Localization of dipeptidyl aminopeptidase yscIV in the plasma membrane of \textit{Saccharomyces cerevisiae}

CARMEN BORDALLO, ROSARIO CUEVA and PAZ SUÁREZ RENDUELES*

Area de Bioquímica y Biología Molecular, Departamento de Biología Funcional, Facultad de Medicina, Universidad de Oviedo, E-33071 Oviedo, Spain

(Received 4 October 1989; accepted 5 December 1989)

The subcellular distribution of dipeptidyl aminopeptidase activity was studied in protoplast lysates of \textit{Saccharomyces cerevisiae} that were virtually free from vacuolar contamination. Dipeptidyl aminopeptidase yscIV, the \textit{STE13} gene product, was found to be associated with plasma membrane vesicles after sucrose gradient isopycnic centrifugation. Another dipeptidyl aminopeptidase activity, not yet fully characterized, was localized in a microvesicular population co-sedimenting with chitosomes.

Introduction

The yeast \textit{Saccharomyces cerevisiae} contains a number of proteinases which play a vital role in cellular control (for recent reviews see Suárez-Rendueles & Wolf, 1988; Hirsch et al., 1989).

We have previously described a membrane-bound X-prolyl dipeptidyl aminopeptidase activity in yeast (Suárez-Rendueles et al., 1981; Garcia Ramos et al., 1983), later reported to be due to at least two different enzymes (Julius et al., 1983; Bordallo et al., 1984). One of the enzymes, which is inactivated by heating crude extracts at 60°C for 20 min, was termed dipeptidyl aminopeptidase yscV. This enzyme is associated with vacuolar membranes (Bordallo et al., 1984) from which it was purified and characterized as a serine peptidase acting on proline-containing substrates (García-Alvarez et al., 1985). Genetic studies have shown that the gene designated \textit{DAP2} is the structural gene for dipeptidyl aminopeptidase yscV, the biological role of which is unknown as \textit{dap2} mutants do not show any obvious difference in phenotype (Suárez-Rendueles & Wolf, 1987).

The synthesis and processing of \textit{pre-pro-\alpha-factor} have been studied with the aid of mutants blocked at various stages in the yeast secretory pathway (Julius et al., 1984). The first stage of proteolytic cleavage of the precursor into smaller units was detected in a mutant strain (\textit{sec7}) that accumulates Golgi-body-like structures at the restrictive temperature; processing is then completed within secretory vesicles (Julius et al., 1984). These findings led to the hypothesis that the specific proteinase responsible for the last processing step of \textit{\alpha-factor} precursor, i.e. dipeptidyl aminopeptidase yscIV, is located in secretory vesicles, plasma membrane or both (Julius et al., 1984).

In this paper we describe the subcellular distribution of dipeptidyl aminopeptidase activity in \textit{S. cerevisiae} lysates prepared by disrupting protoplasts in conditions minimizing organelle rupture.

Methods

\textit{Yeast strains and growth conditions.} \textit{S. cerevisiae} 1022 (ETH, Zurich, Switzerland; ATCC 32167), a wild-type strain kindly supplied by Dr A. Wiemken, and \textit{S. cerevisiae} A2S3 (\textit{MATa ste13-1 ade6 leu2 trp5 his5 met1 can1 trm1 gal2}) (Sprague et al., 1981), defective in dipeptidyl aminopeptidase yscIV activity, were used. Cells were grown in liquid mineral medium (0·57% yeast nitrogen base without amino acids, 1% w/v, glucose) supplemented, when necessary, with the amino acids required by the auxotrophic strain. Cells were grown aerobically at 28°C and harvested at the exponential phase of growth.

\textit{Preparation, labelling and lysis of protoplasts.} Protoplasts from \textit{S. cerevisiae} 1022 were prepared and purified according to the method of Schwencke et al., 1983. When strain A2S3 was used, exponential-phase
yeast cells pre-treated as above were resuspended in 0·6 M-sorbitol, 10 mM-EDTA, 20 mM-MES/Tris (pH 6·0) containing 0.5% (w/w) Zymolyase 20000. Conversion to protoplasts was usually complete within 60 min with both strains. Two different methods of protoplast preparation were used because we found that the cell wall of strain A283 was more sensitive to Zymolyase than to Helicase.

