Complications in cell-surface labelling by biotinylation of *Candida albicans* due to avidin conjugate binding to cell-wall proteins

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Initial contact between the opportunistic fungal pathogen *Candida albicans* and host tissue occurs at the cell surface. Biotin derivatives have been used to label the cell-surface proteins of yeasts, with labelled proteins subsequently detected by avidin–reporter conjugates. Previous work has indicated that avidin can bind to *C. albicans* proteins in the absence of biotin, suggesting a possible host-cell-recognition mechanism by fungal cell-surface proteins. To investigate this mechanism, Western blots of proteins extracted from biotinylated and mock-treated cells were probed with avidin or modified-avidin reagents. Each avidin reagent bound to cell-wall proteins extracted from non-biotinylated cells. Binding did not appear to be due to the lectin-like activity of the cell-wall proteins of *C. albicans* or to the presence of biotin in the sample itself. Binding was inhibited by added biotin, by the chaotrope KSCN and by NaCl in a concentration-dependent manner, although inhibition varied among the avidin conjugates tested. Thus, the non-specific binding of avidin to the cell-wall proteins of *C. albicans* appears to involve hydrophobic and electrostatic interactions, depending on the particular avidin species. These observations demonstrate potential pitfalls in the use of avidin–biotin complexes to identify cell-surface molecules and could provide insights into protein–protein interactions at the *C. albicans* cell wall.

Keywords: yeast, adhesion

INTRODUCTION

*Candida albicans* is found in the normal fungal microbiota of humans. Although usually an obligate commensal, this organism is opportunistically pathogenic and has become an increasing problem over the past two decades, particularly in immunocompromised patients. *Candida* species account for nearly 80% of nosocomial fungal infections, of which nearly 60% are due to *C. albicans* alone (Beck-Sague et al., 1993; Sternberg, 1994). Colonization by *C. albicans*, initiated by the adhesion of the fungal cell surface to the host tissue, occurs early in pathogenesis. Because of this, there has been interest in the characterization of the cell-wall proteins of *C. albicans* in general, and cell-surface proteins in specific, since the late 1960’s [for reviews see Cassone (1989), Odds (1988) and Reiss et al. (1992)].

Our work focuses on one particular characteristic of the cell surface that influences *C. albicans* pathogenesis, namely cell-surface hydrophobicity, and on its influence on the onset of disease. A current model states that cell-surface hydrophobicity in *C. albicans* is due to the presence of hydrophobic proteins at the cell surface (Hazen & Hazen, 1992). However, in order for the cells to be observed as hydrophobic their proteins must be exposed at the surface. This exposure occurs via changes in the conformation of surface fibres (Hazen & Hazen, 1992, 1993). We are therefore interested in labelling cell-surface groups to identify hydrophobic cell-wall proteins, and to identify fibril proteins or glycans potentially involved in hydrophobic protein exposure.

Previous labelling studies have identified differences in the cell-surface proteins of hydrophobic and hydrophilic cells, using iodination in the labelling procedure (Hazen & Hazen, 1992, 1993; Hazen et al., 1990). Although

Abbreviations: HRP, horseradish peroxidase; LYT, supernatant produced following treatment with lyticase; SDS extract, supernatant produced following SDS treatment of disrupted cell pellet; SUP, supernatant produced following cell disruption by glass beads.
radioisotopic labelling was useful for analytical purposes, subsequent purification schemes were complicated by the presence of radioactive material. The use of reactive biotinyl derivatives and avidin conjugates for the non-radioactive detection of proteins and glycoconjugates on blots has been described (Bayer & Wilchek, 1980; Bayer et al., 1987). These reagents were subsequently adapted for labelling the surface of intact mammalian (Hurley et al., 1985) and yeast cells (Alexandre et al., 2000; Casanova et al., 1992; Kandasamy et al., 2000; Marot-Leblond et al., 2000; Mrsa et al., 1997).

Our previous experience with avidin–biotin systems has involved using biotinylated antibodies and horseradish peroxidase (HRP)-conjugated avidin detectors in Western blots of the cell-wall proteins of *C. albicans*. We observed what appeared to be non-specific avidin binding in control lanes where the biotinylated antibody had been omitted. Although Duhamel & Whitehead (1990) have discussed possible contributors to this apparently non-specific binding and methods for blocking it, we were more interested in identifying which particular interactions were involved in the binding of avidin to the cell-wall proteins of *C. albicans*. Understanding these interactions might provide insights into the binding of *C. albicans* to host tissues. For example, avidin is known to possess sequences similar to cell-recognition domains (Alon et al., 1990). If these domains are involved in the observed non-biotin-mediated binding, then the non-specific avidin binding in our control lanes may represent integrin-like proteins, such as the one described by Gale et al. (1996). If hydrophobic interactions are driving non-specific binding, then it might be possible to identify additional proteins that are responsible for cell-surface hydrophobicity (Singleton et al., 2001). Thus, proteins that are both biotinylated during surface labelling and bound by avidin in the absence of a biotin tag might prove relevant in the adhesion of *C. albicans* to host tissues.

