Distinct domains of the *Candida albicans* adhesin Eap1p mediate cell–cell and cell–substrate interactions

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The adhesion of *Candida albicans* to host tissues contributes to its virulence, and adhesion to tissues or medical devices is a necessary step in biofilm formation. *EAP1* encodes a glycosylphosphatidylinositol (GPI)-anchored glucan-cross-linked cell wall protein that mediates adhesion of *C. albicans* to various materials and cells, and appears to be required for *C. albicans* biofilm formation *in vitro* and *in vivo*. In this study, we demonstrated that the Eap1p N-terminal signal peptide and C-terminal GPI-anchor sequences result in similar protein localization in *Saccharomyces cerevisiae* and *C. albicans*. To investigate the contribution of different Eap1p domains to adhesion, we expressed Eap1p domain deletion mutants in non-adherent *S. cerevisiae* strains. The N-terminal domain mediates yeast cell–cell adhesion and invasive growth. Two Ser/Thr-rich domains containing tandem repeats were required to project the N-terminal region into the extracellular environment and to mediate adhesion to polystyrene. The N-terminal tandem repeat domain mediated adhesion to mammalian epithelial cells and promoted *S. cerevisiae* pseudohyphal growth. These results suggest a modular structure of Eap1p in which each domain serves multiple, often distinct, functions.

**INTRODUCTION**

*Candida albicans* is the leading cause of candidiasis, most often manifesting as superficial mucosal infections. *Candida* spp. are also major agents of systemic bloodstream infections, causing 8% of all such nosocomial infections in the USA (Edmond et al., 1999; Jarvis, 1995; Pfaller et al., 1998; Pfaller & Diekema, 2007). *C. albicans* is also the most common fungus associated with biofilm formation on bioprosthetic materials (Lopez-Ribot, 2005). Cells in biofilms are much more resistant to antifungal agents, including amphotericin B and azoles (Al-Fattani & Douglas, 2004; Hawser & Douglas, 1995).

Adhesion between *C. albicans* cells and materials or host cells has been implicated as an early step in biofilm formation. Adhesion of *C. albicans* to mammalian tissues is also considered to be a very important determinant of pathogenesis (Calderone & Fonzi, 2001). *C. albicans* glycosylphosphatidylinositol-dependent cell wall proteins (GPI-CWP) are characterized by the presence of an N-terminal signal peptide and a C-terminal sequence signalling GPI anchor attachment (Dranginis et al., 2007). Many of these proteins are involved in mediating the adhesion of *C. albicans* cells to host materials and/or inert surfaces (Fu et al., 2002; Gaur & Klotz, 1997; Phan et al., 2007; Staab et al., 1999; Zhao et al., 2005, 2007). Previous studies suggest that the N-terminal domains of the adhesins that are members of the GPI-CWP family mediate binding to substrates and the GPI anchor is required for incorporating these proteins into the cell wall (Frieman et al., 2002; Loza et al., 2004). These adhesins, in many cases, also contain serine/threonine-rich tandem repeat domains essential for proper presentation of the N-terminal substrate-binding sites (Frieman et al., 2002; Loza et al., 2004). However, the conserved Als tandem repeats in Als5p facilitate adhesion to fibronectin and greatly increase cell–cell aggregation (Raucoo et al., 2006).

