Identification and Characterization of a Novel Trehalose Synthase Gene Derived from Saline-Alkali Soil Metagenomes

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Abstract

A novel trehalose synthase (TreS) gene was identified from a metagenomic library of saline-alkali soil by a simple activity-based screening system. Sequence analysis revealed that TreS encodes a protein of 552 amino acids, with a predicted molecular weight of 63.3 kDa. After being overexpressed in *Escherichia coli* and purified, the enzymatic properties of TreS were investigated. The recombinant TreS displayed optimal activity at pH 9.0 and 45 °C, and the addition of most common metal ions (1 or 30 mM) had no inhibition effect on the enzymatic activity evidently, except for the divalent metal ions Zn²⁺ and Hg²⁺. Kinetic analysis showed that the recombinant TreS had a 4.1-fold higher catalytic efficiency (Kcat/Km) for maltose than for trehalose. The maximum conversion rate of maltose into trehalose by the TreS was reached more than 78% at a relatively high maltose concentration (30%), making it a good candidate in the large-scale production of trehalose after further study. In addition, five amino acid residues, His172, Asp201, Glu251, His318 and Asp319, were shown to be conserved in the TreS, which were also important for glycosyl hydrolase family 13 enzyme catalysis.

Introduction

Trehalose is a naturally occurring non-reducing disaccharide in which the two glucose units are linked via an α,α-(1,1)-glycosidic bond. Although there are three different anomers of trehalose (i.e. α,α-1,1-,α,β-1,1- and β,β-1,1-), the only known biologically active form is α,α-1,1-glucosyl-glucose [1]. This disaccharide has been isolated from a large number of prokaryotic and eukaryotic cells including mycobacteria, streptomycetes, enteric bacteria, archaea, yeast, fungi, algae, low orders of the animal kingdom and higher orders of the plant kingdom, especially those living in harsh environments [2,3]. Initially, trehalose was thought to act solely as a reserve energy and carbon source in a manner similar to that of starch and glycogen, but a growing number of studies indicate that this sugar instead has important biological function of playing a major role in cell protection against various physical and chemical stresses, such as heat, cold, dehydration, desiccation, oxygen radicals, and acidic/alkali environmental conditions [4-6]. Moreover, the ability of the microorganisms to survive in these environments correlates with their trehalose content [7,8]. In yeast, the most ancient actor of biotechnological transformation, trehalose was found to accumulate, in certain physiological conditions, up to 10%-15% of cell dry weight [9]. Investigation on the cell membranes of anhydrobiotic yeast has unraveled that intracellular trehalose can stabilize proteins in their native state and to reduce their heat-induced denaturation and aggregation [10,11]. As a matter of fact, the build-up of trehalose upon heat shock has been shown to be more important than the induction of heat-shock proteins as a thermotolerance response element [12]. Also, trehalose was shown to decrease oxidative damage to...
cell proteins by oxygen radicals and thus to increase the tolerance of organisms to reactive oxygen species [13]. In particular, trehalose also has the same protection effect in vitro, which opens a new field for its application in food, cosmetic, and pharmaceutical industries, ranging from serving as a sweetener to a biomaterial stabilizer [14]. Besides acting as a protectant, trehalose is also an important component of the cell walls of many mycobacteria and corynebacteria in the form of glycolipids. A well-known example is trehalose dimycolate (or cord factor), which was composed by a trehalose core with mycolic acid esterified at the 6-OH and 6'-OH positions [15]. The cord factor is the most toxic lipid produced by many mycobacteria and corynebacteria, and dramatically increases the impermeability of the cell wall to various antibiotics and thus was identified as a virulence factor [16].

Hayashibara Co. Ltd. has isolated a two-step enzymatic route for the production of trehalose, which has led to a major decrease in the commercial price of trehalose to 5-6 US$·kg$^{-1}$, and for the first time, successfully exploited in industrial production of trehalose. However, further decrease in the production cost of trehalose could be achieved no other than the application of the brand-new enzymatic route [19].

