

O-Mannosylation in *Candida albicans* Enables Development of Interkingdom Biofilm Communities

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ABSTRACT *Candida albicans* is a fungus that colonizes oral cavity surfaces, the gut, and the genital tract. *Streptococcus gordonii* is a ubiquitous oral bacterium that has been shown to form biofilm communities with *C. albicans*. Formation of dual-species *S. gordonii*-*C. albicans* biofilm communities involves interaction of the *S. gordonii* SspB protein with the Als3 protein on the hyphal filament surface of *C. albicans*. Mannoproteins comprise a major component of the *C. albicans* cell wall, and in this study we sought to determine if mannosylation in cell wall biogenesis of *C. albicans* was necessary for hyphal adhesin functions associated with interkingdom biofilm development. A *C. albicans* *mnt1* Δ *mnt2* Δ mutant, with deleted α -1,2-mannosyltransferase genes and thus defective in O-mannosylation, was abrogated in biofilm formation under various growth conditions and produced hyphal filaments that were not recognized by *S. gordonii*. Cell wall proteomes of hypha-forming *mnt1* Δ *mnt2* Δ mutant cells showed growth medium-dependent alterations, compared to findings for the wild type, in a range of protein components, including Als1, Als3, Rbt1, Scw1, and Sap9. Hyphal filaments formed by *mnt1* Δ *mnt2* Δ mutant cells, unlike wild-type hyphae, did not interact with *C. albicans* Als3 or Hwp1 partner cell wall proteins or with *S. gordonii* SspB partner adhesin, suggesting defective functionality of adhesins on the *mnt1* Δ *mnt2* Δ mutant. These observations imply that early stage O-mannosylation is critical for activation of hyphal adhesin functions required for biofilm formation, recognition by bacteria such as *S. gordonii*, and microbial community development.

IMPORTANCE In the human mouth, microorganisms form communities known as biofilms that adhere to the surfaces present. *Candida albicans* is a fungus that is often found within these biofilms. We have focused on the mechanisms by which *C. albicans* becomes incorporated into communities containing bacteria, such as *Streptococcus*. We find that impairment of early stage addition of mannose sugars to *C. albicans* hyphal filament proteins deleteriously affects their subsequent performance in mediating formation of polymicrobial biofilms. Our analyses provide new understanding of the way that microbial communities develop, and of potential means to control *C. albicans* infections.

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Candida albicans is an opportunistic fungal pathogen found in the microbiota of the gut, oral cavity, and genital tract (1). In healthy individuals, *C. albicans* growth is kept in check by a combination of the resident microbiota and both the innate and acquired immune systems. Overgrowth of *C. albicans*, associated with administration of broad-spectrum antibiotics or immune dysfunction, may lead to superficial infections, such as oropharyngeal candidiasis (thrush) and vulvovaginal candidiasis (vaginal candidiasis). Systemic infections have become increasingly prevalent because more individuals are immunocompromised. *C. albicans* biofilm infections are common in patients with urinary or intravascular catheters or artificial joints or voice boxes (2). Indeed, *C. albicans* is associated with >90% of human oral fungal diseases and is becoming an increasingly serious problem in hospital infections (3).

C. albicans forms biofilms on a range of surfaces *in vivo* (4–6) and is well adapted to conditions in the oral cavity. Biofilm for-

mation occurs in three or four phases, starting first with the deposition of yeast-form cells onto the substratum and then the formation of hyphae (7). The latter are critical for biofilm formation because mutants that do not form hyphae under biofilm-promoting conditions are unable to form robust biofilms (8). Biofilm formation is under genetic regulation by the transcription factors Bcr1 (9, 10), Efg1 (11), and Ace2 (12), among others. In particular, Bcr1 is known to control expression of cell surface proteins, such as Als3 and Hwp1 (13), that are involved in adherence of hyphal filaments to each other (14) in the building of biofilms. A number of techniques may be utilized to grow *C. albicans* biofilms *in vitro* on surfaces of acrylic, rubber disks, catheter strips, or glass coverslips (15). Static biofilm models provide good *in vitro* data but do not take into account conditions *in vivo*, where *C. albicans* biofilms growing on medical implants, such as catheters, prosthetic joints and heart valves, would be exposed to flow forces and replenishment of nutrients from body fluids. In addi-

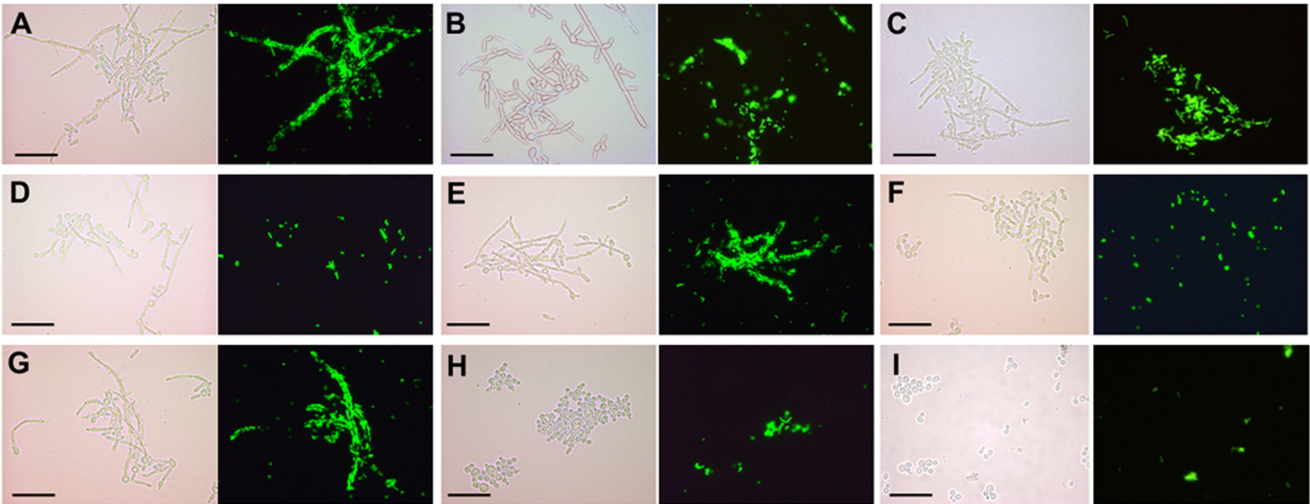


FIG 1 Interactions of *S. gordonii* DL1 cells with *C. albicans* wild-type or glycosylation-deficient mutants induced to form hyphae in YPT-glucose medium in suspension culture (planktonic conditions). Bacteria were fluorescently labeled with fluorescein isothiocyanate (FITC) and incubated with hypha-forming cells of *C. albicans* in YPT-Glc medium for 1 h at 37°C with gentle agitation. Samples (10 μ l) were then removed and placed on glass microscope slides, coverslips were applied, and cells were visualized with a Leica microscope under phase contrast or fluorescence at magnification $\times 400$. The panels each show a light microscopic image and the corresponding fluorescence image. (A) *C. albicans* wild type; (B) *mnt1* Δ mutant; (C) *mnt2* Δ strain; (D) *mnt1* Δ *mnt2* Δ strain; (E) *mnt1* Δ *mnt2* Δ + *MNT1* strain; (F) *mnt1* Δ *mnt2* Δ + *MNT2* strain; (G) *mnn4* Δ strain; (H) *och1* Δ strain; (I) *Saccharomyces cerevisiae* BY4742 (negative control). Scale bar = 50 μ m.

tion, biofilms formed under flow have been shown to develop architectures different from those of static biofilms grown under the same conditions (16).

The cell surface of *C. albicans* is the major point of contact between fungus and host, along with its constituent microbiota, and provides adhesive and immunomodulatory functions (17). The outer cell wall layer is comprised principally of mannoproteins which are embedded in a polysaccharide skeleton provided by β -(1,3)- and β -(1,6)-linked glucan chains and covalently linked chitin (18). The protein and carbohydrate components of the outer wall layers have both been implicated in adhesion to host surfaces (19) and in biofilm formation (20). It is predicted that ~ 115 proteins encoded by the *C. albicans* genome are glycosylphosphatidylinositol (GPI)-modified proteins (21). Among these are adhesins, such as Hwp1, Rbt1, Eap1, and the Als family of proteins (22), some of which have established roles in biofilm formation (13, 23, 24) and in interactions with host tissues (25). Glycosylation of these and other cell wall proteins is thought to be important for their functions (26). The N-linked glycans carry a conserved core structure and highly branched outer mannose chains (27). The O-linked glycans are, conversely, linear oligosaccharides of one to five α -1,2-linked mannose residues attached to serine or threonine residues (28). The N- and O-glycosylations are completed in the Golgi apparatus, mediated by a set of multifunctional mannosyltransferases (29). Mnt1 and Mnt2 are partially redundant α -1,2-mannosyltransferases that catalyze the addition of the second and third mannose residues in an O-linked mannose pentamer (30). The Mnt1 enzyme may also add the fourth and fifth mannose residues (31). *C. albicans* mutants deficient in the Mnt1 and Mnt2 proteins are modified in cell wall structure (30, 32), diminished in adherence to human buccal epithelial cells (28), and attenuated in virulence (33). In addition, *mnt1* Δ , *mnt2* Δ , and *mnt1* Δ *mnt2* Δ mutants form hyphae that are hypersensitive to killing by *Pseudomonas aeruginosa* (34).

In polymicrobial infections (35), *C. albicans* has been found in close association with *P. aeruginosa* (36), *Staphylococcus aureus* (37), and oral streptococci in denture- and mucosa-related diseases (38). The interactions of oral streptococci, such as *Streptococcus gordonii*, with *C. albicans* are proposed to facilitate oral carriage and persistence of *C. albicans* in mixed-species biofilms on natural or prosthetic surfaces (39, 40). The cell surface protein SspB expressed on *S. gordonii* is responsible at least in part for mediating adherence to *C. albicans* hyphal filaments (41, 42). *S. gordonii* appears to promote hyphal filament formation by *C. albicans* (42, 43), in contrast to *P. aeruginosa* and *Salmonella enterica* serovar Typhimurium, which kill hyphae (44, 45). The aims of this work were to determine if *C. albicans* cell wall O-mannosylation was necessary for interactions of hyphae with *S. gordonii* and for formation of dual-species biofilms.

