Release of Wall-bound Invertase and Trehalase in \textit{Neurospora crassa} by Hydrolytic Enzymes

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(Accepted for publication 17 September 1971)

SUMMARY

About 25\% of the total cellular invertase and trehalase activity were found in a purified \textit{Neurospora} wall preparation. Attempts were made to dissociate these enzymes from the walls with chemical reagents and hydrolytic enzymes. A detergent (Triton X-100), a sulphydryl reducing agent (\(\beta\)-mercaptoethanol), a chelating agent (ethylenediaminetetraacetate), a concentrated salt solution (1 M-KCl), and buffers ranging in pH from 3 to 10 did not release them significantly. Snail-gut juice released more than 90\% of both enzymes. \(\beta\)-1,3-Glucanase, prepared from \textit{Bacillus circulans} WL-12, also released similar amounts. Chitinase released about 80\% of invertase and 60\% of trehalase. Cellulase did not release any significant amount of either enzyme. Trypsin released only a few per cent of invertase and severely inactivated trehalase. Thus it appears that these two wall-bound enzymes are released only when covalent bonds of the \textit{Neurospora} wall constituents are disrupted.

INTRODUCTION

According to Pollock (1962), the locations of enzymes in micro-organisms such as bacteria can be grouped into three categories: intracellular (inside the cytoplasmic membrane), surface-bound (outside the cytoplasmic membrane but cell-bound), and extracellular (easily separated from the cells by mild procedures). Considerable confusion exists in their designation because the surface-bound enzymes have been variously referred to as mural enzymes (Eberhart & Beck, 1970), exocellular enzymes (Weimberg & Orton, 1966), external enzymes (Lampen, Neumann, Gascon & Monteneceur, 1967), exoenzymes (Eberhart, 1961) and extracellular enzymes (Matile, 1964). The last three terms have also been used interchangeably to delineate any enzyme occurring in the milieu exterior to the wall. For a clearer operational definition, we propose to restrict the use of "extracellular" to describe only the enzymes that are free in the medium or are easily separated from the cells by processes such as centrifugation or washing with water. Enzymes that are liberated on destruction of walls, e.g. during sphaeroplast formation, and whose activities are governed by factors in the medium, such as pH and the presence of substrates or inhibitors that do not penetrate the cell membrane, will be specifically described as 'cell-bound' or 'surface-bound' exoenzymes, or 'mural' exoenzymes (Trevithick & Metzenberg, 1966) if they are exclusively found in the wall fraction. 'Exoenzymes' will refer generally to all the enzymes outside the permeability barrier (i.e. plasma membrane), regardless of whether they are still cell-bound or truly extracellular.

Many cell-bound exoenzymes have been identified in bacteria (Pollock, 1962; Malamy & Horecker, 1964; Coles & Gross, 1967) and fungi (Mandels, 1953; Matile, 1964; Lampen, 1968; Eberhart & Beck, 1970; Scott & Metzenberg, 1970). Those that have been localized in the fungal wall fraction, i.e. mural exoenzymes, include trehalase (Hill & Sussman, 1964),

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In our laboratory about 25% of the total mycelial activities of invertase (EC. 3.2.1.26, β-D-fructofuranoside fructohydrolase) and trehalase (EC. 3.2.1.28, α,α'-glucoside 1-glucohydrolase) were found in the purified wall fraction of Neurospora in the post-log phase of growth. Consequently, it was of interest to investigate the forces keeping these enzymes in their mural location. Attempts were made to dissociate these two enzymes from the wall fraction with a variety of chemical reagents and hydrolytic enzymes. The chemical reagents chosen were known to affect specific types of binding force, such as hydrogen bonding, hydrophobic bonding, salt linkage, or metal-ligand co-ordination bonding. Each hydrolytic enzyme used was known to hydrolyse a specific component of the Neurospora wall, i.e. protein, chitin, and glucan [mostly in β-1,3-glycosidic linkages with some β-1,4- and α-1,4-linkages (de Terra & Tatum, 1963; Mahadevan & Tatum, 1965; Galsworthy, 1966; Burnett, 1968)].