Trehalose synthase (TreS, EC 5.4.99.16) was first demonstrated in Pimelobacter sp. R48 through an extensive screening of 2,500 strains of soil bacteria [20]. It can catalyze the intramolecular rearrangement of maltose into trehalose in a single step, which represented a simple, fast, and low-cost method for the future industrial production of trehalose [1]. So far, a number of TreSs from several bacterial strains have been identified and characterized (Table 1). However, these TreSs were still not satisfying in a practical application with regard to either their activities or conversion efficiency. Furthermore, all these TreSs were from cultured microorganisms, and little attention had been paid to those from uncultivable microorganisms, which may account for more than 99% of microorganisms in the environment [21]. It is imaginable that there is a large number of industry-potential TreSs for the production of trehalose in the uncultivable microorganisms of environment.

To expand the range of TreSs discovery, culture-based methods have been complemented or replaced by culture-independent metagenomic approaches, which theoretically provide access to the collective nucleic acids from the uncultivable organisms of various environmental samples [22]. Functional metagenomics based on the direct isolation of DNA from soil environmental samples, generation of metagenomic libraries from the isolated DNA, and function-driven screening of the constructed libraries has been successfully employed in the identification and characterization of enzymes with special biocatalytic activities [23]. In the present study, a metagenomic library from saline-alkali soil sample of Lop Nur in Xinjiang Uigur Autonomous Region of north-west China was constructed for the screening new TreSs. Finally, one novel TreS with high activity and conversion efficiency was identified and subsequently expressed in Escherichia coli (E. coli). The specific enzymatic properties of the recombinant enzyme were also characterized after purification. Furthermore, the functional

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### Table 1. Summary of several identified and characterized TreSs from bacterial strains.

| Strain name                  | Gene size (bp) | Molecular size (kDa) | Optimum temperature (°C) | Optimum pH | Optimum substrate | Conversion rate | Citation                  |
|------------------------------|----------------|----------------------|--------------------------|------------|-------------------|-----------------|--------------------------|
| Mycobacterium ameghinae ATCC 14468 | 1781          | 71                   | 37                       | 7.2        | 0.5M              | 45%             | Pan et al., 2004          |
| Thermobifida fusca DSM 43792  | 1830          | 66                   | 25                       | 6.5        | 800mM             | 60%             | Wei et al., 2004          |
| Pseudomonas stutzeri Cj 38    | 2070          | 76                   | 35                       | 8.5        | 600mM             | 72%             | Lee et al., 2005          |
| Picrophilus torridus DSM 9790 | 1677          | 65                   | 45                       | 6          | 150mM             | 71%             | Chen et al., 2006         |
| Thermus thermophilus ATCC 33923 | 2898         | 106                  | 65                       | 6.5        | 3mM               | 62%             | Wang et al., 2007; Anna et al., 2005 |
| Deinococcus radiodurans ATCC 13939 | 1659       | 61                   | 15                       | 6.5        | 800mM             | 65%             | Wang et al., 2007; Wei et al., 2004 |
| Arthrobacter aurescens CGMCC1.1892 | 1797       | 68                   | 35                       | 6.5        | 90mM              | 60%             | Wu et al., 2009           |
| Enterobacter hormaechei        | 1626          | 65                   | 37                       | 6          | 100mM             | 48%             | Yue et al., 2009          |
| Methanothermus ruber CBS-01    | 2889          | 106                  | 50                       | 6.5        | 60mM              | 65%             | Zhu et al., 2010          |
| Corynebacterium glutamicum ATCC 13032 | 1812      | 70                   | 35                       | 7          | 3mM               | 69%             | Kim et al., 2010          |
| Deinococcus geothermalis DSMZ 11300 | 1692       | 65                   | 40                       | 7.6        | 300mM             | 60%             | Pawel et al., 2012        |
| Thermomonospora curvata DSM 431383 | 1806      | 60                   | 35                       | 6.5        | 3mM               | 70%             | Liang et al., 2013        |
| Rhodococcus opacusATCC 41021   | 1857          | 79                   | 25                       | 7          | 90mM              | 67%             | Yan et al., 2013          |
| Deinococcus sp.                | 1656          | 63.3                 | 45                       | 9          | 800mM             | 78%             | This study                |

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amino acid residues have been predicted by the site directed mutation based on homology modeling and structure analysis.

**Materials and Methods**

**Bacterial strains and cultivation**

*E. coli* DH5α was used for construction of recombinant plasmids and *E. coli* BL21(DE3) was used as expression host. *Thermus thermophilus* ATCC 33923 was purchased from the China General Microbiological Culture Collection Center (CGMCC). The pUC118 (TaKaRa, Dalian, China) and pET-22b(+) were used to construct metagenomic libraries and express the target protein, respectively. *E. coli* transformants were grown at 37 °C in Luria-Bertani medium containing 50 μg/mL ampicillin [24].

