Effect of Ethanol on Activity of the Plasma-membrane ATPase in, and Accumulation of Glycine by, *Saccharomyces cerevisiae*

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The pH optimum of the ATPase activity in plasma membranes from *Saccharomyces cerevisiae* NCYC 431 from 8 h cultures was around 6.5 and that in membranes from organisms from 16 h cultures near 6.0. The $K_m[ATP]$ of the enzyme was virtually unaffected by the age of the culture from which organisms were harvested, although the $V_{max}$ of the enzyme in membranes from organisms from 8 h cultures was higher than that for organisms from 16 h cultures. Ethanol non-competitively inhibited ATPase activity in membranes, although the inhibition constant for the enzyme from organisms from 8 h cultures was lower than that from organisms from 16 h cultures. Glycine accumulation by the general amino acid permease was non-competitively inhibited by ethanol. Inhibition constants were virtually the same for glycine uptake by de-energized organisms from 8 h and 16 h cultures, but under energized conditions the value was greater for organisms from 16 h rather than 8 h cultures. The data indicate that inhibition of plasma-membrane ATPase activity by ethanol could account, at least in part, for inhibition of glycine accumulation by ethanol.

INTRODUCTION

Three distinct ATPase enzymes are synthesized by *Saccharomyces cerevisiae*. In addition to the plasma-membrane enzyme, both mitochondrial (Schatz et al., 1967) and vacuolar membranes (van der Wilden & Matile, 1978) possess ATPase activity. The properties of the plasma-membrane ATPase of *S. cerevisiae* have been well documented (Goffeau & Slayman, 1981). The enzyme has a relatively low pH optimum of 5.5–6.5 (Willsky, 1979; Borst-Pauwels & Peters, 1981), an absolute requirement for Mg$^{2+}$ (Ahlers et al., 1978; Willsky, 1979) and a high substrate specificity (Serrano, 1978; Peters & Borst-Pauwels, 1979). In addition, it shows differential sensitivity to a number of inhibitors. Oligomycin, an inhibitor of the mitochondrial ATPase (Serrano, 1978), has little effect on the plasma-membrane ATPase, whereas orthovanadate (Willsky, 1979) and diethylstilboestrol (Serrano, 1980), which do not inhibit the mitochondrial ATPase, are potent inhibitors of the plasma-membrane enzyme. DCCD inhibits all three ATPases in *S. cerevisiae*, although the mitochondrial and vacuolar enzymes are approximately ten times more sensitive to this inhibitor than the plasma-membrane ATPase (Serrano, 1978; Okorokov et al., 1982). The plasma-membrane ATPase in *S. cerevisiae* functions as an electrogenic proton pump (Malpartida & Serrano, 1981; Dufour et al., 1982) coupling ATP hydrolysis to translocation of protons across the membrane, resulting in the establishment of a transmembrane proton electrochemical gradient ($\Delta p$).

The $\Delta p$ has been shown to energize active transport of solutes by *S. cerevisiae* (Borst-Pauwels, 1981; Eddy, 1982). When grown in the presence of ammonium ions this yeast synthesizes at least

Abbreviations: DCCD, *N*,*N*'−dicyclohexylcarbodiimide; GAP, general amino acid permease; $\Delta p$, proton-motive force.

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10 amino acid transport systems, each of which is specific for one or a small number of L-amino acids (Eddy, 1982). Very little has been reported on transport of glycine by a specific system, and it is generally assumed that transport of this amino acid is effected mainly by the general amino acid permease (GAP), a high-velocity broad-specificity system capable of transporting L- and D-amino acids (Rytka, 1975) and subject to nitrogen catabolite repression (Grenson et al., 1970). Two kinetically distinct systems for glycine accumulation by \textit{S. cerevisiae}, a high-affinity system which was assumed to be the GAP and a low-affinity system, have been described (Eddy et al., 1970; Ballarin-Denti et al., 1984). Accumulation of glycine by the GAP has been extensively studied. This amino acid can accumulate in \textit{S. cerevisiae} to a concentration as high as 1400 times that outside the organism (Eddy et al., 1970), a proportion of this being accumulated in the vacuole (Indge et al., 1977). Two or more protons are transported intracellularly with each glycine molecule (Seaston et al., 1973), although under some conditions only one extracellular proton-binding site dissociates (Ballarin-Denti et al., 1984).