This work presents a more detailed study of the binding of avidin to the cell-wall proteins of *C. albicans*, and a characterization of the interactions between the two components. Our characterization of the interactions between avidin and the cell-wall proteins of *C. albicans* considered avidin–biotin components, structural components (such as cell recognition sequences and glycosylation) and physico-chemical forces (electrostatic and hydrophobic).

**METHODS**

**Reagents.** HRP–avidin (2.5 mg ml⁻¹), HRP–Z-avidin (1.25 mg ml⁻¹), HRP–streptavidin (1.25 mg ml⁻¹); mouse anti-biotin monoclonal antibody were purchased from Zymed Laboratories. HRP–ExtrAvidin (2.1 mg ml⁻¹) and gelatin (teleostean, no. G-7765) were purchased from Sigma. Based on information from the manufacturers, the molar ratio of HRP:avidin was 2:1. Sulfo-N-hydroxysulfosuccinimide-(biotinamido) hexanoate (sulfo-N-hydroxysulfosuccinimido-LC-biotin) was purchased from Pierce. All other chemicals were of reagent grade.

**Cultures and growth conditions.** *C. albicans* LGH1095 has been previously described (Hazén & Hazen, 1987). Cultures were maintained as frozen stocks and were prepared for use by three passages in yeast nitrogen base (Difco) buffered to pH 7.0 with sodium phosphate and supplemented with 2% (w/v) glucose, as previously described (Hazén & Hazen, 1987). Cultures were grown aerobically at 37°C. The final passage culture volume was 500 ml.

**Biotinylation.** Biotinylation was carried out according to Mrsa et al. (1997). Cells were harvested by centrifugation and washed three times with cold, sterile distilled water; the cell pellet was divided in two. Both pellets were washed twice with buffer A (50 mM KPO₄, pH 8.0). One pellet was suspended in 10 ml of buffer A (mock-treated); the other was suspended in 10 ml buffer A containing 10 mg sulfo-N-hydroxysulfosuccinimide-LC-biotin. Both the mock-treated and the treated sample were incubated on ice for 90 min, with occasional inversion. The samples were then centrifuged for 5 min at 1500 g. Both pellets were washed twice in buffer B (50 mM Tris, pH 7.5; 50 mM MgCl₂) followed by a final wash in buffer A.

**Release and isolation of cell-wall proteins.** Cell walls were prepared by glass-bead breakage, as modified from Hazen & Couter (1982). The cell pellets were re-suspended in 20 ml buffer A and 5 ml of the suspension was transferred to each of four 30 ml Corex tubes containing 10 g of glass beads (Glaspelren; 0.45–0.5 mm diameter; B. Braun Instruments). The tubes were subjected to alternating 15 s cycles of vortex mixing and ice incubation until cell breakage, determined microscopically, reached 80–90%. Corresponding supernatants were removed and combined. The glass beads were washed three times with buffer A and the washes were added to their respective supernatants. Cell walls were pelleted by centrifugation (10 min at 3000 g) and the supernatant (SUP) was removed and reserved.

Proteins were extracted from isolated cell walls by sequential treatment with hot SDS and β-1,3-glucanase. The pellets were suspended in SDS sample buffer (60 mM Tris, pH 6.8; 2%, w/v, SDS; 0.5%, v/v, β-mercaptoethanol) and heated at 95°C for 5 min. Samples were centrifuged (5 min at 2000 g) and the supernatant (SDS extract) was removed and resolubilized. The SDS-extracted pellets were washed five times in buffer B and finally re-suspended in 3 ml 50 mM NaPO₄, pH 7.4. A protease inhibitor cocktail, consisting of PMSF (0.2 mM final concentration; Roche), EDTA (1 mM; Sigma), leupeptin (1 µM; Sigma), pepstatin A (1 µM; Sigma) and 4-(2-aminoethyl)-benzenesulfonyl fluoride (1 mM; Roche), was added to the suspension. A β-1,3-glucanase was added at 250 U (ml digest)⁻¹ (Lyticase; Sigma, no. L-5263; 10000 U ml⁻¹ stock) and the digests were incubated on a rotary mixer at 37°C overnight. The digests were centrifuged at 3000 g for 10 min and the supernatant (LYT) was removed and reserved. The protein concentration of all supernatants was determined by using the Coomassie Plus-200 assay (Pierce).