Previous studies have demonstrated that the *C. albicans* Eap1p is a member of the GPI-CWP family (Li et al., 2007). The *C. albicans* Eap1p adhesin was originally identified because of its ability to mediate adhesion to polystyrene when the *EAP1* gene was expressed in a flocculin-deficient *Saccharomyces cerevisiae* strain. *EAP1* expression enhanced attachment of *S. cerevisiae* to HEK293 kidney epithelial cells and played an additional role in mediating interactions between *S. cerevisiae* and *C. albicans* cells (Li & Palecek, 2003, 2005). Expression of *EAP1* in a *C. albicans* efg1/efg1 mutant was able to restore its partially reduced...
adhesion to HEK293 epithelial cells (Li & Palecek, 2003). Deleting EAP1 in C. albicans reduced cell adhesion to polysytrene and epithelial cells in a gene dosage-dependent manner. C. albicans eap1/eap1 mutant cells were also defective in cell–cell adhesion when those cells were grown in a parallel-plate flow chamber under shear flow (Li et al., 2007). The defect in Eap1p-mediated adhesion was also associated with a defect in C. albicans biofilm formation in an in vitro and an in vivo model (Li et al., 2007). In C. albicans, EAP1 is expressed at similar levels in both yeast and filamentous forms (Li et al., 2007); however, the relative expression of EAP1 compared to other adhesins and the localization of Eap1p remain unknown. Many insights into the functions of C. albicans adhesins have been derived in C. albicans (Fu et al., 2002; Li et al., 2007; Phan et al., 2007; Staab et al., 1999; Zhao et al., 2005, 2007). Heterologous expression of C. albicans adhesins in S. cerevisiae has also proven valuable in exploring C. albicans adhesion function (Loza et al., 2004; Rauceo et al., 2006). Eap1p mediates adhesion to many substrates, including synthetic materials, yeast cells and mammalian cells, when expressed in both S. cerevisiae and C. albicans, suggesting that S. cerevisiae may be used as a relevant heterologous system to study the adhesive function of Eap1p in C. albicans.

In the experiments described in this paper, we constructed deletion mutants of EAP1 and expressed wild-type Eap1p and Eap1p mutants in an adhesin-deficient S. cerevisiae strain to study the mechanisms by which Eap1p mediates adhesion and define the critical functions of the domains of Eap1p in the context of adhesion. We demonstrated that different regions of Eap1p mediated adhesion to distinct substrates.

**METHODS**

**Strains and media.** Escherichia coli strain DH5α was used for general recombinant techniques according to protocols described by Sambrook et al. (1989). S. cerevisiae haploid strain SPY308 (Sigma278b MATα::MATα/ura3-52 his3::hisG leu2::hisG flo8::kan’, diploid strain SPY311 (Sigma278b MATα/α::ura3-52/ura3-52 his3::hisG/his3::hisG leu2::hisG/LEU2::flo8::kan’/flo8::HIS3) and strain BJ5464 (Bj2) (MATα ura3-52 trp1 leu2A1 his3A200 pep4::HIS3 prb1Δ16R can1 GAL) were used as the hosts for all mutants. Yeast strains were routinely cultured in YPD medium (1% yeast extract, 2% peptone, 2% glucose) at 30 °C. Synthetic complete media lacking specific nutrients and filamentous growth media have been described previously (Ahn et al., 1999; Kron et al., 1994). Synthetic low-ammonium (SLAD) medium contained 50 μM ammonium sulfate. Uracil was added to SLAD medium to a concentration of 0.2 mM to make SLAD + Ura. Galactose was added to media to replace glucose in order to express genes within plasmids containing the S. cerevisiae GAL1 promoter. Yeast cells were transformed using lithium acetate transformation (Gietz et al., 1992). Pseudohyphal growth assay and agar invasion assay were performed as previously described (Li & Palecek, 2003).

**Plasmid construction.** Sequences of all oligonucleotides used in cloning are provided in Table 1. Plasmids designed to express GFP fusions to the N- and truncated C-termini of EAP1 were constructed in a similar manner to that described by Mao et al. (2003). A partial ORF encoding the N-terminal 42 amino acids of Eap1p was amplified from C. albicans SC5314 genomic DNA. This PCR product was digested with PvuI and Spel and ligated into pRHp1Sig.GFP.GPI (Mao et al., 2003) to yield pEap1Sig.GFP.Hwp1GPI. A partial ORF encoding the C-terminal 47 amino acids of Eap1p was amplified from C. albicans SC5314 genomic DNA. This PCR product was digested with BamHI and Smal and ligated into pRHP1Sig.GFP.GPl to generate pEAP1.Sig.GFP.GPl. pEAP1.Sig.GFP.NOGPI is essentially identical to pEAP1.Sig.GFP.GPl except that 21 amino acid residues from the C-terminus of Eap1p encoding the GPI anchor signal were deleted. The sequences between the PvuI and Smal sites encoding Eap1p/GFP fusion proteins were also PCR-amplified from pEAP1.Sig.GFP.GPl and pEAP1.Sig.GFP.NOGPl then cloned into pCRT302 (Boder et al., 2000) cut with EcoRI and XhoI to generate pCTEAP1.Sig.GFP.GPl and pCTEAP1.Sig.GFP.NOGPl, respectively.

