Purification and Characterization of a Chloride ion-Dependent α-Glucosidase from the Midgut Gland of Japanese Scallop (P. yessoensis)
Abstract

Marine glycoside hydrolases hold enormous potential due to their habitat-related characteristics such as salt tolerance, barophilicity, and cold tolerance. We purified an α-glucosidase (PYG) from the midgut gland of the Japanese scallop (*Patinopecten yessoensis*) and found that this enzyme has unique characteristics. The use of acarbose-affinity chromatography during the purification was particularly effective, increasing the specific activity 570-fold. PYG is an interesting chloride ion-dependent enzyme. Chloride ion causes distinctive changes in its enzymatic properties, increasing its hydrolysis rate, changing the pH profile of its enzyme activity, shifting the range of its pH stability to the alkaline region, and raising its optimal temperature from 37°C to 55°C. Furthermore, chloride ion altered PYG’s substrate specificity. PYG exhibited the highest $V_{\text{max}}/K_m$ value toward maltooctaose in the absence of chloride ion and toward maltotriose in the presence of chloride ion.

Keywords

scallop, α-glucosidase, chloride-ion dependent, substrate specificity
Introduction

Marine organisms are attracting attention as a biological resource because they have adapted to the diversity of the oceanic environment, such as high salinity, high pressure, and low temperature, which allows enzymes from marine sources to hold enormous potential [1, 2]. Knowledge of these enzymes is certainly beneficial from a biotechnological perspective. Most of the useful materials obtained from marine resources thus far have been derived from microbes [1]; however, animals, plants, and algae may have hidden potential.

Marine glycoside hydrolases can have just as much potential as any other enzymes. However, studies of the glycoside hydrolases of marine organisms have been inadequate compared with the rigorous studies of the enzymes from land-dwelling creatures. Shelled invertebrates in the marine environment have been shown to possess various polysaccharide-degrading enzymes, such as alginate lyase, cellulase, and mannanase, in their digestive fluids [3-8]. This is because they feed on seaweed and need the ability to digest the abundant polysaccharides, such as alginic acid, cellulose, laminarin, and mannan present in seaweed. Thus, studies on alginate lyase, cellulase, laminarinase, and mannanase can often be found in literature, whereas there is little information on enzymes that digest α-glucans [9].

α-Glucosidase (α-glucoside glucohydrolase, EC 3.2.1.20) catalyzes the liberation of α-glucose from the non-reducing termini of substrates such as maltooligosaccharides and α-glucan. In mammals, α-glucosidase, existing in the small intestine as sucrase-isomaltase and the maltase–glucoamylase complex, is a terminal enzyme in the degradation of starch and its related compounds. These carbohydrates are digested into maltooligosaccharides by salivary and pancreatic α-amylases, and
maltooligosaccharides are digested into glucose by α-glucosidases localized to the small intestine. Thus, α-glucosidase is one of the most important enzymes of α-glucan utilization in mammals.

The digestive glands of several bivalves, including Pecten maximus, Placopecten magellanicus, and Perna viridis, possess α-glucosidase as well as other carbohydrate metabolic enzymes such as amylase, laminarinase, and cellulose [10-12]. Accordingly, α-glucosidase likely plays an important role in the degradation of α-glucan in bivalves. However, to date, there has been no detailed examination of α-glucosidases from bivalves. In this study, we purified an α-glucosidase from the midgut gland of the Japanese scallop Patinopecten yessoensis (PYG) and characterized its enzymatic properties. Kinetic studies indicated that PYG is a chloride ion-dependent enzyme. Chloride ion significantly increases the catalytic activity and alters the substrate specificity of PYG.

Materials and methods

Materials. Midgut glands of scallop (P. yessoensis) were collected in Mutsu Bay of Aomori prefecture, Japan in 2005. Maltose, maltotetraose, maltopentaose, maltohexaose, maltoheptaose, and maltooctaose were kindly gifted by Nihon Shokuhin Kako (Tokyo, Japan). Maltotriose, kojibiose, and nigerose were purchased from Wako Pure Chemical Industries (Osaka, Japan). Other chemicals used were analytical grade and obtained from Nacalai Tesque (Kyoto, Japan) if not otherwise specified.

