Isomaltulose Synthase from *Klebsiella* sp. Strain LX3: Gene Cloning and Characterization and Engineering of Thermostability

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The gene (palI) encoding isomaltulose synthase (PalI) from a soil bacterial isolate, *Klebsiella* sp. strain LX3, was cloned and characterized. PalI converts sucrose into isomaltulose, trehalulose, and trace amounts of glucose and fructose. Sequence domain analysis showed that PalI contains an α-amylase domain and (β/α)7-barrel structures, suggesting that it belongs to the α-amylase family. Sequence alignment indicated that the five amino acid residues of catalytic importance in α-amylases and glucosyltransferases (Asp134, Glu296, Asp309, His145, and His309) are conserved in PalI. Purified recombinant PalI displayed high catalytic efficiency, with a *K*~m~ of 54.6 ± 1.7 mM for sucrose, and maximum activity (approximately 328.0 ± 2.5 U/mg) at pH 6.0 and 35°C. PalI activity was strongly inhibited by Fe3+ and Hg2+ and was enhanced by Mn2+ and Mg2+. The half-life of PalI was 1.8 min at 50°C. Replacement of selected amino acid residues by proline significantly increased the thermostability of PalI. Simultaneous replacement of Glu396 and Arg310 with proline resulted in an 11-fold increase in the half-life of PalI at 50°C.

Isomaltulose (6-O-α-d-glucopyranosyl-d-fructose, commonly referred to as palatinose) is a sucrose isomer with physical and organoleptic properties similar to those of sucrose. It has been suggested as a noncariogenic alternative to sucrose (1) and has now been widely used as a sugar substitute in food. Unlike ingestion of sucrose, ingestion of isomaltulose has only a minor effect on the concentration of glucose in blood, indicating its potential as a parenteral nutrient acceptable to diabetics and nondiabetics (6). Several microorganisms have been found to form isomaltulose and trehalulose from sucrose, for example *Protaminobacter rubrum* (39), *Serratia plymuthica* (5), *Erwinia rhapontici* (4), *Klebsiella planticola* CCRC 19112 (7), *Pseudomonas mesoacidophila* MX-45 (18), and *Agrobacterium radiobacter* MX-232 (19).

Isomaltulose synthase, also known as sucrose isomerase, has been purified from *S. plymuthica* (14, 33), *E. rhapontici* (4), a *Klebsiella* sp. (24), and *Pseudomonas mesoacidophila* MX-45 (18). In addition to isomerizing sucrose to produce isomaltulose and trehalulose, the enzyme reaction also releases small amounts of glucose and fructose as by-products (7, 14, 31, 33). The isomaltulose synthase from *S. plymuthica* even converted sucrose to isomaltose and isomelezitose (33). The product composition varies depending on the bacterial strain used. *Protaminobacter rubrum* (15), *S. plymuthica* NCIB 8285 (5), *E. rhapontici* NCPPB 1579 (4), and *K. planticola* CCRC 19112 (7) produce mainly isomaltulose (75 to 85%), whereas *Pseudomonas mesoacidophila* MX-45 (17) and *A. radiobacter* MX-232 (19) produce more trehalulose (90%) than isomaltulose. The reaction of these enzymes is strongly influenced by temperature. The optimum temperature for isomaltulose production by the enzyme from *S. plymuthica* ATCC 15928 is around 30°C, and no product is produced at 55°C (33), indicating the thermolability of isomaltulose synthase. Isomaltulose synthase sequence information is required to clarify the molecular mechanism of isomerization and assess the feasibility of engineering thermostability.

We recently identified a new isomaltulose-producing bacterial isolate, *Klebsiella* sp. strain LX3 (L.-H. Zhang, unpublished data). Biochemical analyses showed that this isolate converts sucrose mainly into isomaltulose, with small amounts of trehalulose and trace amounts of glucose and fructose. In the present study, we report the cloning and characterization of the gene (palI) encoding isomaltulose synthase (PalI) from *Klebsiella* sp. strain LX3. The recombinant enzyme was overexpressed in *Escherichia coli*, purified, and characterized. In addition, site-directed mutagenesis was performed based on protein domain and secondary structure analysis to improve the thermostability of PalI by proline substitution.