RESULTS

Mannosyltransferase Mnt1 is necessary for *C. albicans* to bind streptococci. Previous work has demonstrated that *S. gordonii* cells adhered to hyphae formed by *C. albicans* SC5314 (43). In the present study, *C. albicans* CAI4+Clp10 has been designated wild type, this being a *Ura3*⁺ reintegrant generated from *C. albicans* CAI4 to control for any effects of *ura3* gene deletion and transposition in mutagenesis (46). The strain behaved identically to SC5314, from which it was initially derived (see Table S1 in the supplemental material). Hyphal filament formation by the wild-type strain was induced in planktonic phase under our standard conditions with glucose (YPT [1 \times Difco yeast nitrogen base, 20 mM NaH₂PO₄-H₃PO₄ buffer, pH 7.0, and 0.1% Bactotryptone]-Glc). Hyphae were shown to avidly bind *S. gordonii* DL1 cells along their lengths (Fig. 1A). A homozygous *mnt1* Δ mutant, deficient in production of α -1,2-mannosyltransferase (30), which adds the second mannose residue to growing O-linked glycan chains (30), showed reduced binding of *S. gordonii*

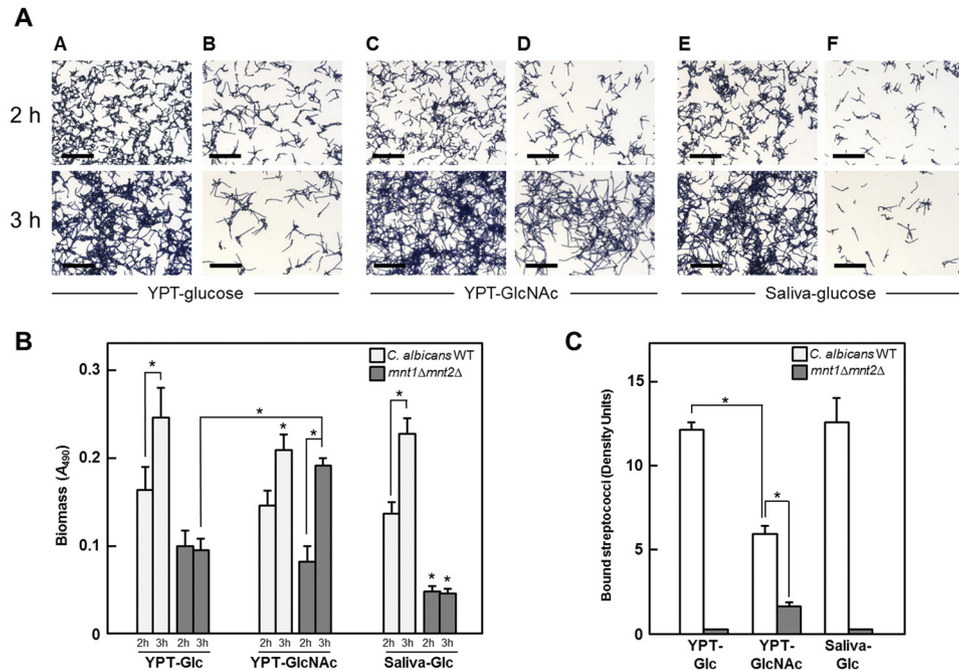


FIG 2 Effects of different hyphal filament induction media on biofilm formation and intermicrobial adherence by *C. albicans* wild type and *mnt1Δ mnt2Δ* mutant strains. (A) Biofilms were grown on saliva-coated coverslips (as described in Materials and Methods) for 2 h or 3 h in YPT-Glc, YPT-GlcNAc, or saliva-Glc medium, as indicated, and the cells were then stained with crystal violet and visualized by light microscopy. Biofilms of the wild-type strain showed increased density and extent of hyphal filament formation over 2 to 3 h (columns A, C, and E). Between 2 h and 3 h, the biofilms formed by the *mnt1Δ mnt2Δ* mutant in YPT-Glc (column B) and saliva-Glc (column F) ceased to develop further, while biofilms in YPT-GlcNAc continued to grow (column D). Bar = 50 μ m. (B) Corresponding biofilm biomass values for panel A images (*, $P < 0.05$; $n = 3$). (C) Extent of FITC-labeled *S. gordonii* DL1 cells binding to hyphal filaments formed by wild-type or *mnt1Δ mnt2Δ* cells after a 3-h incubation in suspension cultures in the three different induction media. Density units of bound streptococci were calculated from image capture software measurements, as described in Materials and Methods (*, $P < 0.05$; $n = 3$).

(Fig. 1B). Hyphae formed by a homozygous *mnt2Δ* mutant were not noticeably impaired in streptococcal cell attachment (Fig. 1C). In the homozygous double mutant strain UB1933 *mnt1Δ mnt2Δ*, hyphal filaments were abrogated in binding *S. gordonii* (Fig. 1D). Reintroduction of *MNT1* into the double mutant resulted in restoration of *S. gordonii* attachment (Fig. 1E), while reintroduction of *MNT2* did not (Fig. 1F). These results suggested that expression of *MNT1* was essential in order to provide hyphal cell wall receptors for adherence of *S. gordonii* to *C. albicans*.

We also tested *mnn4Δ* and *och1Δ* *N*-linked glycosylation-impaired mutants (47, 48) for the ability to recognize *S. gordonii*. Deficiency in *Mnn4*, which normally regulates mannosyl phosphorylation of *N*- and *O*-linked glycans, had no effect on *S. gordonii* binding to hyphae (Fig. 1G). The mutant strain UB1939 *och1Δ* (48), deficient in production of α -1,6-mannosyltransferase, which initiates *N*-glycan outer chain branch addition, did not support adherence of *S. gordonii* (Fig. 1H). However, this mutant had a highly pleiotropic phenotype, including being impaired in growth rate and hypha production. Under the conditions of these experiments, *Saccharomyces cerevisiae* cells did not bind *S. gordonii* (Fig. 1I).

Effect of growth medium on development of biofilms. To investigate in more detail the role of *O*-mannosylation in biofilm-forming and *S. gordonii*-binding properties of *C. albicans*, we compared phenotypes of the wild type and the *mnt1Δ mnt2Δ* double mutant under different growth conditions. In biofilm studies with YPT-Glc medium, a high proportion of deposited *C. albicans* wild-type cells had formed hyphal filaments at 2 h

(Fig. 2A). After a 3-h incubation, an intensified network of hyphae was apparent, with a concomitant increase in biofilm biomass (Fig. 2B). Although at 2 h, fewer *mnt1Δ mnt2Δ* mutant cells were associated with the substratum (Fig. 2A, column B), a proportion of cells similar to that for the wild type had formed hyphae. However, after 2 h, the mutant cells appeared to become arrested in biofilm formation (Fig. 2A, column B), and no further significant increase in biomass had occurred at 3 h (Fig. 2B). When *N*-acetyl-D-glucosamine (GlcNAc), a known inducer of hypha formation (49), was substituted for glucose, the previous inability of the *mnt1Δ mnt2Δ* mutant to further develop a biofilm and increased biomass over the 2- to 3-h period was reversed (Fig. 2A, columns C and D, and 2B). The numbers of cells forming hyphae at 3 h in YPT-GlcNAc were similar ($65\% \pm 6.0\%$) for the wild type and double mutant (Fig. 2A). Fewer *S. gordonii* cells bound to wild-type hyphae in YPT-GlcNAc than in YPT-Glc (Fig. 2C). This we attributed to a deficiency in metabolism of GlcNAc by *S. gordonii*. We also utilized 10% saliva containing 0.01% glucose (saliva-Glc) as a growth medium, and in this medium, the *C. albicans* wild type showed patterns of biofilm formation and biomass production similar to those with YPT-Glc (Fig. 2A, column E, and 2B). With saliva-Glc medium, the *mnt1Δ mnt2Δ* mutant formed only sparse biofilms (Fig. 2A, column F) and showed no increase in biomass after 2 h (Fig. 2B).

Although the presence of GlcNAc rescued the *mnt1Δ mnt2Δ* mutant from deficiency in biofilm production in YPT-Glc medium, the ability of these double mutant hyphae to bind *S. gordonii* was not restored to wild-type levels (Fig. 2C). Confocal scan-