Enzyme activity assay. Reaction mixtures containing 0.2% (w/v) maltose, 60 mM
sodium acetate buffer (pH 5.0), 0.03% (v/v) Triton X-100, and 300 mM NaCl were
incubated at 37°C. Reactions were terminated by the addition of two volumes of 2 M
Tris-HCl buffer (pH 7.0). Glucose liberated from the substrate was quantified using the
glucose oxidase-peroxidase method [13] performed with the Wako Glucose CII Test
(Wako Pure Chemical Industries). One unit of maltase activity was defined as the
amount of enzyme hydrolyzing 1 µmol of maltose per min under the conditions
described above.

Preparation of acarbose-Sepharose 4 Fast Flow. Acarbose (0.1 mmol) (Sigma-
Aldrich, St. Louis, MO, USA) was stirred overnight with 3.5 mmol 2-(4-
aminophenyl)ethylamine (Tokyo Chemical Industry, Tokyo, Japan). The resulting
aminoethylated acarbose was reduced with sodium borohydride (0.63 mmol) in ethanol
(1.5 mL) for 5 h at room temperature. The mixture was titrated with 1 M acetic acid to
pH 5.6 and then applied to a Sephadex G-25 column equilibrated with 1 M sodium
acetate buffer (pH 5.0). Upon isocratic elution with 1 M sodium acetate buffer (pH 5.0),
the fractions containing modified acarbose were identified by using thin-layer
chromatography on silica gel 60 (nitromethane/1-propanol /water, 4/10/3). The plates
were visualized by charring with a methanolic solution of sulfuric acid containing
0.02% α-naphthol. The modified acarbose was coupled with NHS-activated Sepharose
4 Fast Flow (GE Healthcare, Buckinghamshire, England) by following the supplier’s
instruction.

Purification of PYG. Midgut glands (52 g) were stirred in 10 mM sodium acetate
buffer (pH 5.0) containing 5 mM MgCl₂ and 500 mM NaCl overnight at 4°C to extract
protein. The supernatant (crude enzyme) was recovered by centrifugation at 18,700 \times g for 20 min at 4°C. Ammonium sulfate was added to the crude enzyme until its concentration reached 70% saturation, and the suspension was left overnight at 4°C. The precipitate was collected by centrifugation at 18,700 \times g for 20 min. The precipitate was dissolved in 10 mM sodium acetate buffer (pH 5.0) containing 5 mM MgCl₂ and 30% saturated ammonium sulfate and then loaded onto a TOYOPEAL Butyl-650M (Tosoh, Tokyo, Japan) column equilibrated with the same buffer. Absorbed proteins were eluted with a linear gradient of 30%–0% saturated ammonium sulfate in 10 mM sodium acetate buffer (pH 5.0) containing 5 mM MgCl₂. The active fractions were pooled and subjected to acarbose-affinity chromatography at room temperature. An acarbose-Sepharose 4 Fast Flow column (1.5 cm × 11 cm) was equilibrated with 10 mM sodium acetate buffer (pH 5.0) containing 5 mM MgCl₂ and 200 mM NaCl. Bound enzyme was eluted with 100 mL of 4 M Tris-HCl buffer (pH 7.0). The eluted fractions were pooled and dialyzed against 10 mM sodium acetate buffer (pH 5.0) containing 5 mM MgCl₂. The dialysate was concentrated using Amicon Ultracel-30K Ultra Centrifuge Filters (Merck Millipore, Billerica, MA, USA). The concentrated solution was loaded onto a Sephacryl S-200HR (GE Healthcare) column (1.6 cm × 55 cm) equilibrated with 10 mM sodium acetate buffer (pH 5.0) containing 5.0 mM MgCl₂ and 200 mM NaCl. The active fractions were pooled and dialyzed against a 10 mM sodium acetate buffer (pH 5.0). Purification procedures, except acarbose-affinity chromatography, were performed at 4°C. The protein concentration of the purified enzyme was calculated from its absorbance at 280 nm in a 1-cm cuvette using an extinction coefficient of (1.0 g/L)⁻¹·cm⁻¹.