The signal peptide of EAP1 was cloned in pCTEAP1.Sig.GFP.GPl as an EcoRI–Spel fragment generated by PCR with the oligonucleotides EAP1.Sprimer.EcoRI and EAP1.HA.Sig3, which contains the sequence encoding the haemagglutinin (HA) epitope of the influenza virus, to generate pCTEAP1.HA.Sig.GFP.GPl. The PCR products produced by amplifying the genomic DNA of C. albicans strain SC5314 with oligonucleotides EAP1.Sprimer.XhoI and EAP1-1, EAP1-2, EAP1-3,

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’–3’)</th>
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</tr>
<tr>
<td>EAP1.3primer</td>
<td>GAGGACCTGAGTCATACAAAGTGACTAATGACGCCC</td>
</tr>
<tr>
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<tr>
<td>EAP1-2</td>
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<tr>
<td>EAP1-3</td>
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<tr>
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<tr>
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Table 1. Oligonucleotide primers used in this study

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EAP1-4, and EAP1-5, respectively, were digested with SpeI and XhoI and ligated into pCTEap1.HA.Sig.GFP.GPI to yield HA-Eap1p, HA-Eap1pΔ1N, HA-Eap1pΔ2N, HA-Eap1pΔ3N and HA-Eap1pΔ4N, respectively. The PCR products produced by amplifying the genomic DNA of C. albicans strain SC5314 with oligonucleotides EAP1-1 and EAP1-1-Right, EAP1-2-Right, EAP1-3-Right, EAP1-4-Right, and EAP1-5-Right, respectively, were digested with SpeI and BamHI and ligated into pCTEap1.HA.Sig.GFP.GPI to yield HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI, HA-Eap1pΔ3C.GPI and HA-Eap1pΔ4C.GPI, respectively. pCTEap1.HA.Sig.GFP.GPI was digested with SpeI and BamHI. The fragments amplified by PCR with oligonucleotides EAP1-2 and EAP1-2-Right and oligonucleotides EAP1-4 and EAP1-2-Right were cloned into this digested vector to generate HA-Eap1p.Sig.TR1.GPI and HA-Eap1p.Sig.TR2.GPI, respectively. For all constructs in which fragments were generated by PCR, the final constructs were verified by DNA sequence analysis.

**Microscopy.** To detect Eap1p-GFP fusion proteins and HA-tagged Eap1p mutant proteins, constructs were transformed into S. cerevisiae BJ5464 (B1α) cells and ura3 prototrophs were selected on minimal glucose plates. Cells carrying these plasmids were cultured overnight at 30 °C in minimal medium containing galactose. The cells were then washed in PBS and immunofluorescence staining was performed as previously described (Li et al., 2007). To visualize yeast cell aggregates, cells were resuspended in PBS buffer and placed onto a microscope slide for photographing.

**Western analysis.** Yeast cells expressing each Eap1p-GFP fusion protein were cultured as described above and harvested by centrifugation. To analyse the Eap1p-GFP fusion proteins in medium, the supernatant was concentrated using 10 kDa molecular mass cutoff Microcon centrifugal filters. Yeast cell walls were isolated according to the method of Mao et al. (2003). The proteins released into the supernatant by laminarinase digestion or from concentrated cell-free medium were subjected to SDS-PAGE and analysed by Western blotting using the ECL Western blot kit (Amersham) and an anti-GFP monoclonal antibody.

**Polystyrene binding assay.** Yeast cells were cultured overnight at 30 °C in minimal medium containing galactose. After brief sonication to break cell clumps, equal number of cells was added to the wells of 24-well plates and incubated in sodium phosphate buffer (pH 6.0) for 2 h. Plates were washed a few times and the number of cells remaining attached was quantified by an XTT [2,3-bis(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide]-reduction assay (Ramage et al., 2001).

