A cell surface/plasma membrane antigen of *Candida albicans*

REN-KAI LI and JIM E. CUTLER*

Department of Microbiology, Montana State University, Bozeman, MT 59717, USA

(Received 7 August 1990; revised 26 October 1990; accepted 14 November 1990)

Antibody from BALB/cByJ mice immunized against a membranous fraction of *Candida albicans* agglutinated whole cells as well as the membranous fraction. Hybridoma techniques were used to isolate an IgM monoclonal antibody (mAb) designated 10G which agglutinated whole cells and reacted with the subcellular fraction. Yeast cells of 15 additional *C. albicans* strains and isolates of *C. stellatoidea*, *C. tropicalis*, *C. intermedia* and *C. lusitaniae* were also agglutinated by mAb 10G. The antigen was not detected on other fungi, including *Candida krusei*, *C. utilis*, *Cryptococcus neoformans*, *Cr. albida*, *Torulopsis glabrata*, *Rhodotorula* spp. and *Saccharomyces cerevisiae*. To determine the cellular location of the epitope to which mAb 10G is specific, freeze-substitution was compared with traditional chemical fixation methods in preparation of samples for immunocolloidal gold electron microscopy (IEM). With both fixation procedures, the antigen recognized by mAb 10G was found randomly and densely concentrated on the plasma membrane on exponential-phase yeast-form cells and had a patchy distribution on the cell wall surface. Association of the antigen with the plasma membrane was confirmed by IEM of isolated membranes. On developing hyphal cells, antigen appeared first on the plasma membrane and later on the cell wall surface. Treatment of yeast cells with β-mercaptoethanol and Zymolyase before fixation removed the antigen from the surface but left the cytoplasmic antigen undisturbed. Treatment of yeast cells or solubilized antigen with heat or proteolytic enzymes (trypsin, Pronase B, proteinase K) did not remove or destroy the antigen, suggesting a non-protein nature of the epitope.

**Introduction**

The cell wall and plasma membrane of *Candida albicans* constitute the barrier between its cytoplasm and the external environment. These layers are intimately involved in the shape of the cell, antigenicity and secretion of hydrolytic enzymes. The cell wall of *C. albicans* has thus been a major area of investigation on candidal-host interactions mainly because of the role in adherence to its host (Levitz *et al.*, 1987; Calderone *et al.*, 1984; Maisch & Calderone, 1981), and interactions with host immune systems (Calderone *et al.*, 1988; Kuruganti *et al.*, 1988). The plasma membrane, on the other hand, is of particular importance in homeostatic control and susceptibility to antifungal agents (Shepherd, 1987).

The mechanism of transportation and secretion of *C. albicans* wall precursors from the cytoplasm to their destined wall location is unclear (Andaluz *et al.*, 1986; Marriott, 1977; Rajasingham & Cawson, 1984; Elorza *et al.*, 1983). Although the biosynthesis of the wall of *C. albicans* is not fully understood, protoplast regeneration studies, electron microscopy and chemical determinations have revealed the chemical composition of the wall. These observations have confirmed the importance of the plasma membrane in the synthesis of cell wall layers (Duran *et al.*, 1975; Tronchin *et al.*, 1984; Marriott, 1977; Djaczenko & Cassone, 1971; Korn & Northcote, 1960; Takamiya *et al.*, 1985; Elorza *et al.*, 1983). Further understanding of the role of the plasma membrane in cell wall biosynthesis requires detailed studies of critical cell wall epitopes which should be associated with the plasma membrane during various stages of wall synthesis. In spite of profound chemical, structural and functional differences between the cell membrane and wall of *C. albicans*, it has been suggested that certain epitopes may in fact be shared at various times in the cell cycle (Hopwood *et al.*, 1986; Ollert & Calderone, 1990).