**Electrophoresis and Western blotting.** The cell-wall proteins released in each fraction were separated by SDS-PAGE (12.5%, w/v, acrylamide; Laemmli, 1970). Typically, 10 µg total protein were loaded into each lane. Following electrophoresis, the proteins were transferred onto nitrocellulose (Osmonics) membranes using a Trans-Blot SD apparatus (Bio-Rad) run at a constant current (1.0 mA cm⁻²) for 1 h, in buffer containing 25 mM Tris, pH 8.3, 192 mM glycine and 20% (v/v) methanol (Towbin et al., 1979).

Membranes were blocked with TBST (50 mM Tris, pH 7.5; 150 mM NaCl; 0.1%, v/v, Triton X-100) containing 4%
RESULTS

Binding of avidin conjugates to cell-wall proteins of C. albicans

HRP–Z-avidin bound to various proteins extracted from the cell wall of C. albicans (Fig. 1). Although the signal in the mock-treated samples was weaker than that in the biotinylated samples, the bands were clearly evident. The most prominent bands in the SUP fraction from mock-treated cells were approximately 116, 77, 53, 43 and 31 kDa (Fig. 1). The most prominent bands in the SDS extracts were approximately 116, 77, 43 and 14 kDa. Weakly staining bands were also observed in the LYT extract (approx. 17 and <13.1 kDa, Fig. 1). Non-specific binding was also observed with samples from other yeast strains (C. albicans CAH and Saccharomyces cerevisiae X2180, not shown).

Replica blots were carried out and these were probed with HRP–streptavidin, HRP–ExtrAvidin and HRP–avidin. Similar binding profiles to those of the mock-treated SDS extracts were observed with all avidin conjugates tested (Fig. 2, TBST lanes). The binding profiles of the SUP and LYT extract fractions were also the same, regardless of the avidin conjugate used (not shown). Control experiments, in which the membranes were probed with unconjugated HRP or with the colorimetric reagents alone, were negative (not shown).

We tested the possibility that the observed binding was due to endogenous biotin, by probing blots of mock-treated cell-wall proteins with an anti-biotin antibody. A single intensely staining band (116 kDa, apparent molecular mass) was observed in the extract from mock-treated cells (Fig. 2, right panel, far right). A few weakly staining bands were also visible, particularly after prolonged development. These bands may represent proteolytic products of the larger band or may be due to non-specific antibody binding.

Experimental parameters affecting the observed non-specific binding

Along with TBST+BSA, BLOTTO and gelatin are commonly used to block unoccupied membrane sites. Blots blocked with gelatin were identical to those blocked with TBST+BSA. BLOTTO almost completely blocked the binding of Z-avidin, ExtrAvidin and avidin to the cell-wall proteins of C. albicans, even in biotinylated samples, but did not block the binding of streptavidin to proteins from mock-treated cells (Fig. 2). BLOTTO effectively blocked non-specific binding by the anti-biotin monoclonal antibody without significantly reducing binding to the biotinylated proteins (Fig. 2).

Blots of SDS extracts from both biotinylated and mock-treated cells (10 µg lane−1) were probed with increasing dilutions of Z-avidin. In most cases, increased dilution of the avidin conjugate resulted in loss of signal in mock-treated lanes while still allowing visible staining of the biotinylated proteins. However, several bands in the mock-treated samples (apparent molecular masses of 116, 33 and 27 kDa) were still visible at the highest Z-
avidin dilution (1:160000). The 116 kDa band maintained equivalent staining in treated and mock-treated samples, regardless of the Z-avidin dilution. The converse experiment was carried out by probing twofold serial dilutions of SDS extract (10–0-3125 µg) with a constant dilution (1:5000) of Z-avidin. Again, there was a diminution in the signal for both the biotinylated and mock-treated samples. At 1-25 µg of sample, most of the binding to unlabelled proteins was no longer apparent, whereas binding to biotinylated samples was still seen. However, even at 1-25 µg, a few of the non-specific bands were observed. In addition, the less intensely stained biotinylated bands began to be lost at the higher sample dilutions.

Characterization of non-specific binding

Several potential competitors were mixed, in parallel, with the avidin conjugates prior to probing of the blot. Binding by avidin, Z-avidin and ExtrAvidin to proteins from mock-treated cells was completely blocked by biotin down to a biotin concentration of 1 nM. Biotin inhibition of streptavidin binding was incomplete at 1 nM. Biotin also inhibited binding of the avidins to biotinylated proteins in a concentration-dependent manner (not shown). These results suggest that the site of binding to unlabelled proteins includes or is adjacent to the avidin–biotin binding site.

