Improved Expression and Optimization of Trehalose Synthase by Regulation of P_{glv} in *Bacillus subtilis*

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Trehalose synthase (TreS) converts maltose to trehalose, which has several important functions; therefore, enhancing TreS expression is desirable. Here, a recombinant *Bacillus subtilis* W800N (ΔamyE)-P_{glv} strain was constructed to achieve enhanced expression of TreS. Process optimization strategies were developed to improve the expression level of TreS in *B. subtilis* W800N (ΔamyE)-P_{glv}. Intracellular activity of TreS was induced using 60 g/L of maltose in shake flask culture. The protein activity reached 5211 ± 134 U/g at 33 °C and pH 7.0 in Luria-Bertani medium. A fed-batch fermentation strategy was applied in a 30 L fermenter containing 18 L terrific broth to achieve high cell density by replacing glycerol with high maltose syrup as a carbon source and an inducer. After 32 h of fermentation, recombinant *B. subtilis* W800N (ΔamyE)-P_{glv} activity reached 6850 ± 287 U/g dry cell weight. Our results demonstrate the efficiency of the P_{glv} promoter in increasing the expression of TreS in *B. subtilis* W800N (ΔamyE)-P_{glv}.

Trehalose is a non-reducing disaccharide comprising two glucose units joined by an α-1,1-glycosidic bond⁴. Functions of trehalose include protection from desiccation⁵,⁶, stabilization of vaccines⁷–⁹, and protection of mammalian cells against desiccation⁹. Trehalose is widely used in agricultural, cosmetic, and pharmaceutical industries due to its stability with most chemically nonreactive sugars⁸,⁹. Trehalose is industrially produced directly from maltose using trehalose synthase (TreS, EC 2.4.1.245) because of the low cost of the substrate and the simplicity of the process, which may be beneficial for industry-scale production¹⁰,¹¹.

Dual- and single-enzymatic methods are commonly used in the production of trehalose. The drawbacks of the dual-enzymatic method are that it requires the cultivation of two types of recombinant bacteria. These bacteria must be lysed to obtain crude enzymes, which are used for conversion. Thus, the process is associated with the risk of contamination with endotoxins. The single-enzymatic method involves the use of *Escherichia coli* as the expression vector. The removal of endotoxin is also difficult with this method. In addition, isopropyl β-D-1-thiogalactopyranoside (IPTG), the inducer used for the expression of the two enzymes, is expensive and difficult to remove.

TreS is produced by several strains of bacteria, including *Bacillus licheniformis*, *Thermobifida fusca*, and *Yarrowia lipolytica*, which have been considered for use in trehalose production. However, the yield of TreS from bacteria is usually very low, limiting the application of bacterial systems for TreS production. Genetic engineering of bacteria has been carried out with the goal of improving the expression of TreS, including cloning of *treS* gene in *E. coli*¹²–¹⁴. This approach is time-consuming, and the removal of pathogenic enterobacteria prior to the use of trehalose in the food industry is an expensive process.

*B. subtilis* is generally recognized as safe (GRAS) and has been used as a standard host strain because of the ease of cultivation and control over production, which are beneficial factors for large-scale production. *B. subtilis* is used in the food industries in many countries because of its GRAS designation. Several inducible expression

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systems containing inducer-specific promoters, including those for T7, grac, spac, xylA, sacB, and α-amylose promoter are widely used in B. subtilis. Many proteins have been successfully overexpressed in B. subtilis using these inducible promoters. However, the high cost of inducer compounds such as IPTG limits their industrial application. The promoter of glv operon (P\textsubscript{glv}) in B. subtilis is positively regulated by maltose, which is inexpensive and widely available. Thus, P\textsubscript{glv} has been considered as a potential promoter system with industrial applications. However, P\textsubscript{glv} promoter is markedly repressed by glucose via a catabolism repression element (Cre) located downstream of the transcription origin site of P\textsubscript{glv} promoter.

