Endosomal and AP-3-Dependent Vacuolar Trafficking Routes Make Additive Contributions to Candida albicans Hyphal Growth and Pathogenesis

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Candida albicans mutants deficient in vacuolar biogenesis are defective in polarized hyphal growth and virulence. However, the specific vacuolar trafficking routes required for hyphal growth and virulence are unknown. In Saccharomyces cerevisiae, two trafficking routes deliver material from the Golgi apparatus to the vacuole. One occurs via the late endosome and is dependent upon Vps21p, while the second bypasses the endosome and requires the AP-3 complex, including Aps3p. To determine the significance of these pathways in C. albicans hyphal growth and virulence, aps3ΔΔ, vps21ΔΔ, and aps3ΔΔ vps21ΔΔ mutant strains were constructed. Analysis of vacuolar morphology and localization of the vacuolar protein Mlt1p suggests that C. albicans Aps3p and Vps21p mediate two distinct transport pathways. The vps21ΔΔ mutant has a minor reduction in hyphal elongation, while the aps3ΔΔ mutant has no defect in hyphal growth. Interestingly, the aps3ΔΔ vps21ΔΔ double mutant has dramatically reduced hyphal growth. Overexpression of the Ume6p transcriptional activator resulted in constitutive hyphal growth of wild-type, aps3ΔΔ, and vps21ΔΔ strains and formation of highly vacuolated subapical compartments. Thus, Ume6p-dependent transcriptional responses are sufficient to induce subapical vacuolation. However, the aps3ΔΔ vps21ΔΔ mutant formed mainly pseudohyphae that lacked vacuolated compartments. The aps3ΔΔ strain was virulent in a mouse model of disseminated infection; the vps21ΔΔ mutant failed to kill mice but persisted within kidney tissue, while the double mutant was avirulent and cleared from the kidneys. These results suggest that while the AP-3 pathway alone has little impact on hyphal growth or virulence, it is much more significant when endosomal trafficking is disrupted.

The fungal vacuole is an acidic cellular compartment containing a variety of hydrolytic enzymes. It has a central role in the turnover of cellular macromolecules, is a storage site for many metabolites, and is crucial for resistance to osmotic, ionic, pH, and oxidative stresses. A number of previous studies have attempted to define the importance of vacuolar function in pathogenic fungi, including the mucosal and systemic opportunistic Candida albicans. Several C. albicans mutants deficient in vacuolar trafficking have been reported which are sensitive to various stresses and defective in polarized hyphal growth (28, 39). Furthermore, several of these mutants have profound defects in virulence. The vacuole’s role in stress tolerance is likely to be crucial for colonization of host tissue and survival of the ensuing immune response. In addition, C. albicans virulence requires transition between yeast and highly polarized hyphal modes of growth (28, 39). Thus, the fungal vacuole may act on two levels during infection: to enhance C. albicans survival within host tissue and to support polarized hyphal growth required for tissue damage (23, 35). However, the relative contribution of either facet of vacuolar function to pathogenesis is yet to be determined.

During C. albicans hyphal growth, the fungal vacuole expands dramatically to produce subapical cells devoid of cytoplasm, while the cytoplasm migrates within the hyphal tip compartment (2, 21). This increase in volume necessitates a significant redistribution of cellular membrane to the subapical vacuoles. However, the mechanism by which subapical vacuoles expand is unknown, as is the mechanism by which vacuolar transport supports polarized hyphal growth. Furthermore, how subapical vacuole expansion is coordinated with apical extension during hyphal growth is unknown. Finally, while several distinct vacuolar trafficking pathways are known to exist (9), the importance of each to C. albicans hyphal growth and pathogenesis has not been determined.

Many vacuolar enzymes are delivered via the early stages of the secretory network. In Saccharomyces cerevisiae, at least two distinct biosynthetic trafficking routes deliver proteins from the Golgi apparatus to the vacuole (6). One delivers proteins, including carboxypeptidase Y, via a late endosome (also known as the prevacuolar compartment [PVC]). This endosomal route is dependent upon the Rab GTPase Vps21p (18). A second pathway sorts cargo, including alkaline phosphatase (ALP), into a distinct set of vesicles which bypass the late endosome and are delivered directly to the vacuole. The ALP trafficking pathway is facilitated by the AP-3 coat complex, which has four subunits (Aps3p, Apm3p, Apl5p, and Apl6p) and functions in the formation of ALP vesicles at the trans-
Golgi apparatus (12, 40). The purpose of this study was to determine the relative contributions of endosomal and AP-3-dependent trafficking routes to \textit{C. albicans} hyphal growth and virulence.

