# Recombinant production and characterization of full-length and truncated $\beta$-1,3-glucanase PgIA from Paenibacillus sp. S 09 

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#### Abstract

Background: $\beta$-1,3-Glucanases catalyze the hydrolysis of glucan polymers containing $\beta$-1,3-linkages. These enzymes are of great biotechnological, agricultural and industrial interest. The applications of $\beta-1,3-\mathrm{g}$ lucanases is well established in fungal disease biocontrol, yeast extract production and wine extract clarification. Thus, the identification and characterization of novel $\beta-1,3$-glucanases with high catalytic efficiency and stability is of particular interest. Results: A $\beta$-1,3-glucanase gene designated Pg|A was cloned from a newly isolated strain Paenibacillus sp. S09. The gene Pg/A contained a 2631-bp open reading frame encoding a polypeptide of 876 amino acids which shows $76 \%$ identity with the $\beta-1,3-\mathrm{glucanase}(\mathrm{Bg} \mid \mathrm{H})$ from Bacillus circulans IAM1165. The encoded protein PgIA is composed of a signal peptide, an N-terminal leader region, a glycoside hydrolase family 16 (GH16) catalytic domain and a C-terminal immunoglobulin like (Ig-like) domain. The Escherichia coli expression system of Pg|A and five truncated derivatives containing one or two modules was constructed to investigate the role of catalytic and non-catalytic modules. The pH for optimal activity of the enzymes was slightly affected (pH 5.5-6.5) by the presence of different modules. However, the temperature for optimal activity was strongly influenced by the C-terminal domain and ranged from 50 to $60^{\circ} \mathrm{C}$. Deletion of C-terminal domain resulted in obviously enhancing enzymatic thermostability. Specific activity assay indicated that PgIA specifically hydrolyzes $\beta-1,3-$ glucan. Insoluble $\beta-1,3$-glucan binding and hydrolysis were boosted by the presence of N -and C -terminal domains. Kinetic analysis showed that the presence of N -and C -terminus enhances the substrate affinity and catalytic efficiency of the catalytic domain toward laminarin. Carbohydrate-binding assay directly confirmed the binding capabilities of the N -and C -terminal domains. Conclusions: This study provides new insight into the impacts of non-catalytic modules on enzymatic properties of $\beta-1,3-g$ lucanase. Activity comparison of full-length PglA and truncated forms revealed the negative effect of C-terminal region on thermal stability of the enzyme. Both the N -and C -terminal domains exerted strong binding activity toward insoluble $\beta-1,3$-glucan, and could be classified into CBM families.


Keywords: $\beta$-1,3-glucanase, Paenibacillus, Carbohydrate-binding, Ig-like domain, Thermostability

## Background

$\beta-1,3$-Glucanase catalyzes the hydrolysis of $\beta-1,3$-glucosidic bonds existing in $\beta-1,3$-glucan, which is the main cell wall component of yeast and filamentous fungi, and a major structural and storage polysaccharides (laminarin) in marine macroalgae, and produced as exopolysaccharide by some bacteria [1]. $\beta-1,3$-Glucanases are widely distributed among from plants, marine animals, fungi and bacteria.

[^0]According to the divergence of amino acid sequence, $\beta-1,3-$ glucanases from plants are divided into glycosyl hydrolase (GH) family 17 while the bacterial enzymes are grouped into GH16 family [2]. $\beta-1,3$-Glucanases synthesized in plants are highlighted because of the plant defense against fungal pathogens. Additionally, they also play roles in the normal biological and developmental processes in plants and fungi. In bacteria, $\beta-1,3$-glucanases participate in the degradation of polysaccharides which are utilized as an nutrient source $[3,4]$.
There has been a great deal of interest in research of bacterial $\beta$-1,3-glucanases. They have shown potential
biotechnical applications in structural analysis of yeast and fungal cell wall [5], and preparation of protoplasts [6]. $\beta-1,3$-Glucanase could also act as a potential biocontrol agent against pathogenic fungi in agriculture [7], and a useful tool to prevent slime production and undesirable yeast growth during vinification in the wine-making process [8]. In relation to pharmaceutical application, the soluble $\beta$-1,3-glucan oligosaccharides digested by $\beta-1,3$-glucanase are potential immuneactivators [9]. Based on the substrates hydrolyzing mechanisms, $\beta-1,3$-glucanases are classified into two types: exo-1,3-$\beta$-D-glucanases (EC 3.2.1.58) which release glucose residues from the non-reducing end of a substrate, and endo-1,3- $\beta$-D-glucanases (EC 3.2.1.6 and EC 3.2.1.39) which catalyze the hydrolysis of internal $\beta-1,3$-glucosidic bonds at random sites and release short oligosaccharides [10]. There are many $\beta$ - 1,3 -glucanases that have been purified and characterized from different bacterial species, including Bacillus [11], Paenibacillus [12,13], Micromonospora [6], Streptomyces [7] and Cellulosimicrobium [14]. Proposed physiological functions of bacterial $\beta-1,3$-glucanases vary depending on the source and structure of enzyme [15]. In view of the industrial and biotechnological importance of $\beta-1,3$-glucanase, there is of great potential interest in identifying and characterizing novel $\beta-1,3$-glucanases.
In the present study, a novel $\beta$-1,3-glucanase gene was identified by from Paenibacillus sp . S09, which was isolated from the rhizosphere of oat plants. The deduced amino acid sequence analysis and homology search revealed that this protein contained an N-terminal leader region, a GH16 catalytic domain and a C-terminal domain with Ig-like fold. The important role of Ig-like domain has been revealed in many cellulases, but little is known about the biochemical properties of Ig-like domain in bacterial $\beta-1,3$-glucanases. To investigate the role of the N - and C -terminal domains in enzymatic function, the Escherichia coli (E. coli) expression system of PglA and its truncated mutants was constructed in this study. Here, we report the cloning of a new $\beta$-1,3-glucanase gene pglA from S09, the construction and purification of the truncated enzymes, the comparison of the enzymes in terms of their biochemical characteristics and affinities to insoluble $\beta$-1,3-glucans. To the best of our knowledge, this is the first report that describes function activity relationship between catalytic and non-catalytic domains of a bacterial $\beta$-1,3-glucanase which containing a C-terminal Ig-like fold.