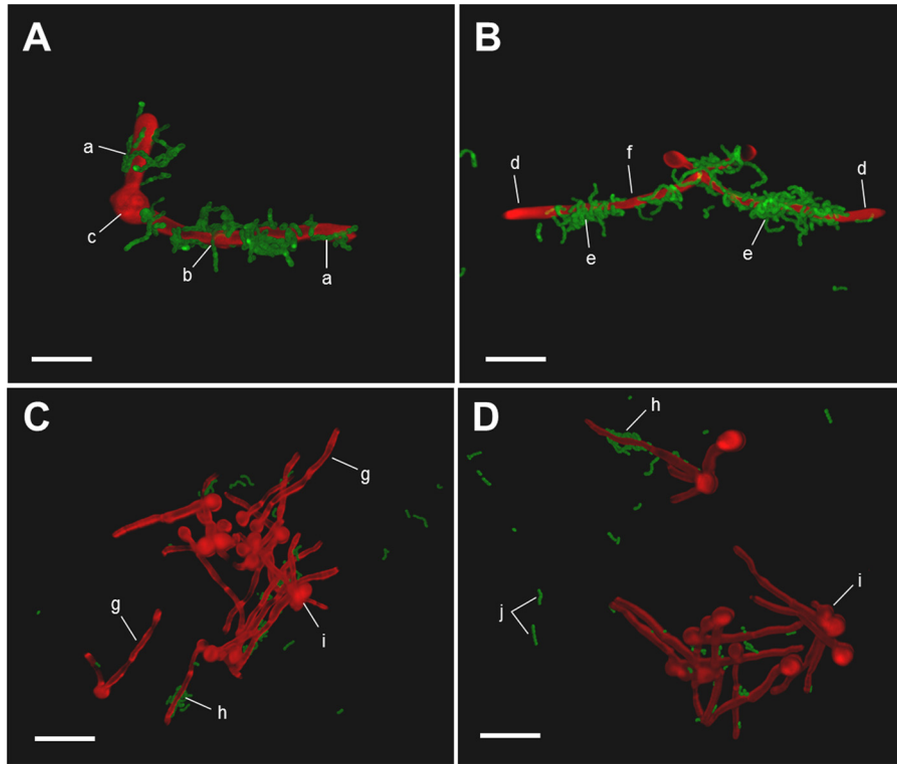


FIG 3 CSLM images of *S. gordonii* cells interacting with hyphal filaments formed by *C. albicans* wild-type or *mnt1Δ mnt2Δ* mutant strains in YPT-GlcNAc suspension culture. *S. gordonii* DL1 cells (green) expressing GFP (Table 1) were incubated for 1 h at 37°C with filament-forming *C. albicans* cells stained with calcofluor white (images recolored red). *C. albicans* wild type (A and B) or *mnt1Δ mnt2Δ* mutant (C and D) is shown. (A) short chains of streptococci may be seen aligning along the hyphal filaments (a) or bound to the filament by one end of a streptococcal cell chain (b) but not associated with yeast morphology mother cells (c). (B) the tips of hyphal filaments and their adjacent regions were not usually seen to bind *S. gordonii* cells (d); some parts of filaments were heavily populated by streptococcal communities (e), while other regions were not bound by streptococci (f). (C and D) Hyphal filaments of the *mnt1Δ mnt2Δ* mutant were sometimes unevenly stained with calcofluor white (g) and were generally not recognized by *S. gordonii* (g), but some individual filaments among the population showed low levels of bacterial cell binding (h). The *mnt1Δ mnt2Δ* mutant mother cells often emanated multiple hyphal filaments from the one cell (i). Nonadhered streptococci (j) were clearly visible in the *mnt1Δ mnt2Δ* strain preparations. Bar = 20 μm .

ning laser microscopy (CSLM) images showed that streptococci in pairs or short chains interacted avidly with hyphae formed by wild-type *C. albicans* (Fig. 3A and B), accumulating as localized clusters of cells but tending not to bind to the hyphal tip regions (Fig. 3B). Conversely, *S. gordonii* cells bound only sparsely to hyphae produced by the *mnt1Δ mnt2Δ* mutant (Fig. 3C), although some hyphal filaments seemed to support localized *S. gordonii* attachment (Fig. 3D).

Effect of *mnt1Δ* and *mnt2Δ* mutations on formation of monospecies or dual-species biofilms. In monospecies biofilms, *C. albicans* wild-type cells formed a confluent mat of blastospores and hyphae after 6 h at 37°C in YPT-Glc or YPT-GlcNAc growth conditions (Fig. 4). When saliva-Glc was utilized to grow *C. albicans* biofilms, aggregates or clusters of cells were visible and consequently overall coverage of the substratum was lower (Fig. 4). Saliva-Glc medium did not support such extensive hyphal filament and biofilm formation. The *mnt1Δ mnt2Δ* mutant cells were unable to form biofilms in YPT-Glc or saliva-Glc medium (Fig. 4). However, the deleterious effects of the *mnt1Δ* and *mnt2Δ* mutations on hypha formation and biofilm formation in YPT-Glc and saliva-Glc media were not evident in YPT-GlcNAc medium (Fig. 4). Although the *mnt1Δ mnt2Δ* mutant biofilms still showed reduced coverage compared to findings for the wild type (Fig. 4), there was extensive hyphal filament formation (more detail can

be seen in Video S1 and Video S2 in the supplemental material).

C. albicans wild-type cells formed dual-species biofilms with *S. gordonii* under the three different growth conditions, with more abundant growth and hypha formation in YPT-GlcNAc (Fig. 4). In YPT-Glc medium, *S. gordonii* cells formed an evenly distributed biofilm over the substratum surface between the deposited *C. albicans* cells, and hyphae appeared integrated within the streptococcal cell community (Fig. 4). More extensive *C. albicans* hypha formation in YPT-GlcNAc medium was accompanied by hyphae appearing to emanate from the *S. gordonii* community. In saliva-Glc medium, both *C. albicans* and *S. gordonii* formed patches of growth (Fig. 4) associated with saliva-mediated aggregation of the microorganisms. In dual-species biofilm experiments, the *mnt1Δ mnt2Δ* mutant was unable to form biofilms in YPT-Glc or saliva-Glc in the presence of *S. gordonii* (Fig. 4). In YPT-GlcNAc medium, a dual-species biofilm with *S. gordonii* was formed (Fig. 4), similar in architecture to the monospecies biofilm. Taken collectively with the results shown in Fig. 2, it is evident that the *mnt1Δ mnt2Δ* deletions result in the following: (i) reduced levels of adhesion to a salivary pellicle substratum, (ii) a depressed rate of hyphal filament extension, and (iii) a curtailed ability (in YPT-Glc or saliva-Glc medium) to form a robust biofilm.

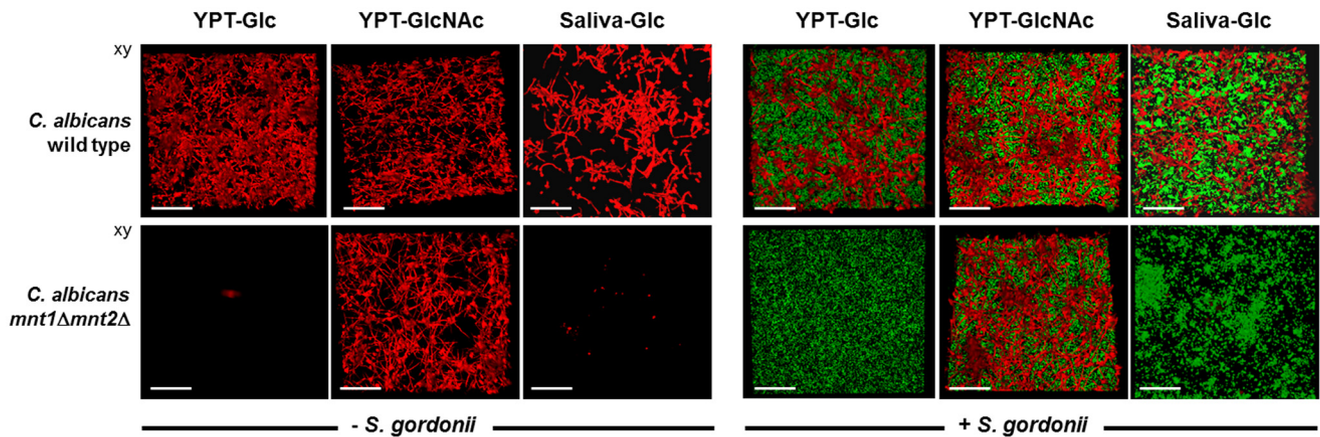


FIG 4 CSLM images of biofilms formed by *C. albicans* wild type or *mnt1Δ mnt2Δ* mutant strains in the absence (monospecies) or presence (dual species) of *S. gordonii* DL1. Monospecies or dual-species biofilms were prepared as described in Materials and Methods and were grown for 6 h at 37°C in the three different growth media indicated. The *mnt1Δ mnt2Δ* mutant was unable to form monospecies or dual-species biofilms in YPT-Glc or saliva-Glc but produced robust monospecies and dual-species biofilms in YPT-GlcNAc. Bar = 50 μ m.

Architecture of monospecies or dual-species biofilms. We then investigated in finer detail the architecture of *C. albicans* and *S. gordonii* in mono- or dual-species biofilms. In YPT-Glc biofilms, *C. albicans* was present as a mixture of blastospores and filaments integrated within and on top of a dense layer of streptococci (see Fig. S1A in the supplemental material). A vertical section (see Fig. S1B) showed *C. albicans* attached to regions of the pellicle substratum and to the streptococcal layer. In the presence of YPT-GlcNAc, *C. albicans* formed a more robust and filamentous biofilm (see Fig. S1, center panels) either alone (see panel E) or with *S. gordonii* (see panel D). The *S. gordonii* component of the dual-species biofilm was much less compact (panel C), and *S. gordonii* cells were more clearly interdigitated with the extending hyphal filaments. In saliva-Glc medium, a highly integrated dual-species biofilm was produced (see Fig. S1), with streptococci and hyphal filaments extending up to 115 μ m from the substratum. Both of the microbial components grew more luxuriantly under these conditions, which was especially apparent when comparing *C. albicans* dual- and monospecies biofilms (see Fig. S1D and E).

Biofilm formation under flow. Under flow conditions, which might more closely mimic those in the human oral cavity, there were some remarkable changes in morphology and composition of single- or dual-species biofilms. *C. albicans* monospecies biofilms that formed in YPT-Glc under flow contained more hyphae than when grown under nonflow conditions, and hyphae extended further from the surface (107 μ m) (Fig. 5B) than under nonflow (73–80 μ m) (see Fig. S1D and E in the supplemental material). *C. albicans* wild-type biofilms formed in YPT-GlcNAc produced hyphal filaments that protruded far out into the environment (to 239 μ m), giving a brush-like appearance (Fig. 5B; see also Video S3). In saliva-Glc medium, extensive hypha formation was also observed, but the hyphae tended to lie flatter across the surface of the biofilm (Fig. 5B). Conversely, the *mnt1Δ mnt2Δ* mutant was unable to form robust biofilms under flow conditions, even in YPT-GlcNAc (Fig. 5C and D; see also Video S4).

In dual-species biofilms, patterns of hypha formation by *C. albicans* wild-type cells were observed that were similar to those of

monospecies biofilms (see Video S5 in the supplemental material). Hyphae formed by the *C. albicans* wild type in saliva-Glc were less protruding and lay flatter over the *S. gordonii* (Fig. 5B). The presence of *S. gordonii* did not promote incorporation of *mnt1Δ mnt2Δ* mutant cells into the flow biofilm except in YPT-GlcNAc medium (Fig. 5). In this medium, the presence of *S. gordonii* stimulated hypha formation by the *mnt1Δ mnt2Δ* mutant (Fig. 5C; see also Video S6), suggesting that metabolic signaling between these two strains was occurring.