**Abbreviations:** IFA, indirect immunofluorescence assay; IEM, immunocolloidal gold electron microscopy; PBS, phosphate-buffered saline; mAb, monoclonal antibody; GYE, glucose yeast extract peptone; BSA, bovine serum albumin.
In the work described here, we determined the presence of an epitope found on the plasma membrane and also on the cell wall surface of *C. albicans* blastoconidia and germination tubes. The antigen distribution was observed at the ultrastructural level in cells fixed by the disparate techniques of traditional blastoconidia and germination tubes. The antigen and also on the cell wall surface of *C. albicans* was assayed by the Summerson photoelectric colorimeter, no. 66 filter) and harvested in spheroplasts, yeasts were grown in GYEP for 22-24 h at 37 °C with and in all experiments unless indicated otherwise. This strain and other and grown on modified Sabouraud dextrose agar (Difco) slants. For electron microscopy (IEM) of intact yeast cells and preparation of media were used for induction of germination. In most experiments, Torulopsis glabrata, Rhodotorula Saccharomyces cerevisiae and grown on modified Sabouraud dextrose agar (Difco) slants. For most experiments, cells were transferred from slants and grown in glucose (2%, w/v)/yeast extract (0-3%, w/v)/peptone (1%, w/v) broth (GYEP) at 37 °C under constant aeration by rotation of flasks at 180 r.p.m. (Gyrotryor incubator; New Brunswick Scientific). For immuno-electron microscopy (IEM) of intact yeast cells and preparation of spheroplasts, yeasts were grown in GYEP for 22-24 h at 37 °C with aeration, reincubated into fresh GYEP to a Klett value of 100 (Klett-Summerson photoelectric colorimeter, no. 66 filter) and harvested in the exponential phase (Klett value 400). In some experiments, stationary-phase cells were harvested (Klett value 550). Two different media were used for induction of germination. In most experiments, hyphae were induced in a synthetic medium originally described by Lee et al. (1975) as modified by Sundstrom & Kenny (1984) and referred to as modified Lee medium. Cells were grown in the modified medium without bovine serum albumin (BSA) for 48 h at 25 °C to obtain yeast-form cells and transferred to warmed medium at 37 °C containing 1% (w/v) BSA (ImmunoBiologicals) to promote germination. Cultures were incubated at 37 °C and aerated by rotation at 180 r.p.m. An alternative medium was used in the IEM studies of germinating cells in which yeast-form cells were grown to stationary phase in GYEP at 37 °C, inoculated into GM-2 medium (Hazen & Cutler, 1983) at 2 × 10⁶ cells ml⁻¹ and incubated at 37 °C under aeration for hypal induction. Germination was assayed by direct light microscopy and the cells were used if germination exceeded 90% by criteria defined previously (Hazen & Cutler, 1979).

*Spheroplasts and subcellular fractions.* Spheroplasts were prepared following procedures described by others (Hudsprth et al., 1980; Wills et al., 1984) with slight modifications. Exponential-phase yeast cells were washed in 0-1 M-EDTA (pH 7-5, disodium salt, Sigma) suspended in 0-1 M-EDTA [2 ml (g wet weight)⁻¹] with 0-3 M-β-mercaptoethanol (pH 9-0) and incubated at 24 °C for 15 min. Cells were pelleted by centrifugation at 2000 g for 10 min and suspended to the same concentration in 1-0 M-sorbitol/0-1 M-EDTA (pH 7-5). Zymolyase 20T (ICN) was added at 80 units per g wet weight of yeast cells and incubated at 37 °C for 60 min with frequent agitation. In some preparations for IEM, the β-mercaptoethanol-treated cells were incubated with Zymolyase for only 10 min to remove surface layers. The efficiency of spheroplast formation was determined by counting (haemocytometer) intact cells before and after lysis in 1% (w/v) SDS (Sigma) as described by Glee et al. (1987). Spheroplasts were washed in 1 M-sorbitol/0-1 M-EDTA and either fixed for electron microscopy or processed as described below for mitochondrial fractions.

