Molecular cloning and extracellular expression of cyclodextrin glycosyltransferase gene from Bacillus sp. NR5 UPM

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The cloning of a polymerase chain reaction (PCR) gene fragment from Bacillus sp. NR5 UPM isolated from the soil in Malaysia into an Escherichia coli expression vector was successfully carried out. Analysis of the nucleotide sequences revealed the presence of an open reading frame of 2112 bp which encoded a protein containing 704 amino acids with a putative molecular weight of 78.6 kDa. The deduced amino acids sequence showed about 98% homology with the CGTase from Bacillus sp. KC201. Compared to the wild type, the CGTase that was produced in E. coli cells only required one-fourth of culture time and neutral pH to produce CGTase. After 12 h of cultivation, the CGTase activity in the culture medium reached 29.6 U/ml, which was approximately 2.5-fold higher than the CGTase from the parental strain. The CGTase was produced extracellularly by E. coli (94%) indicating the signal peptide was functional in E. coli.

Key words: Molecular cloning, nucleotide sequence, cyclodextrin glycosyltransferase, Bacillus sp. NR5 UPM.

INTRODUCTION

Cyclodextrin glycosyltransferase (CGTases, 1,4-α-D-glucopyranosyltransferase (cyclizing), EC 2.4.1.19) is an important enzyme that catalyzes the formation of α-CD, β-CD and γ-CD, containing 6, 7 and 8 glucose residues linked with α-1,4-glucosidic bonds, respectively. Due to their unique abilities to form inclusion complexes with a variety of hydrophobic materials and to entrap volatile compounds, these CDs have found extensive applications in food, pharmaceuticals, agricultural chemicals, cosmetics, industrial chemicals and others (Hashimoto, 2002).

Recently, many researchers have studied the molecular cloning of CGTase genes and analysed the genetic information in order to provide a better CGTase production method. The over expression of CGTase genes could enhance the enzyme activity, reduce cultivation time and produce less contaminating proteins compared to wild type (Charoensakdi et al., 2007). In this study, we have succeeded in isolating the CGTase gene from Bacillus sp. NR5 UPM. The isolated CGTase gene was cloned into an E. coli expression vector and over expressed to study the improved properties of the enzyme.

MATERIALS AND METHODS

Bacterial strain, plasmids and media

Bacillus sp. NR5 UPM was grown in Horikoshi medium II, containing 1% (w/v) soluble starch, 0.5% (w/v) yeast extract, 0.5% (w/v) peptone, 0.1% (w/v) K₂HPO₄, 0.02% (w/v) MgSO₄·7H₂O and
Table 1. Primers used for amplification of upstream and downstream adjacent sequences based on known sequences of the CGTase gene.

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequence (5' to 3' direction)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DW2-ACP2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>ACPCTCGA</td>
</tr>
<tr>
<td>DW2-ACP3&lt;sup&gt;a&lt;/sup&gt;</td>
<td>ACPCCTACG</td>
</tr>
<tr>
<td>DW2-ACPn&lt;sup&gt;b&lt;/sup&gt;</td>
<td>ACPN</td>
</tr>
<tr>
<td>Universal Primer, UniP2</td>
<td>GAGTTTAGGTCACGGCGTTGG</td>
</tr>
<tr>
<td>TSP1R&lt;sup&gt;c&lt;/sup&gt;</td>
<td>GCCAAGTGGTGTGTTTGG</td>
</tr>
<tr>
<td>TSP2R&lt;sup&gt;c&lt;/sup&gt;</td>
<td>GTTTGCGACCTTCTGACATA</td>
</tr>
<tr>
<td>TSP3R&lt;sup&gt;c&lt;/sup&gt;</td>
<td>GACACCATTCTAAACATAGTTAGGG</td>
</tr>
<tr>
<td>TSP1F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>AAAAAATTAGTGGTATCCG</td>
</tr>
<tr>
<td>TSP2F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>CGAAATCTACTGCTGATAACCTG</td>
</tr>
<tr>
<td>TSP3F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>CCCCCAAAACCTCCCTTG</td>
</tr>
<tr>
<td>nTSP3F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>GGCTATGCGACTACACTGAAC</td>
</tr>
<tr>
<td>n2TSP3F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>ACAATGGCCAAACAGGCCAG</td>
</tr>
<tr>
<td>n3TSP3F&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>AGGCTAAGCGGTGGGAGAAAG</td>
</tr>
</tbody>
</table>

<sup>a</sup> Annealing control primer, provided by the manufacturer;  
<sup>b</sup> Target specific primer, designated on the basis of known sequence of CGTase gene;  
<sup>c</sup> Target specific primer, designated on the basis of the newly determined sequence in the third PCR operation.