Labelling and lysis of protoplasts were done as described by Flores Martinez & Schwenccke (1988). A suspension of protoplasts (107 ml−1) in 0·6 M-sorbitol, 10 mM-MgCl2, 20 mM-MOPS/Tris, pH 7·0 (buffer A) was mixed with 0·25 μCi (9·25 kBq) [3H]concanavalin A dissolved in buffer A. After 10 min at 30°C, protoplasts were collected by centrifugation (swinging-bucket rotor) at 3500 g for 5 min and washed twice with ice-cold buffer A. The pellet was carefully resuspended in 1 M-sorbitol, 25 mM-triethylamine (TEA)/HCl, pH 8·0, 1 mM-EDTA and kept on ice for 10 min. The protoplast suspension was diluted 1:4 with pre-warmed (30°C) disrupting buffer (0·36 M-sorbitol, 25 mM-TEA/HCl, pH 8·0, 50 mM-glucose, 2·5 mM-K2CO3) and incubated for 10 min at 30°C. Rupture of protoplasts and vacuole liberation was assessed by phase-contrast microscopy.

Isopycnic sucrose gradient centrifugation. The protoplast lysate was centrifuged at 500 g for 15 min (swinging-bucket rotor) to remove vacuoles, intact protoplasts and debris. The supernatant was filtered through Sepharose-2B as described by Flores Martinez & Schwenccke (1988). Void volume fractions were pooled, sucrose was added to adjust the concentration to 20% (w/w) and used in conjunction with a 20–60% (w/w) linear sucrose gradient (32 ml). The gradients were centrifuged in a Beckman SW-25 rotor at 70000 g (r, 9·27 cm) for 40 h. Fractions (1 ml) were collected from the bottom upwards by means of a peristaltic pump and analysed for enzyme activities as described below. Density was determined by refractometry.

Enzyme assays. Dipeptidyl aminopeptidase activity was determined with Ala-Pro-4-nitroanilide as substrate as described previously (Suárez-Rendueles et al., 1981). Chitin synthetase was assayed according to Ruiz-Herrera & Bartnicki-Garcia (1976), but using 60 μg α-chymotrypsin ml−1 to activate zymogenic chitin synthetase as indicated by Flores Martinez & Schwenccke (1988). Carboxypeptidase yscY activity was assayed by the method of Aibara et al. (1971) with benzoyl-tyrosine-4-nitroanilide as substrate; as a modification, 0.5% sodium deoxycholate was added (Bordallo et al., 1984). α-Mannosidase activity was determined with p-nitrophenyl α-D-mannopyranoside as substrate as described by Opheim (1978).

Protein determination. Protein was determined according to the method of Bradford (1976).

Substrates and chemicals. Ala-Pro-4-nitroanilide was supplied by Bachem, Switzerland. Crystallized α-chymotrypsin was purchased from Serva. Uridine diphospho-N-acetyl-α-[U-14C]glucosamine [ammonium salt; 247 mCi mmol−1 (9·14 GBq mmol−1)] and N-acetyl-[3H]concanavalin A [47·6 Ci mmol−1 (1·76 TBq mmol−1)] were obtained from Amersham. Sepharose-2B was from Pharmacia. All other chemicals, which were of the highest purity available, were purchased from either Merck or Sigma. Zymolyase was obtained from Kirin, Japan.

Results

Subcellular distribution of dipeptidyl aminopeptidase yscIV

The use of a mild disruption procedure for yeast protoplasts, as described by Flores Martinez & Schwenccke (1988), allowed us to investigate the subcellular localization of the membrane-bound dipeptidyl aminopeptidase yscIV free from the interference caused by dipeptidyl aminopeptidase yscV, an enzyme with overlapping substrate specificity, which is located in the vacuolar membrane (Bordallo et al., 1984). Protoplasts from the wild-type strain S. cerevisiae 1022 were labelled with [3H]concanavalin A and lysed as indicated in Methods. After filtration through Sepharose-2B, the particulate fraction eluting in the void volume of the column was subjected to isopycnic sucrose gradient centrifugation (20–60%, w/w, sucrose); two peaks of dipeptidyl aminopeptidase activity were reproducibly obtained (Fig. 1a). The main peak (peak II) contained vesicles having a buoyant density (1·145 g cm−3) typical of chitosomes (Bracker et al., 1976; Ruiz-Herrera et al., 1984; Flores Martinez & Schwenccke, 1988). The other peak (peak I) contained plasma membrane vesicles, identified on the basis of their buoyant density (1·203 g cm−3) and by the fact that the [3H]concanavalin A co-sedimented with them (Scarborough, 1975; Boller et al., 1976; Flores Martinez & Schwenccke, 1988).
Localization of yeast dipeptidyl aminopeptidase