Inhibition of growth of \textit{S. cerevisiae} by ethanol is thought to be caused, at least in part, by interaction of ethanol with the plasma membrane (Ingram & Buttke, 1984). This interaction probably accounts for the non-competitive inhibition, by ethanol, of solute accumulation by this organism (Thomas & Rose, 1979; Leão & van Uden, 1982, 1983, 1984; Loureiro-Dias & Peinado, 1982). Since most transport processes are active and require a \( \Delta p \) (Borst-Pauwels, 1981; Serrano, 1977), the effect of ethanol on the ability of \textit{S. cerevisiae} to maintain a \( \Delta p \) across the plasma-membrane was examined. Cartwright et al. (1986) have shown that ethanol, up to a concentration of 2.0 M, partially dissipates the \( \Delta p \) across the plasma membrane of \textit{S. cerevisiae} NCYC 431 and, under conditions where the plasma-membrane ATPase is active, this dissipation is more pronounced in organisms from mid-exponential phase (8 h) compared with early stationary phase (16 h) cultures.

The present paper reports on the effect of ethanol on the activity of the plasma-membrane ATPase in \textit{S. cerevisiae} NCYC 431, and on the possible relationship between ethanol-induced inhibition of this enzyme and retardation of glycine accumulation caused by ethanol.

**METHODS**

\textbf{Organism.} The strain of \textit{S. cerevisiae} used in this study, NCYC 431 (Beavan et al., 1982), was maintained on slopes of malt extract/yeast extract/glucose/mycological peptone (MYGP) medium (Wickerham, 1951) solidified with 2\% (w/v) agar. A mutant (CC1) lacking GAP activity was isolated by a modification of the method of Rytka (1975) as described by Gregory et al. (1982). It was maintained on slopes of defined medium (Gregory et al. 1982) supplemented with L-methionine (20 mg l\(^{-1}\)).

\textbf{Experimental cultures.} All experiments were done with organisms from self-induced anaerobic cultures grown as described by Beavan et al. (1982). Organisms were harvested as described by Cartwright et al. (1986) after approximately 8 h (0.28 mg dry wt ml\(^{-1}\)) or 16 h (2.4 mg dry wt ml\(^{-1}\)) incubation.