DNA manipulation and amplification of known sequences of CGTase gene

Chromosomal DNA was isolated from Bacillus sp. NR5 UPM by the method of Ish-Horowicz and Burke (1981). Other general molecular experiments involving PCR reaction and transformation were performed according to the methods described by Sambrook and Fritsch (1989). Genomic DNA was extracted and used as the template for the preparation of the CGTase gene by PCR technique. A pair of degenerate primers; forward primer C1, 5'-GGN GAY TGG CAR GGN-3' and reverse primer C2, 5'-CAT RTC RTG RTT RTC DAT RAA-3' designed based on the conserved sequences of the CGTase gene was used as PCR primers (Ong et al., 2008). The reaction contained 1 μg of Bacillus sp. NR5 UPM DNA, 100 pmol of each forward and reverse primer, 0.01 U/μl Taq Polymerase in 1X reaction buffer and 0.2 mM of each dNTPs. The DNA was initially denatured at 94°C for 5 min, followed by denaturation at 94°C for 20 s. Then, the annealing step was carried out at 50°C for 20 s and extension at 72°C for 5 s. The total number of cycles was 30.

Sequence analysis of the adjacent region of the known sequences

The adjacent region of known sequences of the CGTase gene was amplified by PCR using DNA walking SpeedUp™Premix Kit II (Seegene, Inc). The PCR reaction was performed with the following primers (Table 1) and Bacillus sp. NR5 UPM DNA as the template. The primers used were designed to amplify the upstream and downstream adjacent sequences based on known sequences of the CGTase gene. For the first PCR reaction, the primers used were a combination of either DW2-ACP2 or DW2-ACP3 together with TSP1R and TSP1F to amplify the upstream and downstream regions, respectively. Each of the first PCR products was used as a template for the second PCR. The second PCR was performed with DW2-ACPn together with TSP2R for amplification of the upstream region and TSP2F for amplification of the downstream region.

The same operation was repeated with the other primers to determine the nucleotide sequence of the extending region. The primers used were TSP3R for the amplification of the upstream region and TSP3F for amplification of the downstream region together with Universal Primer, UniP2 for each of them. Other primers were designed on the basis of the newly determined sequence in the third PCR reaction, namely nTSP3F, n2TSP3F and n3TSP3F for amplification of the downstream region. DNA sequencing of the resulting third PCR was performed by First Base Laboratories, Biosyntech Malaysia on the ABI PRISM 377 DNA sequencer.

Cloning of CGTase gene

The expression plasmid for the new ORF was constructed as follows. The new ORF of the CGTase gene was amplified by PCR. The chromosomal DNA of Bacillus sp. NR5 UPM was used as a template with 5'-AGCGGATCCCTTTATTTATACGTGTT-3' (BamHI site underlined) as the forward primer and 5'-GTCAGCTTTTACCAATTACGACCTGTT-3' (HindIII site underlined) as the reverse primer. The amplified fragment was digested with both BamHI and HindIII, and then ligated into the corresponding sites on the vector pUC 19. The resultant plasmid was designated as pCGT3D. The ligation products were used for transformation into E. coli JM109 cells. All the white colonies were tested by the agar plate assay method which contained 1% soluble starch and 100 μg/ml ampicillin in LB agar. After growth at 37°C for 24 h, the ability of the possible white colony to form halo zone after exposure to KI-I<sub>2</sub> was observed.