**RESULTS**

**Eap1p/GFP fusion protein is localized to the cell wall in S. cerevisiae**

EAP1 encodes a GPI-anchored, glucan-cross-linked cell wall protein in C. albicans (Li et al., 2007). Mao et al. (2003) demonstrated that green fluorescent protein (GFP) fused to the N- and C-termini of the C. albicans GPI-anchored cell wall proteins Hwp1p, Als3p and Rbt5p was localized to the C. albicans cell wall. We used this approach to verify the cell wall localization of Eap1p in S. cerevisiae. A synthetic GFP (Cormack et al., 1997) was fused between 42 amino acid residues from the N-terminus and 47 amino acid residues from the C-terminus of Eap1p. The resulting fusion protein was expressed in S. cerevisiae under regulation of the GAL1 promoter. The pCTEAP1.Sig.GFP.GPI transformants expressing this fusion protein showed localized fluorescence at the cell surface (Fig. 1a), demonstrating that the EAP1-derived sequences were capable of targeting the fusion protein to the cell surface. We next used GFP fused to the 42 amino acids from the N-terminus but lacking the C-terminal sequence (pCTEAP1.Sig.GFP.NOGPI) to investigate if the GFP fusion was secur ed to the cell surface via a GPI anchor. This fusion did not localize to the cell wall, but was found inside the cell, appearing concentrated in extranuclear organelles (Fig. 1a). We tested the cell-free medium of the pCTEAP1.Sig.GFP.NOGPI transformants for secretion of the GFP fusion by Western blotting with an anti-GFP monoclonal antibody. The pCTEAP1.Sig.GFP.NOGPI transformants secreted a protein that was detected by the anti-GFP antibody and whose size corresponded to that of the fusion protein, whereas GFP was not detected in culture supernatants of the pCTEAP1.Sig.GFP.GPI transformants (Fig. 1a). To address whether the Eap1p-GFP fusion proteins anchored to the plasma membrane or were incorporated in the cell wall, purified cell walls from S. cerevisiae strains transformed with GFP constructs were digested with β-glucanase. The GFP fusion protein was released from the cell walls of the pCTEAP1.Sig.GFP.GPI transformants and detected with anti-GFP antibodies (Fig. 1c). In control experiments, immunoreactive GFP proteins were not found in β-glucanase-treated cell walls of cells transformed with pCTEAP1.Sig.GFP.NOGPI nor in samples treated with buffer alone (Fig. 1c). The N- and C-termini of Eap1p were also required to localize Eap1p in C. albicans cell wall (Li et al., 2007). Based on these results, we concluded that Eap1p N-terminal and C-terminal sequences result in similar protein localization at the cell surface in S. cerevisiae and C. albicans.

**Two Eap1p Ser/Thr-rich domains containing tandem repeats are required for projecting the N-terminal region into the extracellular environment**

Two Ser/Thr-rich regions containing a variable number of tandem repeats divide Eap1p into five regions (Fig. 2). The N-terminal region contains a signal peptide and may also contain a substrate-binding domain based on the common structures among adhesins. Two alleles of EAP1 exist in C. albicans. The segment following the N-terminal region consists of ~90 and ~17 copies of an imperfect peptide sequence, STPATE (Fig. 2, boxes with vertical lines), in the long and short allele, respectively. The long allele was used in this study. The region following this segment is homologous to the putative N-terminal adhesion domain (Fig. 2, grey boxes). The C-terminal tandem-repeat domain consists of ~12 copies of an imperfect peptide sequence, TPAAPGTPVESQPVIPGTET (Fig. 2, hatched boxes). The C-terminal region contains a GPI anchor addition signal involved in cross-linking Eap1p to the yeast cell wall (Fig. 2). To investigate the function of each domain in Eap1p, we constructed a series of
HA-tagged Eap1p deletion mutants by sequentially deleting one domain from the N-terminus or the C-terminus while leaving the signal peptide and the GPI anchor signal intact to ensure localization to the cell wall (Fig. 2). The HA epitope was engineered into the EAP1 coding sequence C-terminal to the 30th amino acid residue.