## Methods

## Strains, media, vectors and chemicals

The strain Paenibacillus sp. S09 (CCTCC accession no: M2012196; China Center for Type Culture Collection, Wuhan, China) was originally isolated from the rhizosphere of oat plants. Luria-Berntani (LB) broth supplemented with $0.5 \%$ pachyman was used for the production of
$\beta-1,3$-glucanase by strain S09. E. coli competent cell DH5 $\alpha$ (Takara, Otsu, Japan) and the plasmid pCR2.1 vector (Invitrogen, Carlsbad, CA, USA) were used for gene cloning. E. coli BL21 (DE3) and the pET-29a(+) vector (Novagen, San Diego, CA, USA) were used for gene expression. Recombinant E. coli was cultured in LB broth supplemented with kanamycin ( $50 \mu \mathrm{~g} / \mathrm{ml}$ ) at $37^{\circ} \mathrm{C}$. Recombinant E. coli BL21 (DE3) harboring the expression vector was cultivated in $2 \times$ YT medium which containing $1.0 \%(\mathrm{w} / \mathrm{v})$ yeast extract, $1.6 \%(\mathrm{w} / \mathrm{v})$ peptone and $0.5 \%(\mathrm{w} / \mathrm{v})$ NaCl . The $\mathrm{His}_{6}$-tagged protein was purified by Ni -sepharose 6 fast flow column (GE healthcare, Sweden). The DNA purification kit, restriction endonucleases, T4 DNA ligase, $p f u$ and LA Taq DNA polymerase with GC buffer and dNTPs were purchased from Takara. Isopropyl- $\beta$-Dthiogalactopyranoside (IPTG) and 5-bromo-4-chloro-3-indolyl $\beta$-D-glucuronide (X-Gal) were also purchased from Takara.
Laminarin (from Laminarium digitatum), barley $\beta$-glucan, zymosan A were purchased from Sigma-Aldrich (Co., St. Louis, USA). Pachyman was purified from commercial fruiting bodies of the basidiomycete Poria cocos. Carboxymethyl cellulose-sodium (CMC) and dextran were also purchased from Sigma. Salecan, a novel soluble linear $\beta$-1,3-glucan with $\alpha-1,3$-linkages, were prepared from Agrobacterium sp. ZX09, as described by Xiu et al. [16]. All other reagents were of analytical grade.

## Molecular cloning of the $\beta-1,3$-glucanase encoding gene

The genomic DNA of strain S09 was extracted as described by Sambrook and Russell [17]. In order to obtain partial of the $\beta-1,3$-glucanase gene sequence, degenerate primers ( $\mathrm{Pdg}-\mathrm{F}$ and Pdg-R) was designed based on the two conserved amino acid sequences of GH16 $\beta-1,3$-glucanases (Table 1, see Additional file 1: Figure S1). the PCR was performed to amplify a partial $\beta-1,3$-glucanase gene as follows: $94^{\circ} \mathrm{C}$ for $5 \mathrm{~min} ; 30$ cycles of $94^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 55^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 72^{\circ} \mathrm{C}$ for 1 min ; one final extension at $72^{\circ} \mathrm{C}$ for 10 min. The genomic DNA of strain S 09 was used as the template. The products of PCR reaction were gel-purified, ligated into pCR2.1 vector, transformed into DH5 $\alpha$ and sequenced by Invitrogen. Inverse PCR (I-PCR) and SelfFormed Adaptor PCR (SEFA-PCR) were employed to obtain the full sequence of $\beta-1,3$-glucanase gene according to the manufacturer's instructions [18]. The primers used in I-PCR and SEFA-PCR were listed in Table 1. The primers "gIF1, gIF2, gIR1, gIR2" constructed according the partial $\beta-1,3$-glucanase gene were used in I-PCR. Briefly, genomic DNA was digested with Hind $W$ for 3 h . The digested genomic DNA fragments were gel-purified and subsequently circularized by use of the DNA ligation kit (Takara). I-PCR amplification was conducted with the circulated DNA as template. The primers $5^{\prime}$-sp1, $5^{\prime}$-sp2, and $5^{\prime}$-hemi-sp3 were used to amplify the 5 ' flanking

Table 1 Primers used for the cloning of $\beta$-1,3-glucanase gene

| Primer name | Primer sequence $\left(\mathbf{5}^{\prime} \rightarrow \mathbf{3} \mathbf{3}^{\prime} \mathbf{a}^{\mathbf{a}}\right.$ |
| :--- | :--- |
| Pdg-F | TAY ACS CGS ACR TAR TC |
| Pdg-R | GAY TAY GTS CGS GTR TA |
| gIF1 | GTAGCGGTTGACCGGCCATTGG |
| gIF2 | GCTGCTGTTCGCCTTGGAGT |
| gIR1 | CGACCAGCCGTTCTATCTGATCAT |
| gIR2 | GGCGACGATGCAGGTAGACTATGT |
| 5'-sp1 | CGTTTCATTGACATTCTGGAAGACGAGCGT |
| $5^{\prime}-$ sp2 | TGTAGTTGAGTGTGACGCTGCTGTTCGC |
| 5'-hemi-sp3 | AATGTCGACCCAGCTNNNNNNNNNCTTGAC |
| 3'-sp1 | AGAGATCGACATCATGGAGGCGAAGGG |
| 3'-sp2 | TACCTCCCGCAAGGTACGACATTCG |
| 3'-hemi-sp3 | ACCAGCCGTTCTATCNNNNNNNNAACCTG |