### MATERIALS AND METHODS

**Growth conditions.** Strains were routinely grown on YPD (1% yeast extract, 2% Bacto peptone, 2% dextrose) at 30°C, supplemented with uridine (50 μg ml^{-1}) when necessary (22). For growth curves, overnight cultures were subcultured into 20 ml fresh YPD medium to an optical density at 600 nm (OD_{600}) of 0.2 and incubated at 30°C with shaking. The OD_{600} was determined from samples taken hourly. Transformants were selected on minimal medium (YNB [6.75 g liter^{-1} yeast nitrogen base plus ammonium sulfate and without amino acids, 2% dextrose, 2% Bacto agar]) supplemented with the appropriate auxotrophic requirements, as described for \textit{S. cerevisiae} (10), or uridine at 50 μg ml^{-1}.

**Plasmid construction.** Plasmids pGEMURA3, pGEMHIS1, and pRSARGSpEc (45) were provided by A. Mitchell (Carnegie Mellon University). Plasmid pLUX (38) was provided by W. Fonzi (Georgetown University). Plasmid pLUXPS21 was previously constructed (23), and pLUXAPS3 was made as follows. The APS3 open reading frame (ORF) with 5' and 3' untranslated region (UTR) sequences was amplified from S. cerevisiae gDNA using HiFi Platinum Taq (Invitrogen) and primers APS3AMPF2 and APS3AMPF (Table 1). Purified products were then cloned into the BamHI site of vector pLUX. pACTI was made by amplifying 1,000 bp of the ACTI 5' UTR using primers ACT1PF and ACT1PR with HiFi Taq. The product was then cloned between the KpnI and SalI sites of CIp10 (31).

**Strain construction.** Strains used in this study are described in Table 2. Gene deletion strains were constructed by the PCR-based approach described by Wilson et al., using the ura3 \textit{his1} \textit{arg4} strain BWP17 (45), kindly provided by A. Mitchell (Carnegie Mellon University). \textit{C. albicans} was transformed using the lithium acetate procedure (19). The \textit{ura3Δ} mutant was constructed in a previous study (23). APS3 deletion cassettes were amplified by PCR using pARGSpEc (ARG4 marker), pGEMHIS1 (HIS1 marker), or pDDB57 (reusable UR3-3pl200 cassette) (44, 45) as a template with the primer pair APS3DISF and APS3DISR. \textit{C. albicans} was diploid; thus sequential deletion of each APS3 allele was achieved using first \textit{aps3aΔ::ARG4} and then \textit{aps3aΔ::HIS1} disruption cassettes to generate a \textit{ura3Δ} strain. An \textit{aps3aΔ vps21Δ} double mutant was made by sequentially deleting both APS3 alleles from a \textit{vps21Δ} \textit{ura3Δ} strain previously constructed (23). This strain was transformed with the \textit{psu3Δ::URA3-dpl200} cassette and then the \textit{aps3aΔ::ARG4} cassette. The regeneration of \textit{ura3Δ} recombinants was selected on YNB medium supplemented with uridine and 1 μg ml^{-1} 5-fluoroorotic acid (5-FOA) (5). Correct integration of either cassette was confirmed at each step by diagnostic PCR using primers ARG4DET2 plus APS3AMPF (ARG4 integration), H11032 plus APS3AMF (HIS1 integration), or HIS1F1268 plus ARG4DET2 (URA3 integration). Southern blot analysis was also performed using a specific probe to the 3' UTR of \textit{APS3}, amplified using primers APS3AMPF plus APS3PBR. Correct gene deletion resulted in replacement of the entire 480-bp \textit{APS3} ORF. Finally a wild-type copy of \textit{ARG4} or \textit{VPS1}, including 5' and 3' flanking sequences, was introduced to the deletion strains on pLUXAPS3 or pLUXVPS21 (23) to produce prototrophic “reconstituted” strains. Prototrophic gene deletion strains were produced by transforming the mutant strains with plasmid vector (pLUX) alone. Either plasmid was digested with NheI to target integration into (and reconstitute) the \textit{URA3} loci. This strategy circumvents the well-described positional effects of \textit{URA3} integration (26).