In this study, the Cre gene sequence on P\textsubscript{glv} promoter was mutated by site-directed mutagenesis. The recombinant plasmid P\textsubscript{glv}-pHT01-treS was constructed as an expression vector. The P\textsubscript{glv} promoter regulated the expression of TreS in recombinant B. subtilis W800N (ΔamyE)-P\textsubscript{glv}. Moreover, the conditions for the TreS-catalyzed production of trehalose were optimized.

### Methods

#### Bacterial strains, plasmids, primers, and culture media.

The bacterial strains, plasmids, and primers used in this study are listed in Table 1. Plasmid pHT01 was purchased from Hangzhou Biosci Biotech Co., Ltd. (Hangzhou, China). E. coli DH5α and B. subtilis 168 from our laboratory culture collection were used as hosts for gene cloning. The engineered B. subtilis WB800N was used for gene expression. Luria-Bertani (LB) medium containing 10 g/L tryptone, 5 g/L yeast extract, and 10 g/L sodium chloride (NaCl) was used as the culture medium. Terrific broth (TB) medium containing 12 g/L tryptone, 24 g/L yeast extract, 4 mL/L glycerol, 2.4 g/L monopotassium phosphate (KH\textsubscript{2}PO\textsubscript{4}), and 16.5 g/L dipotassium phosphate (K\textsubscript{2}HPO\textsubscript{4}) was used as the fermentation medium. Growth medium (GM) containing 10 g/L tryptone, 5 g/L yeast extract, 10 g/L NaCl, and 0.5 mol/L sorbitol was used as the proliferation medium, while regrowth medium RM containing 10 g/L tryptone, 5 g/L yeast extract, 10 g/L NaCl, 0.5 mol/L sorbitol, and 0.38 mol/L mannitol was used as the recovery medium. The electroporation buffer comprised 0.5 mol/L sorbitol, 0.5 mol/L mannitol, 0.5 mol/L trehalose, and 10% glycerol. The antibiotics used for selection were 25 μg/mL chloramphenicol and 50 μg/mL spectinomycin.

### Construction of the recombinant expression vector pHT01-P\textsubscript{glv}-treS.

The Cre gene sequence (GTAAACGTTATCA) was embedded in the P\textsubscript{glv} promoter. In the presence of the Cre gene sequence, a glucose metabolism (CcpA) inhibits the expression from the P\textsubscript{glv} promoter. The site-directed mutagenesis of Cre gene with CG to AT change may alleviate the repression of glucose and improve the expression and activity of the protein. Therefore, mutant fragments of P\textsubscript{glv}+1 (269 bp) and P\textsubscript{glv}+2 (121 bp) were synthesized by the primer pairs P\textsubscript{glv}+1-F/P\textsubscript{glv}+1-R-C and P\textsubscript{glv}+2-F-C/P\textsubscript{glv}+2-R-C using the chromosome of B. subtilis 168 as the template. A BamHI restriction site was introduced at the 5'-end of fragment P\textsubscript{glv}+1 and the 57 bp homologous sequence fragment upstream of glvA gene, and the TAA stop codon and Shine-Dalgarno (SD) sequence of Pglv gene from B. subtilis 168 were introduced into the fragment of P\textsubscript{glv}+2. Overlapping polymerase chain reaction (PCR) connected P\textsubscript{glv}+1 and P\textsubscript{glv}+2 fragments to obtain P\textsubscript{glv}+1+2 fragments, wherein the overlap section of P\textsubscript{glv}+1 and P\textsubscript{glv}+2-F introduced the mutation. The fragment P\textsubscript{glv}+1+2 was created with site-directed mutations in Cre sites was obtained through the fusion of P\textsubscript{glv}+1 and P\textsubscript{glv}+2 overlap PCR fragments using primer pairs P\textsubscript{glv}+1-F and P\textsubscript{glv}+2-R. The complete fragment comprised a 6-bp enzyme digestion site BamHI sequence added to the corresponding gene sequence on Pglv promoter was mutated by site-directed mutagenesis. The recombinant plasmid P\textsubscript{glv}-pHT01-treS was constructed as an expression vector. The P\textsubscript{glv} promoter regulated the expression of TreS in recombinant B. subtilis W800N (ΔamyE)-P\textsubscript{glv}. Moreover, the conditions for the TreS-catalyzed production of trehalose were optimized.
site sequence. The $P_{\text{glv}}$-treS gene fragment was obtained by overlapping PCR with $P_{\text{glv}}$-1 + 2 and two treS gene fragments using $P_{\text{glv}}$-1 – F and treS-R primer pairs. The resulting 2.5-kb fragment was digested with BamHI and AutII and cloned into pH101 digested with the same enzymes, resulting in $P_{\text{glv}}$-P101-treS (Fig. S1).