These interactions under flow are shown in greater detail in Fig. S2 in the supplemental material, in which the dual fluorescence channels are separated. Hyphae produced by the *C. albicans* wild type in YPT-Glc medium grew within, and protruded through the *S. gordonii* biofilm (see Fig. S2B in the supplemental material). In YPT-GlcNAc, hyphal filaments extended into the environment by at least 250 μ m and streptococcal cells were seen to be associated with the hyphae through the depth of the biofilm (see Fig. S2B and C and Video S5). In saliva-Glc medium, hyphal filaments tended to grow within and across the surface of the *S. gordonii* biofilm (see Fig. S2B, C, and D), the biofilms being more compact than the corresponding monospecies *C. albicans* flow biofilm (see Fig. S2E). The *mnt1Δ mnt2Δ* mutant formed a dual-species flow biofilm with *S. gordonii* in YPT-GlcNAc medium (see Fig. S2). However, the *S. gordonii* component of the biofilm appeared to be reduced compared to that for the wild type (see Video S6 for greater detail). Taken collectively, these results show that Mnt1/Mnt2 deficiency can severely restrict *C. albicans* biofilm development but that this restriction can be at least partly negated in the presence of GlcNAc. Importantly, *C. albicans* wild-type monospecies biofilms or dual-species biofilms with *S. gordonii* appeared to be more substantive and vigorous when formed under flow conditions. This suggests that suppressing factors under nonflow conditions or shear forces under flow may play a role in modulating biofilm development.

Expression of *C. albicans* Als3. We have shown that a major adhesin for *S. gordonii* on the hyphal cell surface is Als3 (42), a filament-specific glycoposphatidylinositol (GPI)-modified cell wall protein with multiple adhesin functions (50, 51). Therefore, a possible explanation for the inability of *mnt1Δ mnt2Δ* mutant

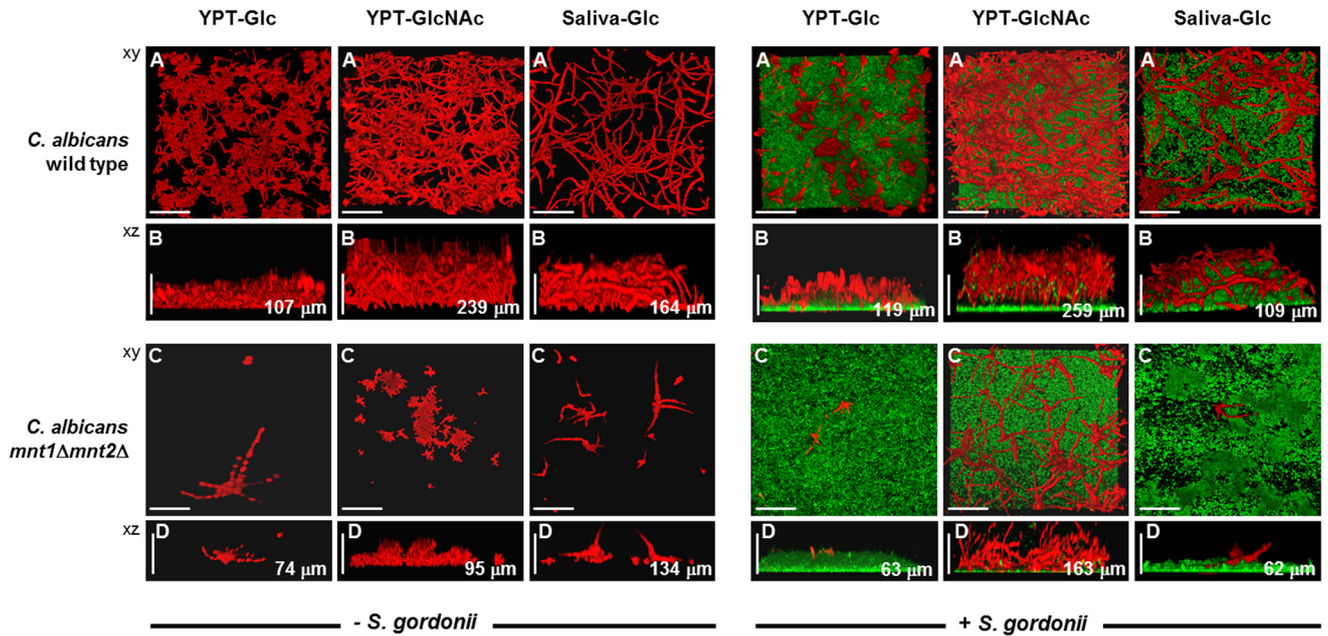


FIG 5 CSLM images of mono- or dual-species biofilms formed under flow conditions by the *C. albicans* wild type or *mnt1Δ mnt2Δ* mutant in three different growth media as indicated. Biofilms were prepared as described in Materials and Methods and grown for 6 h at 37°C at a medium flow rate of 6 ml/h. In the left-side group of 12 panels, monospecies biofilms of *C. albicans* wild-type and *mnt1Δ mnt2Δ* mutant strains are shown; the right-side group of 12 panels shows dual-species biofilms with *S. gordonii*. The image at the top of each group of 6 is a representative *xy* stack of the biofilm assembled from the top down (A and C). The lower image in each column (B and D) is the corresponding *xz* image showing organization and thickness (depth) of the biofilm. Red, *C. albicans*; green, *S. gordonii*. The significant structural and architectural differences associated with the different growth conditions employed are described in the text. The values shown in μm are average thickness (depth) measurements calculated from across the sections of the biofilms shown. Bar = 100 μm .

hyphae to bind *S. gordonii* was that expression of Als3 was affected. To test this, we reacted hyphal filaments formed by wild-type or *mnt1Δ mnt2Δ* mutant cells after a 3-h incubation in YPT-GlcNAc with a monoclonal antibody specific to an epitope within the N-terminal region of Als3 (52). Immunofluorescence microscopy showed that hyphal filaments of the wild type expressed Als3 evenly along their lengths (Fig. 6A and B). However, hyphal filaments formed by the *mnt1Δ mnt2Δ* mutant cells also reacted with the antibody along their lengths (Fig. 6C and D), with no evidence

for any differences in distribution or intensity of antibody reactivity compared to findings for the wild type.

To investigate the functionality of Als3, hypha-forming cells of the wild type or mutant were incubated with fluorescently labeled cells of *Lactococcus lactis* expressing the *S. gordonii* adhesin SspB. This streptococcal cell wall-anchored protein has been shown to interact with Als3 (42). *L. lactis* SspB⁺ cells adhered to wild-type hyphae (Fig. 6E and F) but not to hyphae produced by the *mnt1Δ mnt2Δ* mutant (Fig. 6G and H). *L. lactis* MG1363 control cells did

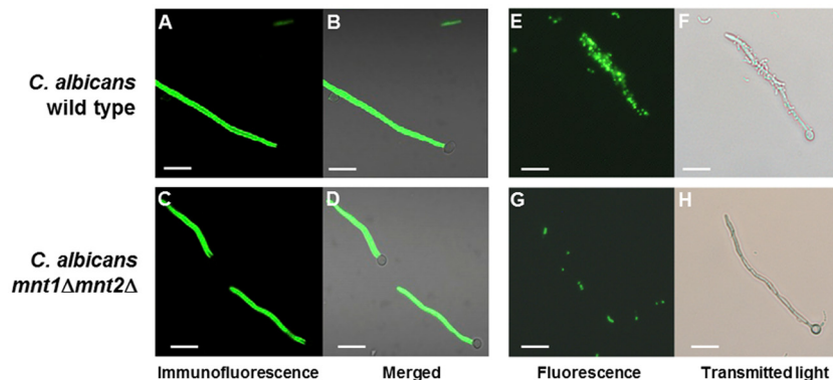


FIG 6 Hypha-forming cells of the *C. albicans* wild type or *mnt1Δ mnt2Δ* mutant immunolabeled with monoclonal antibody to the hyphal cell wall protein Als3 or reacted with *L. lactis* cells expressing SspB. *C. albicans* cells were induced to form hyphae in YPT-GlcNAc medium suspension culture, as described in Materials and Methods, for 3 h at 37°C, harvested by centrifugation, and then incubated with anti-Als3 MAb 3-A5 (52). (A to D) Antibody binding was detected with FITC-conjugated goat anti-mouse F(ab)₂ fragment-specific antibody, and wet mounts were observed microscopically by phase contrast and fluorescence (52). (E and F) Light and corresponding fluorescence microscopic images of FITC-labeled *L. lactis* cells expressing SspB binding to hyphae. (A, B, E, and F) *C. albicans* wild type. (C, D, G, and H) *mnt1Δ mnt2Δ* mutant. Bar = 20 μm .

TABLE 1 Cell wall proteomes for the *C. albicans* wild type or *mnt1Δ mnt2Δ* mutant undergoing hyphal filament formation in three different media