${ }^{\text {a }}$ Restriction sites in the primers are italicized. $Y=C / T, S=C / G, R=A / G$.
sequence, $3^{\prime}$-sp1, $3^{\prime}$-sp2, and $3^{\prime}$-hemi-sp3 were used to amplify the $3^{\prime}$ flanking sequence of the partial $\beta-1,3-$ glucanase gene. The detailed procedures for SEFA-PCR were shown in Additional file 2. The amplified PCR products were purified, cloned into the pCR2.1 plasmid, and sequenced.

## Sequence analysis

Nucleotide sequences were assembled using the DNAMAN software package (Version 5.2.2, Lynnon BioSoft, USA). The signal peptide was predicted using SignalP 4.0 server (http://www.cbs.dtu.dk/services/SignalP) [19]. The alignments of DNA and protein sequences were performed at National Center for Biotechnology Information (NCBI) using the blastn and blastp program (http://www.ncbi.nlm. nih.gov/BLAST/), respectively. The conserved domains and classification of GH family were identified with the website http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml [20]. Protein secondary structure was predicted using PSIPRED 3.0 [21] (http://bioinf.cs.ucl.ac.uk/psipred). Structures of endo-1,3- $\beta$-glucanase from Cellulosimicrobium cellulans (PDB ID: 3atgA) and the C-terminal Ig-like domain of the bacteriophage $\lambda$ tail tube protein (PDB ID: 2104A) were used as templates for protein structure homology modeling using the automated mode of SWISS-MODEL [22]. Multiple sequence alignments were performed with ClustalX2 [23]. Distance matrix for nucleotides was calculated according to the Kimura's two-parameter model [24]. Phylogenetic tree, assessed by 1,000 bootstraps, was constructed using neighbor-joining algorithms with MEGA 5.0 software [25].

## Construction of expression vectors

Five truncated mutants containing one or two modules were designed according to the conserved domain analysis (Figure 1). The corresponding coding regions were amplified using respective pairs of primers by PCR (Table 2).

The amplification reactions performed with pfu DNA polymerase consisted of 30 cycles of $94^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 60^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 72^{\circ} \mathrm{C}$ for 1-3 min, followed by a 10 min extension at $72^{\circ} \mathrm{C}$. The PCR-amplified fragments were purified and digested with KpиI and HindШ and then ligated into the expression vector pET-29a(+) with a C-terminal His ${ }_{6}$-tag. After transformation into E. coli $\mathrm{DH} 5 \alpha$, positive transformants were screened and checked by DNA sequencing. The recombinant plasmids were used to transform E. coli BL21 (DE3) competent cells for protein expression.

## Gene expression and protein purification

For recombinant protein production, recombinant E.coli BL21 cells harboring the expression vectors were first inoculated into LB medium supplemented with kanamycin $(50 \mu \mathrm{~g} / \mathrm{ml})$ and grown overnight at $37^{\circ} \mathrm{C}$. Then, the culture was transferred into fresh $2 \times$ YT medium (5:100 dilution) containing $50 \mu \mathrm{~g} / \mathrm{ml}$ kanamycin and grown at $37^{\circ} \mathrm{C}$ to an $\mathrm{OD}_{600} \approx 0.8$, induced with 0.1 mM IPTG and further cultivated for $6-8 \mathrm{~h}$ at $25^{\circ} \mathrm{C}$. The culture medium was centrifuged at $12,000 \times \mathrm{g}$ for 10 min at $4^{\circ} \mathrm{C}$. Cells were washed and suspended in PBS buffer, lysed by sonication on ice and centrifuged at $15,000 \times \mathrm{g}$ for 20 min at $4^{\circ} \mathrm{C}$ to remove the cell debris. The supernatant was applied to Nisepharose 6 fast flow column (GE healthcare, Sweden) that had been equilibrated with binding buffer ( 20 mM sodium phosphate, $500 \mathrm{mM} \mathrm{NaCl}, \mathrm{pH} 7.2$ ). The protein was eluted using a linear gradient of elution buffer ( 20 mM sodium phosphate, $500 \mathrm{mM} \mathrm{NaCl}, 15-300 \mathrm{mM}$ imidazole, pH 7.2 ). Fractions having the $\beta-1,3$-glucanase activity were pooled and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE).

## Enzyme activity assay and protein determination

The $\beta$-1,3-glucanase activity assay was carried out by measuring the amount of reducing sugars released from a $\beta-1,3$-glucan substrate (laminarin) according to the 3,5-dinitrosalicylic acid (DNS) method [26]. Unless otherwise stated, the reaction mixture was incubated at $55^{\circ} \mathrm{C}$ for 20 min using laminarin ( $5 \mathrm{mg} / \mathrm{ml}$ ) as the substrate in 20 mM sodium acetate buffer ( pH 6.0 ). The amount of reducing sugar was then determined spectrophotometrically at 550 nm . One unit (U) of enzyme activity was defined as the amount of enzyme required to release reducing sugars that equivalent to $1 \mu \mathrm{~mol}$ glucose per min under the test conditions. Hydrolytic activities toward insoluble substrates were performed by gently mixing reaction mixtures during incubation. Then the mixture was centrifuged at $12,000 \times \mathrm{g}$ at $4^{\circ} \mathrm{C}$ for 10 min . The reducing sugars in the supernatant were measured by the DNS method. The protein concentration was measured by the bicinchoninic acid (BCA) method using the BCA protein assay reagent (Thermo Fisher Scientific, MA, USA), Bovine serum albumin (BSA) was used as the standard protein.