\textbf{Isolation of plasma membranes.} Plasma membranes were isolated from spheroplasts that had been surface-labelled with cationic silica microbeads (Schmidt et al., 1983). To prepare spheroplasts, organisms were washed once in water, then in buffered sorbitol (20 mm-Tris containing 10 mm-MgCl\(_2\), 1.2 mm-sorbitol and 0.1 mm-sodium metabisulphite; pH 7.2) and suspended in the same buffer to 10 mg dry wt ml\(^{-1}\). Zymolyase 10000 was then added [0.1 mg (10 mg dry wt organism\(^{-1}\))] and the suspension incubated at 30°C with reciprocal shaking (120 r.p.m.). After incubation for 1 h, a check that formation of spheroplasts was complete was made by diluting 0.1 ml portions of the suspension into 2.9 ml of either 1.2 mm-sorbitol or water and measuring the optical density at 600 nm. Spheroplasts were washed three times in coating buffer (1.2 mm-sorbitol containing 25 mm-sodium acetate and 0.1 mm-KCl; pH 6.0), the population counted and then suspended in coating buffer to 1.5 \( \times 10^8 \) spheroplasts ml\(^{-1}\). Suspensions of spheroplasts and microbeads (3\%, w/v, in coating buffer) were mixed in the ratio 2:1. After incubation for 3 min at 4°C the suspension of coated spheroplasts was centrifuged (500 g; 4 min), the spheroplasts washed once in coating buffer and then suspended in the same buffer to a concentration of \( 10^8 \) spheroplasts ml\(^{-1}\). This suspension was diluted with an equal volume of coating buffer containing 2 mg polycryllic acid (\( M, 90000 \)) and spheroplasts were washed once with coating buffer. The spheroplasts were then resuspended in lysis buffer (5 mm-Tris/HCl containing 1 mm-EGTA; pH 8.0) to \( 10^8 \) spheroplasts ml\(^{-1}\) and the suspension vortexed for 5 min causing at least 95\% lysis of spheroplasts. The lysate was centrifuged (1000 g; 5 min), the pellet of plasma membrane washed three times in lysis buffer and then suspended in assay buffer (100 mm-MES/Tris containing 80 mm-KCl; pH 6.5). The protein content of the plasma-membrane preparation was assayed using the Bio-Rad protein assay, based on the dye-binding technique of Bradford (1976). Portions (0.1 ml) of plasma-membrane
preparation were solubilized by boiling for 5 min after addition of an equal volume of 0-1 M-NaOH. The solution was then neutralized by addition of 0-1 M-HCl (0-1 ml) and 0-2 ml acetic acid/sodium acetate buffer (0-2 M; pH 5-0). The amount of protein in samples was calculated using bovine serum albumin as a standard.

**Assay of ATPase activity.** Plasma-membrane ATPase activity was assayed by following release of P, from ATP. The reaction mixture consisted of 1 ml 100 mM-MES/Tris buffer containing 80 mM-KCl, 0-25-6-0 mM-MgCl₂ and plasma-membrane preparation containing 25-50 μg protein. Except when otherwise indicated, the pH of the buffer was 6-0 for membrane preparations from 16 h cultures and 6-5 for preparations from 8 h cultures. ATP concentrations were varied over the range 0-25-6-0 mM and, in all reaction mixtures, the ATP: Mg²⁺ ratio was maintained at unity. The reaction was started by adding ATP (sodium salt) and the mixture incubated for 10-40 min at 30 °C. The amount of P, liberated was determined as described by Serrano (1978). The reaction was stopped by adding 2 ml of 20% (v/v) conc. H₂SO₄ containing 0-5% (w/v) ammonium molybdate and 0-5% (w/v) SDS. Ascorbic acid (0-02 ml; 10%, w/v) was then added and the colour allowed to develop over 5 min at 30 °C. The absorbance of the solution was measured at 750 nm and the value related to P, concentration by a standard curve. ATPase activity is quoted as μmol P, released (mg protein)⁻¹ min⁻¹. To examine the effect of ethanol on ATPase activity, the alkanol (95%, v/v) was included in suspensions of plasma membrane in assay buffer to give final concentrations of 0-5, 1-0, 1-5 and 2-0 M. After 5 min incubation at 30 °C, the reaction was started by addition of ATP. ATPase inhibitors were similarly included in the membrane suspensions over a suitable range of concentrations and incubated for 10 min before starting the reaction.