Subcellular fractions were prepared as described by Wills et al. (1984) for isolation of mitochondria. Spheroplasts, induced by treatment of yeast cells with Zymolyase for 60 min as above, were washed and suspended in 0-25 M-sucrose (Sigma), 1-0 mM-EDTA and 5 mM-Tris/HCl (Sigma) pH 7-5, and homogenized in a Dounce tissue homogenizer at 4 °C. After removal of cell debris and intact cells by centrifugation at 3000 g for 10 min, the supernatant material was centrifuged at 15000 g for 20 min to pellet the mitochondrial fraction. Fractions enriched for mitochondria obtained from continuous sucrose gradients (25-50%, w/v) run at 41000 g for 90 min (Wills et al., 1984) (SW 41Ti rotor, Beckman) were used for immunization of animals. This fraction contained mitochondria and other membranous vesicles as determined by electron microscopy.

**Plasma membrane fraction.** Plasma membrane fractions were obtained from yeast-form cells as described by Fuhrmann et al. (1974). Stationary-phase yeast cells were washed and suspended in an osmotic stabilizer (0-4 M-KCl, 20 mM-triethanolamine, Sigma, pH 7-0). Cells were broken in a heavy-walled centrifuge tube (Corex tubes, Corning Glass Works) by mixing with 0-45-0-5 mm glass beads on a vortex mixer (Hazen & Cutler, 1982). The suspension was spun at 2000 g for 5 min to remove cell debris and the supernatant material was centrifuged at 5000 g for 10 min to pellet membranes. The crude membrane pellet was washed three times in the osmotic stabilizer, fractionated by sucrose density gradients (10-60%) at 23500 g for 70 min and the resultant bands were collected.

**Hybridoma and monoclonal antibody (mAb).** Preliminary observations showed that mitochondria-rich fractions induced mice to produce antibodies which reacted with the mitochondrial fraction and with the cell surface of *C. albicans*. Thus, fractions enriched for mitochondria and other membranous vesicles, and hence referred to as the membranous fraction, were used for six weekly intravenous 0-1 ml injections into BALB/cByJ mice. Lowry protein concentration was 4-6 mg ml⁻¹. Three days after the last booster injection, mouse sera were checked for agglutinins against intact *C. albicans* blastoconidia, and splenocytes from agglutinin-positive animals were prepared for cell fusion with a non-secreting myeloma cell line. Detection of antibody produced by hybrids, cloning by limiting dilution and production of ascites in mice was done as previously described (Brawner & Cutler, 1984).

**Agglutination and ELISA.** Qualitative agglutination tests were done on heat-killed yeast cells as described before (Brawner & Cutler, 1984). ELISA tests (from either Bethesda Research Laboratories or Boehringer Mannheim Biochemicals) were used for detection and titrating of mAb in either culture fluid or ascites. Microtitre plates (96-well; Corning) were coated with 150 μl of yeast cell suspension (2 × 10⁶ cells ml⁻¹) in modified Lee medium at 37 °C for germination. After 4 h incubation, the wells, now coated with germinating cells, were washed four times with 0-01 M-phosphate buffered saline (PBS, pH 7-2), and 100 μl of appropriately diluted test fluid or negative control material were added and incubated at room temperature for 1 h. ELISA detection of mouse mAb was done by following the manufacturer's instructions. In addition to negative controls as indicated above, fresh RPMI medium was used for detecting inappropriate binding of secondary antibody to the cells. An IgM mAb described previously as C6 was used as a positive control and identical results were obtained as before (Brawner & Cutler, 1986).