Figure 1 Schematic overview of PgIA and its truncated derivatives. A, Schematic representation of the module composition of truncated proteins. Abbreviations: sp: singal peptide; $\Delta X$ : deletion derivative, with $X$ referring to deleted domain; $N$ : $N$-terminal leader region ( N -domain); CD: GH16 catalytic domain of PgIA (GH16); C, C-terminal region containing the Ig-like domain. Every truncated protein has a histidine tag fused to the C-terminus. The scale at the top of the figure indicates amino acid residues. B, SDS-PAGE analysis of the purified derivatives of PgIA. SDS-PAGE was done using a $12 \%$ polyacrylamide gel. Proteins were stained with Coomassie Brilliant Blue R250. Lane M: protein molecular weight marker; Lane 1: rPglA; Lane 2: rPglA $\Delta \mathrm{C}$; Lane 3: rPglA $\Delta \mathrm{N}$; Lane 4: rPglA-CD; Lane 5: rPglA-N; Lane 6: rPglA-C.

## Enzyme characterization

With laminarin as the substrate, the optimal temperature for $\beta-1,3$-glucanase activity was determined over the range of $30-80^{\circ} \mathrm{C}$ in sodium acetate buffer (pH 6.0). Enzyme thermostability was determined by measuring the residual activity after pre-incubation of the enzyme in sodium acetate buffer ( pH 6.0 ) at $55^{\circ} \mathrm{C}$ without substrate for up to 60 min . The optimal pH for $\beta-1,3$-glucanase activity was determined at $55^{\circ} \mathrm{C}$ in buffers with pH ranging from 2.0 to 10.0. The enzyme stability at different pH was estimated by measuring the residual enzyme activity after incubating the enzyme solution in buffers with pH of $2.0-12.0$ at $4^{\circ} \mathrm{C}$ for 4 h . The buffers ( 50 mM ) used were glycine- HCl ( pH 2.0-4.0), sodium acetate ( $\mathrm{pH} 5.0-7.0$ ), Tris- HCl ( $\mathrm{pH} 7.5-8.5$ ) and glycine- NaOH ( $\mathrm{pH} 9.0-12.0$ ).

Table 2 Primers used for the construction of the expression plasmid

| Construct | Primer sequence ( $\left.5^{\prime} \rightarrow 3^{\prime}\right)^{\text {a }}$ |
| :---: | :---: |
| PgIA | F1 CGGGGTACCGCGACGACCGTGACGTCGAT |
|  | R3 CCCAAGCTITTACTCAACAACACCTTGTGTTACC |
| PgIA $\triangle$ C | F1 CGGGGTACCGCGACGACCGTGACGTCGAT |
|  | R2 CCCAAGCTITAGCCCGCGCCCTCTITGTAC |
| PgIA $\triangle$ N | F2 CGGGGTACCTGGAATCTGTATTGGCAGGATG |
|  | R3 CCCAAGCTITTACTCAACAACACCTTGTGTTACC |
| PgIA-CD | F2 CGGGGTACCTGGAATCTGTATTGGCAGGATG |
|  | R2 CCCAAGCTITAGCCCGCGCCCTCTITGTAC |
| PgIA-N | F1 CGGGGTACCGCGACGACCGTGACGTCGAT |
|  | R1 CCCAAGCTITTATCCTGCGATGGCCGGATCG |
| PgIA-C | F3 CGGGGTACCAACGTCGCGGTGACCGGC |
|  | R3 CCCAAGCTITTACTCAACAACACCTTGTGTTACC |

[^1]The substrate specificity was assayed in reactions with $1 \%$ laminarin, pachyman, zymosan A , barley $\beta$-glucan, CMC, dextran and $0.5 \%$ salecan under the optimal conditions for each enzyme. The reaction mixture was often oscillated slightly during incubation for insoluble polysaccharides.
Kinetic study was performed with laminarin as the substrate under the optimal conditions for each enzyme. The initial velocities under laminarin concentrations ( $0.5-5 \mathrm{mg} / \mathrm{ml}$ ) were measured, and the values of $K_{\mathrm{m}}$ and $V_{\max }$ were obtained from Lineweaver-Burk plots.

## Binding assay

Binding strength of the enzymes to insoluble polysaccharides was estimated by mixing $15-20 \mu \mathrm{~g}$ of the purified proteins with various substrates ( 10 mg ) in sodium acetate buffer (200 $\mu \mathrm{l})$ at $4^{\circ} \mathrm{C}$ for 1 h with gentle shaking. Binding mixture without substrate was set as control. The binding ability of enzymes containing catalytic domain to insoluble polysaccharides was determined by measuring the residual enzyme activity in the supernatant of the binding mixture. The amount of the enzyme bound to the polysaccharide was estimated by subtraction of the activity recovered from the original activity [27]. The binding ability of non-catalytic domain to insoluble substrates was also determined [28]. After 1 h of binding reaction, the fraction of protein in the supernatant and the pellet was estimated by $12 \%$ SDS-PAGE analysis. BSA was used as the negative control and was handled similarly to the other samples.

## Accession number

The nucleotide sequence for the GH16 $\beta$-1,3-glucanase gene pglA were deposited in GenBank database under the accession number JX070086.