**Measurement of velocity of glycine accumulation.** Organisms from 8 h or 16 h cultures were washed twice with citric acid/trisodium citrate buffer (20 mM; pH 4-0) suspended at 5 mg dry wt ml⁻¹ in the same buffer and stored in an ice-water mixture. The suspension (5 ml) used to measure the velocity of glycine accumulation consisted of citrate buffer (20 mM; pH 4-0) containing 2-5 mg dry wt organisms ml⁻¹ and was maintained at 30 °C in a water-bath and stirred continuously. Glucose (100 mM) or 2-deoxy-D-glucose (20 mM) was included in the suspension as indicated, as was ethanol to a final concentration of 0-5, 1-0, 1-5 or 2-0 M. After 5 min incubation, the experiment was started by addition of [U-¹⁴C]glycine [10⁻³-10⁻⁶ M; 0-1-1-0 μCi μmol⁻¹ (3-7-37 kBq μmol⁻¹)] after which portions (0-75 ml) were removed at predetermined times, rapidly filtered through membrane filters (0-45 μm pore size; 25 mm diameter; Millipore) and the filter and cells immediately washed with ice-cold citrate buffer (5 ml; pH 4-0) containing glycine at the concentration included in the suspension. Filters were then transferred to scintillation vials containing 7-5 ml Optiphase Safe (Fisons) and radioactivity measured in a liquid scintillation spectrometer (LKB Rackbeta, model 1217). Velocities of accumulation were determined from the linear segments of plots of the time-course of accumulation for up to 60 s.

**Determination of the phospholipid composition of plasma membranes and of the fatty-acyl composition of phospholipids.** A plasma membrane–cationic microbead preparation from 500 mg dry wt organisms was taken up in 15-20 ml methanol, to which chloroform was added to give a chloroform/methanol ratio of 2:1 (v/v). Non-specific lipase activity was inhibited by inclusion of 1 mm-p-chloromercuribenzoate, and the suspension was stirred for 2 h at room temperature. Extracts were washed with 0-25 vol. 0-88% (w/v) KCl and allowed to separate overnight at −20 °C. The lower phase was removed, dried on a rotary evaporator and the residue immediately dissolved in a minimum of chloroform. Phospholipids were separated from other lipid classes by TLC on plates of silica gel G (Whatman; 0-25 mm thick), using a solvent system of light petroleum (b.p. 60-80 °C)/diethyl ether/acetic acid (70:30:1, by vol.). Bands of lipid classes were visualized by spraying with 0.25% (w/v) BF₃ in ethanol, and then viewing plates under UV light (254 nm). Individual lipid classes were identified by reference to simultaneously run standards. Bands containing phospholipid classes were scraped off and suspended in 10 ml light petroleum/methanol (1:1, v/v). The suspension was vigorously shaken, 5 ml 30% (w/v) NaCl added, the suspension allowed to separate at room temperature, and the upper layer removed and retained. A further 5 ml light petroleum was added, and the extraction procedure repeated. The combined extracts were dried under nitrogen, the residue taken up in 0-3 ml chloroform, and individual phospholipid classes separated by TLC using a solvent system of chloroform/methanol/acetic acid (65:25:8, by vol.; Kramer et al., 1978). Individual phospholipid classes were visualized by spraying plates with 4:2 M-H₂SO₄ containing 0-65% (w/v) molybdenum oxide (Dittmer & Lester, 1964) and identified by reference to standards. The relative proportions of the major phospholipid classes were determined using a scanning densitometer (Joyce Loebl Chromoscan 3; wavelength 626 nm; aperture width 0-1-0-3 mm). To determine the fatty-acyl composition of phospholipid classes, bands were visualized and identified as described above, and scraped off. Fatty-acyl methyl esters were prepared by heating (80 °C, 60 min) the silica gel with 5 ml methanol containing 14% (w/v) BF₃ in a Reactivial (Pierce). After cooling, the solution was extracted twice with light petroleum/methanol/30% (w/v) NaCl (1:1:1, by vol.). Fatty acid methyl esters were analysed by GLC on a column (1-5 m) containing 10% S2330 supported on 100-200 mesh Chromosorb W-AW at 165 °C. The injection port was at 210 °C, the detector temperature was 190 °C and the carrier gas (N₂) flow rate was 60 ml min⁻¹. Percentage fatty acid compositions were calculated using a Pye Unicam CDPI integrator.