**MATERIALS AND METHODS**

**Bacterial strains and chemicals.** The bacterial strains and plasmids used in this study are listed in Table 1. *Klebsiella* sp. isolate LX3 was isolated from a soil sample collected in Singapore. Cosmid vector pLAFR3 and cloning vector pBluescript II SK(+) (Stratagene) were used for construction of a genomic library and for subcloning, respectively. *E. coli* DH5α was used as the host for cloning and overexpression of palI. *E. coli* was grown at 37°C in Luria-Bertani (LB) medium containing 100 μg of ampicillin/ml. *Klebsiella* sp. strain LX3 was grown at 30°C in SPY medium (4% sucrose, 0.5% peptone, and 0.4% yeast extract, pH 6.5).

**DNA manipulation.** DNA manipulation, agarose gel electrophoresis, and transformation of *E. coli* were performed according to standard procedures. DNA fragments were isolated from agarose gels with the QIAEX II gel extraction kit (Qiagen), and plasmids were purified with the Qiagen plasmid minikit. A DNA library was constructed by partially digesting DNA from *Klebsiella* sp. strain LX3 with BamHI, cloning the fragments into the BamHI site of cosmids vector pLAFR3, and transfecting into *E. coli* DH5α after in vitro packaging with Gigapack II (Stratagene). Following transformation, individual bacterial colonies were incubated in liquid LB medium supplemented with sucrose (0.4%) for 15 h, and the reducing sugar released in the supernatant was tested using the dinitrosalicylic acid method (16). The presence of isomaltulose was further verified by thin-layer chromatography (TLC), using authentic isomaltulose as control.
pGEK:E498P was produced using pGEK as a template. Double mutant pGEK was introduced by using the QuickChange site-directed mutagenesis kit (Stratagene). Mutant DNA was verified by DNA sequencing.

For the kinetic analysis, 2-min reaction velocities were determined using the Bradford method (3) with bovine serum albumin as a standard. The protein concentration was determined by the method of Lowry et al. (35) using a 0.5% Bio-Rad reagent at 550 nm.

Computer analysis. DNA and peptide sequence homologies were analyzed using the BLAST program to search major databases (GenBank, EMBL, and SWISSPROT databases). Sequence alignment was performed using the Genetics Computer Group program (http://www.bic.nus.edu.sg:8888/gcg/java/). Domain analysis was carried out with the Pfam program (http://pfam.wustl.edu). Signal peptide prediction was performed with the SignalP program (23). The secondary structure of PalI was predicted using the PredictProtein program (25) (http://dodo.cpmc.columbia.edu).

Construction of GST-PalI. The palI gene without the coding sequence for the first 28-amino-acid signal peptide was amplified by PCR using the forward primer 5'-TTGGATCCGCACCATCCTTGAATCAGGATATTCAC-3' and the reverse primer 5'-TTGAATCTTACCGCAGCTTATACACACCTGCCTG-3'. After digestion with BamHI and EcoRI, the fragment was fused to the glutathione-S-transferase (GST) gene in the same open reading frame (ORF) in fusion vector pGEX-2T to generate an expression construct, pGEK, which was verified by DNA sequencing.

Site-directed mutagenesis. A point mutation was introduced into the palI gene by using the QuickChange site-directed mutagenesis kit (Stratagene). Mutant pGEK:E498P was produced using pGEK as a template. Double mutant pGEK:E498P/R310 was created using pGEK:E498P as a template. Mutations were verified by DNA sequencing.