### Screening of recombinant B. subtilis WB800N ($\Delta$amyE).

Single-cross integration was used to excise the amyE gene from the chromosome of B. subtilis WB800N. The sequence fragment comprising a 6-bp BamHI digestion site and a 500-bp homologous fragment upstream of amyE gene was cloned in B. subtilis 168 using amyE-1-F and amyE-1-R primers. The spectinomycin resistance gene was amplified from plasmid pHIL2 using primers Spec-F and Spec-R. These two fragments were fused by overlapping PCR and transformed into the competent B. subtilis WB800N in LB agar containing 50 μg/mL spectinomycin to screen for spectinomycin-sensitive recombinants. The obtained strain B. subtilis WB800N ($\Delta$amyE) was analyzed with PCR using primers Spec-F and Spec-R, and subjected to sequencing to confirm the deletion of amyE gene (Fig. S2). The integration of spec gene at amyE locus disrupted the amyE gene, resulting in the manifestation of amylase-negative phenotype on LB medium supplemented with 1% starch. After incubation at 37°C overnight, the plates were stained with iodine to examine amylase activity.

### Preparation of competent recombinant B. subtilis WB800N.

Single colonies of B. subtilis WB800N from LB agar medium were individually inoculated in 5 mL LB liquid medium for 24 h at 37°C with constant stirring at 200 rpm. A 500-μL aliquot of culture was transferred to 50 mL GM proliferation medium. After the optical density at 600 nm (OD$_{600}$) wavelength reached 1.0, the culture was transferred to a 100-mL centrifuge tube in an ice water bath for 10 min, followed by centrifugation at 4°C and 5,000 rpm for 8 min to collect bacterial cells. The bacteria were washed thrice with pre-cooled ETM electroporation buffer and finally suspended in 500 μL of ETM buffer to obtain competent B. subtilis. The competent bacteria were stored at −80°C (60 μL per tube).

### Construction of recombinant strains.

One tube each of 60 μL of B. subtilis WB800N and B. subtilis WB800N ($\Delta$amyE) was recovered from −80°C freezer and uniformly mixed with 6 μL of pH101-P$_{glv}$-treS and pH101-P$_{glv}$-treS plasmid solution. After pre-cooling for 5 min, the mixed liquid was added to a 2-mm electroporation cup. The samples were subjected to electric shock at 1,500 V and 5 ms with an Eppendorf electric rotating apparatus. After electroporation, the electroporation mixture was rapidly mixed with 1 mL of RM medium at 37°C and 180 rpm for 4 h. The recombinant bacteria were recovered by centrifugation and grown on LB agar medium containing chloramphenicol (25 μg/mL) at 37°C for 48 h. Chloramphenicol-resistant bacteria were recovered and the recombinant plasmid in the bacteria was extracted for use as template. The treS gene was amplified by treS-F-C/treS-R primer to obtain a positive clone of B. subtilis WB800N.

### Induced expression and optimization of shake flask induction conditions.

Each recombinant plasmid was transformed into B. subtilis W800N and B. subtilis W800N ($\Delta$amyE), yielding different TreS-producing strains. A single colony of each recombinant strains selected using LB agar containing antibiotics was inoculated into a 250-ml shaking flask containing 50 mL LB liquid medium with chloramphenicol. The primary seed culture was obtained by incubation at 37°C for 12 h. A total of 1 mL of primary seed culture of each recombinant strain was transferred to a 500-ml shaking flask containing 100 mL LB medium supplemented with 25 μg/mL chloramphenicol and cultured at 37°C overnight on a rotating oscillator (200 rpm/min) until the OD$_{600}$ value reached 1.2. Maltose was added at a final concentration of 60 g/L to induce protein expression during the subsequent 18 h cultivation at 27°C, 30°C, 33°C, and 37°C to examine the temperature of protein expression. In another experiment, final maltose concentrations of 20–100 g/L were used to induce the expression of target protein at 37°C. Maltose was added (final concentration of 60 g/L) at various time points during the cultivation of the recombinant strains at 33°C to examine the influence of time of maltose addition on growth and TreS production. Samples were collected at certain intervals, and OD$_{600}$ and enzyme activities were measured. Each reported value represents an average of two or three independent measurements and does not vary from the mean value by over 10%.