We monitored the surface expression of the Eap1p mutants expressed in S. cerevisiae by fluorescence microscopy. A monoclonal antibody directed against the HA epitope recognized HA-Eap1p, HA-Eap1pΔ1N, HA-Eap1pΔ2N, HA-Eap1pΔ3N (Fig. 3a), HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI and HA-Eap1pΔ3C.GPI (Fig. 3b) on the surface of yeast, but the antibody did not bind to yeast transformed with the plasmid encoding HA-Eap1pΔ4N and HA-Eap1pΔ4C.GPI (Fig. 3a, b). The HA epitope could also be detected by the anti-HA antibody when only the N-terminal or the C-terminal tandem repeat domain was expressed in the presence of the signal peptide and the GPI anchor addition signal (Fig. 3c). These results suggest that both the N-terminal and the C-terminal tandem repeat domains can project the N-terminus of the protein away from the cell and into the extracellular environment, where it is accessible to antibody binding. We also verified the surface expression of all the constructs using fluorescence-activated cell sorting (FACS) after sonication to break cell clumps, and a similar amount of each Eap1p mutant protein was found on the surface of yeast cells in which the construct was accessible to the anti-HA antibody (data not shown).

**Fig. 1.** Cell wall localization of Eap1p-GFP fusion proteins. (a) Epifluorescence microscopy analysis of S. cerevisiae strain BJ5464 transformed with empty vector, plasmid pCTEAP1.Sig.GFP.GPI and pCTEAP1.Sig.GFP.NOGPI. Plasmid pCTEAP1.Sig.GFP.GPI encodes a yeast-enhanced GFP fused to 42 amino acid residues from the N-terminus of Eap1p and 47 amino acid residues from the C-terminus of Eap1p. Plasmid pCTEAP1.Sig.GFP.NOGPI encodes a protein identical to pCTEAP1.Sig.GFP.GPI except that 21 amino acid residues from the C-terminus of Eap1p encoding the GPI anchor signal were deleted. The transformants were grown in minimal medium containing galactose for 20 h at 20 °C prior to imaging. Images were taken with identical exposure. (b) Analysis of the cell-free supernatant from S. cerevisiae strain BJ5464 transformed with plasmid pCTEAP1.Sig.GFP.GPI, pCTEAP1.Sig.GFP.NOGPI, and empty vector pCT302. The fractions were run on a 12 % SDS-PAGE gel and visualized by Western blotting with an anti-GFP monoclonal antibody. (c) Cell wall extractions of S. cerevisiae strain BJ5464 transformed with plasmid pCTEAP1.Sig.GFP.GPI, pCTEAP1.Sig.GFP.NOGPI, and empty vector pCT302. Cell walls were extracted twice in boiling SDS and digested with β-1,3-glucanase (+) or no enzyme (−), then separated into pellet and supernatant fractions. The glucanase-treated and untreated supernatant fractions were loaded on a 12 % SDS-PAGE gel and visualized by Western blotting with an anti-GFP monoclonal antibody. Positions of size standards (kDa) are shown on the left in (b) and (c).
carrying an empty vector was unable to invade agar on synthetic medium lacking uracil, but wild-type EAPI expression induced invasive growth in the flo8Δ strain (Fig. 4). The expression of any of the EAPI mutants lacking the N-terminal domain was not able to restore invasiveness to the haploid flo8Δ strain, suggesting that the N-terminal domain of Eap1p was required for agar invasion (Fig. 4a). Expression of all C-terminal EAPI mutants, except HA-Eap1pΔ4C.GPI, was able to restore invasiveness to the haploid flo8Δ strain (Fig. 4b). Presumably the N-terminal domain of HA-Eap1pΔ4C.GPI was not exposed to the extracellular environment; the HA tag on this protein was not accessible to an anti-HA antibody (Fig. 3b). To further investigate the role of the N-terminal domain of Eap1p in...
haploid invasive growth, we expressed a construct encoding a chimera protein of the N-terminal domain of Eap1p and the C-terminal Ser/Thr-rich tandem repeat domain of Candida glabrata Epa1p. The C-terminal Ser/Thr-rich domain of Epa1p is able to project the N-terminal domain of Epa1p through the permeability barrier of cell wall and into the extracellular environment (Frieman et al., 2002). Expression of the C-terminal domain of Epa1p alone did not restore the ability of the haploid flo8A strain to invade agar, but the strain carrying the construct encoding the Eap1p-Epa1p chimera was able to invade agar (Fig. 4c), suggesting that the N-terminal domain of Eap1p is able to mediate invasive growth if projected beyond the cell surface.