**Electron microscopy.** The purity of plasma-membrane preparations was examined by transmission electron microscopy using a modification of the method of Henschke et al. (1983).
RESULTS

Properties of the plasma-membrane ATPase

Plasma-membrane preparations appeared essentially free from contamination with nonmembraneous and mitochondrial material when examined in the electron microscope. ATPase activity of membranes was inhibited by sodium orthovanadate, diethylstilboestrol and DCCD to an extent virtually identical with membranes from organisms from 8 h (Fig. 1) and 16 h cultures (data not shown). Oligomycin, at concentrations up to 300 μg ml⁻¹, caused less than 5% inhibition of ATPase activity in membranes from organisms from 8 h and 16 h cultures. The pH optimum of the ATPase activity in membranes from organisms from 8 h cultures was around 6-5, while the optimum for the activity in membranes from organisms from 16 h cultures was near 6-0 (Fig. 2). Values for $K_m$, derived from Hanes (1932) plots, for ATPase activity of membranes from 8 h and 16 h cultures were virtually identical, whereas $V_{max}$ values differed (Table 1). Ethanol, at concentrations up to 2-0 M, inhibited ATPase activity in membranes from organisms from both 8 h and 16 h cultures, although the inhibition constant $K_i$ derived from Dixon (1953) plots was different for the two preparations (Table 1).

Phospholipid and fatty-acyl compositions of plasma-membranes

Phosphatidylethanolamine and phosphatidylcholine were the principal phospholipids in plasmamembranes, the former being present in greater proportion in membranes from organisms from 8 h compared with 16 h cultures (Table 2). Phosphatidylinositol was present in about the same proportion in membranes from organisms from 8 h and 16 h cultures while the proportions of phosphatidylserine differed. The fatty-acyl composition of individual plasma-membrane phospholipids differed in organisms from 8 h as compared with 16 h cultures, the proportion of unsaturation decreasing with culture age (Table 3). Of the four major phospholipids detected, phosphatidylinositol showed greatest conservation of fatty-acyl unsaturation with culture age.
organisms. The inhibitory effect on glycine accumulation closely paralleled the decrease in glycine. Values for caused by ethanol (Cartwright et al., 1986) are greater for organisms from cultures grown for 8 h cultures (data not shown) and in suspensions containing 50–100 μM ethanol. This correspondence was observed with organisms grown for 8 h or 16 h incubation

<table>
<thead>
<tr>
<th>Phospholipid</th>
<th>Percentage of total phospholipid*</th>
</tr>
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<tbody>
<tr>
<td>Phosphatidylethanolamine</td>
<td>46.32 ± 5.44</td>
</tr>
<tr>
<td>Phosphatidylcholine</td>
<td>32.01 ± 3.60</td>
</tr>
<tr>
<td>Phosphatidylinositol</td>
<td>18.49 ± 3.11</td>
</tr>
<tr>
<td>Phosphatidylserine</td>
<td>3.06 ± 0.33</td>
</tr>
</tbody>
</table>

* Values quoted are ± SE.

<table>
<thead>
<tr>
<th>Fatty-acyl residue</th>
<th>8 h</th>
<th>16 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>12:0</td>
<td>1.98 ± 0.46</td>
<td>0.23 ± 0.08</td>
</tr>
<tr>
<td>14:0</td>
<td>2.14 ± 0.14</td>
<td>1.92 ± 0.33</td>
</tr>
<tr>
<td>16:0</td>
<td>22.70 ± 2.27</td>
<td>28.98 ± 1.21</td>
</tr>
<tr>
<td>18:0</td>
<td>4.41 ± 0.90</td>
<td>3.92 ± 0.24</td>
</tr>
<tr>
<td>18:1</td>
<td>21.69 ± 1.53</td>
<td>30.40 ± 2.04</td>
</tr>
<tr>
<td>Δ mol⁻¹</td>
<td>0.69</td>
<td>0.73</td>
</tr>
</tbody>
</table>

* Values quoted are ± SE.