Plasmids harboring wild-type (pEM-31) or “dominant active” (pEM16-1 and pEM16-3) \textit{RIM101} alleles (15) were linearized with HpaI to target integration into the \textit{RIM101} locus; \textit{ura3Δ} recipient strains were transformed, and \textit{Ura}+ transfectants were selected. pCT7UM65 was made by amplifying the \textit{UME6 ORF} and 3' UTR sequences with \textit{UME6ORF} and \textit{UME6AMPMLUR} primers and cloning downstream of the \textit{ACT1} promoter in pACTI, between the Sall and MluI sites.

**Stress phenotypes.** \textit{C. albicans} strains were grown overnight in YPD broth at 30°C, cell density was adjusted to 10^7 ml^{-1} in sterile water, and serial 1:5 dilutions were performed in a 96-well plate. Cells were then applied to agar using a sterile multipronged applicator. Resistance to temperature stress was determined on YPD agar at 37 and 42°C, and resistance to osmotic stress was determined on YPD agar plus 2.5 M glycerol or 1.5 M NaCl. Other stresses included YPD plus 5 mM caffeine, 10 mM rapamycin, 5 mM sodium orthovanadate.

### Table 1. Oligonucleotides used in this study

| Primer | Sequence (5'→3') |
|--------|------------------|
| APS3DISF | AATTGTGTTCCGAGCTAATAAGATCCATTTATACAAAGATTCGATAT |
| APS3DISR | CCAACACAGAAATTCTGCTGGTGGATTTGTGGGGAAT |
| APS3AMPF2 | TCATCGAGTCGACTTTGGAATTATTTTTTTAATATTAATATCGAG |
| APS3AMPR | TCATCGAGTCGAGGAACGAGAAATAGG |
| APS3PBR | TGCAACAGGAATATACAAATCC |
| APS3DET | AAGATTTGGATGATATCAAGG |
| APS3DTR | CCTAACCCAGAACCCCTTCTATC |
| UME6ORRF | ATCACGTCAGCTGATATAATGAATCCATCATG |
| UME6AMPMLUR | ATCACGTCAGCTGATATAATGAATCCATCATG |
| ACT1PF | TCATCGAGTCGACTTTGGAATTATTTTTTTAATATTAATATCGAG |
| ACT1PR | TCATCGAGTCGACGTATAAATGATTACCCATATGG |
| MLT1GFPF | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| MLT1GFPF | GATATATATATCGCACCGATATATATCTAGAAGGACCACCTTTGATT |
| MLT1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| ARG4DET | GATATATATATCGCACCGATATATATCTAGAAGGACCACCTTTGATT |
| ARG4DETR | CCGAGCGTTGCGTAATACG |
| HIS1F1268 | CCAGCTACTGTCTCTACTTTG |
| HIS1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| HIS1DETR | CCGAGCGTTGCGTAATACG |
| MLT1GFPF | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| MLT1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| ARG4DET | GATATATATATCGCACCGATATATATCTAGAAGGACCACCTTTGATT |
| ARG4DETR | CCGAGCGTTGCGTAATACG |
| HIS1F1268 | CCAGCTACTGTCTCTACTTTG |
| HIS1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| HIS1DETR | CCGAGCTACTGTCTCTACTTTG |
| ARG4DET | GATATATATATCGCACCGATATATATCTAGAAGGACCACCTTTGATT |
| ARG4DETR | CCGAGCTACTGTCTCTACTTTG |
| HIS1F1268 | CCAGCTACTGTCTCTACTTTG |
| HIS1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| HIS1DETR | CCGAGCTACTGTCTCTACTTTG |
| MLT1GFPF | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| MLT1DET | ATCAAAAAAAATGGTATTGATAATATAAATATCAATAT |
| MLT1DETR | CATCACCTTACCCCAAGG |
| MLT1DETR | CATCACCTTACCCCAAGG |