### Fed-batch fermentation.

The recombinant B. subtilis W800N ($\Delta$amyE)-P$_{glv}$ strain was used to scale-up fermentation in 30 L fermenters (Bailun Biological Technology Co. Ltd., China) containing 20 L of fermentation medium and 100 μg/mL of chloramphenicol. Seed culture in LB medium was inoculated (5% v/v) into TB medium for cultivation. The dissolved oxygen concentration was maintained between 10% and 30% by stirring (0–650 rpm) and a constant sterile air flow rate (0.2–1 vessel volume/min) maintained. A pH value of 7.0 ± 0.2 was maintained using automatic pH control with the simultaneous addition of 250 g/L sodium hydroxide solution and 25% hydrochloric acid solution. Temperature was controlled at 37°C for cell growth and expression. TreS was expressed using a heating tube and integrated cooling system from 37°C to 33°C. High concentration of maltose syrup containing 88–90% maltose, 2.5–4% glucose, and 4–6% polysaccharide was used as inducer because of its low cost and easy availability. Samples were obtained every 2 h to analyze the dry cell weight and TreS enzyme activity after induction. After fermentation, the culture was centrifuged (8,000 rpm for 20 min), washed twice with distilled water, and dried to a constant weight at 80°C to determine the biomass of the culture in terms of dry cell weight.

### Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and TreS activity.

The cultured bacteria were harvested by centrifugation at 8,000 rpm for 20 min at 4°C. The OD$_{660}$ of every sample was measured, and a volume corresponding to approximately the same OD$_{660}$ was harvested and centrifuged. The pelleted cells were resuspended in phosphate-buffered saline (PBS, pH 8.0) and disrupted by sonication on ice using 5 s pulses at an interval of 5 s for 15 min. The crude cell extract was separated by centrifugation and subjected to...
SDS-PAGE according to the standard procedure. The amount of TreS produced was determined by staining the gels with Coomassie Brilliant Blue R250.

**Preparation, separation, and crystallization of TreS.** To identify the enzymatic characteristics of TreS, the enzyme was purified by nickel-nitrilotriacetic affinity chromatography for 1 h using maltose (100 g/L final concentration) as substrate at different pH and temperature values. The enzyme activity of TreS at different temperatures was studied in 100 mM phosphate buffer (pH 8.0) during a 60-min incubation using 100 g/L maltose as substrate. To examine the thermal stability of TreS, TreS was pre-incubated at temperatures from 10 °C to 60 °C for 60 min at pH 8.0. Residual TreS activity was measured at 25 °C. The optimum temperature for enzyme activity was 25 °C, and the enzyme activity was relatively stable from 10 °C to 40 °C. However, TreS activity rapidly decreased with time as the reaction temperature exceeded 40 °C (Fig. S3). The enzyme activity of TreS at various pH values was studied at a constant temperature of 25 °C in 100 mM phosphate buffer (pH 3.0–10.0) for 60 min using 100 g/L maltose as substrate. To examine the temperature and pH stability of TreS, the enzyme was pre-incubated in buffers with pH varying from 3.0 to 10.0 for 60 min at 25 °C. Residual TreS activity was measured at pH 8.0. The optimal pH for enzyme activity was 8.0, and the enzyme activity was relatively stable between pH 6.5 and 9.0 (Fig. S4).