Enzyme purification. E. coli strains containing different constructs were incubated in LB medium at 28°C. The GST-PalI fusion protein and its derivatives were overexpressed by the addition of isopropyl-β-D-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM after the optical density at 600 nm of the bacterial culture reached 0.6. Purification of fusion proteins was carried out by a method described previously (40). Further purification was performed by fast protein liquid chromatography with a Protein Pak 300SW semipreparative column at a flow rate of 0.5 ml/min with sodium phosphate buffer (50 mM, pH 7.4). Purified enzymes were analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (10% polyacrylamide).

Enzyme assay. A sample (100 μl, 2 μg of protein) of enzyme was mixed with 400 μl of 0.1 M citrate-phosphate buffer (pH 6.0) containing sucrose (40 g/liter) and incubated at 35°C for 15 min with gentle agitation. The reaction was stopped by boiling for 5 min, and the total reducing sugar was assayed using the dinitrosalicylic acid method (16). One unit of enzyme is defined as the amount of protein that forms 1 μmol of reducing sugar (with isomaltulose as a standard) per min under the assay conditions specified. The protein concentration was determined using the Bradford method (3) with bovine serum albumin as a standard. For the kinetic analysis, 2 μg of the purified enzyme was mixed with different concentrations of sucrose (10 to 150 mM) dissolved in 0.1 M citrate-phosphate buffer (pH 6.0) at a final reaction volume of 500 μl. The initial reaction velocity was determined after incubation at 35°C for an appropriate reaction time. The kcat and Km values were calculated based on the Michaelis-Menten equation.

Influence of pH on activity and stability. Reaction solutions were prepared by adding sucrose to a final concentration of 4% and 2.0 μg of PalI in 0.1 M citrate-sodium phosphate buffer at a range of pH values, except that 0.05 M Tris buffer was used at pH 8.5. To investigate the effect of pH on stability, the same amount of enzyme was added to each reaction buffer and maintained at room temperature for 1 h before addition of sucrose. Enzyme activity was assayed as described above.

Effect of temperature on enzyme activity. The effect of temperature on enzyme activity was determined by incubating the reaction solution (pH 6.0) at the temperatures indicated for 15 min. To determine the effect of temperature on enzyme activity, the enzyme solution (pH 6.0) was subjected to different temperatures for 10 min and the residual activity was assayed. Enzyme thermostability was determined by incubating the enzyme at 50°C for various periods, and the remaining activity was determined at different intervals as stated. The first-order constant, k, of irreversible thermostabilization was obtained by linear regression in semilog coordinates. The enzyme half-life was calculated using the equation t1/2 = ln 2/k.

Effects of metal ions and other reagents on PalI activity. Purified PalI was extensively dialyzed against a 50 mM citrate-phosphate buffer (pH 6.0). Samples (2.0 μl) were preincubated with metal ions and other reagents in various concentrations at room temperature for 10 min (in a 200-μl final volume). Aliquots (2.0 μl) were withdrawn and tested for activity as described above.

Effect of temperature and pH on product formation. To investigate the influence of temperature on product formation, the conversion reaction was performed for 4 h in a test tube containing 2 ml of 4% sucrose solution (pH 6.0) and 10 μg of PalI at the temperatures indicated. The reaction mixture was boiled for 5 min and subjected to high-pressure liquid chromatography (HPLC) analysis. The pH experiments were conducted at a fixed temperature of 35°C.

Analytical methods. Qualitative analysis of isomaltoolose, glucose-fructose, and trehalulose was carried out using TLC in an ethyl acetate-acetic acid-water (4:3:1 by volume) solvent system. Sugar spots were visualized by spraying TLC plates with diphenylamine–aniline–85% phosphate reagent and incubating for 5 min at 80°C (27). Quantitative analysis of sugars was performed by HPLC (Waters 2690 separation modules) on a Symmetry C18 column (4.6 by 250 mm) coupled to a Waters 2410 refractive index detector. Samples were eluted isocratically with water at a flow rate of 1 ml/min. Under these conditions, the retention times of 1kb.  