*a* Engineered restriction enzyme sites are underlined.
date, 200 μg ml⁻¹ Congo red, and 0.05% SDS. Carbon source utilization was tested using 3% ethanol, glycerol, or potassium acetate instead of glucose. Secreted protease activity was examined on bovine serum albumin (BSA) plus YE agar at 37°C (13). Resistance to nitrogen starvation was determined in SD-N date agar at 37°C (13). The number of colonies on each plate was determined after 48 h, and the CFU g⁻¹ of tissue were calculated.

### RESULTS

#### Identification of Candida albicans AP-3 orthologs.

BLAST searches of the CGD database identified homologs of all four *S. cerevisiae* AP-3 subunits. The most highly conserved was the small subunit encoded by *APS3*, which was 57% identical to ScAsp3p. In order to determine if *C. albicans* possesses a functional AP-3 pathway, *aps3ΔΔ* deletion and *aps3ΔΔ* plus *AP3* “reconstituted” strains were constructed. In *S. cerevisiae*, ALP, as well as the vacuolar t-SNARE Vam3p, is delivered to the vacuole through the AP-3 pathway. However, it has been shown that when AP-3-dependent transport is blocked, both Vam3p and ALP still reach the vacuole, as they are redirected into the Vps21p-dependent endosomal trafficking route (12).

#### Mouse model of hematogenously disseminated candidiasis.

All procedures were approved by the Institutional Animal Care and Use Committee of LSUHSC-New Orleans and complied with all relevant federal guidelines. *C. albicans* was grown overnight in YPD cultures at 30°C (200 rpm). Cells were washed twice in sterile PBS, and cell density was determined using a hemocytometer. Each strain was then diluted to 5 x 10⁶ cells ml⁻¹ in sterile PBS. Then, 0.1 ml of each cell suspension was inoculated into the lateral tail vein of BALB/c mice (Charles River Laboratories, Maryland). Viable cell counts of each inoculum were confirmed by plating appropriate dilutions onto YPD agar plates and counting the colonies formed after 48 h. Mice were monitored for 28 days, and those showing distress were euthanized. Kaplan-Meier survivor curves were plotted, and statistical significance was determined using the log rank test. At day 28, surviving mice were euthanized. The kidneys from each mouse were weighed and homogenized in PBS, and serial dilutions of homogenate were plated to YPD agar. The number of colonies on each plate was determined after 48 h, and the CFU g⁻¹ of tissue were calculated.

#### Table 2. *C. albicans* strains used in this study

| Strain        | Relevant genotype | Source or reference |
|---------------|-------------------|---------------------|
| SC5314        | VPS21/VPS21 APS3/APS3 | 20                  |
| YJB6284       | VPS21/VPS21 APS3/APS3 | 3                   |
| BWP17         | VPS21/VPS21 APS3/APS3ΔΔ his1ΔΔ arg4ΔΔ | 45                  |
| CA14          | VPS21/VPS21 APS3/APS3ΔΔ | 21                  |
| APD1/2        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ/ARG4/psa3ΔΔ/URA3 | This study |
| APR1/3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ/ARG4/psa3ΔΔ/URA3 | This study |
| 21F1          | vps21ΔΔ/His1/vps21ΔΔ/dpl200 APS3/APS3ΔΔ | 23                  |
| VDI1/4        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 | 23                  |
| VR1/4         | ura3ΔΔ/URA3/VPS21 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 | 23                  |
| VDI1/4        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 | 23                  |
| AVU1-3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ/ARG4/psa3ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
| VU1-3         | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
| CRM1-3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ/ARG4/psa3ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
| ARM1-3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ/ARG4/psa3ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
| AVR1-3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
| AVR1-3        | ura3ΔΔ/URA3 his1ΔΔ arg4ΔΔ vps21ΔΔ/ARG4/vps21ΔΔ/His1 UME6/UME6::ACT1-UME6::URA3 | This study |
tant also had large intact vacuoles but exhibited the same morphological defects described for the \textit{aps3}/H9004 mutant. As expected, reintroduction of \textit{APS3} (but not \textit{VPS21}) into the double mutant resolved these defects. The \textit{aps3}/H9004 \textit{vps21}/H9004 mutant also had a delay in dye internalization, which was restored by reintroduction of \textit{VPS21} but not \textit{APS3}. This clearly established that \textit{C. albicans} \textit{APS3} is involved in vacuolar membrane transport and that its functions are distinct from those of \textit{VPS21}.