**Isolation of DNA from environmental sample**

For the construction of a metagenomic library, an environmental sample was obtained from the soil in Lop Nur. The total DNA was extracted based on a method described previously [25]. Soil sample (4 g of wet weight) was mixed with 13.5 mL of DNA extraction buffer, which composed of 100 mM Tris-HCl (pH 8.0), 100 mM sodium EDTA (pH 8.0), 100 mM sodium phosphate (pH 8.0), 1.5 M NaCl, 1% cetyltrimethylammonium bromide (CTAB), and 1.5 mL of 20% sodium dodecyl sulfate (SDS). The mixture was incubated in a 65 °C water bath for 2 h with gentle inversion every 15 to 20 min. The supernatants were collected after centrifugation (16,000 × g, 20 min) at 4 °C, washed twice with cold 70% ethanol and suspended in an appropriate volume of sterile deionized water.

**Construction of a metagenomic library**

To construct the metagenomic library, the purified DNA was partially digested with BamHI. DNA fragments of 2.5-10 kb were ligated into BamHI-digested pUC118, and the ligated products were transformed into *E. coli* DH5α. The transformed cells were plated onto LB agar plates containing 50 μg/mL ampicillin (Amp), 0.5 mM isopropyl-β-D-thiogalactopyranoside (IPTG) and 100 μM 5-Bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-gal). After incubation at 37 °C for 24 h, clones with white color were selected and further tested by colony polymerase chain reaction (PCR).

**Subcloning and gene sequence analysis**

Several bacterial trehalose synthase sequences published in the NCBI database were collected and analyzed by the online multiple sequence alignment program CLUSTAL W2 (http://www.ebi.ac.uk/Tools/clustalw2). As shown in Figure 1, two degenerate primers TF1 (5'-AGYCCNCTNCNNGAYGRNGGNT-3') and TF2 (5'-AGNGTNAYTCRTCTGRT-3') were synthesized based on the conserved domains. Then colony PCR was carried with primer TF1 and TF2, clones in white color were used as templates, the genome of *Thermus thermophilus* ATCC 33923 was acted as a positive control. PCR products were detected on a 2% agarose gel. Only the clones containing recombinant plasmids (pUC118-treS) as well as positive control could produce detectable band. Then the recombinant plasmids were extracted and sequenced. The deduced amino acid sequence analysis and open reading frame search were performed with BLAST program provided by NCBI (http://www.ncbi.nlm.nih.gov/). The phylogenetic tree was constructed by the neighbor-joining method using Molecular Evolutionary Genetics Analysis software (MEGA, version3.1).

**Cloning of the treS gene and construction of the expression vector**

Primers TR1 (5'-CCCATATGATGATCCAGACCACCACCACTACAG-3') and TR2 (5'-CCAAGGCTTGTTCAGGCGCCGCAAGTATAGT-3') were synthesized according to the open reading frame (ORF) sequence to introduce Ndel and HindIII sites into the 3' and 5' ends of treS ORF, respectively. The stop codon of treS was eliminated to in-frame read a His(6)-tag on the C-terminus for one step purification. The treS gene was amplified by PCR with the pUC118-treS as template. The PCR product was then digested with restriction enzymes Ndel and HindIII, and inserted into pET22b(+) vector to generate pET22b(+)-treS. After confirmed by DNA sequencing, the recombinant plasmid was transformed to the *E. coli* BL21(DE3).

**Protein expression and purification**

The *E. coli* BL21(DE3) harboring pET22b(+)-treS was inoculated into LB medium supplemented with 50 μg/mL Amp and then grown at 37 °C in a shaker at 200 rpm. When the OD₆₀₀ of the culture reached 0.8, IPTG was added to a final concentration of 1.0 mM, and then the incubation was continued for another 8 h at 30 °C. The cells were harvested by centrifugation and resuspended in lysis buffer (50 mM KH₂PO₄, 500 mM NaCl, pH 6.0) followed by sonification and centrifugation at 12,000 × g for 20 min at 4 °C to remove insoluble cell debris. The supernatant was filtered through a 0.45-μm filter and loaded onto a Ni-NTA affinity chromatography column according to the manufacturer’s purification protocol manual (Novagen, Ni-NTA His Bind Resins). The cell extracts and purified enzyme were analyzed by 12.5% (w/w) SDS-polyacrylamide gel electrophoresis (SDS-PAGE). Protein concentrations were determined by the method of Bradford using bovine serum albumin as a standard [26].

