WGA-TRITC); violet, 405 nm (for CFW and DyLight 405). Data were analyzed using BD FACSDiva or FlowJo v10.1 software. A total of 10,000 events were gated in the forward scatter/side scatter (FSC/SSC) plots and represented as histograms with mean fluorescence intensity (MFI) on the x axis and cell counts on the y axis. Unstained cells were used as negative controls. For fluorescence microscopy, the following channels were used: fluorescein isothiocyanate (FITC) (EY and DyLight 488), DAPI (4′,6-diamidino-2-phenylindole) (CFW and DyLight 405), and DiSer (Texas Red and TRITC). Imaging was performed at 100× magnification in a Zeiss Axio Observer or a Zeiss Axiovert 200M (Thornwood, NY). The same exposure time was used to image young and old cells. The images were processed using ImageJ software.

β-1,3-glucan quantification. Cn cells (4 × 10⁶) were washed with Tris-EDTA buffer (pH 8.0) and suspended in 500 µL of Tris-EDTA and 56 µL of NaOH (10 M) to achieve a final concentration of 1 M. To
solubilize (1,3) β-D-glucan, cells were incubated at 80°C for 30 min. Then, 2.1 mL of aniline blue solution (0.03% aniline blue, 0.18 M HCl, 0.49 M glycine/NaOH, pH 9.5) was added to the cells to interact with linear (1,3) β-D-glucan. Following homogenization, cells were incubated for 30 min at 50°C and for 30 min at RT. The fluorescence was measured using a plate reader SpectraMax i3X (Molecular Probes), with excitation at 400 nm and emission at 460 nm (95).

**Ultramicroscopy.** Cells were fixed in 3% glutaraldehyde in 0.1 M sodium cacodylate buffer, pH 7.4, at 4°C overnight. Then, cells were washed 3 times with 0.1 M sodium cacodylate buffer and dispersed in ultra-low-temperature agarose at 4°C for 30 min. Subsequently, the agarose samples were cut into small cubes, and the pieces were postfixed in aqueous 1% potassium permanganate (KMnO₄) for 1 h at RT. Blocks were rinsed with distilled water (dH₂O) and treated with 0.5% sodium meta-periodate for 15 min at RT to allow infiltration. Next, blocks were rinsed with dH₂O and dehydrated through a graded ethanol series. Samples were treated with 100% propylene oxide, embedded with Spur resin, and polymerized in an oven at 60°C for 24 h. Ultrathin sections (80 nm) were cut using a Leica EM UC7 ultramicrotome, placed on 300-mesh copper grids, and counterstained with uranyl acetate and lead citrate. Images were acquired at magnifications of 18,500× and 49,000× using an FEI TeCnai 12 BioTwinG² TEM. Digital images were acquired with an AMT XR 60 CCD digital camera system. At least 30 young and old cells were analyzed, and the thickness of cell wall layers was measured with ImageJ software.

**Intracellular glucuronoxylomannan (GXM) of Cn.** Cn cells were fixed for 45 min at RT in fixate solution (synthetic medium + 4% PFA + 0.1% glutaraldehyde + 0.1 M cacodylate buffer). Samples were washed twice with 0.1 M cacodylate buffer, suspended in 0.1 M cacodylate buffer, and left overnight at 4°C. The cells were washed with PBS, placed in gelatin, cut into cubes of 1 mm, and postfixed in 0.1 M sodium cacodylate buffer, 2.0% paraformaldehyde. Samples were then dehydrated through a graded series of ethanol with a progressive lowering of the temperature to −50°C in a Leica EM AFS, embedded in Lowicryl HM-20 mono-step resin (Electron Microscopy Sciences), and polymerized using UV light. Ultrathin sections were cut on a Leica EM UC7, and immunolabeled with MAb 1887 (50 μg/mL) and gold-labeled anti-mouse IgG. Control systems included ultrathin sections incubated without the MAb 1887, followed by the gold-labeled anti-mouse IgG. Samples were stained with uranyl acetate and viewed on a JEOL 1400 Plus transmission electron microscope at 80 kv. Gold particles in electron micrographs were quantified as described (96). For quantitative analysis of GXM labeling, 50 cells of the young group and 47 cells from the old group were analyzed. For normalization, the average number of gold particles counted in control systems (no MAb 1887 labeling) was subtracted from the total number of gold particles of each MAb 1887-labeled section, as described reference 97.

**Quantification and pH analysis.** Vacuoles from 10⁵ fungal cells were stained at 37°C with FM 4-64 (Thermo Fisher) (10 μg/mL) for 10 min following manufacturer’s protocol. Another set of 10⁵ fungal cells were labeled with quinacrine in separate tubes at a final concentration of 200 μM and incubated for 20 min at 37°C. Cells were washed three times with PBS and imaged at 100× using a deconvolution microscope (Zeiss Axiovert 200M), using the filters Texas Red (for FM4-64) and green fluorescent protein (GFP; for quinacrine). Then, 25 cells from each group and strain were analyzed using ImageJ software. Also, 200 μL of yeast suspensions (10⁵) labeled with quinacrine were plated in triplicates in a black 96-well plate, and the fluorescence was measured with a Spectramax i3X spectrophotometer (Molecular Probes), using excitation at 436 nm and emission at 525 nm. The values of unstained cells were subtracted from the values obtained from the stained cells. All experiments were repeated three times independently.

**Cell wall stressor assay.** Serial dilutions of yeast suspensions (final concentrations 10⁴ to 10² cells) were spotted on YPD agar containing each of the following stressors: fluorescent brightener 28 (FB 28), 1.5 mg/mL; Congo red, 0.5%; caffeine, 1 mg/mL or 0.5 mg/mL; sodium dodecyl sulfate (SDS), 0.01% or 0.0025%. Next, the plates were incubated at 30°C and 37°C for 48 h. Control conditions included cells added on YPD agar without the stressors. For Congo red, the stressor was added to the YPD medium prior to autoclaving, whereas for caffeine, FB 28, and SDS, stock solutions were prepared, filter-sterilized, and added to YPD after autoclaving (28).

**Statistical analysis.** Statistical differences between young and old cells were performed using a t test with Welch’s correction. All statistical tests were performed with Prism 9.0 software (GraphPad Software, Inc.).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1,** TIF file, 1.4 MB.

**FIG S2,** TIF file, 1.6 MB.

**FIG S3,** TIF file, 0.5 MB.

**FIG S4,** TIF file, 0.6 MB.

**TABLE S1,** PDF file, 0.1 MB.

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