In \textit{S. cerevisiae}, ALP is delivered to the vacuole as an inactive precursor, which is proteolytically activated upon delivery to the vacuole (24). In order to assess ALP trafficking in the \textit{aps3}/H9004 and \textit{vps21}/H9004 mutant strains, ALP activity was assayed. While neither single mutant was significantly affected, the \textit{aps3}/H9004 \textit{vps21}/H9004 mutant had reduced ALP activity compared to that of the control strain YJB6284 (Fig. 1B). ALP activity was also largely restored by reintroduction of either \textit{APS3} or \textit{VPS21} to the double mutant. This is consistent with ALP trafficking in \textit{S. cerevisiae}, where ALP can transit to the vacuole through either AP-3- or \textit{VPS21}-dependent pathways.

We also examined the localization of an Mlt1-GFP fusion protein, previously localized to the vacuolar membrane (41), in each mutant. In “wild-type” cells, Mlt1-GFP colocalized with FM4-64 in large ring structures, as expected for a vacuolar membrane protein (Fig. 2). Both \textit{aps3}/H9004 and \textit{vps21}/H9004 mutants showed largely similar distributions, indicating that the fusion protein reached the vacuole, except that the \textit{aps3}/H9004 mutant also had significant Mlt1-GFP localization to the previously noted intravacuolar vesicles. In the \textit{aps3}/H9004 \textit{vps21}/H9004 double mutant, Mlt1-GFP colocalized with FM4-64 on the vacuole, but significant amounts were also mislocalized within the cytoplasm as punctate spots and a diffuse “halo” surrounding the vacuole. These data are consistent with \textit{APS3} and \textit{VPS21} functioning in two independent trafficking pathways to the vacuole, with the Mlt1-GFP cargo being able to utilize
either. It also suggests that Mlt1-GFP is able to utilize a third delivery pathway to reach the vacuole when the Aps3p and Vps21p pathways are blocked.

As previously reported, the vps21ΔΔ/H9004/H9004 mutant had increased secreted protease activity compared to the control strains (23), while loss of APS3 had no impact on this phenotype (data not shown).

AP-3- and VPS21-dependent pathways play distinct roles in resisting cellular stress. The fungal vacuole has established roles in resisting cellular stress. Therefore, to determine the biological significance of AP-3 trafficking in C. albicans, resistance of each mutant to various cellular stresses was examined.

FIG. 2. Mlt1-GFP can transit to the vacuole by either Aps3p- or Vps21p-dependent pathways. An MLT1-GFP fusion was introduced into control, aps3ΔΔ, vps21ΔΔ, or aps3ΔΔ vps21ΔΔ strain backgrounds. Each strain was pulse-labeled with FM4-64 to label the vacuole. Cells were then observed by deconvolution fluorescence microscopy to determine Mlt1-GFP (left panel) and FM4-64 (center left) distribution and by Nomarski optics (center right). Merged images are also shown (right panel). Arrowheads indicate punctate Mlt1-GFP, and arrows indicate diffuse cytoplasmic Mlt1-GFP distributions in the double-mutant strain. Scale bar, 10 μM.

Both single mutants were sensitive to rapamycin, with the double mutant even more sensitive, suggesting that each pathway makes an independent contribution to rapamycin resistance. The aps3ΔΔ mutant seems otherwise unaffected in stress tolerance. No growth defects were observed at 37°C for either mutant strain, and both single mutants grew at 42°C. However, the double mutant showed a synthetic growth defect at 42°C (Fig. 3A), under ionic stress (1.5 M NaCl), and in the presence of caffeine. The vps21ΔΔ mutant was also found to be hypersensitive to Congo red and SDS, agents known to disrupt cell wall integrity. However, loss of APS3 had no impact on the sensitivity to these agents. Interestingly, the
aps3ΔΔ mutant was consistently more resistant to nitrogen starvation than the wild-type control strain (Fig. 3B). The vps21ΔΔ mutant was slightly sensitive to nitrogen starvation, as was the aps3ΔΔ/vps21ΔΔ mutant. Thus, while loss of APS3 alone has little effect on stress resistance, the synthetic stress phenotypes of the aps3ΔΔ/vps21ΔΔ mutant suggest that there is some functional interplay between AP-3 and endosomal trafficking routes.