To obtain trehalose at high purity, the pH of the conversion system containing the mixed syrup was adjusted as substrate. To examine the temperature and pH stability of TreS, the enzyme was purified by nickel-nitrilotriacetic affinity chromatography for 1 h using maltose (100 g/L final concentration) as substrate at different pH and temperature values. The enzyme activity of TreS at different temperatures was studied in 100 mM phosphate buffer (pH 8.0) during a 60-min incubation using 100 g/L maltose as substrate. To examine the thermal stability of TreS, TreS was pre-incubated at temperatures from 10 °C to 60 °C for 60 min at pH 8.0. Residual TreS activity was measured at 25 °C. The optimum temperature for enzyme activity was 25 °C, and the enzyme activity was relatively stable from 10 °C to 40 °C. However, TreS activity rapidly decreased with time as the reaction temperature exceeded 40 °C (Fig. S3). The enzyme activity of TreS at various pH values was studied at a constant temperature of 25 °C in 100 mM phosphate buffer (pH 3.0–10.0) for 60 min using 100 g/L maltose as substrate. To examine the temperature and pH stability of TreS, the enzyme was pre-incubated in buffers with pH varying from 3.0 to 10.0 for 60 min at 25 °C. Residual TreS activity was measured at pH 8.0. The optimal pH for enzyme activity was 8.0, and the enzyme activity was relatively stable between pH 6.5 and 9.0 (Fig. S4).

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**High-performance liquid chromatography (HPLC) analysis of trehalose production.** The maltose solution (300 g/L) was treated with crude enzyme (5 mL) at 25 °C for 1 h, followed by incubation in boiling water for 15 min. The solution was centrifuged at 13,000 rpm for 15 min, and the supernatant was analyzed by HPLC, as previously described22. The transition rate of trehalose was calculated as follows:

\[
\alpha = \frac{m_{\text{trehalose}}}{m_{\text{trehalose}} + m_{\text{glucose}} + m_{\text{maltose}}} \times 100\%
\]

where \(m_{\text{trehalose}}\), \(m_{\text{glucose}}\), and \(m_{\text{maltose}}\) are weights of trehalose, glucose, and maltose, respectively.

**Results**

**Expression of TreS.** The recombinant clones (B. subtilis W800N (ΔamyE)-Pglv, M1; B. subtilis W800N-Pglo, M2; and B. subtilis W800N (ΔamyE)-Pgac, M3) were subjected to fermentation followed by sonication to extract intracellular proteins. Comparison of enzyme activity of the recombinant B. subtilis strains (Fig. 1) showed that the enzyme activity of M1 reached 4,870 ± 189 U/g dry weight with 60 g/L maltose as the induction substrate after 18 h, higher than that of M2 (3,750 ± 185 U/g) using 60 g/L maltose as the induction substrate and M3 (2,490 ± 87 U/g) in the presence of 0.5 mM IPTG as inducer. The expression of recombinant TreS was confirmed by 10% SDS-PAGE. The recombinant proteins had a molecular weight of nearly 75 kDa (Fig. 2), consistent with the molecular weight deduced from the amino acid sequence of TreS. These data suggest that the Pglo promoter was superior to Pgac promoter in enhancing TreS expression and that the knockout of amyE gene further increased the expression of TreS in the recombinant group. As the highest yield was obtained from B. subtilis W800N (ΔamyE)-Pglv, the construct (ΔamyE)/pHT01-Pglv-treS was chosen for further studies.

**Optimization of conditions for recombinant enzyme production.** To identify the optimal conditions for recombinant enzyme production, pH, induction temperature, maltose induction concentration, and time of induction were assessed.

Temperature is an important parameter for recombinant protein expression in host bacteria. In general, the growth of host bacteria at low temperatures lowers the rate of synthesis of recombinant proteins and increases exogenous protein solubility. To determine the optimal temperature for TreS production, M1 was induced at 27 °C, 30 °C, 33 °C, 37 °C, and 40 °C in LB medium. The highest cell density (OD600) of 5.6 ± 0.23 was obtained at 40 °C, while the maximum activity of 4918 ± 193 U/g was reported at 33 °C (Fig. 3). Although temperatures of 37 °C and 40 °C may be more suitable for the growth of bacteria, the lower enzyme activity per gram weight limits the total enzyme activity. Therefore, M1 was more conducive for expression at a temperature of 33 °C.