**Immunocolloidal gold electron microscopy (IEM).** This technique was used to visualize the antigen expression on intact cells, spheroplasts and membranous fractions. Two procedures for fixation of samples were adopted for IEM. Samples were either chemically fixed in glutaralde-
hyde (0.5%, v/v)/osmium tetroxide (1%, w/v), and dehydrated through a graded ethanol series before embedding (Brawner & Cutler, 1987), or freeze-fixed and substituted with anhydrous acetone as described by others (Hoch & Howard, 1980, 1981; Hoch, 1986). In the latter procedure, a thin layer of yeast cells was very rapidly frozen at -192°C in a mixture of liquid propane and ethane, and the frozen samples were transferred to anhydrous acetone substitution fluid containing 2% osmium tetroxide and 0.05% uranyl acetate at -80°C. Samples were dehydrated and fixed at -80°C for 2-3 d, and gradually brought up to room temperature (22-24°C) (2 h at -20°C, 2 h at 4°C and then room temperature). This freeze-substitution method was used in fixing yeast cells and spheroplasts. Samples were washed three times in anhydrous acetone and embedded in Spurr's resin (Spurr, 1969) for thin sectioning. Thin sections on nickel grids were reacted with 10-times concentrated culture fluid from a cloned hybridoma cell line, blocked with 3% BSA, reacted with goat anti-mouse secondary antibody conjugated with colloidal gold (10 nm) (Janssen Biotech), and finally contrast-stained with lead citrate and uranyl acetate.

Indirect immunofluorescence assay (IFA). The assay was modified from that described by Ruff et al. (1976). Mycelium-form cells induced in modified Lee medium were harvested after different incubation times and fixed in 3% (v/v) formaldehyde at 4°C for 12 h. Cells were washed and suspended in PBS, cyto centrifuged onto a glass slide and treated for 5 min at 4°C in 5% (v/v) acetic acid in 95% (v/v) ethanol. Slides were washed in PBS and air dried before adding antibody. Undiluted ascites fluid containing the mAb or 10-times concentrated culture fluid from the chosen hybridoma clone was added to the fixed cells on the slides and incubated in a moist chamber for 15 min at 37°C. Slides were washed by dipping in PBS several times at room temperature and air dried before adding fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse immunoglobulin (Cappel) (diluted as suggested by the manufacturer). After incubation for 15 min at room temperature and washing in PBS, cells were examined with a fluorescence microscope (Nikon Episcopic-fluorescence microscope, type 104, equipped with a high-pressure mercury lamp 100W/2). The assay was modified by Ouchterlony double diffusion in which goat antibodies specific for mouse light or heavy chain immunoglobulin determinants were used (Sigma).

Results
mAb preparation and characterization

Four stable hybridoma cell lines were obtained which reacted strongly against C. albicans strain 1 as determined by ELISA. One which was chosen for further investigation was cloned by limiting dilution, recloned and designated mAb 10G. The ascites induced in BALB/cByJ mice by the cloned hybrid gave an ELISA titre of 2400. The mAb 10G belongs to the IgM class with kappa light chains as determined by specific antisera (not shown).

Expression of antigen on yeast cells

Surface expression of the epitope recognized by mAb 10G (i.e. Ag 10G) was detected by agglutination and IFA of whole cells of C. albicans strain 1. Antigen distribution at the ultrastructural level was further investigated by IEM in cells fixed by two different procedures. Similar results were obtained from cells fixed by either glutaraldehyde or freeze-substitution. Ag 10G was usually found randomly distributed on the plasma membrane but had a patchy distribution on the wall surface (Fig. 1a, b). The ultrastructural preservation of plasma membranes, nuclei, walls and most cytoplasmic contents was improved by freeze-substitution techniques compared to traditional chemical fixation.

To investigate the strain and species specificity of expression of the antigen, other isolates of C. albicans as well as other Candida spp. and other fungal species were examined for surface expression of the antigen by the agglutination test. The following organisms gave positive agglutination reactions: all 15 C. albicans clinical isolates and one isolate each of C. stellatoidea, C.
tropicalis, C. lusitaniae and C. intermedia. No antigen was detectable by agglutination of other fungi, which included Candida utilis, C. krusei, Cryptococcus neoformans, Cr. albidos, Torulopsis glabrata, Rhodotorula spp. and Saccharomyces cerevisiae. Two of the agglutination-positive isolates of C. albicans (strains 207 and 325) were examined by IEM and a similar pattern of antigen distribution was observed (data not shown).