Binding by Z-avidin and ExtrAvidin was also completely inhibited by the addition of 1 M NaCl. NaCl had only a slight effect on the binding of avidin and streptavidin. KSCN (1 M), which acts as a chaotrope (Hatefi & Hanstein, 1969), inhibited the binding of Z-avidin and ExtrAvidin in a concentration-dependent manner (Fig. 3). There was also a slight concentration-dependent KSCN inhibition of avidin binding. Streptavidin binding was not inhibited by 1 M KSCN.

Cell recognition (RGD) sequences have been shown to inhibit the binding of C. albicans cells to surfaces and to host cells (Klotz & Smith, 1991; Negre et al., 1994). Furthermore, streptavidin was shown to possess a similar sequence, RYD, which mimics the RGD sequence of fibronectin (Alon et al., 1990). RGD (20, 20 and 0-2 µg ml⁻¹) and RGD-containing peptides from fibronectin (1358Fn, 50 µM) or from laminin (1343Ln, 50 µM) were tested as potential inhibitors of avidin binding to the cell-wall proteins of C. albicans. None of the three peptides had an effect on binding by the avidin conjugates at the concentrations tested.

Fig. 2. Representative comparison of, and effect of blocking agent on, the binding of the different avidin conjugates to SDS extracts of C. albicans from biotinylated (B) or mock-treated (M) cells. Membranes were blocked with BLOTTO or TBST › 4% BSA and then probed with HRP-Z-avidin, HRP-streptavidin or HRP-ExtrAvidin (left panel), and HRP-Avidin or an anti-biotin antibody (right panel). Bound complexes were detected with 3,3'-diaminobenzidine H₂O₂. Molecular mass markers are shown at the left of both images.

Fig. 3. Representative Western blot of KSCN inhibiting the binding of the different avidin conjugates to proteins from unlabelled cells. The first lane of each set of four contains no KSCN, the ‘wedge’ above the subsequent three lanes of each set represents the decreasing KSCN concentration. Bound avidin complexes were detected by 3,3'-diaminobenzidine H₂O₂. Molecular mass markers are shown at the left of the image.
DISCUSSION

Several groups have used biotin derivatives to label cell-surface proteins in yeasts (Alexandre et al., 2000; Casanova et al., 1992; Kandasamy et al., 2000; Marot-Leblond et al., 2000; Mrsà et al., 1997). Our results show that avidin can bind to the cell-wall proteins of *C. albicans* in the absence of a biotin tag. This non-biotin-mediated binding may complicate data interpretation by leading to unwarranted conclusions of surface exposure. Supplemental data from biotin-labelled *S. cerevisiae* showed non-specific binding to SDS extracts of cell-wall proteins (I. Hagen and W. Tanner, personal communication). The relative signal strength of non-specific to specific binding was comparable to the results presented in this study. Non-specific binding of streptavidin to human B cell and myeloid cell-surface proteins has also been previously reported (Cole et al., 1987). However, prior to this point, there have been no reports in the literature of non-specific binding by avidin to yeast cell-wall proteins or analyses of what mechanism is responsible for the interactions.

One study (Kandasamy et al., 2000) demonstrated another potential problem for biotin labelling. In this study, biotinylation was carried out in a Tris buffer, even though free amine species (e.g. Tris or glycine) will compete for the biotinylation reagent (Bayer & Wilchek, 1980). Kandasamy et al. (2000) performed subsequent experiments using Western, Southern and Northern blots which clearly demonstrated that the proteins under investigation were expressed and present in the cell wall of *C. albicans*. Due to the biotinylation conditions, however, the conclusion that these proteins are present on the cell surface remains unsupported.

Marot-Leblond et al. (2000) reported the extraction of cellular components, and the separation of these components by hydrophobic-interaction chromatography, from unlabelled and biotinylated cells. Since only the results of the binding of streptavidin to proteins from biotinylated cells were reported, the reader is left to infer that no streptavidin staining occurred with extracts from unlabelled cells. They did, however, report the results of two important controls. Cell-wall protein biotinylation affected neither the protein extraction efficiency nor the hydrophobic-interaction chromatography profile (confirmed in our laboratory, unpublished results).