Further evidence for the identity of the two fractions was obtained by measuring chitin synthetase activity along the gradient (Fig. 1a). The distribution of chitin synthetase was similar to that described above for dipeptidyl aminopeptidase. In both peaks chitin synthetase was present mostly in zymogen form: without proteolytic activation by chymotrypsin less than 5% of the total chitin synthetase activity could be measured. The same results concerning zymogenic chitin synthetase distribution were reported by Flores Martínez & Schwencke (1988) using the same yeast strain. These authors identify peak 1 as plasma membrane particles and peak II as chitosome vesicles. In agreement with those authors we were unable to detect carboxypeptidase yscY activity (a soluble vacuolar-enzyme marker) in either the material excluded from the Sepharose-2B column or in the fractions collected after the isopycnic sucrose gradient centrifugation, thus ruling out the possibility of interference by soluble vacuolar proteinases. We also assayed α-mannosidase activity as a measure of cross-contamination by vacuolar membranes: about 15% of the total α-mannosidase activity remained in the 500 g supernatant after protoplast lysis. However, no α-mannosidase activity could be detected after filtration through the Sepharose-2B column, as this process inactivated the enzyme for unknown reasons. Therefore, the possibility of a low level of contamination of the gradient fractions by vacuolar membranes and hence by the tonoplast-bound dipeptidyl aminopeptidase yscV could not be ruled out completely.

Preliminary evidence that dipeptidyl aminopeptidase yscIV, the enzyme responsible for α-factor precursor processing, was the activity detected in plasma membrane vesicles (Fig. 1a, peak 1) came from kinetic studies. No significant loss of activity was observed after heating gradient fractions (under the bar in peak I, Fig. 1a) at 60 °C for 20 min. This behaviour is characteristic of dipeptidyl aminopeptidase yscIV (Julius et al., 1983; Bordallo et al., 1984). Also, the affinity for the substrate Ala-Pro-4-nitroanilide was very similar to that reported for dipeptidyl aminopeptidase yscIV (Bordallo et al., 1984), the apparent Kₐ value being 0.9 mM (data not shown). Further evidence that dipeptidyl aminopeptidase yscIV is bound to the plasma membrane was obtained by using a yeast strain devoid of dipeptidyl aminopeptidase yscIV activity due to a single chromosomal gene mutation affecting its structural gene (STE13) (Julius et al., 1984; Suárez-Rendueles & Wolf, 1987). Protoplasts from strain A2S3 (ste13) were labelled with [3H]concanavalin A, lysed and filtered through a Sepharose-2B column following the protocol used for the wild-type strain. When the void volume fractions from this column were subjected to isopycnic sucrose gradient centrifugation (20–60%, w/w), the distribution of dipeptidyl aminopeptidase activity was as shown in Fig. 1(b). Only the peak of activity (peak II) at specific gravity 1.145 g cm⁻³ (chitosomes) was detected, confirming that the plasma-membrane-bound activity detected in the wild-type strain was due to dipeptidyl aminopeptidase yscIV.

Characterization of the dipeptidyl aminopeptidase activity co-sedimenting with chitosomes

The thermal stability of dipeptidyl aminopeptidase activity was measured both in a whole membrane fraction (prepared as described by Suárez-Rendueles et al., 1981) and in purified chitosomes prepared from the mutant strain lacking dipeptidyl aminopeptidase yscIV (fractions under the bar in Fig. 1b). As expected, in the whole membrane fraction a heat-labile dipeptidyl aminopeptidase activity, corresponding mainly to dipeptidyl aminopeptidase yscV (Bordallo et al., 1984), was detected (Fig. 2). A quite different picture was obtained when the heat stability of chitosome-bound activity was studied. Heating chitosomes at 60 °C for 5 min inactiva-
showed that both activity peaks had a PI of 4-9 (Garcia-}
focusing in the pH range
ed by PMSF, a specific inhibitor of serine proteinases,
yscV and the activity co-sedimenting with chitosomes
substrate
isoelectric point. Both activities were completely inhibit-
ed chitosomal dipeptidyl aminopeptidase activity com-
pletely (Fig. 2), whereas dipeptidyl aminopeptidase yscV
retains almost 50% activity under such conditions
(Bordallo et al., 1984).

The affinity for the substrate Ala-Pro-4-nitroanilide was
determined using either a whole membrane fraction or
purified chitosomes from strain A2S3 as enzyme
sources (Fig. 3). Two components with different sub-
strate affinities ($K_m$ values 0-09 and 0-44 mM) were
present in both membrane preparations but their relative
contribution to the total dipeptidyl aminopeptidase
activity measured depended strongly on the enzyme
source. Thus, dipeptidyl aminopeptidase yscV ($K_m$
0-09 mM) (Bordallo et al., 1984) was the major activity
detected in the whole membrane preparation (Fig. 3a),
while a different enzyme, with lower affinity for the
substrate ($K_m$ 0-44 mM) was the major component in the
chitosome vesicles (Fig. 3b). Dipeptidyl aminopeptidase
yscV and the activity co-sedimenting with chitosomes
could not be distinguished by sensitivity to inhibitors or
isoelectric point. Both activities were completely inhib-
ited by PMSF, a specific inhibitor of serine proteinases,
and by various divalent cations such as Hg$^{2+}$, Cd$^{2+}$, Zn$^{2+}$
and Ni$^{2+}$. The isoelectric point determined by isoelectric
focusing in the pH range 4-6-5 using a glycerol gradient
showed that both activity peaks had a pI of 4-9 (Garcia-
Alvarez et al., 1985).