**Activity assay of recombinant TreS**

The activity of TreS was assayed by measuring the amount of trehalose produced from maltose. The reaction was performed in a mixture containing the TreS solution and 100 mM maltose in 50 mM phosphate buffer (pH 9.0) at 45 °C for 30 min, then boiled for 10 min to terminate the reaction. The amount of trehalose, glucose, and maltose of each reaction mixture was measured using a high-performance liquid chromatography (HPLC) system equipped with an RID...
Properties of recombinant TreS

The optimum pH of TreS was assayed by incubating the purified enzyme with 200 mM maltose substrate in 50 mM potassium phosphate buffer at pH 3.0 to 11.0 and 45 °C for 30 min, respectively. The optimum temperature for TreS activity was determined at 10 to 80 °C using the same buffer at pH 9.0 for 30 min, respectively. To determine the effect of metal ions and different chemical reagents on TreS, its activity was also assayed in the presence of these ions or compounds at 1 mM, respectively. To determine the conversion efficiency of TreS, the 5-L reaction system consisted of a 5-L stirred-tank fermentor (B. Braun, B. Braun Biotech International, Melsungen, Germany) containing 2-L of a medium was employed [27].

Determination of kinetic parameters

The Michaelis-Menten ($K_m$) and maximum activity ($V_{max}$) constant for recombinant TreS were determined under conditions of pH 9.0 and 45 °C for 30 min in 200 mM sodium phosphate buffer containing substrate (maltose and trehalose) at various concentrations. The resulting data were plotted with Origin 6.0 software (Microcal, Northampton, MA). All experiments were carried out in triplicate.

Construction and analysis of protein models for TreS

Models were built through an online Automatic Modeling Mode server at http://swissmodel.expasy.org. Obtained models were analyzed through Swiss-Pdb Viewer [28] and PyMOL (http://www.pymol.org). Structure predictions for TreS were made by the development of the homology model using the resolved X-ray structure of α-amylase with Protein database entry code 1SMA as template.

Site-directed mutagenesis of TreS

Mutants were obtained through a cloning method with two complementary primers containing mutation bases. The pET22b(+)–treS plasmids were amplified by PCR with PrimeSTAR HS DNA polymerase (Takara, Dalian, China). To remove the templates, Dpn I was added to PCR reactions for 1 h at 37 °C. The digested products were then directly transformed into competent E. coli DH5α to obtain mutation recombinant plasmids. After identified by sequencing, the recombinant plasmid containing mutation site was transformed to the E. coli BL21(DE3).
Results

Construction and screening of the metagenomic library

A metagenomic library of ca. 85,000 clones was constructed for obtaining trehalose synthase genes. Restriction analysis of 20 randomly selected clones indicated that all the clones harbored insertion DNAs ranging from 2.5 to 5 kb in size, with an average of approximating 3.5 kb. The metagenomic library processed a capacity ~300 Mb of soil microbial genomic DNA. One clone expressing trehalose synthase activity was isolated after all the clones in the library had been screened.

Sequence analysis of treS gene

The plasmid of pUC118-treS was extracted and the complete insert DNA was sequenced. The length of the insert DNA was 5,356 bp. An ORF-finder and BLAST analysis revealed the presence of an ORF consisting of 1,656 bp, encoding a full-length treS gene, which further encoded a protein of 552 amino acids with a predicted molecular mass of 63.3 kDa. The deduced amino acid sequence of treS was used to perform a BLAST research of the NCBI and SwissProt databases. This search revealed that the protein has the highest similarity (84%) with trehalose synthase from Deinococcus radiodurans R1 [29]. As can be seen in Figure 1, multiple sequence alignments of this treS gene with 10 reported trehalose synthase revealed that they share several highly conserved amino acid motifs, such as SPLRDG/DGYDV/I, FPL/VMPRI/LF/Y, NHDELTLE, GIRRRLA/MPL, and so on. However, the region SRVPAPNTVF in this treS gene of the present study was significantly different from the conserved regions YYGDEIGMGD in other listed treS genes (Figure 1). Such a result suggests that the cloned fragment may originate from an uncultured organism, and the identified gene-encoding products possibly had a unique function. The phylogenetic tree based on amino acid sequence was further constructed to verify the evolutionary relationship of this treS gene to other known trehalose synthases, and 24 trehalose synthase proteins were selected for the phylogenetic tree analysis. As shown in Figure 2, this recombinant TreS has a close relationship to Deinococcus genus regarding sequence homology.