Effect of ethanol on glycine accumulation

Woolf–Hofstee (Hofstee, 1959) plots showed that organisms possessed two separable glycine accumulation systems, which differ in affinity for glycine and in velocity of solute accumulation (Table 4). The high-affinity system was not detectable in the GAP-less mutant. Ethanol inhibited glycine accumulation by the high-affinity system in energized and de-energized organisms. The inhibitory effect on glycine accumulation closely paralleled the decrease in Δp caused by ethanol (Cartwright et al., 1986). This correspondence was observed with organisms from 8 h (Fig. 3) and 16 h cultures (data not shown) and in suspensions containing 50–100 μM glycine. Values for Kₐ (Dixon, 1953) were virtually the same for de-energized organisms from 8 h (0.71 ± 0.09 M) and 16 h (0.74 ± 0.08 M) cultures, but with energized organisms the value was greater for organisms from 16 h cultures (1.55 ± 0.11 M) compared with 8 h cultures (0.97 ± 0.10 M). (Values are means ± SE.)
Fig. 3. Relationship between the effect of ethanol on the velocity of glycine accumulation by the GAP and on the Δp generated by S. cerevisiae NCYC 431 harvested from 8 h cultures. ●, Response of de-energized organisms; ○, response of energized organisms. Data are for glycine accumulation from a suspension containing 100 μM-glycine. Values for the Δp are from Cartwright et al. (1986). The correlation coefficient for the data shown is 0.95, and that for data on organisms from 16 h cultures (not shown) 0.81.

Table 4. Kinetic constants for accumulation of glycine by S. cerevisiae NCYC 431

<table>
<thead>
<tr>
<th>Value</th>
<th>Affinity system</th>
<th>8 h</th>
<th>16 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>KT (mM)</td>
<td>High</td>
<td>0.06 ± 0.02</td>
<td>0.09 ± 0.04</td>
</tr>
<tr>
<td></td>
<td>Low</td>
<td>2.25 ± 0.04</td>
<td>2.86 ± 0.08</td>
</tr>
<tr>
<td>Vmax [pmol (mg dry wt)^{-1} s^{-1}]</td>
<td>High</td>
<td>7.80 ± 0.96</td>
<td>8.21 ± 1.52</td>
</tr>
<tr>
<td></td>
<td>Low</td>
<td>275.9 ± 18.6</td>
<td>1352.8 ± 26.9</td>
</tr>
</tbody>
</table>

* Values were corrected as described by Neal (1972), and are quoted ± se.

DISCUSSION

Electron microscopy confirmed that plasma membranes of S. cerevisiae, isolated using cationically charged microbeads, were free from contaminating material derived from other yeast organelles. Furthermore, the effects of inhibitors on the activity of the plasma-membrane ATPase, an enzyme often taken as a marker for the yeast plasma membrane, attest the purity of the preparations obtained. One of the main advantages of using microbeads to isolate plasma membranes from eukaryotic micro-organisms is that the time taken to obtain reasonably pure preparations is much shorter than when differential centrifugation and density-gradient methods are employed. Shortening of the preparation time might decrease the extent to which membrane lipids are degraded during isolation. It was therefore of interest to compare the lipid composition of membranes obtained using microbeads with those reported for preparations obtained using alternative techniques. The proportions of phospholipid classes in plasma membranes were very similar to those discovered by Longley et al. (1968), and Suomalainen & Nurminen (1970). Moreover, the changes that occurred in the overall fatty-acyl composition of plasma-membrane phospholipids as cultures progressed from the exponential to the stationary phase of growth were similar to those reported by Beavan et al. (1982) for whole-cell
phospholipids, suggesting that culture age has a similar effect on the degree of unsaturation of phospholipids in most types of yeast membrane. Our study also revealed that changes in the relative proportions of phospholipids and in the fatty-acyl composition of individual classes of plasma-membrane phospholipid accompany the transition from the exponential to the stationary phase of growth. The relative proportions of phosphatidylethanolamine and phosphatidylserine, but not of phosphatidylcholine or phosphatidylinositol, in plasma membranes varied with culture age. Interestingly, only phosphatidylinositol retained the same degree of fatty-acyl unsaturation in plasma membranes from organisms from 8 h and 16 h cultures. This suggests that this phospholipid supports some plasma-membrane function which, associated with a critical degree of fatty-acyl unsaturation, is of physiological importance to the yeast.