**FIG. 1.** Cloning of the palI gene from Klebsiella sp. strain LX3. B, BamHI; EV, EcoRV; C, ClaI; X, XhoI; P, PstI; +, PalI activity detected; −, no activity. The arrow indicates the relative position of the palI coding region. The size of the insert in each clone is listed in parentheses.
trehalose, glucose, fructose, isomaltulose, and sucrose were 2.20, 2.67, 2.87, 3.25, and 3.87 min, respectively. Each sugar was quantified based on peak area.

Nucleotide sequence accession number. The nucleotide and peptide sequences of the Klebsiella sp. strain LX3 isomaltulose synthase gene (palI) were deposited in the GenBank database under accession number AY040843.

RESULTS

Cloning and sequence analysis of the palI gene, encoding isomaltulose synthase. The cosmId library of a Klebsiella sp. strain was screened for isomaltulose biosynthesis activity. Two positive clones, designated pPLAK164 and pPLAK169, were selected from about 1,500 colonies for their ability to produce isomaltulose. Restriction enzyme analysis of the two clones revealed that they contained the same BamHI fragment of ~6.0 kb. Deletion analysis narrowed down the region encoding isomaltulose biosynthesis to a 2.5-kb EcoRV fragment in the subclone pBKEE (Fig. 1).

DNA sequence analysis of pBKEE identified an ORF of 1,797 nucleotides. Deletion analysis confirmed that it was the coding region of isomaltulose synthase (data not shown). The ORF encodes a protein of 598 amino acids with a calculated molecular mass of 69.94 kDa and an isoelectric point at 6.62. The first 28 amino acids (Met1 to Ala28) at the N-terminal constitute a signal peptide, as predicted by the SignalP computer program (23). Deletion of the signal peptide resulted in accumulation of isomaltulose synthase in the cytoplasm of bacterial cells (data not shown).

Homology comparison of the isomaltulose synthase from Klebsiella sp. strain LX3. The best homolog of the isomaltulose synthase from Klebsiella sp. strain LX3 is the isomaltulose synthase from Enterobacter sp. strain SXZ62, reported in a patent publication (15). The two enzymes share 99.3% homology at the nucleotide sequence level and 99.7% homology at the peptide sequence level. There are differences in the C-terminal regions of the enzymes. The Klebsiella sp. strain LX3 isomaltulose synthase contains an extra amino acid, Ala297, and differs in two residues (Ala578 and Val594) from isomaltulose synthase from Enterobacter sp. strain LX3.

Pall of Klebsiella sp. strain LX3 exhibits 47% identity to the oligo-1,6-glucosidase (OGL) from Bacillus thermoglucosidasius (35) and the α-glucosidase from a Bacillus sp. (22) and 45% identity to the OGL from Bacillus cereus (36). Despite high sequence similarity (62 to 65%) between α-glucosidase and the two OGLs, the enzymes exhibit significant differences in substrate specificity (22). OGL and α-glucosidase are hydrolases with α-1,6 and α-1,4 bond specificities, respectively, whereas isomaltulose synthase catalyzes a two-step reaction with sucrose as the substrate, i.e., hydrolysis of the α-1,2 bond and then formation of an α-1,6 bond between the glucose and fructose moieties.

The Pall protein is a member of the α-amylase family. A domain search with the Pfam program revealed that Pall contains an α-amylase domain spanning residues 52 to 518, with an E value of 4.2e−125. Secondary-structure prediction indicated that Pall contains a (βα)8-barrel structure and a subdomain 8-barrel structure and a subdomain 8-barrel structure and a subdomain 8-barrel structure.

FIG. 2. Sequence comparison and assignment of secondary structure. The peptide sequence of Pall of Klebsiella sp. strain LX3 (K. si) is compared with the sequences of the thermolabile OGL of B. cereus ATCC 7064 (B. ce) (accession no. P21332) and the thermostable OGL of B. thermoglucosidasius KP1006 (B. th) (accession no. P29094). The secondary structural features, i.e., α (helix) and β (sheet), of B. cereus OGL, based on crystal structural analysis (11), are indicated above the sequences. The predicted secondary structural regions of Pall are underlined and labeled at the bottom (H, α-helix; E, β-sheet). The selection marker for the α-amylase family, QPDLN, is indicated by blue lines. The potential amino acid residues for probe substitution are shown in boldface, and the two mutated residues, R310 and E498, are indicated by . The crucial invariant residues (D241, E295, D369, H145, and H368 in Pall) conserved in α-amylase and glucosyltransferase are indicated by +. Boxes I to IV show the regions containing the five essential residues.