Protein	Description or function	Glucose (Glc) medium emPAI value ^a		GlcNAc medium emPAI value		Saliva-Glc medium emPAI value	
		Wild type	<i>mnt1Δ mnt2Δ</i>	Wild type	<i>mnt1Δ mnt2Δ</i>	Wild type	<i>mnt1Δ mnt2Δ</i>
Als1	GPI-modified yeast cell wall and hyphal filament base adhesin	0.18 ± 0.03	— ^c	0.22 ± 0.08	0.05 ± 0.09	0.09 ± 0.05	0.10 ± 0.05
Als3	GPI-modified hyphal cell wall adhesin and invasin	0.34 ± 0.09	0.03 ± 0.06	0.44 ± 0.08	0.41 ± 0.02	0.13 ± 0.08	0.02 ± 0.02
Als10	GPI-modified, related to Als3	0.03 ± 0.01	—	—	—	—	—
Pga4/Gas1	GPI-modified cell surface β-1,3-glucanosyltransferase	0.48 ± 0.17	0.12 ± 0.22	0.29 ± 0.09	0.19 ± 0.09	0.47 ± 0.09	0.29 ± 0
Phr1	Cell wall glycosidase, glucan remodeling, pH regulated	0.31 ± 0.21	—	0.17 ± 0.06	0.22 ± 0.19	1.07 ± 0.17	0.51 ± 0.17
Phr2	Cell wall glycosidase, pH regulated	0.09 ± 0.03	0.06 ± 0	0.06 ± 0 ^b	0.08 ± 0.09	0.31 ± 0.11	0.22 ± 0.03
Rbt1	GPI-modified hyphal cell wall adhesin similar to Hwp1	0.15 ± 0.03	0.04 ± 0	0.19 ± 0.05	0.09 ± 0.05	0.01 ± 0.02 ^b	0.04 ± 0 ^b
Rbt5	GPI-modified hyphal cell wall protein	0.14 ± 0	0.14 ± 0	0.19 ± 0.09 ^b	0.14 ± 0 ^b	0.14 ± 0 ^b	0.14 ± 0 ^b
Ssr1	GPI-modified glucan-associated role in cell wall structure	0.38 ± 0.20	0.20 ± 0.09	0.38 ± 0.20	0.26 ± 0.09	0.26 ± 0.09	0.31 ± 0
Sap9	GPI-modified aspartyl proteinase, adhesion and cell surface integrity	0.04 ± 0.06	—	0.04 ± 0.07	—	0.20 ± 0.03^b	0.04 ± 0.03
Rhd3/Pga29	GPI-modified yeast cell wall protein, induced by high iron	0.58 ± 0.36	0.11 ± 0.20 ^b	0.22 ± 0.10	0.05 ± 0.09 ^b	0.17 ± 0.17	0.17 ± 0.17
Utr2	GPI-modified cell wall glycosidase, role in adhesion	0.13 ± 0.06	0.01 ± 0.02^b	0.02 ± 0.03 ^b	0.04 ± 0.03 ^b	0.30 ± 0.09	0.20 ± 0.08
Hyr1	GPI-modified hyphal cell wall protein, induced by macrophages (Bcr1 activated)	0.01 ± 0.02	—	0.05 ± 0.06	0.33 ± 0.09	—	—
Scw1	Cell surface O-glycosylated mannoprotein, glucan metabolism and adhesion	0.72 ± 0.30	0.24 ± 0.12	0.20 ± 0.10	0.15 ± 0.14	0.20 ± 0.11	0.05 ± 0.05
Cht2	GPI-modified chitinase, required for filamentous growth	0.55 ± 0.05	0.48 ± 0.04	0.34 ± 0.04	0.50 ± 0.17	0.32 ± 0.08	0.02 ± 0.03^b
Ecm33.3	GPI-modified hypha-specific protein, adhesion	0.53 ± 0.17	0.74 ± 0.31	0.38 ± 0.06	0.55 ± 0.34	0.77 ± 0.07	1.00 ± 0.17
Crh11	GPI-modified transglycosylase	0.55 ± 0.28	0.33 ± 0.18	0.35 ± 0.10	0.38 ± 0.20	0.72 ± 0.12	0.90 ± 0.32
Sod4	Superoxide dismutase (Cu/Zn)	0.19 ± 0.09	0.05 ± 0.08 ^b	—	—	—	—
Sod5	Superoxide dismutase (Cu/Zn)	0.19 ± 0.09	0.05 ± 0.08 ^b	0.25 ± 0.09	0.32 ± 0.12	0.30 ± 0	0.36 ± 0.10
Pir1	1,3-β-Glucan-linked cell wall protein, highly glycosylated, full virulence	—	0.03 ± 0.05 ^b	0.03 ± 0.05 ^b	—	0.12 ± 0.06	0.09 ± 0 ^b
Pga24/YWp1	Secreted yeast form specific protein, dispersal <i>in vivo</i>	0.04 ± 0.01	0.02 ± 0.35 ^b	—	0.02 ± 0.03 ^b	0.04 ± 0.03	0.06 ± 0 ^b
Pga31	GPI-modified cell wall protein	—	0.03 ± 0.05 ^b	—	0.03 ± 0.05 ^b	—	0.19 ± 0.10
Ihd1/Pga36	GPI-modified protein, hypha induced	—	—	—	0.03 ± 0.05 ^b	—	—

^a Absolute protein abundance emPAI values (83, 84) are means for 3 independent biological replicates ± relative SE. Bold type indicates a statistically significant difference ($P < 0.05$).

^b Data based on only one peptide.

^c —, no peptides detected.

not bind to hyphae, as previously shown (42). Thus, it was concluded that although Als3 was present on the surface of *mnt1Δ mnt2Δ* mutant hyphae, as detected by antibody binding, the protein was not expressed in a conformation that was recognized by the *S. gordonii* SspB protein.

C. albicans WT and *mnt1Δ mnt2Δ* cell wall proteomes. To determine if there were significant differences in the expression of cell wall proteins (CWPs) by *C. albicans* wild-type and mutant

strains under the different growth conditions, cell wall preparations (see Materials and Methods) were subjected to trypsin digestion and the resulting peptides were identified by matrix-assisted laser desorption/ionization-time of flight (MALDI-TOF) mass spectrometry. The experiments allowed us to reproducibly identify 23 CWPs (Table 1). Thirteen of these were produced by both strains under each of the hypha-forming conditions, while 10 CWPs were not expressed under all conditions. In YPT-Glc me-

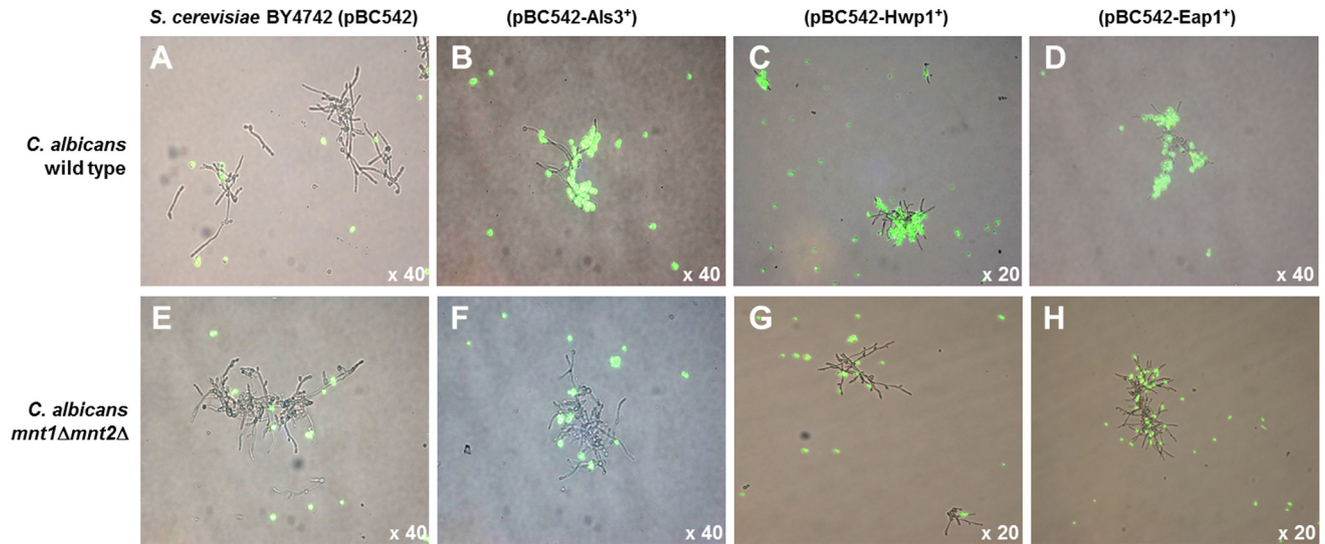


FIG 7 Interactions of *C. albicans* wild type or *mnt1* Δ *mnt2* Δ mutants with *S. cerevisiae* strains expressing *C. albicans* cell wall proteins. *S. cerevisiae* cultures were grown to mid-exponential phase in CSM medium, and the cells were FITC labeled and then mixed with *C. albicans* cells that had been induced to form hyphal filaments for 3 h at 37°C in YPT-GlcNAc medium (see Materials and Methods). After a 2-h incubation of cultures at 30°C with gentle agitation, samples (10 μ l) were removed and placed on glass microscope slides, coverslips were applied, and the cells were visualized using a Leica microscope under phase contrast or fluorescence. The images show phase micrographs merged with respective green fluorescence images of *S. cerevisiae* cells. (A to D) *C. albicans* wild type; (E to H) *C. albicans* *mnt1* Δ *mnt2* Δ mutant. *S. cerevisiae* BY4742 empty vector pBC542 control (A and E), *S. cerevisiae* expressing Als3 (B and F), *S. cerevisiae* expressing Hwp1 (C and G), or *S. cerevisiae* expressing Eap1 (D and H) is shown. Magnification, $\times 20$ or $\times 40$, as indicated.

dium, there were significant ($P < 0.05$) reductions in abundances of the proteins Als1, Als3, Phr1, Rbt1, Sap9, Utr2, and Scw1 for the *mnt1* Δ *mnt2* Δ mutant compared with those for the wild type (Table 1). The apparent reductions for the *mnt1* Δ *mnt2* Δ mutant of most other proteins, except for Rbt5 and Ecm33.3, were not statistically significant. Thus, deficiency in Mnt1 and Mnt2 affected incorporation of glycan-modifying enzymes, e.g., glycosidases and adhesins, into the cell wall when hyphal induction was performed in YPT-Glc medium.

Conversely, in YPT-GlcNAc medium, there were no significant decreases in cell wall protein abundance between the wild type and the mutant except for Sap9 (aspartyl proteinase), which was absent from *mnt1* Δ *mnt2* Δ cell walls (Table 1). The presence of GlcNAc appeared to restore wild-type levels of Als3, Phr1, and Rbt1 in the *mnt1* Δ *mnt2* Δ mutant cell walls, while Hyr1 was significantly ($P < 0.05$) increased compared with that for the wild type. However, in saliva-Glc medium, there were contrasting differences, with the Pga4/Gas1, Phr1, Phr2, Sap9, Utr2, Ecm33.3, and Crh11 proteins being more abundant than in YPT-GlcNAc, and a significant reduction ($P < 0.05$) in levels of Als3, Pga4/Gas1, Phr1, Sap9, and Cht2 was seen for the mutant (Table 1). The apparent increases in Ecm33.3 and Crh11 for the *mnt1* Δ *mnt2* Δ mutant were not statistically significant within the data set. In summary, the *mnt1* Δ and *mnt2* Δ deletions can have pleiotropic effects on hyphal cell wall protein composition depending upon the medium and conditions employed to induce hypha formation. Since the cell wall proteome data indicated that Sap9 was the only protein consistently reduced in abundance from cell wall preparations of the *mnt1* Δ *mnt2* Δ mutant, the possibility that SAP9 expression might be necessary for activating receptors, e.g., Als3 for *S. gordonii*, was considered. To test this, we examined a *sap9* Δ mutant (53) for ability to form biofilms and to support adhesion of *S. gordonii*. The incidence of hyphal filament formation by the

sap9 Δ mutant was similar to that for the *C. albicans* wild type, and *S. gordonii* cells bound at similar levels to filaments formed by the wild type or the *sap9* Δ mutant (data not shown). Therefore, absence of Sap9 from the cell wall of *mnt1* Δ *mnt2* Δ hyphae could not account directly for the deficiency in binding *S. gordonii* cells.