We propose that the invasive growth assay as performed in this study reflects the strength of yeast cell–cell and cell–agar adhesion because the patched cells in lower layers adhere to the agar substrate and bind the cells in upper layers, preventing them from washing away. To follow up on these results, we tested the ability of yeast cells expressing various Eap1p mutants to form cell aggregates in suspension. Any strain expressing an Eap1p mutant with an accessible N-terminal domain formed large cell aggregates, while control strains and strains expressing Eap1p mutants lacking the N-terminal domain grew as single cells or small clusters, further implying a role of the N-terminal domain of Eap1p in mediating yeast cell–cell adhesion (Fig. 5).

The two Ser/Thr-rich domains containing tandem repeats are required for adhesion to polystyrene

Expression of EAP1 in S. cerevisiae increased adhesion to polystyrene and deletion of EAPI in C. albicans reduced adhesion to polystyrene (Li & Palecek, 2003; Li et al., 2007). S. cerevisiae haploid flo8A strains expressing the N-terminal and the C-terminal Eap1p mutants that lack one of the two Ser/Thr-rich tandem repeat domains (HA-Eap1pΔ4N and HA-Eap1pΔ4N.GPI) exhibited reduced adhesion to polystyrene compared to the wild-type and the Eap1p mutants that contain both tandem repeat domains (Fig. 6a, b; P<0.05). We also transformed the haploid flo8A strain with constructs encoding the N-terminal (HA-Eap1pΔ4N and HA-Eap1pΔ4N.GPI) and C-terminal (HA-Eap1pΔ4N.GPI) tandem repeat domains of Eap1p and found that the adhesion to polystyrene was restored by expressing these tandem repeat domains bounded by the signal peptide and GPI anchor sequences of Eap1p (Fig. 6c; P<0.01). S. cerevisiae cells expressing the Eap1p–Epa1p chimera did not exhibit enhanced adhesion to polystyrene compared to cells expressing the C-terminal Ser/Thr-rich domain of C. glabrata Epa1p (Fig. 6c). These results suggest that the two Ser/Thr-rich tandem repeat domains are sufficient to mediate adhesion to polystyrene.

The N-terminal tandem repeat domain mediates adhesion to mammalian cells

Expression of EAPI in S. cerevisiae permitted yeast cell attachment to human HEK293 kidney epithelial cells, and deletion of EAPI in C. albicans reduced yeast cell adhesion to HEK293 cells (Li & Palecek, 2003; Li et al., 2007). To identify the domains of Eap1p involved in adhesion to...
mammalian cells, we expressed the N-terminal and C-terminal EAP1 mutants in a haploid flo8Δ strain and quantified the yeast adhesion to mammalian HEK293 cells. HA-Eap1p induced adhesion of the flo8Δ strain to HEK293 cells, but deletion of the Eap1p N-terminal tandem repeat domain dramatically reduced this adhesion (Fig. 7a, b; \( P < 0.01 \)). Interestingly, deletion of the N-terminal domain, which was required for adhesion to yeast cells, did not reduce adhesion to HEK293 cells (Fig. 7a). Likewise, expression of the chimera protein containing the N-terminal domain of Eap1p and the C-terminal domain of

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Fig. 5. Photomicrographs of yeast cell–cell adhesion. S. cerevisiae strain SPY308 (flo8Δ) was transformed with plasmids encoding the proteins indicated and grown overnight in liquid SC–Ura medium containing galactose. Yeast cell aggregates were visualized by photomicroscopy, (a) HA-Eap1p, HA-Eap1pΔ1N, HA-Eap1pΔ2N, HA-Eap1pΔ3N and HA-Eap1pΔ4N. (b) HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI, HA-Eap1pΔ3C.GPI and HA-Eap1pΔ4C.GPI. (c) EPA1C (the C-terminal Ser/Thr-rich domain of C. glabrata Epa1p) and EAP1ADH.EPA1C (the N-terminal putative yeast–yeast adhesion domain of Eap1p and EPA1C fusion protein).