**AP-3- and VPS21-dependent pathways make an additive contribution in supporting hyphal growth.** Loss of APS3 alone has no obvious effect on C. albicans hyphal growth on M199/FBS agar, in liquid M199 or FBS culture, or under embedded conditions (Fig. 4). The vps21ΔΔ mutant was previously shown to have mild defects in hyphal growth under each of these conditions (23). However, the double mutant has profound defects in hyphal growth. The double mutant was unable to produce filaments on M199 or FBS agar and formed few filaments when embedded in an agar matrix (Fig. 4A and D). In liquid M199 and FBS media the double mutant grew mainly as yeast and pseudohyphae, with half of the pseudohyphae having an elongated “hook” shape (Fig. 4B and C). Reintroduction of VPS21 almost completely restored normal hyphal growth un-
under each of these conditions, while reintroduction of APS3 restored hyphal growth to vps21ΔΔ levels. These data imply that AP-3 trafficking becomes much more important in supporting hyphal growth when Vps21p-mediated trafficking is disrupted.

The elongated pseudohyphal cells observed under hypha-inducing conditions could indicate that the double mutant is unable to maintain the highly polarized cell state necessary for true hyphal growth. UME6 encodes a transcription factor required to maintain polarized hyphal growth and hypha-specific gene transcription (1). Overexpression of UME6 has been shown to promote extended hyphal growth, even under conditions which would normally support growth in the yeast form (11). To examine the relationship between APS3- and VPS21-mediated trafficking and the control of hyphal extension, UME6 was overexpressed in each mutant strain. This caused the control and aps3ΔΔ strains to grow exclusively as extended hyphae (Fig. 5A). The vps21ΔΔ mutant formed marginally

FIG. 4. Aps3p- and Vps21p-dependent pathways make additive contributions to polarized hyphal growth. (A) Cell suspensions of each strain were applied as spots to M199 agar and incubated at 37°C for 4 days. (B) Each strain was induced to filament in 10% FBS at 37°C for 5 h. (C) A minimum of 200 cells grown in FBS at 37°C for 5 h were scored for each strain, according to the morphology index (MI) described by Odds and colleagues (29, 33). MI of 1, spherical and nearly spherical cells; MI of 2, ovoid cells with length up to twice the cell width; MI of 3, pseudohyphal cells with obvious constrictions; MI of 4, parallel-sided true hyphal cells. Data are the means of results of three separate experiments; error bars represent standard deviations. Pearson’s chi-square test was used to determine whether the proportion of MI categories observed differed between the control strain (YJB6284) and each of the other strains. *, P < 0.001. (D) Cells of each strain were sandwiched between two layers of YPD agar and incubated at room temperature for 3 days.
less-elongated hyphae, while the hyphae formed in the \(\text{aps3}\Delta/\Delta \text{vps21}\Delta/\Delta\) background were dramatically shorter than those of the control. Thus, normal hyphal growth cannot be restored in the \(\text{vps21}\Delta/\Delta\) and \(\text{aps3}\Delta/\Delta \text{vps21}\Delta/\Delta\) strains by overexpression of Ume6p. Rim101p is a pH-responsive transcription factor required for alkaline-induced hyphal growth (14, 15). Defects in Rim101p activation have been shown to contribute to the hyphal growth defects in several \(\text{C. albicans}\) endosomal trafficking mutants (25, 46). However, introduction of a “dominant active” \(\text{RIM101}\) allele (15) also failed to restore normal hyphal growth in the \(\text{vps21}\Delta/\Delta\) (23) or \(\text{aps3}\Delta/\Delta \text{vps21}\Delta/\Delta\) mutants (data not shown).