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main B located between Nβ3 and Na3 (Fig. 2). A motif search showed a short stretch (D241, E295, D369, H485, and H586) in PalI are shown in Fig. 2. Site-directed mutagenesis of these residues in PalI verified that they are essential for enzyme activity (D. Zhang, unpublished data).

Overexpression and purification of isomaltulose synthase. The pall without the coding sequence for the signal peptide was subcloned into expression vector PGEX-2T to generate the GST-PalI fusion gene under the control of an IPTG-inducible promoter, Ptac. To maximize the yield of the fusion gene, we tested different induction conditions and found that induction with 0.5 mM IPTG at 28°C for 15 h produced the maximum level of soluble active fusion protein (data not shown). A 500-ml culture of E. coli DH5α/pGEX produced a total of 840 U of PalI. Approximately 60% (540 U) of the total activity was found in the cell-free fraction. The GST-PalI fusion protein bound by glutathione-Sepharose affinity was digested by thrombin, and PalI was released from the affinity column. The product was found to be impure, containing one major band of ~67 kDa (PalI) and several minor bands. These minor bands were eliminated after further purification using semipreparative fast protein liquid chromatography. The homogeneity of the final product, purified up to 350-fold with a 29% final yield, was confirmed by SDS-PAGE (data not shown).

Analysis of PalI reaction product. The PalI reaction product was separated by TLC and detected by diphenylamine-aniline-phosphate reagent, which reacts with glucosaccharides of varied linkages to yield spots with different colors (27). Isomaltulose, which appeared as a green spot, was the major product of PalI. The second product showed the red color typical of trehalulose. Glucose and fructose were hardly detected using this method. HPLC analysis of the reaction solution showed four peaks (Fig. 3A). By comparison with the HPLC profiles of the standards and the typical color reaction of each peak on TLC plate, peak 1, with a retention time of 3.25 min was identified as isomaltulose, and peaks 2, 3, and 4 were identified as trehalulose, glucose, and fructose, respectively.

General properties of PalI. The optimal temperature and pH for PalI activity were examined. Maximal activity was observed at pH 6.0, and the enzyme was stable within a narrow pH range of 5.0 to 6.5. PalI rapidly lost activity at pH values higher than 6.5 (data not shown). PalI was most active at 35°C. At above 40°C PalI activity quickly dropped, indicating the poor thermostability of this enzyme under the assay conditions used. The values of K_m (3.18 mM) and k_cat/K_m (46.4 ± 1.7 mM and 0.27 ± 0.02 mM−1 min−1, respectively, and the specific activity was 328.0 ± 25 U/mg at pH 6.0 and 35°C.

Fe^{2+}, Hg^{2+}, and the detergent SDS completely abolished PalI activity, whereas Mg^{2+} and Mn^{2+} enhanced the activity of the enzyme (Table 2). Divalent cations such as Fe^{2+}, Li^{2+}, Ca^{2+}, and Cu^{2+} partially inhibited the enzyme activity, and Zn^{2+} had no effect. EDTA (a chelating reagent for divalent cations) also inhibited PalI, confirming that the enzyme requires certain cations for activity.