_Polyacrylamide gel electrophoresis._ SDS-PAGE and Blue Native-polyacrylamide gel
electrophoresis (BN-PAGE) were performed using a Mini-Protean III cell apparatus (Bio-Rad, Richmond, CA, USA). SDS-PAGE was performed as described by Laemmli [14] and BN-PAGE as described by Schagger and von Jagow [15]. Protein was stained with Coomassie Brilliant Blue (CBB) R-250 using a Rapid CBB staining kit (Kanto Chemical, Tokyo, Japan). Mark12 unstained protein standards and Native mark unstained protein standard (Invitrogen, Life Technologies, Carlsbad, CA, USA) were used as molecular size markers in SDS-PAGE and BN-PAGE, respectively. Activity staining after BN-PAGE was done by incubating the gel in 100 mM sodium acetate buffer (pH 5.0) containing 1 mg/ml 4-methylumebelliferyl α-D-glucopyranoside (Sigma-Aldrich) and 500 mM NaCl at 37°C for 10 min. Fluorescence from the 4-methylumbelliferone produced by hydrolysis was detected using a UV transilluminator.

In-gel digestion with Trypsin. Purified PYG (15 µg) was subjected to SDS-PAGE, and a CBB-stained protein band was excised. The gel piece was washed three times with 100 µL of 50 mM NH₄HCO₃ in 50% (v/v) acetonitrile for 20 min at 37°C then dried using a vacuum centrifugation. The dried gel was swollen in 5 µL trypsin solution (1 mg/mL) containing 1 mM HCl and 20 mM CaCl₂ at 4°C for 10 min. After reswelling the gel, 15 µL of 50 mM NH₄HCO₃ was added and digestion was continued for 24 h at 37°C. Peptides were extracted three times in 100 µL of 60% (v/v) acetonitrile, lyophilized, and redissolved in 0.1 % trifluoroacetic acid containing 50% (v/v) acetonitrile.

Matrix-assisted laser desorption ionization-time of flight mass spectrometric analysis.
The tryptic peptides were desalted using a Zip-Tip C-18 (Merck Millipore, Billerica, MA, USA) according to the manufacturer’s instructions. The elutate (2 µL) was mixed with 0.8 µl of matrix solution [10 mg/ml α-cyano-4-hydroxycinnamic acid (Sigma-Aldrich) in 0.1% trifluoroacetic acid/50% (v/v) acetonitrile], and dried at room temperature on the sample plate. Mass spectra were recorded with a matrix-assisted laser desorption ionization-time of flight mass spectrometer (MALDI-TOF-MS) (Voyager-DE STR, Applied Biosystems, Foster City, CA, USA) with delayed ion extraction technology. All mass spectra were obtained in reflector mode and calibrated with GRAMS software (Applied Biosystems) using angiotensin II as an external mass standard. We used the MALDI-TOF-MS in the facilities of the Research Faculty of Agriculture, Hokkaido University.

**Effect of various salts on enzyme activity.** The enzyme reactions were performed in 60 mM sodium acetate buffer (pH 5.0) containing 0.03% (v/v) Triton X-100, 0.2% maltose, and diluted enzyme (102 ng/mL) at 37°C while varying the concentrations of various salts (0 mM–400 mM MgSO4, NaCl, KCl, MgCl2 or CaCl2). The liberated glucose was measured as described above.