METHODS

Organisms. Wild-type Neurospora crassa containing the Emerson genetic background was grown in Erlenmeyer flasks containing Fries minimal medium (Beadle & Tatum, 1945), 2.7% (w/v) galactose, and 0.04 M-sodium succinate buffer (pH 5.2) for 20 h at 30° in a gyratory shaker bath (New Brunswick Scientific Co., New Brunswick, New Jersey, U.S.A.) agitated at 180 rev./min. The mycelium was harvested by filtration on a Millipore filter and used for preparation of the wall fraction.

Preparation of wall fraction. The procedure was modified from the method of Chung & Trevithick (1970). Each batch of about 1 g mycelium (blotted dry with filter paper) was suspended in 20 ml cold phosphate buffered saline (PBS: 0.9% sodium chloride in 0.01 M-sodium phosphate buffer at pH 7.1) in a metal container and disrupted with a sonifier (Branson Soni Power S 125) at maximum output for a period of 3 to 25 min. More than 95% of the cells were estimated by phase-contrast microscopy to have been broken. The sample was kept below 10° during this treatment by using an ethylene glycol–dry ice bath and turning on the sonifier intermittently. The suspension was centrifuged at 650 g in a model PR-2 Centrifuge (International Equipment Co., Needham Hts, Massachusetts, U.S.A.) at 4° for 30 min. The residue was washed successively with PBS, 1% (v/v) Triton X-100, M-NaCl (twice), PBS, and glass distilled water by centrifugation at 100 g for 15 min. The final wall residue was free from cytoplasmic debris when observed under the phase-contrast microscope, and contained less than 2% of the total cellular ribonucleic acid when measured by a modified Mejabaum reaction according to Merchant, Kahn & Murphy (1964). The wall suspension used for all the following experiments had a dry wt of 1 to 1.5 mg/ml and contained approximately 9 units of invertase activity and 0.03 units of trehalase activity per ml. (See below for definitions of units.)

Treatment with chemical reagents

(i) Buffers with a range of pH values from 3 to 10 were prepared at a total final concentration of 0.14 M: glycine–hydrochloric acid (pH 3-0); sodium acetate–acetic acid (pH 4.0, pH 5.0); sodium monohydrogen phosphate–sodium dihydrogen phosphate (pH 6.2, pH 7.0, and pH 8.0); glycine–sodium hydroxide (pH 9-0, pH 10-0).
For each buffer a wall suspension (1 ml) was mixed with the buffer solution (0.5 ml) in a graduated centrifuge tube and kept at 23° for 30 min with frequent agitation. The mixture was rapidly chilled in ice and immediately centrifuged for 20 min at 900 g. The supernatant and the residual fractions were assayed for invertase and trehalase activities.

(ii) Triton X-100 (octyl phenoxy polyethoxyethanol): a wall suspension was prepared omitting the Triton X-100 washing step in the routine procedure. It was mixed with an equal volume (0.5 ml) of 2% (v/v) Triton X-100 in a graduated centrifuge tube; after standing in an ice bath for 2 h the suspension was centrifuged at 650 g for 10 min. The wall residue was washed with 10 ml distilled water and made up to 1 ml with water.

Invertase activity of the wall residue and supernatant fractions was determined with and without the above Triton X-100 treatment. No loss of invertase activity occurred in controls incubated under identical conditions.

(iii) EDTA (ethylenediaminetetraacetate): a sterilized 8-ml screw-cap vial containing EDTA (0.1 M, 15 µl), wall suspension (2 ml), phosphate buffer (0.6 M, pH 8, 0.25 ml) and water (0.74 ml) was agitated in a gyratory shaker bath at 30° at 280 rev./min for 4 h, then centrifuged at 650 g for 10 min. Invertase activity in the supernatant and residual fractions was determined. Estimations from controls in which EDTA was replaced by water showed that about 5% of invertase activity was lost under the experimental conditions.

(iv) Potassium chloride: procedures were identical to those of EDTA treatment (iii). The incubation mixture consisted of KCl (4 M, 0.75 ml), wall suspension (2 ml) and phosphate buffer (0.6 M, pH 8, 0.25 ml). Only a negligible amount of invertase was inactivated by this treatment.

(v) β-Mercaptoethanol: procedures were identical to those of EDTA treatment (iii). The reaction mixture consisted of β-mercaptoethanol (0.23 M in 0.6 M-phosphate buffer at pH 8, 0.25 ml), wall suspension (2 ml) and water (0.75 ml). About 15% of invertase activity was lost during this treatment.