Effect of temperature and pH on product composition. Changing the temperature or pH affects the ratio of monosaccharides (glucose and fructose) and trehalulose in the PalI reaction products. Analysis of the products formed at different temperatures showed that high temperatures promote monosaccharide release but suppress trehalulose formation (Table 3). An increase of the temperature from 25 to 50°C caused the amount of monosaccharides in the reaction products to double, while trehalulose formation decreased approximately threefold. Neutral pH favors trehalulose formation, with its content increasing from 11.0 ± 2.3% at pH 5.0 to 13.0 ± 3.1% at pH 7.0. In contrast, slightly more monosaccharides were produced at an acidic pH than at neutral pH (6.9 ± 1.4% at pH 5.0 and 4.5 ± 0.9% at pH 7.0). Under these assay conditions, the proportion of isomaltulose in the PalI reaction products was not significantly influenced by either temperature (Table 3) or pH (data not shown).

Identification of potential sites for proline substitution. The proline rule, i.e., the fact that the presence of proline at the second position of a β-turn makes a protein more stable by decreasing the entropy of unfolding of the backbone (29, 30), has been widely applied to enhance thermostability in α-amylases (8, 21, 28, 37). Given that PalI belongs to the α-amylase family and its predicted secondary structure, i.e., (β/α)_n-barrel, is similar to that of B. cereus OGL (Fig. 2), we assumed that the proline rule could also be applied to PalI. We compared
TABLE 2. Influence of different reagents on PalI activity

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<th>Reagent</th>
<th>Concentration (mM)</th>
<th>Pall activity (%)</th>
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* Purified Pall protein was extensively dialyzed against 50 mM citrate-phosphate buffer (pH 6.0). The enzyme (2 μg in a 200-μl final volume) was preincubated with the indicated reagents and ions at room temperature for 10 min, and 20 μl of sample was taken for enzyme assay as described in Materials and Methods.

The primary sequence and the predicted secondary structure of the protein with those of thermostable B. cereus ATCC 7064 OGL and thermostable B. thermoglucosidasius KP1006 OGL, using the proline rule. The comparison identified five residues (L³⁰¹, R³¹⁰, S³³⁰, R⁴¹⁹, and E⁴⁹⁸) in PalI which are potential candidates for proline substitution (Fig. 2). Two residues, R³¹⁰ in the loop and E⁴⁹⁸ in the β-turn, were selected to investigate the effect of proline substitution on the thermostability and catalytic activity of PalI.

Proline substitution increases thermostability. Site-directed mutagenesis was used to replace E⁴⁹⁸ of PalI with proline to produce pGEK:E⁴⁹⁸P, which was further mutated to change R³¹⁰ to proline to form the double mutant pGEK:E⁴⁹⁸P/R³¹⁰P. The two mutant enzymes were overexpressed and purified by the same procedure as for native PalI. Mutation did not affect the expression levels of these mutant versions, as shown by SDS-PAGE (data not shown). Proline substitution changed the protein with those of thermolabile B. thermoglucosidasius OGL, producing pGEK:E⁴⁹⁸P, which was further mutated to change the optimum temperature of PalI (Fig. 4A). The optimum temperature of PalI was 35°C for PalI but increased to 40 and 45°C for PalI:E⁴⁹⁸P and PalI:E⁴⁹⁸P/R³¹⁰P, respectively. In comparison to PalI, the maximum specific activity increased by 7% for PalI:E⁴⁹⁸P and 16% for PalI:E⁴⁹⁸P/R³¹⁰P. The temperatures resulting in 50% loss of activity were estimated to be approximately 45.8°C for PalI, 48.3°C for PalI:E⁴⁹⁸P, and 51°C for PalI:E⁴⁹⁸P/R³¹⁰P (data not shown). The half-lives of PalI, PalI:E⁴⁹⁸P, and PalI:E⁴⁹⁸P/R³¹⁰P were 1.81, 9.45, and 13.61 min at 50°C (Fig. 4B), respectively. The half-life of PalI:E⁴⁹⁸P/R³¹⁰P at 50°C is therefore about 11-fold longer than that of PalI.