Fig. 6. Adhesion to polystyrene of haploid S. cerevisiae strain SPY308 (flo8Δ) expressing Eap1p mutant proteins. (a) SPY308 was transformed with empty vector (control) and plasmids encoding HA-Eap1p, HA-Eap1pΔ1N, HA-Eap1pΔ2N, HA-Eap1pΔ3N and HA-Eap1pΔ4N. (b) SPY308 was transformed with empty vector (control) and plasmids encoding HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI, HA-Eap1pΔ3C.GPI and HA-Eap1pΔ4C.GPI. (c) SPY308 (flo8Δ) was transformed with empty vector (control) and plasmids encoding HA-Eap1Sig.TR1.GPI, HA-Eap1Sig.TR2.GPI, EPA1C and EAP1ADH.EPA1C. Strains were incubated in 24-well plates for 2 h, then plates were washed and adhesion of cells was quantified by an XTT assay. Error bars represent the standard deviations for three independent experiments, with three replicates in each experiment. An asterisk (*) indicates strains that adhered to a greater extent than the control (\( P < 0.05 \) or \( P < 0.01 \) in a two-tailed Student’s t-test).
**C. glabrata** Eap1p had no effect on adhesion to HEK293 cells (Fig. 7c), indicating that the N-terminal domain of Eap1p is substrate specific. To test whether the N-terminal tandem repeat domain was sufficient to mediate adhesion to HEK293 cells, we expressed HA-Eap1Sig.TR1.GPI and found its expression significantly enhanced the adhesion to HEK293 cells (Fig. 7; P<0.01).

**The N-terminal tandem repeat domain promotes *S. cerevisiae* pseudohyphal growth**

Diploid *flo8Δ*/*flo8Δ* cells failed to form any filaments on low-nitrogen SLAD medium (Fig. 8), and EAPI expression restored filamentous growth to a *flo8Δ*/*flo8Δ* strain (Li & Palecek, 2003). The expression of both HA-Eap1p (wild-type) and HA-Eap1pΔ1N restored the ability of a *flo8Δ/*flo8Δ strain to form pseudohyphae, indicating that the N-terminal domain of Eap1p, which was involved in mediating yeast cell–cell adhesion, was not required for pseudohyphal growth (Fig. 8a). Expression of HA-Eap1pΔ2N, HA-Eap1pΔ3N and HA-Eap1pΔ4N was not able to complement the pseudohyphal growth defect of the diploid *flo8Δ*/*flo8Δ* strain (Fig. 8a), while expression of HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI and HA-Eap1pΔ3C did complement this pseudohyphal defect (Fig. 8b). To test whether the N-terminal tandem repeat domain alone is able to activate the pseudohyphal growth of the *flo8Δ*/*flo8Δ* strain, we transformed the diploid *flo8Δ*/*flo8Δ* strain with HA-Eap1Sig.TR1.GPI and found that pseudohyphal growth was restored (Fig. 8c).

**DISCUSSION**

The *S. cerevisiae* genome contains about 60 putative GPI-anchored proteins, of which many possess Ser/Thr-rich C-terminal domains containing tandem repeats (Caro et al., 1997). The availability of the *C. albicans* genome provides a strategy to predict ORFs that potentially encode GPI-CWP proteins based on their common sequence features. Thus far, 104 putative GPI-anchored proteins have been predicted and more than 65% of these proteins have completely unknown functions to date (Richard & Plaine, 2007). Understanding whether these GPI-anchored proteins mediate adhesion to different materials, host cells, or other microbial cells should increase our ability to design strategies to prevent such adhesion and resulting infections.