Hydrolytic enzymes

The hydrolytic enzymes used were: snail-gut juice – Suc d’Helix pomatia (Industries Biologiques Francaises, Gennevilliers, France) processed according to the method of Trevithick & Metzenberg (1964); trypsin (EC 3.4.4.4; Worthington Biochemical Corp., Freehold, New Jersey, U.S.A.) used as a 0.1% solution in tris buffer (0.125 M, pH 8.1) containing 0.03 M-CaCl₂; chitinase (EC 3.2.1.14; Calbiochem Co., Los Angeles, California, U.S.A.) used as a 0.2% solution in phosphate buffer (0.03 M, pH 6.3); cellulase (EC 3.2.1.4; CSEI, Worthington Biochemical Corp.) used as a 0.1% solution in sodium acetate buffer (0.02 M, pH 5); β-1,3-glucanase (EC 3.2.1.6; prepared according to the method of Tanaka & Phaff (1965) from the culture fluid of Bacillus circulans WL-12, a stock culture of which was kindly supplied by Dr H. J. Phaff, University of California, Davis, California; purified Neurospora wall fraction was used as the carbon source instead of yeast wall). A unit of β-1,3-glucanase activity was defined as the amount of enzyme that liberated 1 µmole of reducing sugar/min at pH 5.8 and 37° from laminarin. The β-1,3-glucanase preparation used had a specific activity of 4.8 units/mg protein and a concentration of 0.35 mg protein/ml sodium succinate buffer (0.05 M, pH 5.8).

Substrate specificity of hydrolytic enzymes

Snail-gut juice, glucanase preparations, and chitinase were tested for their glucanolytic specificities with laminarin (Pierce Chemical Co., Rockford, Illinois, U.S.A), pustulan (obtained through the courtesy of E. T. Reese, Quartermaster Research and Engineering
Centre, Natick, Massachusetts, U.S.A.), sodium carboxymethyl cellulose (Hercules Incorp., Wilmington, Delaware) and soluble starch (Fisher Scientific Co., Don Mills, Ontario, Canada) according to the method of Tanaka & Phaff (1965). The chitinolytic activity of these enzyme preparations was assayed according to the procedure in the Worthington Biochemical Enzyme Manual (Worthington Biochemical Corp.) with colloidal chitin prepared from poly-N-acetylglucosamine (Sigma Chemical Co., St Louis, Missouri, U.S.A.) according to the method of Berger & Reynolds (1958). Trypsin, snail-gut juice and glucanase preparations were assayed for proteolytic activities according to the procedure of Drysdale & Fling (1965).

Digestion of Neurospora wall by hydrolytic enzymes

A sterilized and stoppered Erlenmeyer flask (25 ml) containing 10 ml wall suspension, 5 ml hydrolytic enzyme solution, and 50 μl reconstituted penicillin-streptomycin mixture was incubated in a gyratory shaker bath at 30° and 200 rev./min. Samples of 1 to 2 ml were aseptically withdrawn at timed intervals with sterilized pipettes and centrifuged in graduated centrifuge tubes at 650 g for 10 min under refrigeration. Trehalase and invertase activities were measured in the supernatant and residual fractions. In the digestion of wall fraction by β-1,3-glucanase, glucose was released from the wall polymers. This would have interfered with the trehalase and invertase assays which involved a determination of glucose as a hydrolytic product from their substrates (trehalose and sucrose respectively). Therefore the glucose in the digested wall samples was measured before the enzyme assays and subtracted from the total glucose detected after the enzyme assays to yield the experimental values.

Snail-gut juice and cellulase preparations, besides liberating glucose from walls, also had sucrose- and trehalose-cleaving activities. Therefore heat-inactivated (100° for 5 to 10 min) wall suspensions were similarly incubated with these two enzyme preparations. ‘Invertase’ and ‘trehalase’ activities detected in this control were attributed to the snail-gut juice or cellulase preparations and subtracted from the gross activities obtained from the experimental samples.

Preparation of 14C-labelled wall fraction

A conidial suspension (0.1 ml, absorbance at 600 nm = 20) was inoculated into a 125 ml Erlenmeyer flask containing 50 ml Fries minimal medium, sodium succinate buffer (0.04 M, pH 5.2), glucose (0.5 %) and 0.05 mCi D-glucose-14C (U) with a specific activity of 2.8 mCi/mM (Radiochemical Centre, Amersham, Buckinghamshire) for 24 h in a gyratory shaker bath at 30° and 200 rev./min. The mycelium was harvested and 14C-labelled wall fractions prepared according to the procedure already described.