To determine the difference in catalytic activity, we measured Kₘ (millimolar) and kₗₚₜ (minutes⁻¹) at pH 6.0 and 35°C. The kₗₚₜ/Kₘ values for PalI, PalI:E⁴⁹⁸P, and PalI:E⁴⁹⁸P/R³¹⁰P were 0.27 ± 0.02, 0.29 ± 0.04, and 0.31 ± 0.03 min⁻¹ mM⁻¹, respectively. This result indicates that replacement of E⁴⁹⁸ and/or R³¹⁰ by proline slightly increases the apparent catalytic efficiency (kₗₚₜ/Kₘ) of PalI on sucrose.

Proline substitution facilitates isomaltulose formation. The reaction products of PalI and its mutant versions were analyzed to test whether proline replacement affects the percent contents of PalI products. The percent content of monosaccharides decreased from 5.9 ± 1.0% for PalI to 3.4 ± 0.8% for PalI:E⁴⁹⁸P and 3.3 ± 0.8% for PalI:E⁴⁹⁸P/R³¹⁰P, whereas the percentage of isomaltulose increased from 81.9 ± 1.4% for PalI to 84.8 ± 1.0% for PalI:E⁴⁹⁸P and 84.1 ± 2.0% for PalI:E⁴⁹⁸P/R³¹⁰P. The percent content of trehalulose in the reaction products was not significantly influenced by proline substitution. It is likely that proline substitution enhances the stability of the glucosyl-enzyme complex and consequently diminishes the release of monosaccharides.

DISCUSSION

The gene (palI) encoding the isomaltulose synthase (PalI) has been cloned and characterized. palI encodes a peptide of 598 amino acids with a molecular mass of 69.94 kDa. Domain structure analysis indicates that PalI contains an α-amylase domain. The main homologs of PalI, besides several uncharacterized isomaltulose synthases from an Enterobacter sp., Protaminobacter rubrum (15), and E. raphontici (2), are OGLs from B. thermoglucosidasius and B. cereus (35, 36) and the α-glucosidase from a Bacillus sp. (21). The α-amylase domain of PalI shows about 47.7, 47.5, and 48.4% identity to the same domain of α-glucosidase from the Bacillus sp., OGL from B. thermoglucosidasius, and OGL from B. cereus, respectively. The predicted (β/α)₉-barrel secondary structure of PalI is similar to that of B. cereus OGL. These characteristics, together

![Table 3: Influence of temperature on product formation](http://aem.asm.org/)

<table>
<thead>
<tr>
<th>Sugar</th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
<th>35</th>
<th>45</th>
<th>50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose + fructose</td>
<td>ND</td>
<td>ND</td>
<td>5.5 ± 0.7</td>
<td>5.9 ± 0.8</td>
<td>5.9 ± 1.0</td>
<td>9.1 ± 0.4</td>
<td>12.7 ± 2.3</td>
</tr>
<tr>
<td>Trehalulose</td>
<td>20.9 ± 3.2</td>
<td>21.5 ± 2.7</td>
<td>17.0 ± 2.3</td>
<td>14.4 ± 0.8</td>
<td>12.2 ± 0.6</td>
<td>8.3 ± 0.6</td>
<td>5.2 ± 0.8</td>
</tr>
<tr>
<td>Isomaltulose</td>
<td>79.1 ± 2.8</td>
<td>78.9 ± 4.3</td>
<td>77.4 ± 2.7</td>
<td>78.7 ± 3.1</td>
<td>81.9 ± 1.4</td>
<td>82.6 ± 1.7</td>
<td>82.1 ± 3.3</td>
</tr>
</tbody>
</table>

* Samples (2 ml) of sucrose (4%) in citrate-phosphate buffer (pH 6.0) containing 10 μg of PalI were incubated for 4 h at the indicated temperatures. The reaction mixture was boiled for 5 min and analyzed by HPLC on a Symmetry C₁₈ column (4.6 by 250 mm).

* ND, not determined.
with the presence of the marker sequence 209QPDLN, indicate that PalI is a member of the α-amylase family.