Expression and purification of the recombinant TreS

To characterize the biochemical properties of the recombinant TreS, the treS gene was expressed as an N-terminal His-tag fusion protein using pET22b(+) expression system in E. coli BL21(DE3). No inclusion bodies were found in
cell lysates, which suggested that this recombinant TreS was expressed in a soluble form. The cells were harvested and disrupted by sonication on ice. When compared to the sample without induction (Figure 3, lanes 1), only the induced cells containing the recombinant vector expressed an extra protein band migrating at about 65 kDa upon induction (Figure 3, lanes 2). The recombinant protein was about 1.5 kDa heavier than the predicted molecular mass of 63.3 kDa, which was due to the additional 13 amino acids including the His(6)-tag at the N-terminus. After purification with the Ni-NTA column, a single band was shown on the SDS-PAGE gel correlating with the size of enzyme, indicating that the recombinant enzyme was purified to homogeneity (Figure 3, lane 3).

**Effects of pH and temperature on activity of TreS**

The optimum pH of the recombinant TreS was found to be 9.0. The enzyme maintained high activity at a broad pH range of 5.0-10.0 (Figure 4A). The optimum temperature was 45 °C. Meanwhile, the enzyme maintained high activity when reaction temperature ranged from 15 °C to 55 °C. However, relative activity quickly decreased when temperature was above 55 °C or below 15 °C (Figure 4B). Thus, it is probably more efficient to carry out conversions at moderate temperatures. In this study, the recombinant TreS exhibited a stable performance under the wide working conditions (pH 5.0-10.0 and 15-55 °C, Figure 4). Further studies of the effects of temperature, reaction mixtures containing 200 mM maltose were incubated under pH 9.0 at 55 °C for 1 h, the remaining activity of the recombinant TreS was as high as 80% of its initial activity.

**Activity assay of the recombinant TreS**

Enzymatic activity was detected with purified TreS in reactions of the conversion between maltose and trehalose. It was confirmed that the DNA fragment was the intrinsically coding sequence of active TreS. The highest enzyme activity was calculated to be 133.5 ± 4.8 U·mg⁻¹ protein, and the conversion efficiency was test in a 5-L reaction system. After 18 h of TreS-catalyzed reaction, the final yield of trehalose was...
constantly above 75%, with the maximum value of 78% under the optimum condition at a relatively high maltose concentration (30%).

Effects of metal ions and reagents on activity of TreS

The effects of metal ions and reagents were further determined by examining recombinant TreS activity in the presence of 1 and 30 mM of these substances under standard assay conditions (Table 2). As shown in Table 2, the TreS activity was inhibited strongly by Hg²⁺, Zn²⁺, and SDS and
moderately by Cu$^{2+}$, Cd$^{2+}$, Pb$^{2+}$, and Tris at the concentration of 1 mM. More uplifting was the result that, at the concentration of 30 mM, almost all metal ions and reagents, except for Hg$^{2+}$ and Zn$^{2+}$, had no more inhibition effect on the enzyme activity than the concentration of 1 mM.

Kinetic analysis of TreS

Kinetic parameters of the recombinant TreS were investigated at pH 9.0 and 45 °C for 30 min with maltose or trehalose as substrate. When this data was plotted by the method of Lineweaver and Burk, the $K_m$ values for the recombinant TreS were found to be 9 ± 1.2 mM for maltose and 64 ± 3.5 mM for trehalose, the $V_{max}$ values of 1.5 ± 0.2 mM min$^{-1}$ mg$^{-1}$ protein for maltose and 3.1 ± 0.6 mM min$^{-1}$ mg$^{-1}$ protein for trehalose were calculated, respectively. Although a higher $K_m$ for trehalose than maltose was seen (78 ± 6.5 s$^{-1}$ vs. 44 ± 3.8 s$^{-1}$), TreS had a 4.1-fold higher catalytic efficiency ($K_{cat}/K_m$) toward maltose than trehalose (4.9 ± 1.2 M$^{-1}$ s$^{-1}$ vs. 1.2 ± 0.18 M$^{-1}$ s$^{-1}$). With regard to these results, the recombinant TreS had a higher affinity to maltose and a favorite reaction direction toward the synthesis of trehalose. Interestingly, all reported TreS enzymes share the feature of a reversible conversion at different degrees [26,29].