Digestion of 14C-labelled wall by hydrolytic enzymes

14C-Labelled wall suspension (0.6 ml) at a concentration of 1.1 mg of dry wt and 291,682 d.p.m. of radioactivity/ml was added to 6 μl reconstituted penicillin-streptomycin solution (BBL, Div. of BioQuest, Maryland, U.S.A.) and 0.3 ml of a hydrolytic enzyme solution (snail-gut juice, trypsin, chitinase, cellulase, or β-1,3-glucanase) in a sterilized 8 ml screw-cap vial. After 24 h of incubation in a gyratory shaker bath at 30° and 300 rev./min, 100 μl and 200 μl of the sample in each vial were withdrawn separately, spread evenly on glass-fibre filter paper, washed by filtration with 10 ml and 20 ml distilled water, and dried inside a counting vial for over 24 h at 80°. A toluene-based scintillation counting fluid (10 ml: containing 0.1 g I, 4-bis-2-(4-methyl-5-phenyloxazolyl)-benzene (POPOP) and 3 g
Neurospora mural enzymes

2, 5-diphenyloxazole (PPO) per litre of toluene) was added to each vial and the radioactivity of the samples was measured with a Beckman LS-150 Liquid Scintillation System.

Enzyme assays

For all enzyme assays, incubation mixtures containing wall residues were centrifuged at 6000 rev./min for 5 to 10 min with a GLC centrifuge (Sorvall) before the clear supernatant fractions were taken for colorimetric measurements.

(i) Invertase was assayed according to the method of Metzenberg (1962). A unit of activity was defined as the amount of enzyme that released 1 pmole of glucose from sucrose/min at 37°. (ii) Trehalase activity was assayed by incubating 1 ml of reaction mixture containing enzyme solution, potassium phosphate buffer (50 nmole, pH 6) and trehalose (50 pmole) at 37° for 60 min. Incubation was terminated by heating in a boiling water bath for 2 min. The glucose liberated was determined by the glucose oxidase method (Huggett & Nixon, 1957). A unit of activity was defined as the amount of enzyme that hydrolysed 1 pmole of trehalose to glucose/min at 37°.

RESULTS

Treatment with chemical reagents. Buffers from pH 3.0 to pH 10 (about 0.05 M), Triton X-100 (1%, v/v), EDTA (0.5 mM), KCl (1 M), and β-mercaptoethanol (19 mM), in the final concentrations indicated in parentheses, caused a maximum of about 6% of the invertase activity in the wall fractions to appear in the supernatant fractions when compared to controls using water instead of the reagents. Therefore they failed to dissociate invertase from Neurospora wall fractions to any significant extent.

Treatment with hydrolytic enzymes. Snail-gut juice hydrolysed laminarin, pustulan, starch, and chitin very readily; cellulose and casein were hydrolysed to a smaller extent. With such a wide variety of hydrolytic activities, the snail-gut juice released 90% of the invertase and 93% of trehalase from Neurospora wall fractions over 24 h; half of these enzymes were released after 0.25 h and 0.75 h, respectively (Fig. 1, 2). β-1,3-Glucanase, which specifically hydrolysed laminarin, released invertase and trehalase from Neurospora wall fractions more slowly but also more completely than snail-gut juice. After 24 h incubation, 94% of invertase and 96% of trehalase finally appeared in the supernatant fractions; half of each was released within 2 h. Chitinase was rather specific in hydrolysing chitin, with slight hydrolytic activity towards laminarin evident only after 24 h of digestion. Chitinase digestion resulted in 79% of invertase and 63% of trehalase appearing in the supernatant fractions after 29 h of digestion. Trypsin digestion released only 19% of invertase into the supernatant, compared to almost 15% release in the control. The release of trehalase could not be assessed in this experiment since it was severely inactivated by trypsin. However, preliminary investigations showed a pattern of release similar to that obtained for invertase. Cellulase digestion seemed to release about 5% more of invertase and trehalase into the supernatant than was present in the controls after 8 h of incubation. However, after an extended period, up to 24 h, the experimental results coincided with the control values, indicating that no significant amount of enzyme activity was released from the wall fraction. Hence, the effectiveness of the different hydrolytic enzymes in releasing invertase varied rather dramatically, and could be listed in the order of increasing potency as: cellulase, trypsin, chitinase, snail-gut juice and β-1,3-glucanase. The same order also applied to the release of trehalase. The rather high control values may be attributed to autolytic enzymes in walls as demonstrated in Neurospora by Mahadevan & Mahadkar (1970) and in some other fungi by Mitchell & Sabar (1966).
Fig. 1. Release of invertase from Neurospora wall fractions by hydrolytic enzymes. Conditions of the experiment are described in the Methods.