Loss of adhesin function by *mnt1* Δ *mnt2* Δ mutant hyphae. The proteome studies supported the observations shown in Fig. 6 that the hyphal cell wall Als3 receptor for *S. gordonii* (42) in the *mnt1* Δ *mnt2* Δ mutant was present in normal amounts in YPT-GlcNAc (Table 1). We hypothesized, therefore, that the *mnt1* Δ *mnt2* Δ mutations affected cell wall protein structure such that *S. gordonii* receptors, e.g., Als3, were not presented in a functional conformation. Since the hyphal surface proteins Als3 and Hwp1 have been shown to interact with one another (14), we further tested our hypothesis by determining the abilities of Als3 and Hwp1 to interact with hyphae produced by the wild type or the *mnt1* Δ *mnt2* Δ mutant. *Saccharomyces cerevisiae* cells expressing Als3 on their surfaces (24) (Fig. 7B) were able to interact with hyphae formed by the *C. albicans* wild type but not with hyphae formed by the *mnt1* Δ *mnt2* Δ mutant (Fig. 7F). Wild-type *S. cerevisiae* containing empty vector pBC542 (54) did not interact with hyphae from either strain of *C. albicans* (Fig. 7A and E). *S. cerevisiae* cells expressing Hwp1 (Fig. 7C) also bound to hyphae produced by the *C. albicans* wild type but not to those formed by the *mnt1* Δ *mnt2* Δ mutant (Fig. 7G and H). *S. cerevisiae* cells expressing adhesin Eap1 (55) showed partial recognition of *mnt1* Δ *mnt2* Δ mutant hyphae (Fig. 7D). These results suggested that while the hyphal filaments of the *mnt1* Δ *mnt2* Δ mutant appeared to express Als3, this protein was not presented in a conformation that enabled homotypic (Als3-Als3) or heterotypic (Hwp1-Als3) interactions to occur, as they did with wild-type hyphal filaments. The observations provide direct evidence that mannosylation mediated by Mnt1/Mnt2 is essential for determining hyphal cell wall

protein functions and hence interkingdom biofilm formation with *S. gordonii*.

DISCUSSION

A number of studies have demonstrated that mannosylation reactions in *C. albicans* are required for adherence, invasion of host tissues, and virulence (30, 33). The marked reduction in adherence of *mnt1Δ mnt2Δ* mutants to a number of substrata observed previously suggested that *O*-linked glycosylation might indeed be specifically required for adhesion (56). It has not been precisely clear why mannosylation reactions necessarily affect these properties. One possibility is that deficiency in mannosylation directly affects the synthesis, expression, or activities of cell wall proteins and adhesins crucial for invasion and virulence. Oligosaccharide chains are believed to confer stabilizing properties upon the extended regions of fungal extracellular glycoprotein adhesins (57). Thus, mannosylation could be key to formation of a fully functional *C. albicans* cell surface, especially since there is now good evidence that a number of CWP families, e.g., Als and Rbt, crucial to biofilm formation and virulence, carry extensive amino acid residue repeat blocks in their C-terminal regions that are heavily *O*-glycosylated (58). In addition to being attenuated in adherence and invasion, *C. albicans* glycosylation mutants induced lower levels of cytokine production by human peripheral blood monocytes (29). Disruption of the mannosylation processes seems therefore to have a range of damaging effects on the fungus and interactions with the host (28). Despite these observations, the potential for mannan synthesis to be an antifungal target has generally received less attention.

Previously, it was suggested that deletion of *MNT1* or *MNT2* in *C. albicans* might not result in marked changes in the tertiary structure of cell surface mannoproteins, because protein stabilization is thought to be achieved by addition of the first *O*-linked mannose rather than subsequent residues (30). However, our results suggest that the subsequent (second and third) steps in mannosylation are important. Biofilm development and interactions of *C. albicans* with streptococci were dependent upon expression of *MNT1* and *MNT2*, encoding the enzymes adding the second and third mannose residues to the growing mannan chain. Interestingly, the phenotypic effects of the *mnt1Δ mnt2Δ* double deletion on biofilm development were environmentally sensitive. When induced to form hyphal filaments by nitrogen starvation (YPT-Glc) or with salivary glycoproteins, the double mutant produced a mixture of pseudohyphae and true hyphae. However, when induced with GlcNAc, the double mutant formed hyphae similar to wild-type hyphae in structure and proteome.

GlcNAc induces two sets of responses in *C. albicans*. One causes it to switch from budding to hyphal growth, activated by stimulation of adenylyl cyclase and increased cAMP signaling (59). The second activates a pathway independently of cAMP to induce expression of genes necessary to catabolize GlcNAc (60), but this is not required for hypha induction (61). The hyphal-induction signal from GlcNAc is transmitted through Ngt1 and then to the hyphal gene transcription regulator Efg1 (62). Conversely, the induction signals of low nitrogen or salivary (serous) proteins are received through Mep2 or Ras1, with activation of Cyr1 (adenylyl cyclase). Possibly the latter hyphal induction pathway is more sensitive to the effects of *MNT1/MNT2* deletions, such as cell wall stress (63), than the Ngt1 induction pathway.

GlcNAc is not involved directly in *O*-mannosylation, but GlcNAc treatment induces chitin synthesis and could potentially be converted to other monosaccharide cell wall building blocks via its conversion to fructose-6-phosphate. Therefore, the effect of GlcNAc in effectively complementing the *mnt1Δ mnt2Δ* mutant phenotypes may be multifactorial. Despite GlcNAc enhancing biofilm and hypha formation by the *mnt1Δ mnt2Δ* mutant, the hyphae were still deficient in binding *S. gordonii* and in interacting with the *C. albicans* cell wall adhesins Als3 and Hwp1, expressed on the surface of *S. cerevisiae*. This provides biochemical and physiological demonstration that without the functions of *MNT1* and *MNT2*, the intermicrobial cell adhesins are not presented in a way that promotes dual-species community development. Nevertheless, there must be sufficient levels of interaction between *C. albicans mnt1Δ mnt2Δ* cells in GlcNAc to enable monospecies biofilm formation.

The ability of oral streptococci to interact with *C. albicans* may be relevant to longer-term carriage or persistence of *C. albicans* in the oral cavity (29). Evidence suggests that streptococci and *C. albicans* exhibit growth synergy, implying mutual benefit in cocolonization (43, 64). In this article, dual-species communities formed under flow or nonflow conditions, and *S. gordonii* cells could be seen within the biofilms closely associated with hyphal filaments (see Video S5 in the supplemental material). In biofilms formed under medium flow, hyphal filaments extended further into the environment (see Video S3) than they did under nonflow conditions, where the biofilms were structurally more compact (see Video S1). The stimulation of biofilm development under flow could result from better provision of nutrients, removal of inhibitory metabolic end products, shear-force-induced gene expression, or diffusion away from the biofilm of compounds, such as farnesol (65), that inhibit hypha formation.

To determine the effects of *mnt1Δ mnt2Δ* mutation on the production of hyphal cell wall proteins, we prepared cell walls from hypha-forming cells and subjected them to trypsin digestion and proteome analyses. In *C. albicans* cell walls, the more abundant covalently attached proteins are GPI modified, whereas the least abundant are attached via an alkali-labile linkage (66). Our experiments identified peptides from proteins present in the cell wall through covalent linkage. Proteins missing from the cell wall extracts would be either not expressed (transcription) or not properly incorporated (covalently linked) into the cell wall or lacking amenable trypsin cleavage sites. In the *mnt1Δ mnt2Δ* mutant grown in YPT-Glc, there were significant reductions in a wide range of CWPs that would be expected to have major effects on phenotypes such as adhesion, biofilm formation, and invasion of host cells (9, 50, 51). In the presence of GlcNAc, however, the *mnt1Δ mnt2Δ* mutant showed a cell wall proteome much more similar to that of the wild type. Peptides from the GPI-modified hypha-specific adhesin Hwp1 were not detected by liquid chromatography-tandem mass spectrometry (LC-MS/MS), as also reported by others (67). During hypha formation there is enrichment of carbohydrate-active or cell wall-remodeling enzymes Cht2, Crh11, Mp65, Pga4, Phr1, Phr2, and Utr2 (49, 67). We have identified six of these proteins in our studies. Actual relative amounts will be dependent upon the methods employed in cell wall purification and MS analysis and most importantly the precise conditions utilized for growth and hypha formation.

A protein consistently deficient in the *mnt1Δ mnt2Δ* cell wall proteome was Sap9, one of a family of 10 aspartyl proteinases (Sap1 to 10) that are differentially expressed in *C. albicans* (68).

The main effects of *SAP9* deletion are reduced chitinase activity and increased adhesion to epithelial cells (53). It has been suggested that Sap9 is able to trim cell surface proteins, on *C. albicans* or the host, to modulate receptor-ligand interactions (69–71). In a model for Als3-mediated interactions *in vivo*, soluble or membrane-bound Saps are hypothesized to partly digest proteins on the host cell surface, thus providing additional ligands for Als3 to interact with (71). However, it was concluded that this protease activity did not function in providing receptor availability for binding of *S. gordonii*.