Finally, it has been shown previously that a \(\text{vma7}\Delta/\Delta\) mutant, lacking a subunit of the vacuolar \(\text{v}-\text{ATPase}\), has defects in hyphal growth and virulence (37), presumably due to an inability to acidify its vacuole. Thus, it was considered that a

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**FIG. 5.** Overexpression of **UME6** does not restore normal hyphal elongation to the **aps3\Delta/\Delta \text{vps21}\Delta/\Delta** mutant. **UME6** was overexpressed from the **ACT1** promoter in CAI4 (“wild-type” [WT] control), **aps3\Delta/\Delta**, **vps21\Delta/\Delta**, and **aps3\Delta/\Delta \text{vps21}\Delta/\Delta** strain backgrounds. Each strain was also transformed with the **pACT1** vector alone (shown only for the control strain). (A) Strains were streaked to YPD and grown at 30°C. (B) Each strain was grown in YPD broth at 30°C, stained with FM4-64, and observed using a deconvolution microscope. Scale bar, 10 \(\mu\text{M}\).
block in vacuolar trafficking of the v-ATPase may account for the defective hyphal growth in the *aps3ΔΔ* vps21ΔΔ mutant. However, unlike the *vma7Δ*Δ mutant, *aps3ΔΔ* vps21ΔΔ and *aps3ΔΔ* vps211ΔΔ mutants were all able to grow at pH 8 (Fig. 3A) and accumulated the weak base quinacrine within their vacuoles (data not shown). These data suggest that acidification of the vacuoles in each mutant was unaffected.

Overexpression of *UME6* induces hyphal growth with vacuolated subapical cells. *C. albicans* hyphal growth involves the induction of a hypha-specific transcription program, as well as posttranslational modulation of proteins to establish a polarized, or apical mode of growth. However, it is unknown how subapical vacuolation is coordinated with these events. To examine the interrelationship between hypha-specific transcriptional responses and subapical vacuolation, vacuole morphology was examined in the *UME6* overexpression strains. Strains were grown in YPD medium at 30°C, and vacuoles were labeled with FM4-64. In the wild-type background, the elongated hyphae induced by *UME6* overexpression had extensively vacuolated subapical compartments (Fig. 5B). Cell compartments near the hyphal tip had much smaller, rounder compartments within the kidneys of mice surviving to day 28 was determined. The *aps3ΔΔ* vps21ΔΔ mutant was not detected in kidney tissue, suggesting that it is cleared from the inoculated mice. However, while neither vps21ΔΔ or *aps3ΔΔ* vps21ΔΔ plus *APS3* strains caused lethal infection, both were able to persist within the kidney tissue at significant levels (5.5 × 10^5 ± 5.1 × 10^5 and 6.8 × 10^5 ± 5.7 × 10^5 CFU g^-1, respectively; mean ± standard deviation [SD]; n = 5). This suggests that *APS3*-dependent trafficking does play a role in *C. albicans* pathogenesis in the absence of *VPS21*-mediated transport.

**DISCUSSION**

Previous studies have clearly established that disruption of vacuolar trafficking leads to defective hyphal growth and loss of virulence in *C. albicans* (8, 23, 35). However, the specific trafficking events required for each of these processes are unknown. This study has established that *C. albicans* has an AP-3-dependent vacuolar trafficking pathway. Deletion of *APS3* led to the accumulation of large vesicles within the vacuole lumens of many cells, which could be explained by a deficiency in the breakdown of multivesicular bodies or autophagosomes. The Vps21p Rab GTPase was previously localized to the late endosome and has an established role in vacuolar trafficking (23). The distinct vacuolar morphology and stress phenotypes of the *aps3ΔΔ* and vps21ΔΔ mutants suggest that Aps3p and Vps21p operate in distinct trafficking pathways to the *C. albicans* vacuole, as is the case in *S. cerevisiae*. In support of this, the Mlt1-GFP fusion protein localized to the vacuolar membrane in either single mutant but showed significant mislocalization in the double mutant. This suggests that the Mlt1p “cargo” can transit to the vacuole via either Aps3p- or Vps21p-mediated pathways. Nonetheless, despite disruption of both AP-3 and endosomal trafficking routes, a significant amount of Mlt1-GFP reached the vacuole, and the *aps3ΔΔ* vps21ΔΔ mutant had a largely intact vacuole compartment. Furthermore, the modest reduction in ALP activity and normal vacuolar acidification suggest that additional trafficking routes may exist between the Golgi apparatus and vacuole in *C. albicans*.