\[
\% \text{ activity released from wall} = \frac{\text{activity in supernatant}}{\text{activity in (supernatant + wall residue)}} \times 100.
\]

The percentage recovery of initial invertase activity after treatment with the hydrolytic enzyme is indicated in parentheses after the enzyme preparation used: SGJ (snail-gut juice) (95 %), G (β-1,3-glucanase) (92 %), Ch (chitinase) (84 %), T (trypsin) (87 %), Ce (cellulase) (92 %), Control (hydrolytic enzyme preparation replaced by 0.02 M-sodium acetate buffer, pH 5.0) (85 %).

**Release of radioactivity from 14C-labelled wall fractions by hydrolytic enzymes.** The ability of the hydrolytic enzymes to release enzymes from the wall fraction did not seem to bear a direct relationship with their ability to digest 14C-labelled wall preparation (Table 1). Trypsin did not release any significant amount of radioactivity although it was slightly more effective than cellulase in releasing the mural invertase. The release of about 10 % of 14C-radioactivity by cellulase seemed to indicate the presence of cellulose-like glucan (β-1,4-linked) in the Neurospora walls. A most marked disproportionality was observed in the chitinase digestion experiment. A mere 16 % of 14C-radioactivity was released from the wall fraction although the amounts of invertase and trehalase released each reached about 70 %. However, β-1,3-glucanase, being more efficient than snail-gut juice in releasing the mural exoenzymes, also released coincidentally more 14C-radioactivity (82 %, cf. 73 % by snail-gut juice).

**DISCUSSION**

Several chemical reagents have been shown to alter the permeability or binding characteristics of bacterial and fungal walls. In *Aspergillus oryzae* the surface-bound α-amylase was released at alkaline pH (Tonomura & Tanabe, 1964). In the Gram-positive bacteria *Staphylococcus aureus* a portion of its penicillinase bound to the cell walls at specific ionic binding sites so that it could be instantaneously displaced by certain inorganic anions or polyanions.
Fig. 2. Release of trehalase from Neurospora wall fractions by hydrolytic enzymes. The percentage recovery of initial trehalase activity after treatment with the hydrolytic enzymes is indicated in parenthesis after the enzyme preparation used (see Fig. 1 for a comparison of activities of invertase released): SGJ (snail-gut juice) (60%), G (β-1,3-glucanase) (78%), Ch (chitinase) (81%), T (trypsin) (47%), Ce (cellulase) (99%), control (84%).

(Coles & Gross, 1967). EDTA is known to increase the permeability of Gram-negative bacterial cell envelopes (Costerton, 1970). Either KCl or mercaptoethanol could release acid phosphatase and invertase, two cell-bound exoenzymes, from *Saccharomyces fragilis* (Weimberg & Orton, 1966; Kidby & Davies, 1970). However, the chemical reagents used in our experiments, including buffers of pH 3 to 10, 1% (v/v) Triton X-100, EDTA, KCl and β-mercaptoethanol, did not release any significant amount of the invertase from the wall fractions.

Table 1. Release of radioactivity from ¹⁴C-labelled wall fractions by hydrolytic enzymes