Initial adherence in *C. albicans* biofilm formation is mediated at least in part by Als1 (25), and *ALS1* expression is under control of the transcriptional regulator Bcr1. Other adhesin targets of Bcr1 are Als3 and Hwp1, which mediate cell-cell interactions in biofilms (14). The cell wall protein Eap1 is required for biofilm formation *in vitro* and *in vivo* (55) and may be needed for initial layer formation on specific substrata (24). It has been suggested that the Eap1 and Als proteins may play a role in environmental sensing as well as directly as adhesins (72). Genes encoding transglycosidases involved in maintenance of cell wall integrity, such as *CRH11* (Table 1), are coregulated with other hyphal or virulence genes, while *BGL2*, *PHR1*, and *XOG1* encode enzymes that are crucial for delivery of β -1,3-glucan to the biofilm matrix and for accumulation of matrix biomass (73). Levels of a number of these factors were altered in the *mnt1* Δ *mnt2* Δ mutant cell wall proteomes from cells forming hyphae in YPT-Glc and saliva-Glc medium, possibly disabling biofilm formation. The observation that between 2 h and 3 h biofilm development by the *mnt1* Δ *mnt2* Δ mutant effectively ceased is similar to that made for an *als3* Δ mutant (42). This might be consistent with the 80 to 90% reduction in Als3 in the cell wall proteome of the double mutant under YPT-Glc or saliva-Glc medium conditions. On the other hand, GlcNAc-induced hyphae formed by the *mnt1* Δ *mnt2* Δ mutant had a full complement of adhesins (Table 1). We conclude that in the presence of GlcNAc, the functional expression of adhesins on the *C. albicans* cell surface for binding *S. gordonii* but not for initial biofilm formation is affected by the block in mannosylation.

Evidence that adhesins expressed on the surface of GlcNAc-induced hyphae produced by the *mnt1* Δ *mnt2* Δ mutant were not properly functional was provided by the experiments utilizing *S. cerevisiae* strains expressing the adhesin Als3, Hwp1, or Eap1. Strains expressing these proteins bound avidly to *C. albicans* wild-type hyphae but feebly to hyphae produced by the *mnt1* Δ *mnt2* Δ mutant. This suggested that the *mnt1* Δ *mnt2* Δ mutant hyphae did not present a correct configuration of adhesins to interact with these CWP adhesins. Such effects would be extremely difficult to detect by current imaging techniques and are implicated by probing for functionality, such as ability to interact with known ligands. Of note is that some hyphae in all populations (wild type or mutant) appeared not to bind *S. gordonii* cells, thus indicating potential heterogeneity of hyphal adhesin expression or activation within hyphal filament populations.

A current model for the broad and strong adhesive activity of Als3 suggests that Als3 molecules are able to cluster on the cell surface and bind to each other through rapidly dissociable hydrophobic interactions (74). A long glycosylated C-terminal region stalk elevates the N-terminal region and allows flexibility needed for the adhesins to form amyloids. The Als3 ligand binding cleft is located within the N-terminal region (71). We suggest that the initial SspB-Als3 interaction unfolds or extends the Als3 protein

such that further interactions are substantiated, possibly between amyloid-like regions of the proteins. The Als3 protein contains at least one amyloid sequence (32) and SspB carries three potential amyloid sequences within the central region and C-terminal regions of the polypeptide (VVVYTYT, IWFAF, and TTSEVLN), as predicted by the TANGO computer algorithm (75), that could have amyloidogenic activity.

Amyloid interactions between adhesins may provide cohesive strength to *C. albicans* Als proteins. Formation of Als5 adhesin nanodomains on the cell surface was observed in response to mechanical stimuli, which probably caused the T region (Thr-rich N-terminally located sheet) to partially unfold and expose the amyloid-forming sequences (76). The formation of adhesin clusters could thus explain why Als proteins mediate strong adherence (51). These various *in vitro* observations taken collectively with our results in this article provide the basis for a model (see Fig. S3 in the supplemental material) through which we envisage O-mannosylation to play a critical role in development of CWP adhesin functions. In this model, O-mannosylation catalyzed by the enzymes Mnt1 and Mnt2 initially acts by stabilizing adhesin structure and then allowing for subsequent unfolding and extension. The apical N-terminal domains of Als3 protrude from the hyphal cell wall surface and are recognized by *S. gordonii* cells. Binding causes force-induced unfolding of Als3, which would be enhanced in biofilms under flow conditions (as shown in Results). The adhesive (pulling) forces then elicit clustering of Als3 protein molecules to form nanodomains over the cell surface, thus promoting multiple force interactions of Als3 proteins with bacteria. This would be consistent with observations that streptococci can be seen on some hyphae to show localized attachment and accumulations, suggesting that they are interacting with clusters of adhesins. In Fig. S3, Sap9 aspartyl proteinase is shown as modulating the activities of these nanodomains by proteolytically disrupting the intermolecular forces. This would disaggregate hyphal filaments, leading to increased biofilm permeability and more flexibility for recognition of additional binding partner molecules. Dispersal of intermicrobial cell contacts might also be regulated, at least in part, via Sap9-catalyzed hydrolysis of intramolecular peptide bonds of proteins engaged in intermolecular adhesion events. To date, the evidence for adhesin activation has been derived from biophysical *in vitro* analyses (76), but our data provide direct experimental evidence at the physiological level for the hypothesis that Als-mediated adhesion largely depends upon conformational modifications of existing adhesins and that this is dependent upon early-stage O-mannosylation reactions. These reactions are clearly worthy of further consideration as potential targets for control of *C. albicans* colonization and virulence.

MATERIALS AND METHODS

Microbial growth conditions. The microbial strains utilized in this study are listed in Table S1 in the supplemental material. *C. albicans* strains, all derived from CAI4 (77), were cultivated on Sabouraud dextrose agar (Lab M) aerobically at 37°C. *S. gordonii* strains were cultivated anaerobically at 37°C on BHYN agar (per liter: 37 g brain heart infusion broth, 5 g yeast extract, 5 g Neopeptone, and 15 g agar), which was supplemented with erythromycin (5 μ g/ml) for *S. gordonii* expressing green fluorescent protein (GFP). *Lactococcus lactis* strains were grown anaerobically at 30°C on M17-glucose agar (Oxoid) containing 5 μ g erythromycin/ml where appropriate. *Saccharomyces cerevisiae* strains (24, 78) were cultivated aerobically at 30°C on CSM medium (per liter: 6.7 g Difco yeast nitrogen base, 0.77 g complete supplement mixture [CSM] drop-out [CSM-Ura] [For-

medium, Hunstanton, United Kingdom], 20 g glucose, and 30 g agar). Suspension cultures of *C. albicans* were grown in YPD medium (1% yeast extract, 2% peptone, and 2% glucose) in conical flasks at 37°C with shaking (200 rpm). *S. gordonii* strains were grown in BHY medium (brain heart infusion, containing 5 g/liter yeast extract), stationary, at 37°C. YPT medium (1× Difco yeast nitrogen base, 20 mM NaH₂PO₄-H₃PO₄ buffer, pH 7.0, and 0.1% Bacto-tryptone) supplemented with 0.4% glucose (YPT-Glc) or 0.4% *N*-acetyl-D-glucosamine (YPT-GlcNAc) was utilized to support growth of *S. gordonii* and *C. albicans* in planktonic cultures or biofilms.

Generation of *S. gordonii* expressing GFP. Plasmid pAR4 (*rpsT-gfp-mut3b**pJAR2), kindly provided by Patrick Piggot (Temple University School of Medicine), was extracted from *Streptococcus mutans* UA159 (79) by a modified alkaline-lysis technique (41). The plasmid was purified using a Qiaprep spin miniprep kit (Qiagen). *S. gordonii* DL1 cells were made competent for transformation as previously described (80), and transformants containing pAR4 were selected on BHYN agar containing 5 μg erythromycin/ml (see Table S1 in the supplemental material).

Preparation of saliva. Collection of saliva from human subjects was approved by the National Research Ethics Committee South Central Oxford C (no. 08/H0606/87+5). All samples were collected from at least 6 adult subjects, who provided written informed consent. The samples were pooled, treated with 0.25 M dithiothreitol on ice for 10 min, and centrifuged (8,000 × *g* for 10 min). The supernatant was diluted to 10% with sterile water, filter sterilized (0.45-μm-pore-size membrane), and stored at -70°C in 10-ml portions. Saliva (10%) containing 0.1% glucose was utilized as a microbial growth medium (saliva-Glc).

Preparation of microbial cells. *C. albicans* strains were grown for 16 h in YPD. Cells were harvested by centrifugation (5,000 × *g* for 5 min), washed twice in YPT, and suspended at an optical density at 600 nm (OD₆₀₀) of 1.0 (approximately 1 × 10⁷ cells/ml). *S. gordonii* or *L. lactis* cells were grown for 16 h in 10 ml YPT-Glc, harvested by centrifugation (5,000 × *g* for 7 min), and washed twice with YPT (no glucose). Bacteria were suspended in 0.05 M Na₂CO₃-0.1 M NaCl containing 1.5 mM fluorescein isothiocyanate (FITC) and incubated in the dark at 20°C for 30 min with gentle agitation. The bacterial cells were then washed three times with YPT and suspended at an OD₆₀₀ of 0.5 (2 × 10⁸ cells/ml) in YPT.

***C. albicans* interactions with *S. gordonii* in planktonic phase.** Portions (0.2 ml, 2 × 10⁶ cells) of *C. albicans* cell suspension in YPT were added to glass tubes containing warm YPT-Glc, YPT-GlcNAc, or saliva-Glc medium (1.8 ml). The cultures were incubated for 3 h with shaking at 220 rpm; *S. gordonii* (or in some experiments *L. lactis*) cell suspension (1 ml, 2 × 10⁸ cells) labeled with FITC or expressing GFP was then added, and incubation was continued at 37°C for 1 h. Samples of suspension (10 μl) were applied to glass microscope slides and visualized by light or fluorescence microscopy (Leica DMLB) or with a Leica SP5-AOBS confocal laser scanning microscope attached to a Leica DM I6000 inverted epifluorescence microscope. The Volocity software program was utilized to determine density of fluorescence associated with multiple hyphal filaments over at least 10 randomly selected fields of view (expressed as density units per 50 hyphae).

Planktonic-phase interactions of *S. cerevisiae* with *C. albicans*. *S. cerevisiae* BY4742 strains expressing *C. albicans* cell wall proteins (see Table S1 in the supplemental material) were inoculated into warm CSM (10 ml) and incubated for 16 h at 30°C with shaking at 220 rpm. Portions (5 ml) were centrifuged (5,000 × *g*, 5 min), washed twice with YPT, and fluorescently labeled with FITC as described above. The labeled cells were washed thoroughly with carbonate buffer and twice with YPT before being suspended in YPT-Glc at an OD₆₀₀ of 1.0 (1 × 10⁷ cells/ml). A 1-ml suspension of each strain was transferred to a glass tube containing 1 ml *C. albicans* cells preinduced to form hyphae for 3 h in YPT-GlcNAc and incubated at 30°C for 2 h with shaking at 220 rpm. Portions (10 μl) of the cell suspensions were visualized by transmitted light or fluorescence microscopy.