Expression of CGTase gene

A single colony of E. coli JM109 cells harbouring plasmid pCGT3D was inoculated into 10 ml of LB medium containing 100 μg/ml
ampicillin and grown at 37°C overnight. 10% of overnight culture was then diluted into 100 ml of terrific broth (TB) containing 100 μg/ml ampicillin in a 250 ml flask. The culture was incubated at 37°C on a rotary shaker (200 rpm). Samples were withdrawn hourly and analyzed for enzyme activities.

**Nucleotide and protein sequence analysis**

The blastn and blastx programs provided by the National Center for Biotechnology Information (NCBI) were used to search for existing sequences that were similar to the sequences we obtained. The amino acid and nucleotide sequences we obtained were compared with other sequences using the BioEdit 7.01 program.

**Cell fractionation**

Cell fractionation was performed as described by Ding et al. (2010). The culture broth was centrifuged at 12,000 rpm for 5 min at 4°C to obtain the extracellular fraction. Then, 1 ml culture was harvested and washed twice with 1 ml of 30 mM Tris-HCl buffer (pH 7.0). The cells were resuspended in the same buffer containing 25% (w/v) sucrose and 1 mM EDTA to separate the periplasmic fraction. The cell suspension was incubated on ice for 2 h and centrifuged at 12,000 rpm for 5 min at 4°C. The supernatant obtained was collected as the periplasmic fraction.

**Assay of β-CGTase**

The phenolphthalein assay (Kaneko et al., 1987) was used to determine the CGTase activity. The reaction mixture which contained 1 ml of 40 mg of soluble starch in 100 mM phosphate buffer (pH 6.0) and 0.1 ml enzyme solution was incubated at 60°C for 10 min in a water bath. To stop the reaction, 3.5 ml of 30 mM NaOH was added to the mixture. Subsequently, 0.5 ml of 0.02% (w/v) phenolphthalein in 5 mM Na2CO3 solution was added to the reaction mixture and mixed well.

Then, the reduction in colour intensity was measured at 550 nm after leaving at room temperature for 15 min. Control lacking CGTase was analysed simultaneously with each batch of samples. The soluble starch and enzyme were replaced by 0.5 mg of β-CD and 0.1 ml of water, respectively as a standard. One unit of enzyme activity was defined as the amount of the enzyme that formed 1 μmol β-CD per min under the conditions aforesaid.

**RESULTS AND DISCUSSION**

**Amplification of known sequence of CGTase gene**

Approximately, 800 bp of polymerase PCR product was successfully amplified from the genomic DNA of *Bacillus* sp. NR5 UPM. The nucleotide sequence analysis for the amplification of known sequences of CGTase gene resulted in a 752 bp size product. The sequence was subjected to a search against all known sequences in databases using blastn and blastx searches. It showed 95% maximum identity with *Bacillus* sp. KC201 CGTase (GenBank accession no. D13068.1).

**Sequence analysis for amplification of CGTase gene**

The DNA walking strategy was used for the amplification of the CGTase gene with genomic DNA of *Bacillus* sp. NR5 UPM as the template. The nucleotide sequence analysis of the upstream and downstream adjacent sequences based on known sequences of CGTase gene revealed a new ORF of CGTase gene from *Bacillus* sp. NR5 UPM. The nucleotide sequence has been deposited in the GenBank database under the accession number HQ876173. The new ORF of the CGTase gene consisted of 2112 bp and encoded 704 amino acids.