The pH adaptability of M1 was investigated using a standard assay. The enzyme activity was maximum at pH 7.0 and rapidly increased between pH 6.5 and 7.0 (Fig. 4); no significant changes were reported between pH 7.0
and 8.0. In this expression system, both weak acidic conditions (pH < 6.5) and alkaline conditions (pH > 8.8) strongly inhibited the activity of TreS.

The expression of TreS was induced by maltose; different concentrations of maltose were used to identify the optimum concentration of inducer. TreS activity increased with an increase in maltose concentration and an optimum activity of 5189 ± 197 U/g was achieved following protein expression induction using 60 g/L maltose (Fig. 5). However, further increase in maltose concentration reduced the activity of TreS. In contrast, maltose concentration influenced bacterial growth, but no significant changes were evident at the maltose concentrations used in the present study.

A set of induction experiments was performed using M1 to determine the optimum dose of the inducer xylose. All induced samples were analyzed for TreS activity and cell density after fermentation. As shown in Fig. 6, the activity was high upon addition of xylose to the culture in the mid-logarithmic phase of growth. The maximum activity of 5211 ± 134 U/g was obtained by adding xylose at an OD600 of 1.6.

Scale-up production of TreS. To scale-up the production of TreS, M1 culture was induced to express TreS in a 20 L fermenter containing glycerol-free TB medium (Fig. 7). High maltose concentration was added to the medium in three phases. The first phase was started in the presence of an initial high maltose syrup concentration of 10 g/L. After the OD600 reached approximately 1.6, the second dose of high maltose syrup was added at a
**Figure 3.** Effect of temperature on the intracellular activity and growth of cells (n = 3, x ± SD).

**Figure 4.** Effect of pH on the intracellular activity and growth of cells (n = 3, x ± SD).

**Figure 5.** Effect of different maltose concentrations on the enzyme activity of recombinant *B. subtilis* W800N (∆amyE)-P_{glv} (n = 3, x ± SD).
concentration not exceeding 40 g/L. After 6 h of fermentation, the third dose of high maltose syrup was added at a concentration of 64 g/L. In the second phase, the feed medium (TB medium without glycerol) was used until the OD600 value reached 25 and the temperature decreased to 33 °C. Intracellular TreS enzyme activity reached 6850 ± 287 U/g after 32 h fermentation, a value that was 1.3 times the level attained in shake flask cultivation.

Purification of trehalose. As the optimum temperature for TreS expression was determined to be 25 °C, the enzyme exhibited competent transformation activity at low temperatures. To avoid the growth of microorganisms at room temperature (25 °C) or to prevent any effect on transformation, 15 °C was chosen as the reaction temperature. Recombinant *B. subtilis* W800N cell lysate containing TreS was directly used to convert high maltose syrup for 12 h at 15 °C and pH 8.0 with agitation at 60 rpm in the presence of 300 U TreS/g maltose and 300 g/L high maltose syrup. After conversion in the mixed tank for 12 h under optimum conditions, a mixture of glucose, maltose, and trehalose was obtained. The maximal conversion efficiency of TreS was 67% (Fig. 8). After crystallization, the purified trehalose (≥ 99%) was analyzed by HPLC (Fig. 9).

Discussion
Endospore-forming bacteria of the genus *Bacillus* are used to manufacture several industrial enzymes, in particular amylase and protease. In the majority of cases, a constitutive expression pattern that couples growth and expression is employed. Plasmid pHT01 is a versatile *E. coli-B. subtilis* shuttle vector. The full-length of 7,955 bp is produced using Pgrac as the control element and IPTG as the inducer. Although most published studies have focused on the use of IPTG as the inducer, IPTG is a potentially toxic chemical and is expensive. To avoid these drawbacks, we constructed a new recombinant expression vector, pHT01-Pglv, using Pglv as the control element instead of Pgrac. Pglv was fused to a synthetic ribosome-binding site fragment and tandem stop codons were introduced to efficiently avoid read-through transcription. The presence of unique restriction sites downstream of the maltose-inducible promoter allows the proper ligation of coding regions for target protein overproduction. *B. subtilis* WB800N can secrete α-amylase to hydrolyze maltose and form glucose in the medium, resulting in the marked reduction in the induction efficiency of maltose. Therefore, it is necessary to knockout *amyE* gene in the
chromosome of *B. subtilis* WB800N. The spectinomycin gene was integrated into the chromosome of *B. subtilis* WB800N at *amyE* locus by a single crossover event. Amylase activity analysis confirmed the interruption of the complete expression of amylase, resulting in the abrogation of amylase activity. The recombinant *B. subtilis* strains W800N-P<sub>glv</sub> and W800N-P<sub>grac</sub> described in the present study produced functional TreS in the presence of simple fermentation media containing xylose as inducer. Enzyme activity was the highest for the strain *B. subtilis* W800N (ΔamyE)-P<sub>glv</sub>, suggesting that the P<sub>glv</sub> promoter is more efficient in expression the of TreS than the P<sub>grac</sub> promoter.