## Results

## Cloning of the $\beta-1,3$-glucanase gene and sequence analysis

 A $\beta$-1,3-glucanase-encoding gene designated $\operatorname{PglA}$ was identified from a 5053-bp fragment which was obtained by using degenerate PCR, I-PCR and SEFA-PCR. The deduced amino acid sequence of PglA (GenBank No.: AFO67889.1) comprises 876 amino acids with a predicted N-terminal signal peptide (residues 1-38) (Figure 2). The mature protein consists of 838 amino acid residues with a calculated pI of 4.5 and molecular mass of 90.4 kDa , respectively. The deduced amino acid sequence of PglA exhibited the highest identity (86\%) with the hypothetical protein from Paenibacillus daejeonensis (NCBI Reference Sequence: WP_020617344.1), 77\% identity with the glycoside hydrolase family 16 protein from Paenibacillus lactis 154 (GenBank No.: EHB65983.1), and 76\% identity with the reported $\beta-1,3$-glucanase BglH from Bacillus circulans IAM1165 (GenBank No.: AAC60453.1). PglA showed only 45\% identity with the well-characterized endo- $\beta$-1,3-glucanase LamA from Paenibacillus sp. CCRC 17245 (GenBank No.: ABJ15796.1). Combing the results of conserved domain search in the Conserved Domain Database (CDD) of NCBI, secondary structure prediction and tertiary structure homology modeling by SWISS-MODEL (See Additional file 3: Figure S2), PglA contains a GH16 laminarinase-like domain and a bacterial Ig-like domain (Figures 2 and 3A). The domain organization of PglA is not similar to LamA,which include a leader peptide, a threefold repeat of S-layer homologous module, a GH16 catalytic module, four repeats of CBM_4_9 and an analogue of coagulation factor Fa5/8C from N to C terminus [13]. PglA could be considered as a novel $\beta-1,3$-glucanase of Paenibacillus species. Phylogenetic tree showed that the catalytic region of PglA forms a distinct clade with BglH from B. circulans and putative glycoside hydrolases from Paenibacillus species (Figure 3B). Phylogenetic analysis based on the Ig-like domain sequence alignment indicated the relationships between $\operatorname{PglA}$ and other Ig-like domain containing proteins (Figure 3C).

## Expression of full-length and truncated PgIA in E.coli and protein purification

Using basic molecular cloning techniques, the gene fragment of $P g l A$ without the signal peptide sequence and five truncated mutants were cloned into pET-29a(+) vector and successfully expressed in E.coli BL21 (DE3). Figure 1A represents a schematic overview of the recombinantly produced constructs. The $\mathrm{His}_{6}$-tagged enzymes were purified by affinity chromatography on a Ni-sepharose 6 fast flow column. Molecular masses of rPglA and its truncated mutants rPglA $\Delta \mathrm{C}$, rPglA $\Delta \mathrm{N}$, rPglA-CD, rPglA- N and $\mathrm{rPglA}-\mathrm{C}$ were estimated to be $91 \mathrm{kDa}, 70 \mathrm{kDa}, 49 \mathrm{kDa}, 29 \mathrm{kDa}$, 42 kDa and 21 kDa , respectively. SDS-PAGE analysis indicated protein bands with molecular masses corresponding well to the theoretical values. All the enzymes were purified

[^2]Figure 2 The nucleotide and amino acid sequences of PgIA from Paenibacillus sp. S09. A vertical arrow indicates a cleavage site to produce the mature enzyme. The three N-terminal repeats are indicated by arrow line. The catalytic domain and lg-like domain are shadowed by blue and red, respectively. The putative catalytic amino acids are circled. The stop codon is indicated by asterisk.


Figure 3 Sequence analysis of PgIA. A, Schematic representation of PgIA with the functional domains. The numbers indicated amino acid positions. The GH16 catalytic domain was indicated in black, signal peptide was indicated in gray, the bacterial Ig-like domain was shown by a stippled box. B and C, Phylogenetic analysis of the GH16 and Ig-like domains from different proteins, respectively. The amino acid sequence of the GH16 or Ig-like domain was used in BLASTP search and matched sequences were extracted and aligned by ClustalX2. The starting and ending amino acid positions are shown. The unrooted dendrogram was constructed by neighbor-joining method. The percentages of bootstrap values (based on 1000 bootstraps) are shown at the internal nodes. GenBank accession numbers: P. lactis 154, EHB65983.1; B. circulans BgIH, AAC60453.1; P. vortex, EFU40835.1; C. butyricum DKU-01, EMU55784.1; S. frigidimarina NCIMB 400, ABI71172.1; Sphingomonas sp. SKA58, EAT10234.1, M. maris MCS10, ABI64540.1; Enterobacter sp. Ag1, EJF31699.1; Pantoea sp. At-9b, ADU69429.1; F. johnsoniae UW101, ABQ04240.2; E.coli, EMX44285.1; Clostridium sp. CAG:1024, CCX42470.1.
to a homogeneous state on SDS-PAGE (Figure 1B). The steps involved in a typical purification process are summarized in Table 3.