Casanova et al. (1992) showed that the biotinylation reagent does not cross the membrane into the cytoplasm. This confirmed for yeasts what had been seen previously for bovine leukocytes (Hurley et al., 1985). Thus, the presence of a signal in the bead-break supernatant (Fig. 1) was somewhat unexpected. As an alternative sample preparation technique, we prepared sphaeroplasts following biotinylation and compared the cell-wall fraction with the sphaeroplast lysate (see Casanova et al., 1992). Only a few distinct bands were observed in the lysate, confirming the observation made by Casanova et al. (1992). If the biotinylation reagent was crossing the cellular membrane and labelling the cytoplasmic proteins, a continuous smear of signal (the length of the blot) would be expected. Therefore, the observed signal in the SUP fraction is likely to be due to proteins loosely associated with the cell-wall matrix that are released by the breaking action itself (Pastor et al., 1984).

A simple overabundance of avidin does not explain the observed binding, since the concentration of the avidin conjugates was at or was below that suggested by the manufacturer and was similar to those found in the literature. The anti-biotin monoclonal antibody gave a single intense band in mock-treated cell samples (approx. 116 kDa, Fig. 2, right panel, far right). Because of this band, and other bands that were very faint, we must consider the possibility that some of the avidin conjugate binding to mock-treated cell-wall proteins is due to endogenous biotin. If avidin binding to unlabelled proteins is indeed due solely to endogenous biotin, showing a comparison of labelled and unlabelled proteins is even more of an imperative. However, based on the number of bands appearing in the lanes probed with the avidin conjugates, we think it likely that the observed avidin binding is due to interactions other than avidin–biotin.

Binding does appear to occur near the biotin site, because biotin inhibited the binding of avidin in a concentration-dependent manner (see Results). In addition, BLOTTO, used as a blocking agent, almost completely inhibited avidin binding to proteins from both biotinylated and mock-treated cells (Fig. 2). The concentration of biotin (from the dry milk) in BLOTTO (0.04–0.10 nmol ml⁻¹; Jensen, 1995) is two- to fourfold higher than the concentration of biotin-binding sites (0.005 nmol avidin ml⁻¹ gives 0.02 nmol biotin sites ml⁻¹). This concentration is likely to be sufficient to approach saturation of the biotin-binding sites. The reason BLOTTO did not eliminate binding by streptavidin or the monoclonal antibody (Fig. 2) is less clear. The affinity of streptavidin (4 × 10⁻¹⁴ M) is 67-fold lower than that of avidin (6 × 10⁻¹⁵ M). This lower affinity suggests that, during the probing step, streptavidin is more likely than the other avidins to release the biotin into solution and to be available for binding to the cell-wall proteins.

As noted above, avidin has been shown to possess affinity for other ligands. Avidin contains the RYD sequence that mimics the RGD sequence in the cell-wall recognition site of extracellular matrix proteins (Alon et al., 1990). The binding mediated by RYD is independent of the avidin–biotin interaction and thus represents another binding site. Studies by Gale et al. (1996) demonstrated that *C. albicans* possesses integrin-like proteins. Such integrin–extracellular matrix interactions might then explain some of the non-specific binding, although the specific extracellular matrix peptides tested here did not inhibit this binding activity (see Results). In addition, a lectin-like activity of the cell-wall proteins of *C. albicans* appears to be ruled out, since glycosylated (avidin) and unglycosylated (Z-avidin and streptavidin) forms bound in the absence of a biotin tag.
Both NaCl and KCSN, which interfere with electrostatic and hydrophobic interactions, respectively, were able to inhibit the binding of avidin, Z-avidin and ExtrAvidin to proteins from mock-treated cells. NaCl had a smaller effect on streptavidin and avidin binding to these proteins. KSCN did not inhibit binding by streptavidin, but was able to inhibit avidin binding at higher concentrations.

Because the biotin-labelling reagent is not membrane permeable, it is not radioactive and labels under gentle and convenient conditions, it is a powerful tool for determining proteins at the fungal cell surface. It is these proteins that make and maintain first contact with the host tissues, making them important to the understanding of colonization and pathogenesis. Our results emphasize the necessity for an untreated control to aid in data interpretation. Proteins that produce Western blot signals in extracts from both labelled and unlabelled cells require additional tests (such as immunofluorescence microscopy) before they can be shown to be exposed on the cell surface. Our results further suggest that both the hydrophobic and electrostatic characteristics of the avidin conjugates seem to be involved in non-specific binding, except in the case of streptavidin where only charge seems to play a role. Currently, we are extending our studies to carry out electrophoretic protein separation in two dimensions to more clearly identify those proteins which can be both labelled with biotin and bind to avidin (e.g. ExtrAvidin), i.e. proteins which may be responsible for cell-surface hydrophobicity. We will then apply these techniques to examine the differences in the expression of cell-surface proteins between hydrophobic and hydrophilic cells.

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