Monospecies and dual-species biofilms of *C. albicans* and *S. gordonii*. Sterile 19-mm glass coverslips were incubated with 10% filter-sterilized saliva at 4°C for 16 h. Individual coverslips were then transferred to 12-well tissue culture plates containing 1.9 ml YPT-Glc, YPT-GlcNAc, or saliva-Glc, and 0.1 ml *C. albicans* cell suspension (1 × 10⁶ cells) was added. The plates were incubated at 37°C for up to 6 h with gentle motion (50 rpm) in a humid environment. Coverslips were removed at intervals, rinsed gently with phosphate-buffered saline (PBS), dried, and stained with 0.5% crystal violet solution for light microscopy or estimation of biomass after releasing the stain with 10% acetic acid and measuring the OD₅₉₅ (43).

For visualization of *S. gordonii* interactions with *C. albicans* hyphae in early biofilms, coverslips were incubated with *C. albicans* as described above for 3 h, removed, and placed in wells containing fresh YPT-Glc medium (0.5 ml). Suspensions (0.5 ml) of FITC-labeled *S. gordonii* were added, and the cultures were incubated for a further 1 h. For comparative monospecies biofilms of *C. albicans*, culture medium alone (0.5 ml) was added. Coverslips were then removed, gently rinsed once with PBS, inverted onto clean glass microscope slides, and examined by transmitted light or fluorescence microscopy as described above.

Preparation of biofilms for CSLM. Plastic culture dishes (35-mm diameter; Mat Tek) with 14-mm No. 1.0 coverslip base glass windows were incubated with 2 ml 10% saliva at 4°C for 16 h. The saliva was aspirated, and 1.8-ml growth medium (YPT-Glc, YPT-GlcNAc, or saliva-Glc) was added to each dish, followed by 0.2 ml *C. albicans* cell suspension (2 × 10⁶ cells). Dishes were incubated in a humid environment at 37°C for 1 h with gentle motion at 50 rpm. All subsequent incubation steps were carried out under these growth conditions. The culture suspensions were aspirated and replaced with the appropriate medium, and the dishes were incubated for 2 h. For dual-species biofilms, the *C. albicans* suspension was gently aspirated, and 0.2 ml *S. gordonii* UB2549 (expressing GFPmut3b*) suspension in appropriate medium was added. The dishes were incubated for 30 min, the *S. gordonii* culture suspension was aspirated, and 2 ml appropriate growth medium was added for further incubation at 37°C for 4 h. The cell suspensions were then aspirated, the dishes were washed gently with sterile deionized water, and *C. albicans* cells were stained with calcofluor white (0.2 μg/ml) just prior to visualizing by confocal scanning laser microscopy (CSLM) with a Leica SP5-AOBS confocal microscope attached to a Leica DM I6000 inverted epifluorescence microscope. Volocity software was utilized to prepare three-dimensional (3D) images and to calculate biofilm heights (in μm). For monospecies biofilms, exactly the same protocols were applied, minus bacteria or *C. albicans*, substituted with sterile growth medium.

Flow-cell biofilms. Flow cell units, consisting of two parallel chambers sealed with a glass coverslip, were prepared as described by Palmer (81). The growth medium input line was connected, medium was drawn through the cells with a syringe, and the cells were injected with 0.5 ml 10% human saliva and incubated at 4°C for 16 h to coat the inside surfaces of the chambers with salivary glycoproteins. The effluent line to a peristaltic pump was then connected, and appropriate growth medium was drawn through the flow cell for 15 min at a flow rate of 6 ml/h. *C. albicans* cell suspension in growth medium (0.2 ml) was injected into the flow cell chamber and incubated statically at 37°C for 1 h. The growth medium (containing 1 μg/ml calcofluor white) was then drawn through the flow cell chambers at a rate of 6 ml/h for up to 16 h. For dual-species biofilms with *S. gordonii*, the medium flow was stopped after 2 h, and *S. gordonii* UB2549 (expressing GFPmut3b*) cell suspension (0.2 ml) was injected into the chamber and incubated without medium flow at 37°C for 30 min. Flow of growth medium was then recommenced and continued for 4 h at a rate of 6 ml/h. Biofilms were visualized by CSLM.

Immunolabeling of *C. albicans* hyphae. Suspension cultures of *C. albicans* were grown in YPD medium for 16 h at 37°C with shaking (200 rpm). Cells were harvested by centrifugation, washed with YPT medium, suspended at an OD₆₀₀ of 0.5 in YPT-GlcNAc medium, and incubated with shaking for 3 h at 37°C. Cells were harvested by centrifugation,

washed, blocked with goat serum, and then reacted with MAB 3-5A monoclonal antibody to the protein Als3, as previously described (52). Antibody binding was detected with FITC-labeled goat anti-mouse IgG F(ab) fragment-specific antibody, and wet mounts were visualized by light microscopy using an Olympus BX50 microscope.

C. albicans cell wall purification. Cell wall protein samples were prepared by a modification of the method previously described (82). *C. albicans* cells (200-ml cultures) were grown at 37°C for 6 h with shaking to an OD₆₀₀ of 0.7 to 0.9. The cells were harvested by centrifugation (5,000 × g, 5 min), washed with 10 mM Tris-HCl, pH 7.5, at 4°C, suspended in 0.2 ml Tris-HCl buffer, and mixed with 0.5 g cold silica beads (Biospec). Protease inhibitor cocktail (Sigma) and 1 mM phenylmethylsulfonyl fluoride (PMSF) were added, and the cells were disrupted by shaking using a Fast-Prep cell breakage machine (FastPrep-24 bead beater; MP Biosciences). The contents of the tubes were washed out into cold 50-ml tubes with 1 M NaCl, the beads were allowed to settle, and the supernatant was removed (repeated twice). Cell wall suspensions were centrifuged (3,000 × g, 5 min, 4°C), and the pellets were washed 5 times with 1 M NaCl and once with double-distilled water (ddH₂O). The crude cell walls were then twice extracted with SDS-mercaptoethanol buffer (50 mM Tris, 2% SDS, 0.3 M β-mercaptoethanol, and 1 mM EDTA [pH 8.0]) at 100°C for 10 min to remove noncovalently bound proteins. Cell walls were then washed thoroughly by alternate suspension in ddH₂O and centrifugation (5 times) and freeze-dried.

Preparation of cell wall peptides. Freeze-dried cell wall sample (1 to 2 mg) was mixed with 0.5 M ammonium bicarbonate in water containing 3 mM dithiothreitol and heated at 60°C for 20 min. Iodoacetamide (30 μl of 55 mM stock solution) was then added, and the suspension was incubated at 25°C for 10 min in the dark. Trypsin (30 μl of 20 mg/ml stock) was added, and the suspension was incubated at 37°C for 14 h and centrifuged at 14,000 × g for 10 min. The supernatant was freeze-dried and extracted with 10% formic acid, and peptides were purified using ZipTip mC18 pipette tips (Millipore) and dissolved in 0.1% formic acid.

Proteomic analysis. Samples (3 μl) were injected into an LC-MS system which comprised an UltiMate 3000 LC instrument (Dionex Ltd., United Kingdom) fitted with a PepSwift monolithic poly(styrene-codivinylbenzene) (PS-DVB) column (200-μm inside diameter [i.d.] by 5 cm; Dionex) coupled to an HCTultra ion trap mass spectrometer (Bruker Daltonik GmbH, Bremen, Germany) fitted with a low-flow nebulizer in the electrospray ionization (ESI) source and controlled by HyStar software (version 4.0; Bruker Daltonik). Peptides were separated at a flow rate of 2 μl/min using a linear gradient of 0 to 40% acetonitrile-water-formic acid (80:20:0.04) (solvent B) in water-acetonitrile-formic acid (97:3:0.05) (solvent A) over 40 min, followed by a 1-min column wash in 90% solvent B and a 12-min equilibration step in solvent A. MS/MS data (scan range, *m/z* 100 to 2,200; averages = 2) were acquired in positive data-dependent AutoMS(n) mode using the esquireControl software program (version 6.2; Bruker Daltonik). Up to three precursor ions were selected from the MS scan (range, *m/z* 300 to 1,500; averages = 3) in each AutoMS(n) cycle. Precursors were actively excluded after being selected twice within a 1-min window, and singly charged ions were also excluded. Peptide peaks were detected (maximum of 9,999 compounds above an intensity threshold of 50,000) and deconvoluted automatically using DataAnalysis software (version 3.4; Bruker Daltonik). Mass lists in the form of Mascot Generic Format (*.mgf) files were created automatically and used as inputs to Mascot MS/MS ion searches via a local Mascot server (version 2.2; Matrix Science, London, United Kingdom) with a database built from the <CALBI_prot.txt > file (date stamp, 5 February 2002; 6,166 sequences) downloaded from CandidaDB (<ftp://ftp.pasteur.fr/pub/GenomeDB/CandidaDB/FlatFiles>). Search parameters used were the following: enzyme = trypsin; fixed modifications = carbamidomethyl (C); variable modifications = oxidation (M); mass values = monoisotopic; peptide mass tolerance = 1.5 Da; fragment mass tolerance = 0.5 Da; max missed cleavages = 1; instrument type = ESI-TRAP. Search results were displayed by Mascot after selection of the following parameters: standard scoring; re-

quire bold red; ion score or expect cutoff = 0.05. An open-source web application (emPAI Calc) was utilized for estimation of protein abundance (83, 84).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.00911-14/-/DCSupplemental>.

Figure S1, TIF file, 7.2 MB.
Figure S2, TIF file, 8.1 MB.
Figure S3, TIF file, 7.2 MB.
Table S1, DOCX file, 0.1 MB.
Video S1, AVI file, 55.5 MB.
Video S2, AVI file, 47.6 MB.
Video S3, AVI file, 45.1 MB.
Video S4, AVI file, 8 MB.
Video S5, AVI file, 29.2 MB.
Video S6, AVI file, 24.1 MB.

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