Sequence comparison revealed that PalI contains a potential catalytic triad (Asp241, Glu295, and Asp369) and two histidine residues (His145 and His368) that are highly conserved in α-amylases and glucosyltransferases. The importance of the catalytic triad and the two invariant residues has been verified for a range of α-amylases. Watanabe et al. (38) reported that mutation of the catalytic triad (Asp199, Glu235, and Asp269) in the OGL of B. cereus to asparagine or glutamine resulted in a significant loss of enzyme activity. Mutation of the equivalent residues, Asp294 and Asp401, in Neisseria polysaccharea amylo-sucrase inactivated the enzyme activity (26). A crystallographic study of cyclomaltodextran glucanotransferase from Bacillus circulans revealed that D256 functions as a nucleophilic site, E284 functions as a proton donor, and D354 functions as a substrate binding site (12, 32). The two conserved histidine residues may participate in stabilization of the substrate-binding transition state at subsite −1, as in the case of the Taka-amylase of Aspergillus oryzae (10) and cyclomaltodextran glucanotransferase of alkalophilic Bacillus sp. strain 1011 (20, 32). We found that the potential catalytic triad and the two histidine residues are essential for PalI activity (D. Zhang, unpublished data). The similar protein domain structure and function of the conserved catalytic triad suggests that the catalytic mechanism of PalI may resemble that of α-amylases and glucosyltransferases, especially for the first part of the reaction, i.e., cleavage of the glycosidic linkage and formation of a glucosyl-enzyme covalent intermediate.

PalI efficiently converts sucrose to isomaltulose and trehalulose with trace amounts of glucose and fructose. Formation of multiple products seems to be a characteristic common to isomaltulose synthases from various sources (7, 14, 15, 31, 33). The sucrose isomerase of S. plymuthica converts sucrose into isomaltulose, trehalulose, glucose, fructose, isomaltose, and isomelezitose (33). We found no evidence of isomaltose or isomelezitose in the reaction products of PalI by either TLC or HPLC analysis. The optimum pH (6.0) of PalI is similar to that of sucrose isomerase from S. plymuthica (14, 33), a Klebsiella sp. (24), and E. rhapontici (2). The optimum temperature (35°C) for PalI is identical to that for the sucrose isomerase from the Klebsiella sp. (24) and higher than that (30°C) for the sucrose isomerases from S. plymuthica (14, 33) and E. rhapontici (2).

Our data show that formation of trehalulose and monosaccharides, but not isomaltulose, is significantly influenced by temperature. High temperature tends to stimulate release of monosaccharides but suppress formation of trehalulose. It is likely that low temperature favors tautomerization of fructofuranose into fructopyranose (13), a critical step for trehalulose formation. Our results are consistent with the finding that isomaltulose synthase from Pseudomonas mesoacidophila produces 90% trehalulose at 20°C but only 50% trehalulose at 50°C (18).

The thermal stability of a protein is determined by many factors, for example, packing efficiency, hydrophobic interaction, loop stabilization, reduction of entropy of unfolding, and electrostatic interaction (34). Proline replacement in the OGL of B. cereus ATCC 7064 demonstrated that proline residues, especially at the second site of a β-turn, at the first turn of α-helices, or in coils within the loop binding to adjacent secondary structure, contributed to protein thermostability (30, 35, 37). Structural similarities between PalI and OGLs from B. cereus ATCC 7064 and B. thermoglucosidasius KP1006 and differences in thermostability facilitate the identification of potential residues for proline substitution to engineer protein thermostability properties. Our results showed that the proline substitution of two identified residues, Glu408 and Arg310, significantly enhanced both the optimum temperature for enzyme activity and the thermostability. In particular, the half-life of the double mutant PalI:E408P/R310P is 11-fold higher at 50°C than the half-life of PalI. The similarity of the apparent catalytic efficiencies (kcat/Km) of PalI and its mutant versions implies that no extensive conformational change, especially in the active-site cleft, is induced by proline substitution. Our results indicate that comparison of primary and secondary structures of enzymes with similar domain structures is an effective approach for molecular design in protein engineering.
REFERENCES