**Effects of pH and temperature.** The effect of pH on maltose hydrolysis activity was determined using the standard assay conditions with the exception that 60 mM sodium acetate (pH 3.6–pH 5.6), 60 mM sodium phosphate (pH 5.4–pH 8.2) or Britton-Robinson (pH 2.2–pH 6.9) buffers were used. Enzyme concentrations used were 406 ng/mL (in the absence of NaCl) and 81.2 ng/mL (in the presence of 300 mM NaCl). For pH–stability experiments, the purified enzyme (203 µg/mL) diluted with 40 mM Britton-
Robinson buffer (pH 2.0–pH 11.0) containing 0.05% (v/v) Triton X-100 were kept at 4°C for 24 h in either the absence or presence of 300 mM NaCl. The remaining activity was measured under the standard assay conditions except that the remaining activity of samples treated in the absence of NaCl was measured using the NaCl-free buffer. The optimal temperatures for the hydrolytic activity of the enzyme in the absence of NaCl or in the presence of 300 mM NaCl were measured under the standard assay condition except that the temperature was varied from 25°C to 65°C. Enzyme concentrations used were 406 ng/mL (in the absence of NaCl) and 102 ng/mL (in the presence of 300 mM NaCl). For thermal stability experiments, the purified enzyme (508 ng/mL in the absence of NaCl and 102 ng/mL in the presence of 300 mM NaCl) in 10 mM sodium acetate buffer (pH 5.0) containing 0.05% (v/v) Triton X-100 was incubated at 30°C to 65°C for 15 min. The residual activity was assayed under the standard assay conditions except that the remaining activity treated without NaCl was measured using NaCl-free buffer.

Hydrolysis of maltooligosaccharides and various disaccharides. The initial velocities of the hydrolysis reactions performed using 2 mM concentrations of various substrates (maltose, maltotriose, maltotetraose, maltopentaose, maltohexaose, maltoheptaose, maltooctaose, trehalose, kojibiose, and nigerose) were measured in the absence or presence of 300 mM NaCl under standard assay conditions. Enzyme concentrations used were 0.102 µg/mL to 40.6 µg/mL. The initial velocity was expressed as the number of µmol of glucose released from the non-reducing end of the substrate per min per mg protein. When determining the hydrolysis rate for disaccharides, the concentration of liberated glucose was divided by two.
Kinetic parameters. The kinetic parameters ($V_{\text{max}}$ and $K_m$) for kojibiiose, nigerose, and maltooligosaccharides in the absence or presence of 300 mM NaCl were calculated using the initial velocities of reactions containing six to eight different substrate concentrations (0.2 mM–32 mM). These initial rates were fitted to the Michaelis-Menten equation using the computer program Kaleidagraph 3.6. The initial velocities were measured in the standard assay conditions except for the NaCl concentration. The enzyme concentration used was 127 ng/mL.

Results and Discussion

Purification of PYG

PYG was purified using a combination of ammonium sulfate precipitation and several column chromatographic steps, including acarbose-affinity chromatography, which was performed using acarbose covalently cross-linked to a Sepharose 4 Fast Flow column. The enzyme preparation was subjected to electrophoresis to confirm the purity of the enzyme. The enzyme migrated as a single protein band with a molecular mass of about 66 kDa on BN-PAGE, stained by CBB R-250 or visualized by activity staining (Fig. 1A). However, on SDS-PAGE, the enzyme separated into two components (Fig. 1B). The molecular masses of the major and minor components were 51 kDa and 40 kDa, respectively. The results of PAGEs imply that these components do not dimerize, and exist as a monomer. The components could not be separated using further purification procedures including ion-exchange, gel filtration, and hydrophobic interaction chromatographies. MALDI-TOF-MS of the mixture of tryptic fragments generated by in-gel digestion of the 51 kDa and 40 kDa components indicated that nine peptides derived from each component were identical (Fig. 2). It was likely that the 51
and 40 kDa components were derived from the same peptide, which underwent different proteolysis. The midgut gland contains proteases since it is a digestive tract with functions similar to those of the liver and pancreas in mammals [16]. It is possible that PYG undergoes a proteolytic process that splits it into two components. The enzyme preparation was regarded as the purified enzyme (PYG) and was thus used for further analysis. Peptide masses determined by MALDI-TOF-MS were used in database searches by Mascot; however, no glycoside hydrolase was identified as a significant match. The purification process yielded 0.427 mg of purified PYG, whose specific activity was 122 U/mg, from 52 g of midgut glands with a yield of 39.8%. The PYG purification procedures are summarized in Table 1. Acarbose-affinity chromatography was the most effective step, increasing the specific activity 570-fold.