The open reading frame shows unique properties with TTG, rather than ATG as a start codon. The same result has also been reported in CGTase from *Bacillus* sp. G1 (Ong et al., 2008), *Bacillus* sp. TS1-1 (Rahman et al., 2006), and *Bacillus ohbensis* (Sin et al., 1991). Besides TTG and ATG, GTG and CTG are also frequently used by bacteria as start codons. Initially, TTG at nucleotide 1 was thought to be a start codon. However, the identification of Shine-Dalgarno (SD) sequence showed it was located at nucleotide 40. The identification of SD sequence is important as it aided in the recognition of the initiation codon. The SD sequence, which is also known as ribosome binding site, must be located at 6 to 13 bases upstream from the start codon and the sequence should be totally complementary to the 3’ end of the 16S ribosomal RNA of *Bacillus* sp. NR5 UPM. The sequence of GGAGGA is believed to be the SD sequence which was located at 6 bp upstream from the start codon and the sequence homology among Gram-positive bacteria (von Heijne, 1986). According to the amino acid sequence deduced from the DNA sequence, the first 29 amino acid residues maybe a signal peptide which is involved in the secretion of the protein. Indeed, there is no strong sequence homology among the cleavage sites and it is known to have highly variable and rapidly evolving structures. However, the comparison of signal peptide shows consistent characteristics among Gram-positive bacteria. According to the (-3,-1)-rule of signal peptide, there is a common pattern for a probable cleavage site and this pattern shows in agreement with the amino acid residues of signal peptide deduced from the DNA sequence. The presence of Gly at position -3 is compatible with the (-3,-1)-rule where the residue in that position must not be amino acids with electrically charged side chain, aromatic or large and polar. Besides, the presence of Gly, Ile, Glu and Leu at position -3 through +1, is in accordance with the rule that requires the absence of non-polar amino acid with hydrophobic side chain at that position.

**Amino acid sequence analysis with other CGTases and amylase**

The CGTase sequence from *Bacillus* sp. NR5 UPM was
Figure 1. Nucleotide sequence of the cyclodextrin glucanotransferase (CGTase) gene from *Bacillus* sp. NR5 UPM. The two possible start codons (TTG) are shown in the boxes with the second box most likely to be a true initiation codon. The ribosome binding site, SD, is underlined. The signal peptide is shown in italics. The possible signal peptide cleavage site is shown by a vertical arrow. This sequence was submitted to the GenBank with the accession number HQ876173.
Figure 1. Nucleotide sequence of the cyclodextrin glucanotransferase (CGTase) gene from Bacillus sp. NR5 UPM. The two possible start codons (TTG) are shown in the boxes with the second box most likely to be a true initiation codon. The ribosome binding site, SD, is underlined. The signal peptide is shown in italics. The possible signal peptide cleavage site is shown by a vertical arrow. This sequence was submitted to the GenBank with the accession number HQ876173.

compared with other CGTases and amylase sequences using BLAST program. A homology search revealed that CGTase from Bacillus sp. NR5 UPM had highest similarity (98%) with Bacillus sp. KC201 CGTase (GenBank accession no. D13068.1). The other CGTases such as CGTases from Bacillus sp. G1-2004, Bacillus sp. TS1-1 and alkalophilic Bacillus 1-1 (GenBank accession nos. AAV38118.2, AAV38117.1 and P31746.1, respectively) also showed high homology (more than 95%) with the amino acid sequence of CGTase from Bacillus sp. NR5 UPM. According to Figure 2, six highly conserved regions (labelled as I to VI) were identified among CGTases and amylase. The deduced amino acid sequence of CGTases and amylase showed the homology of sites important for catalysis, which suggested a common evolutionary derivation of these two classes of enzymes (Binder et al., 1986).

Cloning and expression of CGTase gene

The sequence analysis of the open reading frame strongly suggests that it belongs to the CGTase gene. Hence, the 2.1 kb-BamHI-HindIII fragment containing the open reading frame was cloned into pUC19 under lac promoter. The CGTase gene was cloned into E. coli JM 109 as a host. The formation of clear halo zone around the colony cultured on a LB-starch plate (containing 1.5% agar, 100 μg/ml ampicillin and 1% potato soluble starch) when stained with iodine solution indicated the successful expression of starch degrading activity.

For extracellular expression of recombinant CGTase, terrific broth (TB) was used as a culture medium due to its pH-buffering capacity which gives beneficial effects on cell growth and enzyme stability. Indeed, the large amounts of yeast extract contained in the TB medium gives a critical effect on the release of the recombinant enzyme from the periplasmic space (Li et al., 2010).