Since ethanol dissipates the Δp across the plasma membrane of \textit{S. cerevisiae} NCYC 431 (Cartwright \textit{et al.}, 1986), one objective of the present study was to assess to what extent this dissipation is caused by an inhibitory effect of ethanol on the plasma-membrane ATPase. Our finding that ethanol non-competitively inhibited ATPase activity in the same range of concentrations at which it dissipated the Δp suggests that inhibition of this enzyme is a major factor in determining the extent of Δp dissipation. That the $V_{\text{max}}$ value for the ATPase in membranes from organisms from 16 h cultures was lower than the corresponding value for membranes from organisms from 8 h cultures, while the $K_m$ value remained unchanged, confirms the report by Tuduri \textit{et al.} (1985) of a decline in the activity of this enzyme during the late exponential phase of growth. This effect, and also the change in the optimum pH value for the membrane-bound ATPase activity, are conceivably results of alterations that take place in the phospholipid and fatty-acyl compositions of the plasma membrane with culture age. In this context, Stadtlander \textit{et al.} (1982) showed that changes in phospholipid head-group orientation as well as in fatty-acyl chain length and degree of unsaturation modify kinetic characteristics of the yeast plasma-membrane ATPase.

To determine the extent to which ethanol-induced inhibition of active uptake of solutes might be attributable to the dissipative effect of ethanol on the Δp, glycine was chosen as the solute to be studied. Ballarin-Denti \textit{et al.} (1984) reported that glycine accumulation by \textit{S. cerevisiae} had no appreciable effect on intracellular pH value, even under de-energized conditions. Consequently, uptake of glycine by \textit{S. cerevisiae} NCYC 431 should not alter the values for the Δp previously determined for this strain (Cartwright \textit{et al.}, 1986). Our data on glycine accumulation confirm the claims by Eddy \textit{et al.} (1970) and Ballarin-Denti \textit{et al.} (1984) that \textit{S. cerevisiae} possesses two distinct uptake systems for this amino acid. However, using the GAP-less mutant, we were able to show conclusively that the high-affinity system, which was examined further in the present study, was the GAP. The nature of the low-affinity glycine-uptake system remains unclear although, given the high $K_c$ and low $V_{\text{max}}$ values for glycine accumulation by this carrier, it probably functions primarily to transport some other amino acid or acids.

Our data on the effect of ethanol on glycine accumulation agree with the report of Leão \& van Uden (1984) that inhibition of GAP activity is non-competitive. Leão \& van Uden (1984) suggested that ethanol inhibits glycine transport by altering the conformation of the transport protein(s). An alternative explanation is that ethanol inhibits glycine accumulation as a result of its dissipative effect on the Δp. Our belief is that this effect at least partly accounts for ethanol-induced inhibition of glycine accumulation by \textit{S. cerevisiae}. The finding that ethanol has a greater inhibitory effect on glycine accumulation by organisms from 8 h rather than 16 h cultures, under energized but not de-energized conditions, supports this hypothesis, since the more pronounced dissipative effect of ethanol on the Δp in organisms cultured for 8 h rather than 16 h is only apparent under energized conditions (Cartwright \textit{et al.}, 1986). The linear relationship demonstrated between rates of glycine accumulation by the GAP and the magnitude of Δp, both in the presence and absence of ethanol, further illustrates the direct effect that the Δp dissipation has on glycine uptake.

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REFERENCES


purification of yeast plasma membranes from Saccharomyces cerevisiae. NCYC 366 Biochimica et biophysica acta 732, 421-427.