Expression of antigen on germinating cells

Surface expression of antigen on hyphae of C. albicans was examined by IFA. Immunofluorescence of germinating cells was typically dense on the mother yeast cells throughout the incubation time. Antigen expression on the hyphal surface was not detectable for the first 2 h of incubation but became evident as germination proceeded (Fig. 2). These observations were subsequently confirmed by IEM.

Stationary-phase C. albicans cells grown in GYEP were transferred to GM-2 medium and allowed to germinate for up to 8 h. After various incubation times, germinating cells were fixed with glutaraldehyde and prepared for IEM as described above. At time 0, stationary-phase cells had greater surface expression of antigen than exponential-phase cells (compare Fig. 3a to Fig. 1). Throughout the entire incubation time allowed for hyphal development, antigen expression on parent cells was more dense on the cell wall surface than on the plasma membrane. The development of antigen on germ tubes was first observed after 2 h incubation. Unlike the distribution in parent cells, the antigen expressed on germ tubes was primarily in the subsurface area near or on the plasma membrane (Fig. 3b). At 6 h, antigen became obvious on the cell surface and remained dense in the subsurface area (data not shown). Control experiments in which cells were treated with culture fluid of the non-secreting cell line did not show colloidal gold above background level (<10 particles per cell) (not shown).

Fig. 1. Distribution of 10G epitope in exponential-phase cells fixed chemically at room temperature as compared to freeze-substituted cells. (a) Cells were fixed with glutaraldehyde and osmium tetroxide. Antigen was equally distributed near or on the plasma membrane (arrow) but unevenly on the cell wall surface (arrowhead). (b) Cells fixed by freeze-substitution show similar patterns of antigen distribution and better detail of membrane association of the antigen (arrow). Bars, 1 μm.
Expression of antigen on spheroplasts

Exponential-phase yeast cells were treated with β-mercaptoethanol and subjected to Zymolyase digestion for either 10 or 60 min. The efficiency of spheroplast formation after Zymolyase digestion for 60 min was consistently over 90%. Samples were freeze-substituted with acetone and prepared for IEM. In cells treated with β-mercaptoethanol and Zymolyase for 10 min, antigen was removed from the surface but remained undisturbed on the plasma membrane. The fibrillar outer cell wall layer was also removed by this treatment (Fig. 4a). Organelles which had been released from lysed cells were occasionally observed. After 60 min incubation in Zymolyase, no cell wall was observed, yet antigen was still associated with the plasma membrane (Fig. 4b). The ultrastructural preservation of spheroplast organelles was generally improved compared with whole yeast cells.

Expression of antigen on subcellular fractions

To further confirm the association of the antigen with the plasma membrane, lysed spheroplasts and plasma membrane fractions were examined for the presence of antigen. Lysis of spheroplasts was induced by suspension in 0.5% glutaraldehyde without 1 M-sorbitol. In glutaraldehyde-fixed samples of spheroplasts, antigen was mostly found in vesicle structures but not on fragmented membranes (Fig. 5a). The vesicles apparently formed spontaneously during glutaraldehyde lysis of spheroplasts. The antigen was also not associated with the plasma membrane in membrane fractions (Fig. 5b). When lysed spheroplasts were freeze-fixed and substituted in acetone, the antigen was observed evenly distributed throughout the membranes (Fig. 5c).

Mitochondria-enriched fractions were also investigated using IEM in an attempt to explain their immunogenic nature. Fractions of mitochondria and membranes which co-purified with mitochondria were observed, but antigen was detectable only within single and multilamellar membrane vesicles (Fig. 5d).