Secretion of extracellular CGTase from E. coli transformant

The CGTase of Bacillus sp. NR5 UPM in E. coli was produced extracellularly into the culture medium, indicating the signal peptide of CGTase was functional in E. coli. The CGTase activities localized extracellularly and in the periplasm were 29.6 and 1.9 U/ml, respectively; that is 94 and 6%, respectively. A comparative study on the characteristics of a signal peptide shows the importance of an amino-terminal positively charged region (n-region, 1-5 residues) and a central-hydrophobic region (h-region, 7-15 residues) in targeting the enzyme into the extracellular medium. The signal peptidases play a critical role in cleaving the amino terminus of the protein once its targeting function has
been carried out. However, many cases show the translocation of the proteins ended up in the periplasm due to the failure of the integral membrane protein to remove the translocation signal which instead remained anchored to the membrane by an uncleaved signal peptide.

Meanwhile, the polar carboxy-terminal domain (c-region, 3-7 residues) is needed to specify the signal peptidase cleavage site for proper removal of the signal peptide from the mature chain (von Heijne, 1990). Many studies on molecular cloning of the CGTase gene have been carried out, in which the translocation of recombinant enzymes among the constructs varied from extracellular medium into periplasmic and intracellular space (Table 2). Among the recombinants constructed, the NR5 transformant showed the best translocation of the CGTase into the extracellular medium (94%) compared to other constructs which were around 68 to 74% (Kim et al., 1998; Yong et al., 1996). The ability of the signal peptide to secrete the enzyme predominantly in vivo outside the cell is beneficial because this will aid in the downstream processing, enhance the in vivo expression, and potentially improve the yield and quality of the recombinant enzyme.

**Table 2.** Comparison of secretion efficiency of CGTases from *Bacillus* sp. NR5 UPM with other CGTases and amylase. The six highly conserved regions in different CGTases and amylase are boxed. The numbers within brackets are GenBank accession numbers. KC201: CGTase from *Bacillus* sp. KC201 (D13068.1), BC: CGTase from *Bacillus* sp. *B. circulans* (X68326.1), BO: CGTase from *B. ohbensis* (D90243.1), NR5: CGTase from *Bacillus* sp. NR5 UPM (HQ876173.1), amylase: Taka-amylase A from *Aspergillus oryzae* (M33218.1).

**Figure 2.** Comparison of the deduced amino acid sequence of CGTase from *Bacillus* sp. NR5 UPM with other CGTases and amylase. The six highly conserved regions in different CGTases and amylase are boxed. The numbers within brackets are GenBank accession numbers. KC201: CGTase from *Bacillus* sp. KC201 (D13068.1), BC: CGTase from *Bacillus* sp. *B. circulans* (X68326.1), BO: CGTase from *B. ohbensis* (D90243.1), NR5: CGTase from *Bacillus* sp. NR5 UPM (HQ876173.1), amylase: Taka-amylase A from *Aspergillus oryzae* (M33218.1).
Table 2. Translocation of recombinant enzymes into extracellular, periplasmic and intracellular spaces.

<table>
<thead>
<tr>
<th>Parental strain</th>
<th>Cloning host</th>
<th>Expression vector</th>
<th>Mature enzyme (amino acids)</th>
<th>Signal peptide (amino acids)</th>
<th>Translocation of enzyme</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus sp. NR5 UPM</td>
<td>E. coli JM109</td>
<td>pUC19</td>
<td>675</td>
<td>29</td>
<td>Extracellular</td>
<td>This study</td>
</tr>
<tr>
<td>Klebsiella pneumoniae M5a1</td>
<td>E. coli RR28</td>
<td>pHE3</td>
<td>625</td>
<td>30</td>
<td>Extracellular</td>
<td>Binder et al. (1986)</td>
</tr>
<tr>
<td>Bacillus sp. Strain no. 8</td>
<td>E. coli MB 406</td>
<td>pTZ18R and pTZ19R</td>
<td>684</td>
<td>34</td>
<td>Extracellular</td>
<td>Nitschke et al. (1990)</td>
</tr>
<tr>
<td>Bacillus sp. TS1-1</td>
<td>E. coli JM109</td>
<td>pUC19</td>
<td>666</td>
<td>46</td>
<td>Extracellular</td>
<td>Rahman et al. (2006)</td>
</tr>
<tr>
<td>Bacillus sp. Strain No. 38-2</td>
<td>E. coli HB101</td>
<td>pBR322</td>
<td>685</td>
<td>27</td>
<td>Periplasmic</td>
<td>Kaneko et al. (1988)</td>
</tr>
<tr>
<td>Bacillus sp. KC201</td>
<td>E. Coli DH5α</td>
<td>pUC18</td>
<td>674</td>
<td>51</td>
<td>Intracellular</td>
<td>Kitamoto et al. (1992)</td>
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</table>