Adherence activity of the GPI-anchored proteins has commonly been ascribed to the N-terminal domains of these proteins because deletions in these regions usually abrogate binding activity (Frieman et al., 2002; Loza et al., 2004; Sheppard et al., 2004; Staab & Sundstrom, 1998; Wojciechowicz et al., 1993). N-terminal globular domains...
adhesins and ligands at the cell surface determine nanomolar affinities. These affinities and the location of bind peptide or sugar ligands, with millimolar to microscopic and macroscopic characteristics of cell–cell associations (Dranginis et al., 2007). Here, we demonstrated that the N-terminal domain of Eap1p contains a ligand-binding domain mediating adhesion to yeast cells. This domain is also capable of rescuing the invasive defect of a haploid flo8A S. cerevisiae strain. The increase in invasion upon expression of Eap1p domains may be the result of the increased cell–cell adhesion. Numerous mutations that increase cell–cell adhesion via FLO11-dependent or FLO11-independent mechanisms in S. cerevisiae also enhance invasive growth (Guo et al., 2000; Palecek et al., 2000; Svarovsky & Palecek, 2005; Vyas et al., 2003).

The tandem repeats in Eap1p appear to have no direct role in yeast–yeast binding, but instead may project the N-terminal ligand-binding domains away from the body of the cell and into the extracellular environment. Previous studies also suggested that tandem-repeat-containing domains are essential for stabilizing the correct conformation of the N-terminal ligand-binding domain and contributing positively to the adherence activity mediated by members of the Als family of adhesins (Loza et al., 2004; Rauceo et al., 2006). Our data also demonstrated that the Eap1p Ser/Thr-rich domain containing tandem repeats was able to mediate adhesion to polystyrene and to mammalian epithelial cells directly, in addition to its function of projecting the N-terminal domain into the extracellular environment. Further, the N-terminal tandem repeat domain of Eap1p was able to promote the pseudohyphal growth of S. cerevisiae, suggesting that this tandem repeat domain was also able to induce morphological change of this fungus. The mechanism by which Eap1p induces pseudohyphal formation in S. cerevisiae is not known, but may be related to its adhesive properties, as the endogenous S. cerevisiae adhesin Flo11p and endochitinase/endoglucanase regulation of mother–daughter separation influence filamentous differentiation (King & Butler, 1998; Lo & Dranginis, 1998; Pan & Heitman, 2000). However, EAPI expression does not appear to strongly influence morphology in C. albicans (Li et al., 2007). Our results indicate that tandem repeats of GPI-anchored proteins possess multiple functions, including modulating protein structure, mediating adhesion to certain substrates, and signalling morphological changes.

The tandem repeats are thought to trigger frequent recombination events in genes or between genes and pseudogenes, causing expansion and contraction of the gene size in S. cerevisiae, and the size variation creates quantitative alterations in phenotypes (Verstrepen et al., 2005). Different clinical isolates of C. albicans have differing numbers of repeats for the same ALS gene (Hoyer, 2001). The two alleles of EAPI in C. albicans strain SC5314 also contain differing numbers of repeats in the N-terminal tandem repeat domain. Our results suggest these tandem repeat domains can directly mediate adhesion in addition to their roles in generating cell-surface diversity through genetic recombination and epigenetic regulation.

Fig. 8. Pseudohyphal formation in diploid S. cerevisiae strain SPY311 (flo8Δ/flo8Δ) expressing Eap1p mutant proteins. (a) SPY311 transformed with empty vector and with plasmids encoding HA-Eap1p, HA-Eap1pΔ1N, HA-Eap1pΔ2N, HA-Eap1pΔ3N and HA-Eap1pΔ4N. (b) SPY311 transformed with empty vector and with plasmids encoding HA-Eap1p.GPI, HA-Eap1pΔ1C.GPI, HA-Eap1pΔ2C.GPI, HA-Eap1pΔ3C.GPI and HA-Eap1pΔ4C.GPI. (c) SPY311 transformed with empty vector and with a plasmid encoding HA-Eap1p.Sig.TR1.GPI. Cells were streaked onto the low-nitrogen SLAD medium containing galactose. Colonies were allowed to grow at 30 °C for 2 days and then photographed. Scale bars, 0.2 mm.
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Roles of the C. albicans Eap1p domains in adhesion


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