Temperature and pH are usually the main factors that determine the production efficiency of TreS in recombinant hosts<sup>26</sup>. The present findings are consistent with those previously reported on the optimum temperature of approximately 33 °C for TreS expression in different hosts. At present, TreS cloned from *B. subtilis* W800N (ΔamyE)-P<sub>grac</sub> showed the highest activity at pH 7.0, similar to the activities observed for the enzymes from *Mycobacterium smegmatis* (pH 7.0)<sup>27</sup> and *Corynebacterium glutamicum* ATCC 13032 (pH 7.0)<sup>14</sup>.

In general, fed-batch culture is used in bio-industrial processes to achieve high cell density in bioreactors<sup>26</sup>. The fermentation period of recombinant *E. coli* BL(DE3) was only 14 h and the time to add the inducer of protein expression was easy to calculate. Lactose, as an inducer instead of IPTG, is more economical and easier to obtain<sup>28</sup>. Furthermore, lactose can be completely consumed as a carbon source during the fermentation process, leaving no residue or product in the medium<sup>28</sup>. The recombinant *B. subtilis* W800N (ΔamyE)-P<sub>glv</sub> described herein produced functional TreS on a simple fermentation medium containing xylose as the inducer. The maximum TreS activity was obtained upon the addition of xylose at an OD<sub>600</sub> value of 1.6. Crude TreS solutions were directly used for conversion, thereby avoiding the process of purification of TreS. TreS was purified by nickel-nitritoltriacetic affinity chromatography for 1 h using maltose a substrate at different pH and temperature values. The pH of the conversion system containing mixed syrup was adjusted to 5.0, resulting in the production of very pure trehalose. The trehalose solution was concentrated by a plate-type evaporator and cooled to crystallize the product. The purity of the produced trehalose exceeded 99%.

**Figure 8.** HPLC analysis of mixed syrup in the process of conversion.

**Figure 9.** Trehalose product analysis. (A). Trehalose product crystallization. (B). Analysis of HPLC chromatogram from trehalose purification.
Conclusions
We obtained a recombinant B. subtilis W800N (ΔamyE)−Pglv for the intracellular expression of TreS using Pglv as the promoter and maltose as the inducer. The recombinant B. subtilis W800N (ΔamyE)−Pglv efficiently expressed TreS, indicating that the mutant Pglv can evidently alleviate the repression caused by glucose to enhance the expression of the Pglv promoter system. In the food and pharmaceutical industries, processes that are inexpensive, simple, and free of endotoxins are key. The direct use of recombinant B. subtilis W800N cell lysates to prepare trehalose from maltose takes into account these criteria. Thus, the expression of TreS by a recombinant B. subtilis W800N using maltose as an inducer not only provides a safe and efficient expression system for TreS but also provides a new direction for the industrial production of trehalose.

Data Availability
All data generated or analyzed during this study are included in this published article (and its Supplementary Information files).

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Author Contributions
L.H.L. and W.T.F. contributed to the conception and design of the study. Y.S.J., W.T.F. and W.R.M. performed the experiment. L.H.L. and L.H. processed the data and performed statistical analysis. W.T.F. revised the manuscript. L.H.L. wrote the paper. All authors read and approved the final manuscript.

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