Characterization of 10G antigen

The phenol extract of C. albicans strain 1 was subjected to chemical and physical treatment and subsequently tested for antigen reactivity against mAb 10G in Ouchterlony double diffusion. Precipitin bands of variously treated
Fig. 3. Distribution of 10G epitope during germ tube development as determined by IEM. Stationary-phase yeasts were placed into germination conditions, harvested at various time intervals and fixed with glutaraldehyde and osmium tetroxide. (a) At time 0, epitope expression was especially heavy on the cell wall surface. (b) After 2 h germination, antigen remained densely distributed on the mother cell wall surface, whereas on germ tubes it was primarily expressed in the cytoplasm near the cell membrane (arrow). Bars, 1 μm.

phenol extract showed identity with untreated material. The antigen reactivity was not altered by treatment with heat at 96 °C for 10 min or with proteolytic enzymes (Pronase B, trypsin, and proteinase K).

Yeast cells grown in GYEP broth were also treated under the same conditions and screened for surface antigen activities by agglutination and IFA. Treated and untreated yeast cells showed the same degree of agglutination and fluorescence. Immunoelectrophoresis of the phenol extract produced a smeared precipitin band from the well of antigen toward the anode at all pH values tested (not shown).

Discussion

Among the many approaches to understanding cell wall structure and development of C. albicans, mAbs allow investigators to follow the expression of a particular determinant. In the present study, an IgM mAb, with kappa light chains, and designated 10G, was isolated which agglutinates C. albicans yeast cells. In addition to the cell surface location of the antigen to which mAb 10G is specific, the appearance of the immunodeterminant also on the plasma membrane was rigorously documented. These findings are not unique to a single strain of C. albicans because the same pattern of expression was found on other isolates of C. albicans. Our results indicate that Ag 10G is a common antigen in C. albicans and is also expressed in certain other Candida species (C. stellatoidea, C. intermedia, C. tropicalis and C. lusitaniae). The antigen is not expressed by C. utilis and C. krusei. In addition, the antigen was not detectable in other yeast genera. However, the absence of reactive epitope on the cell surface of C. utilis, C. krusei and other fungal genera as demonstrated by the agglutination test did not exclude the possibility of intracellular expression.

It has been suggested by other investigators, who used only immunofluorescence microscopy, that the same polysaccharide epitope recognized by a mAb was located on the cell surface and on the plasma membrane of C. albicans (Hopwood et al., 1986; Ollert & Calderone,
Another group (Takamiya et al., 1985), who used ferritin-labelled antibody, localized a mannan antigen on the exterior cell wall surface as well as in the cytoplasm near the cytoplasmic membrane. The latter location was regarded as the site of mannan synthesis and it was suggested that the mannan antigens were transported from the cytoplasm via channel-like structures in the cell wall to the final deposits in the exterior cell wall layer. Similar speculation has been reported by others on cellular excretion of glycoproteins through the cell wall of C. albicans (Poulain et al., 1989). These workers intimated that antigen emergence at the cell surface may correspond to patches of the antigenic material in the fuzzy coat of the cell surface. Although channel-like structures were not observed in our studies, it may be hypothesized that the patchy distribution of the 10G antigen on the cell wall surface represents the site of antigen transportation. However, knowledge of the chemistry of the antigens at both locations is needed to determine whether the plasma membrane antigen is a cell wall precursor or is an unrelated, cross-reactive substance. Treatment of the cells with β-mercaptoethanol and Zymolyase selectively releases the antigen from the cell wall and will allow differential purification of the antigen from both locations for subsequent chemical comparisons.

Information on yeast wall synthesis comes primarily from studies on Saccharomyces cerevisiae (Tsai et al., 1990).
Fig. 5. Antigen 10G in membrane fractions. Lysed spheroplasts (a) and membrane fractions (b) fixed by glutaraldehyde. Antigen was not found on membrane fragments but was associated with vesicles (arrow). (c) Lysed spheroplasts fixed by freeze-substitution showing antigen association with the plasma membranes (arrow). (d) Mitochondria-enriched fractions in which variously sized vesicles co-migrated and trapped the antigen during spontaneous vesicle formation. Bars, 1 μm.