Effect of various salts on the hydrolysis rate

The effect of MgCl₂ and MgSO₄ on PYG activity was investigated by measuring the rate of maltose hydrolysis (Fig. 3). The hydrolysis rate in 60 mM sodium acetate buffer without any salts was 13.1 µmol/min/mg. MgSO₄ doubled the hydrolysis rate: the hydrolysis rate in the presence of 400 mM MgSO₄ was 26.4 µmol/min/mg. MgCl₂ increased more efficiently than the sulfate ion: the hydrolysis rates in the presence of 400 mM MgCl₂ was 124 µmol/min/mg. The effect of MgCl₂, NaCl, KCl, and CaCl₂ on PYG activity was investigated (Fig. 3). Every salt increased the enzyme activity in the same manner and the activity was saturated at the concentration of 40 mM. The hydrolysis rates in the presence of sufficient concentration (400 mM) of salts were 124 (MgCl₂), 122 (NaCl), 123 (KCl), and 128 (CaCl₂) µmol/min/mg, respectively. These results indicate that chloride ion increased the hydrolytic activity efficiently, but the
identity of the cation had little effect on the activity.

\[ pH \text{ and thermal properties of PYG} \]

The optimal pH of maltose-hydrolytic activity and pH stability of PYG in the presence or absence of NaCl were examined. We used 300 mM NaCl, in which the activity was adequately enhanced, as the salt-existence condition. The optimal pH values in the absence and presence of 300 mM NaCl were 3.9 and 4.0, respectively (Fig. 4A). The pH-activity profiles showed a bell-shaped curve with or without of NaCl. The enzyme was incubated at various pHs in the presence or absence of NaCl at 4°C for 24 h, and its residual activity was measured. The stable range was defined as the pH range exhibiting more than 90% residual activity. NaCl shifted the pH-stability range to the alkaline region. PYG without NaCl was stable in a pH range from 3.2 to 6.3, whereas PYG with NaCl was stable in a pH range from 4.1 to 7.3 (Fig. 4B). In addition, when the enzyme was treated at pH 10.0, the residual activity was 50% in the presence of NaCl, whereas the enzyme was almost inactive in the absence of NaCl.

The optimal temperature and thermal stability in the presence or absence of NaCl was examined. The optimal temperature increased from 37°C to 55°C with the addition of NaCl (Fig. 4C). The thermal stability of the enzyme in the presence or absence was investigated based on the measurement of the residual activity after heat-treatment for 15 min. The stable region was defined as the temperature range exhibiting more than 90% residual activity. PYG was stable up to 40°C under both NaCl conditions. However, the residual activity at 45°C was different in the presence or absence of NaCl. The enzyme exhibited 80% residual activity in the presence of NaCl, whereas the residual activity was 10% in the absence of NaCl. (Fig. 4D).
The increasing of pH and thermal stabilities with the addition of NaCl might be explained by facilitating energetic interactions within the protein structure and by the stabilization of surface-exposed residues. *Alteromonas haloplanctis* α-amylase is stabilized by facilitating energetic interactions between the protein structure and the chloride ion [17]. Furthermore, monovalent cation, including potassium and sodium ions, enhances the overall thermal stability of several enzymes by stabilization of surface-exposed residues [18]. The thermal stabilization of PYG might be caused by the similar manner as these enzymes.