## Effects of temperature and pH on the enzymatic activity and stability

The general properties including the optimal pH and temperature of PglA and the truncated derivatives were investigated with laminarin as the substrate. Deletion of the N -or C-terminal domain did not abolish the hydrolytic function of $\beta-1,3$-glucanase, thus, enzymatic properties of $r \operatorname{PglA}, \operatorname{rPglA} \Delta C, r \operatorname{PglA} \Delta \mathrm{~N}$ and rPglA-CD were characterized respectively.
Effect of domains on thermoactivity and thermostability of rPglA was investigated. Side by side comparison of the temperature activity profiles of rPglA and the truncated mutants $(\mathrm{rPglA} \Delta \mathrm{C}, \mathrm{rPglA} \Delta \mathrm{N}$ and $\mathrm{rPglA}-\mathrm{CD})$ showed that, $\operatorname{rPglA} \Delta \mathrm{N}$, the PglA variant containing the catalytic domain and the C-terminal Ig-like domain, obviously is the least thermoactive enzyme. It had its highest activity at $50^{\circ} \mathrm{C}$, while, $\mathrm{rPglA}, \mathrm{rPglA} \Delta \mathrm{C}$ and $\mathrm{rPglA}-\mathrm{CD}$ have a maximum activity at $60^{\circ} \mathrm{C}$ (Figure 4A). This phenomenon indicated that the deletion of the C-terminus increased the optimal temperature. As a dramatic loss of enzymatic stability occurred in the full length enzyme (rPglA) at temperatures higher than $60^{\circ} \mathrm{C}$, the thermal stability was assayed at $55^{\circ} \mathrm{C}$
for up to 60 min . In conjunction with the temperature optima, also the thermal stability of the truncated derivatives of PglA turned out to be module related. As observed by the thermal stability assay, the absence of C-terminal region is of great relevance for higher temperature stability of the proteins. This can be deduced from the unexpected higher stability of rPglA-CD and rPglA $\Delta \mathrm{C}$, and lower stability of rPglA and rPglA $\Delta \mathrm{N}$ (Figure 4B). These results indicated that truncation of C-terminal domain improved the thermostability of PglA.
Deletion of the N -or C-terminal regions had no drastic effect on the optimal pH. For all PglA derived constructs, an apparent pH optimum between pH 5.5 and 6.5 was observed (Figure 4C). rPglA and the truncated enzymes showed broad pH stability. All enzymes retained more than $80 \%$ of their original enzymatic activity after 4 h incubation in buffers with pH values between pH 5.0 and 10.0 (Figure 4D). It should be noted that the C-terminal domain deleted mutant $\mathrm{rPglA} \Delta \mathrm{C}$ retained higher activity in the acidic range of the pH spectrum ( $\mathrm{pH} 2.0-4.5$ ) than the other deletion derivatives.

## Substrate specificity and kinetic analysis

The purified rPglA and truncated enzymes ( $\operatorname{PglA} \Delta C$, $\operatorname{PglA} \Delta \mathrm{N}$ and $\operatorname{PglA}-\mathrm{CD})$ shared similar substrate specificity. They were highly active toward soluble $\beta-1,3$-glucan

Table 3 Purification of rPgIA and its truncated derivatives

| Purification step | Total activity (U) | Total protein (mg) | Specific activity ( $\mathrm{U} / \mathrm{mg}$ ) | Yield (\%) | Purification (fold) |
| :---: | :---: | :---: | :---: | :---: | :---: |
| rPgIA |  |  |  |  |  |
| Crude extract | 2.56 | 9.80 | 0.26 | 100 | 1 |
| $\mathrm{Ni} 2+$-affinity column | 1.43 | 0.72 | 1.99 | 58.1 | 7.62 |
| rPgIADC |  |  |  |  |  |
| Crude extract | 4.01 | 12.4 | 0.32 | 100 | 1 |
| $\mathrm{Ni} 2+$-affinity column | 2.21 | 1.05 | 2.14 | 55.1 | 6.63 |
| rPgIA $\mathrm{S}^{\text {N }}$ |  |  |  |  |  |
| Crude extract | 2.85 | 9.71 | 0.29 | 100 | 1 |
| Ni2 + -affinity column | 1.53 | 0.75 | 2.04 | 53.7 | 6.96 |
| rPgIA-CD |  |  |  |  |  |
| Crude extract | 2.51 | 8.81 | 0.29 | 100 | 1 |
| Ni2 + -affinity column | 1.05 | 0.60 | 1.76 | 42.1 | 6.17 |

(laminarin) and insoluble $\beta-1,3$-glucan (pachyman and zymosan A). Low activity was detected on soluble linear glucan, barley $\beta$-glucan which contains mixed $\beta-1,3$-and $\beta-1,4$ - linkages, and salecan which contains mixed $\beta$ 1,3 -and $\alpha$-1,3-linkages, little activity on CMC, dextran and xylan (Table 4). It was not surprising as $\beta-1,3$-glucan
specificity is the general rule for laminarinases. Side by side comparison of the specific activity of rPglA and the truncated mutants (rPglA $\Delta \mathrm{C}, \mathrm{rPglA} \Delta \mathrm{N}$ and $\mathrm{rPglA}-\mathrm{CD}$ ) showed that, both the N -and C -terminal domains could enhance the hydrolyzing activity of the catalytic module toward insoluble substrates. It was worth noting that


Figure 4 Enzymatic characteristics of purified rPgIA and the truncated enzymes. A, Influence of temperature on $\beta-1,3$-glucanase activity. Reactions were carried out at pH 6.0 and temperatures ranging from 35 to $80^{\circ} \mathrm{C}$ for 20 min . Values were expressed as percentages of maximal activity. B, Thermal stability of rPgIA and the truncated derivatives. The enzymes were incubated at $55^{\circ} \mathrm{C}$ for up to 60 min and the residue enzyme
 optimal temperatures for each enzyme for 20 min in buffers of varying $\mathrm{pH}(2.0-10.0)$. D, pH stability of rPglA and the truncated derivatives. Purified enzymes were first incubated in buffers of varying $\mathrm{pH}(2.0-12.0)$ at $4^{\circ} \mathrm{C}$ for 4 h , and activity was measured under optimal conditions for 20 min . The buffers ( 50 mM ) used were: glycine- $\mathrm{HCl}(\mathrm{pH} 2.0-4.0$ ), sodium acetate ( $\mathrm{pH} 5.0-7.0$ ), Tris- $\mathrm{HCl}(\mathrm{pH} 7.5-8.5$ ) and glycine- NaOH ( $\mathrm{pH} 9.0-12.0$ ). The measurements were repeated in triplicate.