Table 3. Comparison of growth and CGTase production profiles between wild type (NR5) and transformant (pNR5).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>NR5</th>
<th>pNR5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Growth</td>
<td>30 – 40°C, pH 10</td>
<td>37°C, pH 7</td>
</tr>
<tr>
<td>Optimum CGTase production</td>
<td>37°C, pH 10</td>
<td>37°C, pH 7</td>
</tr>
<tr>
<td>Maximum CGTase activity</td>
<td>11.7 U/ml</td>
<td>29.6 U/ml</td>
</tr>
<tr>
<td>Time of cultivation</td>
<td>48 h</td>
<td>12 h</td>
</tr>
</tbody>
</table>

stability of the secreted enzyme, facilitate the folding processes and enable the production of enzyme in soluble and biologically active form which can finally reduce the subsequent purification cost (Mergulhao et al., 2005). The presence of functional signal peptide presents significant advantages in terms of specific secretion of the enzyme of interest, thus minimizing the contamination of the target enzyme with other non-target proteins.

Besides, the efficient secretion of enzyme into the extracellular medium also eliminates the need for the use of other approaches for protein secretion, such as the use of fusion partners, permeablizing proteins, nutrients or other agents that can create “leakage” and increase the permeability of the outer membrane of *E. coli* (Makrides, 1996). The study by Kim et al. (2005) showed the co-expression of folding accessory proteins facilitated the production of active CGTase of *Bacillus macerans* in recombinant *E. coli*. The soluble expression of the target protein was improved with the use of folding accessory proteins and co-expression with molecular chaperons; otherwise the proteins are mainly expressed as inclusion bodies. The formation of inclusion bodies becomes a significant obstacle in gene expression due to the challenging task in the refolding process of the targeted protein. The supplementation of medium additives also has been verified to lead to the secretion of heterologous protein which has the ability to enhance the permeability of the membrane. Aristidou et al. (1993) reported that the supplementation of glycine gave significant effect to the release of the enzyme which resulted from the extensive cell lysis caused by glycine.

The study on the appropriate time for supplying glycine is also important because the addition of glycine might harm cell growth, thus affecting the overall enzyme production (Li et al., 2009). Together with glycine, the addition of Ca²⁺, SDS and Na⁺ also facilitated the secretion of recombinant enzyme from *E. coli* (Ding et al., 2010).

Enzymatic properties of recombinant CGTase

The recombinant CGTase was characterized and compared with native enzyme from *Bacillus* sp. NR5 UPM with respect to their cyclization activities (Table 3).

The result suggested that the CGTase from transformants required a neutral pH to carry out the cyclization activity, while the wild type required alkaline pH as optimum culture condition for CGTase production. The same finding was
reported by Charoensakdi et al. (2007). The 2.5-fold increment in CGTase activities and one-fourth less in cultivation time proved the transformant CGTase as a successful recombinant enzyme. The recombinant CGTase exhibited a putative molecular weight of 78.6 kDa.

Conclusion

A CGTase gene from Bacillus sp. NR5 UPM was successfully isolated, cloned and expressed into pUC19 cloning vector with E. coli as the host. The deduced amino acid sequence showed 2.1 kb mature CGTase with putative molecular weight of 78.6 kDa and exhibited 98% maximum identity with CGTase sequence from Bacillus sp. KC201. The optimum temperature and pH for recombinant CGTase activity were 37°C and neutral pH, respectively. The CGTase was produced extracellularly (94%) indicating the signal peptide was functional in E. coli. The recombinant CGTase activity was enhanced, approximately 2.5-fold higher than the CGTase from the parent strain and it was beneficial due to it needing less culture time for CGTase production.

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REFERENCES