**Substrate specificity of PYG**

The kinetic parameters for the hydrolysis of maltooligosaccharides, kojiibiose, and nigerose in the presence or absence of 300 mM NaCl were determined to evaluate substrate specificity (Table 2). PYG readily hydrolyzed maltooligosaccharides, nigerose, and kojiibiose, but hardly hydrolyzed trehalose and isomaltose. The hydrolytic rates of 2 mM isomaltose and 2 mM trehalose in the presence of 300 mM NaCl were $1.38 \times 10^{-2}$ and $4.15 \times 10^{-2}$ μmol/min/mg, respectively. NaCl affected $K_m$ values more effectively than $V_{max}$ values. The $V_{max}/K_m$ values for disaccharides followed the order, maltose>nigerose>kojibiose in the presence or absence of NaCl. However, NaCl had an effect on the preference for maltooligosaccharide chain length. In the absence of NaCl, $V_{max}/K_m$ for maltooligosaccharides increased with increasing chain length; the highest $V_{max}/K_m$ (57.1 μmol/min/mg/mM) was observed for the hydrolysis of maltooctaose. On the other hand, in the presence of NaCl, PYG exhibited the highest $V_{max}/K_m$ with maltotriose (370 μmol/min/mg/mM). $V_{max}/K_m$ values for maltooligosaccharides longer than maltotriose were similar. On the other hand, the $V_{max}$ and $K_m$ values with
maltohexaose were lower than those with other maltooligosaccharides. Although it is hard to give a feasible reason for the lower $V_{\text{max}}$ and $K_m$, the lower $K_m$ seems to decrease the $V_{\text{max}}$ value. As mentioned above, the $V_{\text{max}}/K_m$ value for maltohexaose is similar to that for other maltooligosaccharides. It suggests that the activation free energies for the reactions of the free enzyme and these maltooligosaccharide substrates may be similar, and the transition state of the enzyme–maltohexaose may be stabilized equally to other maltooligosaccharides. Meanwhile, the lower $K_m$ makes the reaction fall into a thermodynamic pit, and causes a rise in a free-energy barrier from the Michaelis complex to the transition state.

Possible mechanism for chloride ion dependence of PYG

The hydrolytic rate of PYG was significantly increased by chloride ion, suggesting PYG is chloride ion-dependent enzyme. A few chloride ion-dependent glycoside hydrolases have been reported thus far [19-21]. Human pancreatic $\alpha$-amylase (HPA), a well-studied chloride ion-dependent enzyme [22-24], catalyzes the same $\alpha$-glucoside hydrolysis as PYG and could furnish information on the mechanism of the chloride-ion dependency of PYG. Biochemical and structural studies of HPA have indicated that chloride ion makes the orientation of an acid/base catalyst accurate, thereby increasing its p$K_a$ and enhancing activity. The enhancement of PYG catalytic activity by chloride ion was associated with a small change in the bell-shaped pH-rate profile of PYG (Fig. 4A); thus, chloride ion might be connected to a catalytic residue as in HPA.

PYG underwent not only an increase in catalytic activity but also a change of substrate specificity toward maltooligosaccharides in the presence of chloride ion. In the
case of HPA, chloride ion causes the conformational change of substrate-binding loop. It is possible that chloride ion also causes a conformational change in PYG and that the change in specificity is caused by a conformational change in the substrate binding site.

In conclusion, we succeeded in finding a unique α-glucosidase from a marine source, purifying PYG from the midgut gland of the scallop, and determining the unique properties in this enzyme. The catalytic activity of PYG was evidently enhanced by chloride ion. Furthermore, substrate specificity was altered by chloride ion. To the best of our knowledge, this is the first report of a chloride ion-dependent α-glucosidase. Unfortunately, we could not clarify the molecular mechanism of the chloride-ion dependency at this stage because no structural information exists. It will be necessary to determine the structure of PYG to elucidate the molecular mechanism responsible for the chloride-ion dependency.