Structure analysis and site-directed mutagenesis

Using the structure-resolved *Pseudomonas* Mesoacidiphila trehalulose synthase as template (PDB ID: 2PWG), a TreS model was built through SWISS-MODEL. The sequence identity of TreS and trehalulose synthase was 30%, but both belonged to the glycosyl hydrolase family 13 (GH13 family) and had a typical ($\alpha/\beta)_8$ barrel catalytic domain (Figure 5A). In alignment with the amino acid sequences of trehalulose synthase, five conserved key amino acids constituting a catalytic pocket (His172, Asp201, Glu251, His318 and Asp319) of TreS were deduced. Three dimensional structures showed that all five putative active sites were in the center of the barrel catalytic domain (Figure 5B). To verify the importance of these residues, site-directed mutagenesis was used to replace the five residues individually with Ala, and each mutant recombinant protein was purified by Ni column. The drastic reduction in enzyme activity of all mutants suggested that these five residues might play essential roles in TreS catalysis (Table 3). Moreover, similar conservations of active sites were observed in other TreSs (Figure 1), further supporting the catalytic importance of these residues in TreS activity.

Discussion

Up to now, five different enzymatic routes involved in the biosynthesis of trehalose have been discovered and indentified from all kinds of environmental microorganisms. The first pathway utilizes trehalose-phosphate synthase (TPS, EC 2.4.99.15) (OtsA in *E. coli*) that catalyzes the transfer of the glucosyl moiety from UDP-glucose to glucose-1-phosphate, forming the intermediate trehalose-6-phosphate and UDP. The phosphate is then removed by trehalose-phosphate phosphatase (TPP, EC 3.1.3.12) (OtsB in *E. coli*) to give free trehalose [30]. The second pathway involves a trehalose glycosyltransferring synthase (TreT, EC 2.4.1.245) that catalyzes the synthesis of trehalose using nucleoside diphosphate glucose (NDPG), such as UDPG, as a donor and glucose as an acceptor [31]. The third pathway utilizes trehalose phosphorylase (TreP, EC 2.4.1.64) to catalyze a reversible reaction in which it hydrolyzes trehalose in the presence of inorganic phosphate to form glucose-1-phosphate and glucose and, inversely, gives rise to trehalose from both products in vitro [32]. However, all these three pathways are not suitable in trehalose industrial production due to their expensive substrates. The fourth pathway also involves a two-step enzymatic process, which utilizes glycogen or maltooligosaccharides as the starting material. The maltosyl moiety at the reducing end is first isomerized into a trehalosyl moiety by maltooligosyl trehalose synthase (TreY, EC 5.4.99.15). Hydrolytic release of the trehalose is then catalyzed by maltooligosyltrehalase trehalohydrolase (TreZ, EC 3.2.1.141) [33]. Trehalose has mainly been manufactured through this pathway since it was discovered in 1995. The last pathway utilizes only one enzyme, trehalose synthase (TreS, EC 5.4.99.16), to catalyze the intramolecular rearrangement of the α-1,4-linkage of maltose to the α-1,1-linkage of trehalose [34]. Maltose is relatively cheap, and this single step process saves both time and costs in scale-up mode, which could be an alternative method for industrial trehalose production.

In the present study, we constructed a plasmid metagenomic library from uncultivated microorganisms within saline-alkali soil samples and isolated a novel TreS gene by a sequence-based screening strategy. This recombinant TreS showed an optimal pH of 9.0 and an optimal temperature of 45 °C and retained 80% of its initial activity after heat-treatment at 55 °C for 1 h, which was consistent with the extreme environment such as high temperature, high salinity conditions in Lop Nur region. In comparison with other previously reported trehalose synthases,
this recombinant TreS has the most alkali optimum pH. Moreover, the three well-characterized TreSs from *Thermobifida fusca*, *Enterobacter hormachei*, *Arthrobacter aurescens* showed dramatic decreases in enzyme activity at pH above 9.0 and retained <40% of activity [35-37]. Therefore, it is more alkali-tolerant than other TreSs reported so far. Using an enzyme that is stable at high pH value as well as high reaction temperature can reduce the possibility of microbial contamination, lower viscosity, and improve the solubility of substrates for better access to enzymatic attack.