In C. albicans mannosylation reactions might occur at the plasma membrane level (Marriott, 1977), and enzymes associated with the cell wall matrix may modify the wall during growth and hyphal transformation (King et al., 1980; Barnes et al., 1983; Lehrer et al., 1986). In our observations, the early appearance of antigen on the membrane of germ tubes and later development on the surface suggests a growth-dependent expression of the antigen during hyphal development. In addition, the dense expression of the antigen corresponded with the growing apical region. We also noted that in yeast-form
growth in GYEP, antigen expression was predominantly on the plasma membrane in exponential-phase cells and primarily on the cell surface of stationary-phase cells. Expression of cell-surface antigenic determinants may vary as a function of growth phase or morphological states (Brown & Chaffin, 1981; Smail & Jones, 1984; Sundström & Kenny, 1984; Tronchin et al., 1984; Brawner & Cutler, 1986) and our observations suggest that specific sites of antigen location may be related to structural modifications and/or cellular transport of a cell wall-component during yeast growth and early stages of germination.

Ascribing antigen location at the ultrastructural level to specific cellular sites by results obtained from a single fixation procedure is open to criticism. Conventional chemical fixation using glutaraldehyde followed by fixation procedure is open to criticism. Conventional chemical fixation using glutaraldehyde followed by osmium tetroxide may induce artifactual enlargement of dolipore septa in various basidiomycetes (Hoch & Howard, 1981). Using freeze-substitution fixation, we obtained essentially identical patterns of antigen distribution as with glutaraldehyde fixation, which gives strong support for the cell wall/cell membrane locations of antigen 10G. In addition, freeze-substitution gave apparently better preservation of most of the intracellular organelles, plasma membranes and the fluffy outer layer of the cell wall of C. albicans. One limitation of the method was that only a low percentage of intact yeast cells showed acceptable preservation. The majority of spheroplasts, however, were well preserved. Hence, total or partial loss of cell wall layers may allow faster freezing and better penetration of embedding agents.

Comparison between the two fixation procedures on lysed spheroplasts suggests that although the antigen is associated with the plasma membrane, it is not well anchored. The presence of the antigen on plasma membranes fixed by freeze-substitution but not on those fixed by traditional means implies that the antigen becomes dissociated from the membrane during the many washing steps in the glutaraldehyde fixation procedure. A loose association between the antigen and the plasma membrane may explain the absence of antigen on membrane fractions (Fig. 5).

During preparation of mitochondria-enriched fractions, membrane vesicles spontaneously form (data not shown) and can trap antigen released from the cell membrane, as demonstrated by electron microscopy. Spontaneous vesicle formation and non-specific trapping of substances by the vesicles has also been observed by others during investigations on S. cerevisiae (Christensen & Cirillo, 1972; Fuhrmann et al., 1974). In addition to explaining the specific immunogenicity of mitochondria-enriched fractions in our studies, these results indicate that vesicles may be artifactual created during cellular fractionation procedures. Furthermore, such vesicles may erroneously be thought to be associated with various cytoplasmic substances which were, in fact, passively trapped during vesicle formation.

The antigenic reactivity of either phenol extract or intact yeast cells was not altered after treatment with heat or proteolytic enzymes. These data, together with the behaviour of the phenol-extracted antigen in immunoelectrophoresis, indicate that antigen 10G is not a protein. Experiments to isolate and purify the epitope are now in progress, which will allow chemical comparisons between the cell-surface antigen and that of the plasma membrane.

We thank Pati Glee for assistance in preparation of mitochondria-enriched fractions, Diane Brawner for advice on hybridoma techniques and Susan Zaske for electron microscopic technical assistance.

This work was supported by a Public Health Service grant AI24912 from the National Institutes of Health.

References