**Author contributions**


**Acknowledgements**

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References


### Table 1. PYG purification summary

<table>
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<tr>
<th>Procedure</th>
<th>Total protein (mg)$^a$</th>
<th>Total activity (U)</th>
<th>Specific activity (U/mg)</th>
<th>Purification (-fold)</th>
<th>Recovery (%)</th>
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<tr>
<td>Crude extract</td>
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<td>131</td>
<td>0.0110</td>
<td>1.00</td>
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<tr>
<td>Ammonium sulfate 60-30% fractionation</td>
<td>3,350</td>
<td>124</td>
<td>0.0370</td>
<td>3.36</td>
<td>94.6</td>
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<td>TOYOPEARL Butyl-650M</td>
<td>636</td>
<td>150</td>
<td>0.236</td>
<td>21.4</td>
<td>115</td>
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<td>Acarbose Sepharose 4 Fast Flow</td>
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<td>135</td>
<td>134</td>
<td>12,100</td>
<td>93.9</td>
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<td>Sephacryl S-200</td>
<td>0.426</td>
<td>52.1</td>
<td>122</td>
<td>11,100</td>
<td>39.8</td>
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$^a$ calculated under the assumption that $E_{280nm,1cm}^{1\%} = 10.0$
Table 2. PYG kinetic parameters in the absence and presence of 300 mM NaCl

<table>
<thead>
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<th>Substrate</th>
<th>Absence of NaCl</th>
<th>Presence of NaCl</th>
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<tr>
<td></td>
<td>$V_{\text{max}}$</td>
<td>$K_{m}$</td>
</tr>
<tr>
<td></td>
<td>(µmol/min/mg)</td>
<td>(mM)</td>
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<tr>
<td>Maltose</td>
<td>41.3 ± 0.9</td>
<td>11.1 ± 0.9</td>
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<td>Maltotriose</td>
<td>34.3 ± 1.4</td>
<td>7.47 ± 0.5</td>
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<td>Maltotetraose</td>
<td>32.9 ± 0.1</td>
<td>4.48 ± 0.02</td>
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<td>Maltopentaose</td>
<td>27.9 ± 0.9</td>
<td>4.43 ± 0.55</td>
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<td>Maltohexaose</td>
<td>32.2 ± 1.8</td>
<td>7.02 ± 0.44</td>
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<tr>
<td>Maltoheptaose</td>
<td>67.4 ± 2.9</td>
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<tr>
<td>Maltooctaose</td>
<td>79.7 ± 0.04</td>
<td>1.39 ± 0.32</td>
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<tr>
<td>Kojibiose</td>
<td>8.96 ± 0.14</td>
<td>17.9 ± 0.41</td>
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<tr>
<td>Nigerose</td>
<td>18.6 ± 0.39</td>
<td>13.1 ± 0.33</td>
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Figure legends

Fig 1. Electrophoretic analysis of PYG.
   A: BN-PAGE of PYG. Lane M, marker proteins; lanes 1 and 2, PYG (4 µg protein for each lane) visualized by protein-staining and activity-staining, respectively.
   B: SDS-PAGE of PYG. Lane M, marker proteins; lane 1, 5.5 µg PYG.

Fig 2. MALDI-TOF-MS analysis of PYG tryptic peptides.
   A: MALDI-TOF-MS spectra of tryptic peptides obtained from 51 kDa component. B: MALDI-TOF-MS spectra of tryptic peptides obtained from 40 kDa component.

Fig 3. Effect of various salts on enzyme activity.
   Maltose hydrolysis rates measured in the presence of 0–400 mM MgCl₂ (■), CaCl₂ (□), KCl (●) NaCl (○), and MgSO₄ (▲) represent in panel at left. Panel at right shows an enlarged view at low concentration of salts (0–5 mM).

Fig 4. Effect of NaCl on the pH-activity profile (A), the pH stability (B), the temperature-activity profile (C) and the thermal stability (D) of PYG in the absence (○) and presence (●) of 300 mM NaCl. A: Enzyme activity was measured at various pH values and 37°C. B: Enzyme was incubated at various pH values at 4°C for 24 h, and the residual activity was measured at pH 5.0 and 37°C. C: Enzyme activity was measured at the indicated temperature and pH 5.0. D: Enzyme was incubated at various temperature at pH 5.0 for 15 min, and residual activity was measured at pH 5.0 and 37°C.
Fig. 1 Masuda et al
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