Discussion

Determining the subcellular distribution of dipeptidyl
aminopeptidase yscIV was difficult for two reasons.
First, in the wild-type strain the dipeptidyl aminopepti-
idase yscV, which has overlapping substrate specificity,
accounts for approximately 70% of the total activity
measurable in a protoplast lysate (Julius et al., 1983;
Bordallo et al., 1984). Secondly, obtaining pure prepara-
tions of yeast cell organelles, is difficult, especially when
harsh methods of cell disruption are used.

The recent publication of a mild iso-osmotic disrup-
tion method of yeast protoplasts that leaves most of the
vacuoles intact and minimizes cross-contamination of
subcellular fractions (Flores Martinez & Schwencke,
1988), allowed us to study the location of dipeptidyl
aminopeptidase yscIV. The physiological role (o-factor
precursor processing) of this enzyme indicates that it
could be located in the plasma membrane, in secretory
vesicles or in both (Julius et al., 1984).

The results presented in this paper show that
dipeptidyl aminopeptidase activity from a wild-type
strain is associated with two different particulate
fractions after isopycnic sucrose gradient centrifugation
of a partially purified protoplast lysate from which most
vacuoles and thus most dipeptidyl aminopeptidase yscV
activity had been eliminated. The fraction equilibrating
at a buoyant density of 1-203 g cm$^{-3}$ and containing the
bulk of bound [$^3$H]concanavalin A has been identified by
several criteria as plasma-membrane ghost vesicles by
Flores Martinez & Schwencke (1988) who used proto-
plast homogenates from the same yeast strain. The other
fraction, equilibrating at a lower buoyant density
(1-145 g cm$^{-3}$) has been characterized as chitosomes by
these authors. It must be kept in mind, however, that
other vesicular material such as proteasomes (Tanaka
et al., 1988) may sediment in the same fraction. Neverthe-
less, the proteolytic activity detected so far in protea-
omes shows latency, while the dipeptidyl aminopepti-
dase activity described here can be detected without
previous activation.

The data presented clearly demonstrate that dipepti-
dyl aminopeptidase activity associated with plasma-
membrane vesicles is absent in a yeast strain defective in
dipeptidyl aminopeptidase yscV due to a single chromo-
somal mutation (stel3), strongly supporting the proposal
that this enzyme is localized at the yeast plasma
membrane. The fact that we have been unable to detect
dipeptidyl aminopeptidase activity in whole cells that
have not first been permeabilized to permit entry of the
small peptide substrate used for the assay probably
means that dipeptidyl aminopeptidase yscIV is not an
exocellular plasma membrane protein.

Dipeptidyl aminopeptidase activity co-sedimenting

![Fig. 3. Eadie–Hofstee plot of dipeptidyl aminopeptidase activity with Ala-Pro-4-nitroanilide as substrate in samples prepared from S. cerevisiae A2S3 (stel3). Enzyme activity was assayed at 37°C as outlined in Methods. (a) Whole membrane fraction; (b) chitosomes (dialysed pooled fractions indicated by the bar in Fig. 1b).]
with chitosomes was found both in a wild-type yeast strain and in the ste13 mutant strain. This shows that chitosomal dipeptidyl aminopeptidase activity cannot be due to dipeptidyl aminopeptidase yscIV. The preliminary data presented here concerning the identification and characterization of activity sedimenting with the chitosomal fraction suggest that although there is some vacuolar contamination of the chitosomal fraction, as judged by the presence of dipeptidyl aminopeptidase yscIV activity, most of the activity co-sedimenting with these microvesicles might be due to a different enzyme. Further studies are in progress in order to identify and better characterize this activity.

The relative distribution of dipeptidyl aminopeptidase activity between the plasma membrane and the chitosomes found in the study probably does not exactly reflect the situation 'in vivo', because of the possible fusion of plasma membrane and chitosomes which can occur after cell disruption and during the purification, and because the chitosomal fraction was slightly contaminated with vacuolar dipeptidyl aminopeptidase yscV.

We conclude that the product of the STE13 gene, dipeptidyl aminopeptidase yscIV, is localized at the plasma membrane, and that another dipeptidyl aminopeptidase activity, not yet characterized, is localized in a vesicular fraction co-sedimenting with chitosomes.

This work was supported by grant 2582/83 from the Comision Asesora para la Investigacion Cientifica y Técnica of Spain. We extend special thanks to Jaime Schwencke for providing information prior to publication.

References