Table 4 Substrate specificity of rPgIA and its truncated derivatives

| Substrate ${ }^{\text {a }}$ | Solubility | Main linkage | Specific activity (U/mg) ${ }^{\text {b }}$ of: |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | rPgIA | rPgIA $\Delta C$ | rPgIA $\Delta N$ | rPgIA-CD |
| Laminarin | Soluble | $\beta-1,3, \beta-1,6$ (Glc) | $2.016 \pm 0.213$ | $2.506 \pm 0.131$ | $2.000 \pm 0.099$ | $2.525 \pm 0.141$ |
| Pachyman | Insoluble | $\beta-1,3$ (Glc) | $1.695 \pm 0.165$ | $2.396 \pm 0.339$ | $1.883 \pm 0.045$ | $0.898 \pm 0.078$ |
| Zymosan A | Insoluble | $\beta-1,3, \beta-1,6$ (Glc) | $1.364 \pm 0.131$ | $1.518 \pm 0.036$ | $0.835 \pm 0.022$ | $0.567 \pm 0.035$ |
| Barley $\beta$-glucan | Soluble | $\beta-1,3, \beta-1,4$ (Glc) | $0.086 \pm 0.011$ | $0.074 \pm 0.001$ | $0.076 \pm 0.021$ | $0.081 \pm 0.015$ |
| Salecan | Soluble | $\beta-1,3, a-1,3$ (Glc) | $0.049 \pm 0.010$ | $0.039 \pm 0.005$ | $0.037 \pm 0.009$ | $0.032 \pm 0.007$ |
| CMC | Soluble | $\beta-1,4$ (Glc) | 0 | 0 | 0 | 0 |
| Dextran | Soluble | a-1,6 (Glc) | 0 | 0 | 0 | 0 |
| xylan | Insoluble | $\beta-1,4$ (xyl) | 0 | 0 | 0 | 0 |

${ }^{\text {a }}$ All substrates were used at a final concentration of $10 \mathrm{mg} / \mathrm{ml}$, except salecan ( $5 \mathrm{mg} / \mathrm{ml}$ ).
${ }^{\mathrm{b}}$ Activity $\pm$ standard error of triplicate samples.
$\mathrm{rPglA} \Delta \mathrm{C}$ and $\mathrm{rPglA}-\mathrm{CD}$ show slightly higher activity toward laminarin than rPglA and $\mathrm{rPglA} \Delta \mathrm{N}$. It should be related to the improved thermostability of $\operatorname{rPglA} \Delta \mathrm{C}$ and rPglA-CD as the deletion of the C-terminal region.
The kinetic parameters with laminarin as substrate were determined from substrate concentration dependence of velocity. Based on double reciprocal plots, the values for the Michaelis constant $\left(K_{\mathrm{m}}\right)$, turnover number $\left(k_{\text {cat }}\right)$ and catalytic efficiency ( $k_{\mathrm{cat}} / K_{\mathrm{m}}$ ) are calculated and presented in Table 5. $K_{\mathrm{m}}$ values of rPglA, rPglA $\Delta \mathrm{C}, \mathrm{rPglA} \Delta \mathrm{N}$ and rPglA-CD were calculated to be $0.94,1.01,1.36$ and $2.64 \mathrm{mg} / \mathrm{ml}$, respectively, suggesting that deleting N -and C-terminal domains caused significantly increased $K_{\mathrm{m}}$ values. The $k_{\text {cat }} / K_{\mathrm{m}}$ values of $\operatorname{rPglA} \Delta \mathrm{C}, \operatorname{rPglA} \Delta \mathrm{N}$ and rPglA-CD were about $70 \%, 78 \%$ and $35 \%$ of that of rPglA respectively. These kinetic parameters indicate that the N -and C-terminal domains increase the affinity of PglA for the substrate (laminarin) and thus enhance the catalytic efficiency of catalytic module.

## Binding assay

Binding experiments with rPglA and the truncated derivatives and BSA as the tested protein were performed. Two approaches were taken to explore the polysaccharidebinding function of the N -terminal and C -terminal regions. First, as for rPglA and the truncated enzymes containing the catalytic domain (rPglA $\Delta \mathrm{C}$, rPglA $\Delta \mathrm{N}$, rPglA-CD), residual enzymatic activity in the supernatant of binding mixtures was measured. As shown in Figure 5A, the catalytic domain

Table 5 Kinetic parameters ${ }^{\text {a }}$ of rPgIA and its truncated derivatives toward laminarin

| Enzyme | $K_{\mathrm{m}}(\mathrm{mg} / \mathrm{ml})$ | $k_{\text {cat }}\left(s^{-1}\right)$ | $k_{\text {cat }} / K_{\mathrm{m}}\left(\mathrm{s}^{-1} \cdot \mathrm{mg}^{-1} \cdot \mathrm{ml}\right)$ |
| :---: | :---: | :---: | :---: |
| rPglA | $0.94 \pm 0.07$ | $3.56 \pm 0.18$ | $3.79 \pm 0.31$ |
| rPgIA $\triangle$ C | $1.01 \pm 0.05$ | $2.68 \pm 0.07$ | $2.66 \pm 0.09$ |
| rPglA $\triangle$ N | $1.36 \pm 0.11$ | $4.01 \pm 0.10$ | $2.95 \pm 0.07$ |
| rPgIA-CD | $2.64 \pm 0.15$ | $3.54 \pm 0.21$ | $1.34 \pm 0.07$ |