Initial phenotypic analyses of the *aps3ΔΔ* mutant failed to identify any substantial defect in stress resistance or polarized hyphal growth. This suggested that the ALP trafficking pathway was of little consequence in *C. albicans*. In addition to the mild stress phenotypes and increased protease secretion previously reported (23), the vps21ΔΔ mutant was also found to be sensitive to agents which disrupt cell wall integrity. These phenotypes are consistent with those reported by Lee and colleagues (27), and they suggest that disrupting endosomal transport may influence secretion and cell surface composition. Loss of Aps3p had no effect on secretion, endocytosis, or cell wall integrity. Interestingly, loss of both genes resulted in synthetic stress phenotypes. This further supports the notion that Aps3p and Vps21p operate in distinct pathways which share some functional overlap.

While the loss of *APS3* has little impact upon hyphal growth,
Loss of APS3 in the vps21Δ/Δ background has a profound impact. The double mutant grows at an essentially normal rate in the yeast form; thus, this is not merely the result of reduced growth-limiting apical extension. This suggests that Golgi-to-vacuole transport via Aps3p- and Vps21p-dependent routes is particularly important during polarized hyphal growth. Ume6p overexpression enabled us to induce constitutive hyphal growth in a rich medium at 30°C, stress-free conditions under which all strains grow at similar rates in the yeast form. Ume6p overexpression was sufficient to induce hyphal growth with vacuolated subapical cells in the wild-type, overexpression was sufficient to induce hyphal growth with which all strains grow at similar rates in the yeast form. Ume6p growth in a rich medium at 30°C, stress-free conditions under growth-limiting apical extension. This suggests that Golgi-to-
the yeast form; thus, this is not merely the result of reduced pact. The double mutant grows at an essentially normal rate in

Several previously described C. albicans mutants with disrupted vacuole morphology have profound defects in hyphal growth. Despite an apparently normally-sized vacuole compartment, the aps3AΔ vps21Δ/Δ mutant exhibited profound defects in hyphal growth. Eck and colleagues have also reported a vps34AΔ mutant with defects in vacuolar trafficking (8). The vps34AΔ mutant had an enlarged vacuole in the yeast form but still has severely reduced hyphal growth. These results suggest that a controlled flow of membrane between the Golgi apparatus and vacuole is required to support hyphal growth, rather than an intact vacuole morphology per se. The UME6 overexpression data imply that the mechanisms responsible for subapical vacuole expansion lie downstream of Ume6p and are controlled either directly or indirectly by hypha-specific transcriptional responses. To my knowledge, this provides the first link between hypha-specific signaling and control of vacuolar dynamics in C. albicans. Putting these data together suggests a simple model where activation of the transcription factor Ume6p stimulates Golgi-to-vacuole transport to promote subapical vacuole expansion. Thus, overexpression of a single regulator of hyphal growth, Ume6p, is sufficient to induce both an apical mode of growth and subapical vacuolation. This suggests that these processes may be coregulated.

Vacuolar trafficking may support hyphal growth through several possible mechanisms. First, it seems likely that disruption of vacuolar trafficking would prevent vacuolation of subapical cells, and this may limit hyphal elongation. Second, the vacuole has an established role in regulating turgor pressure within the fungal cell (24), which may help support hyphal extension by providing a force for directional growth (30). Thus, defective regulation of turgor pressure in vacuolar mutants may limit apical extension. Finally, it is possible that a specific “factor” is required to localize to the vacuole to support polarized hyphal growth. For example, loss of the vacuolar v-ATPase subunit Vma7p (and consequently vacuolar acidification) results in defective hyphal growth (37). However, vacuolar acidification was apparently intact in the aps3AΔ vps21Δ/Δ mutant. None-
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