<table>
<thead>
<tr>
<th>Hydrolytic enzyme</th>
<th>Residual radioactivity (d.p.m.) in wall fraction</th>
<th>Average % of radioactivity released into supernatant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (0 h)</td>
<td>Sample A: 29,408</td>
<td>0%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 57,857</td>
<td></td>
</tr>
<tr>
<td>Trypsin</td>
<td>Sample A: 31,162</td>
<td>9%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 57,540</td>
<td></td>
</tr>
<tr>
<td>Cellulase</td>
<td>Sample A: 27,098</td>
<td>16%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 51,623</td>
<td></td>
</tr>
<tr>
<td>Chitinase</td>
<td>Sample A: 24,705</td>
<td>74%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 47,951</td>
<td></td>
</tr>
<tr>
<td>Snail-gut juice</td>
<td>Sample A: 7,907</td>
<td>87%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 15,156</td>
<td></td>
</tr>
<tr>
<td>1,3-Glucanase</td>
<td>Sample A: 5,437</td>
<td>3%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 9,723</td>
<td></td>
</tr>
<tr>
<td>Control (24 h)</td>
<td>Sample A: 28,445</td>
<td>0%</td>
</tr>
<tr>
<td></td>
<td>Sample B: 55,366</td>
<td></td>
</tr>
</tbody>
</table>

¹⁴C-Labelled wall fractions were incubated with the various hydrolytic enzymes for 24 h; 100 μl (sample A) and 200 μl (sample B) of the incubation mixture were withdrawn. Control samples were similarly incubated with PBS substituting for hydrolytic enzymes. The difference of radioactivity in the wall residues between the control at 0 h and the enzyme digested samples after 24 h was taken to be the amount of ¹⁴C-radioactivity released from the wall fractions.
fractions. Therefore this mural exoenzyme in *Neurospora* must be associated with the walls in ways different from the cases mentioned above.

Three conclusions can be drawn from the experiments on the release of invertase and trehalase from Neurospora wall fractions by various hydrolytic enzymes. First, invertase and trehalase are probably held in the walls in similar ways, and released only after covalent bonds of the wall components are broken. Secondly, these mural exoenzymes are not found exclusively associated with any single chemical component of the walls but the dissolution of glucan (by β-1,3-glucanase) or chitin (by chitinase) or protein (by trypsin) was effective in releasing a fraction of the exoenzymes. Thirdly, the amounts of enzymes released are not directly proportional to the amounts of 14C-labelled wall material that can be solubilized by the hydrolytic enzymes. These conclusions suggest three possibilities that may account for the mural association of invertase and trehalase in *Neurospora*. The first is that the exoenzymes are only physically confined within the matrix of the walls. If this is so, the various hydrolytic enzymes would cause a breakdown in the structural integrity of the wall which consequently becomes permeable to the exoenzymes. The second is that the exoenzymes are covalently bonded to one or more of the wall polymers. A third closely related possibility is that exoenzymes are bound to binding sites associated with one or more of the polymers of the wall, as seems possible for the α-amylase of *Aspergillus oryzae* (Yabuki & Fukui, 1970). Wrathal has (private communication, 1971) recently informed us that wall proteins of low molecular weight may be involved in the association of invertase with the walls of *Neurospora*. In the second and third possibilities, hydrolytic enzymes would not only disrupt the wall integrity but also ‘dissect’ out fragments of the wall polymers or portions of binding proteins which are linked to the enzyme molecules. These enzyme molecules, together with the attached wall fragments, would be able to diffuse through the partially degraded walls.

It is impossible to make a choice among these possibilities now but the following observations would favour the first one. During routine storage of purified wall fractions, a cycle of freeze-thawing was found to release up to 10% of the invertase activity into a supernatant fraction. This possibly indicates that mere mechanical force is sufficient to liberate some invertase from its association with the wall fraction. Hence, in the fraction of enzyme that is so released, no covalent bonds between the enzyme molecule and the wall could have been involved. Furthermore, if the liberated mural exoenzymes are still associated with wall fragments or binding proteins, their sizes would be larger than the same enzyme purified from the soluble fraction of the Neurospora homogenate. However, during disc-gel electrophoresis, invertase liberated from the wall fraction by enzymatic hydrolysis migrates to the same position as the invertase purified from Neurospora homogenate (unpublished data). Therefore, these two types of invertase are similar both in their charge and size characteristics—a property that can be best explained by the hypothesis that mural exoenzymes are only physically confined within the Neurospora wall.

The authors thank Dr C. F. Robinow, Dr D. B. Smith and Dr A. Wellman for helpful discussions during the course of this work. The assistance provided by a grant from the Medical Research Council of Canada is gratefully acknowledged. The paper formed part of a dissertation submitted by P. L. Y. C. in partial fulfilment of the requirements for the degree of Doctor of Philosophy, University of Western Ontario, London, Ontario, Canada.
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