It is well known that a glycosidase activity is frequently affected by the presence of metal ions [38]. Various divalent
metal ions were therefore examined and consequently found to have inhibition effect on the enzyme activity in different degree. The results showed that the activity of the recombinant TreS was decreased by Zn$^{2+}$ and Hg$^{2+}$, but not as significantly as other known TreSs, especially at a high concentration of the metal ions (up to 30 mM) [39,40]. However, the activity of this recombinant TreS was not affected by EDTA, suggesting that this enzyme did not require metal ions for activity and stability.

To the target substrate maltose, the specific activity of this recombinant TreS was estimated to be about 133 units/mg of protein, which was 1.67-fold higher than that of *Pseudomonas stutzeri* and *Pichobius torridus* trehalose synthases, and roughly equal to that of *Thermus aquaticus* trehalose synthase [20,39,41]. Because TreS catalyzes the interconversion of maltose and trehalose, but converts maltose to trehalose more rapidly than trehalose to maltose, it was therefore of interest to determine the affinity ($K_a$) of this recombinant TreS for these two substrates. The results showed that the TreS had much greater affinity (7.1-fold) for maltose than trehalose. Moreover, the catalytic efficiency ($K_{cat}/K_a$) for maltose was 4.1-fold higher than trehalose, due to the reduced $K_a$ value as well as the increased $K_{cat}$ value, which resulted in the conversion of trehalose. In this study, the maximum trehalose yield from maltose by this recombinant TreS was exceeded 78% under the optimum condition, which was close to the theoretical equilibrium constant for overall reaction of 82% in favor of trehalose using thermodynamic parameters [42]. Although no three-dimensional structures have been obtained to date on trehalose synthase, previous reports on

**Table 3.** Relative specific activity of wild-type TreS and mutant enzymes.

| mutant   | relative specific activity (%) | mutant (%) |
|-----------|-------------------------------|------------|
| wide type | 100.00                        | E251A 1.71 ± 0.12 |
| H172A     | 7.21 ± 0.51                   | H318A 0.00 |
| D201A     | 4.76 ± 0.24                   | D319A 0.00 |

a. Relative activities are represented as the ratio of mutants to wild type. The specific activity of wild-type TreS was 133 units/mg protein.

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Building a Recombinant Trehalose Synthase

TrES have indicated that this enzyme was belonged to the GH13 family containing a common structural feature and conserved residues for catalysis and substrate binding [36,39,43]. From the 3D model of TreS, we speculated that the deep groove in the catalytic pocket served as an entry point of substrate into the catalytic center and as an export site of products (Figure 5B). Our study showed that residues of catalytic importance in GH13 family enzymes were also conserved in TreS (Figure 1), and the importance of the identified five residues in the central catalytic area (His172, Asp201, Glu251, His318 and Asp319) was revealed in a mutagenesis study (Table 3). The same conserved amino acids were found in other similar class of GH13 family enzymes, such as *Bacillus cereus* oligo-1,6-glucosidase (His103, Asp199, Glu255, His328, and Asp329) [44], *Neisseria polysacchareaialis* amylosucrase (His187, Asp286, Glu328, His392, and Asp393) [45], and *Pseudomonas mesoacidophilia* MX-45 trehalose synthase (His104, Asp200, Glu254, His326, and Asp327) [46]. This finding is consistent with previous suggestions and further supported the assumption that TreS employs a similar hydrolysis mechanism as other GH13 family enzymes.

In conclusion, construction and screening of a large-insert soil-derived metagenomic library has led to the discovery and characterization of one novel trehalose synthase. The novelty of this enzyme arises from the enormous genetic diversity of uncultured saline-alkali soil microorganisms. The recovered enzyme displayed several excellent enzymatic properties, such as resistance to most metal ions, high activity over a wide range of temperatures and pH values, with the maximum trehalose yield to be above 78%, which was considered as a good candidate for the large-scale production of trehalose in the near future.

**Author Contributions**

Conceived and designed the experiments: HH. Performed the experiments: LJ ML. Analyzed the data: YZ YL. Contributed reagents/materials/analysis tools: XX SL. Wrote the manuscript: LJ.

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