[^3]of $\operatorname{PglA}(r \operatorname{PglA}-C D)$ had the minimum binding ability compared to the other three enzymes. Both the N -and C-terminal regions of PglA had significantly carbohydratebinding activities. These results were confirmed by the insoluble carbohydrate-binding analysis of $\mathrm{rPglA}-\mathrm{N}$ and rPglA-C. The N -and C -terminal domain bound strongly to insoluble $\beta-1,3$-glucan (pachyman and zymosan $A$ ), as a large fraction of the protein was retained in the pellet fraction, but not to xylan (Figure 5B). Presence of insoluble polysaccharide did not affect the amount of BSA in supernatant, excluding the nonspecific binding of insoluble polysaccharide to protein.

## Discussion

Enzymes that hydrolyze insoluble complex polysaccharide structures are usually multi-modular enzymes and composed of distinct catalytic domain and non-catalytic carbohydrate binding modules (CBMs) that play a pivotal role in the action of these enzymes against recalcitrant substrates [29]. The presently known CBMs are classified into 67 families in the latest update of the CAZy database (http://www.cazy.org/fam/acc_CBM.html). Among the 67 CBMs families classified in CAZy, one or more of CBMs of $4,6,9,13,54,56$ families have been found in bacterial $\beta-1,3$-glucanases [13,27,29-31]. In the work reported here, a novel $\beta-1,3$-glucanase gene PglA was cloned from Paenibacillus sp.S09. PglA contains three functional regions: an N-terminal domain, a GH16 catalytic domain, and a C-terminal domain with an Ig-like structure. Even high sequence similarity was found between PglA and the $\beta-1,3$-glucanase BglH from Bacillus circulans IAM1165, PglA possessed a novel Ig-like domain which was not observed in BglH by conserved domain search in NCBI. Previous reports demonstrated carbohydrate-binding properties of the N -and C-terminal regions of BglH [27,32], while, the effects of the non-catalytic domain on enzymatic properties of BglH were not described. Furthermore, the enzymatic properties of Ig-like domain, which is generally considered being involved in carbohydrate binding and


Figure 5 Binding assay of the truncated enzymes toward insoluble polysaccharides. The purified protein (15-20 $\mu \mathrm{g}$ ) and the indicated substrate ( 10 mg ) were thoroughly mixed at $4^{\circ} \mathrm{C}$ for 1 h . A, Binding assay of the four active enzymes (PglA, rPglA $\triangle \mathrm{C}, \mathrm{rPg} \mid \mathrm{A} \Delta \mathrm{N}$, rPglA-CD). The enzymatic activity against laminarin remaining in the supernatant was determined and compared to the original activity of each enzyme preparation. The binding activity was calculated by subtraction of the activity recovered from the original activity which was considered as 100\%. The values represent the average of the results from triplicate experiments. B, Binding of rPglA-N and rPglA-C to insoluble carbohydrates. The amount of protein remaining in the supernatant $(S)$ and co-precipitating with the substrate (P) were examined by SDS-PAGE.
commonly found in bacterial proteins [33], have not been illustrated in bacterial $\beta$-1,3-glucanases. We were interested to investigate the functional structural elements of PglA.
It is generally believed that each functional module in a polypeptide can function normally even after separation from an adjacent module [34]. The full-length PglA and five truncated derivatives containing one or two modules were constructed, recombinantly produced, purified and characterized. The function of each module referring to carbohydrate-binding and hydrolyzing properties was identified respectively. Specific activities of PglA and truncated derivatives on various substrates were measured. The strict and selective specificity of PglA suggested that it was a strict $\beta-1,3$-glucanase belonging to EC 3.2.1.39 family of glycoside hydrolases. The truncated mutant PglA-CD, which contained only the catalytic domain, showed reduced hydrolytic activity on insoluble substrates, while, its activity remained unchanged on
soluble substrates (Table 4). It could be suggested that the N - or C-terminal domain did not play a direct or critical role in the structural stability of the catalytic domain. The results of binding assay demonstrated that both the N -and C-terminal domains of PglA had significant binding activities toward insoluble linear $\beta-1,3$-glucan (pachyman) and branched-chain $\beta$-1,3-glucan (zymosan A) respectively (Figure 5), suggesting their module functions in the enzymatic activity of PglA. It was in accordance with the $\beta-1,3$-glucanase (BglH) from Bacillus circulans IAM1165 [11]. Amino acid sequence alignment showed no significant similarity between the N -and C -terminal domains. Thus, both of them could be classified into CBM families respectively. Either N-or C-terminal CBM could exert the strong binding toward insoluble $\beta-1,3$-glucan (pachyman and zymosan A), which probably account for the increase of the specific activities of $\mathrm{rPglA} \Delta \mathrm{C}$ or $\mathrm{rPglA} \Delta \mathrm{N}$ against these insoluble substrates, presumably by assisting the localization of catalytic domains to insoluble polysaccharides.
The full-length $\operatorname{PglA}$ and truncated glucanases $(\operatorname{PglA} \Delta C$, $\operatorname{PglA} \Delta \mathrm{N}$ and PglA-CD) exhibited similar enzymatic properties. The pH activity and stability profiles of rPglA and the truncated enzymes were nearly indistinguishable, except that rPglA $\Delta \mathrm{C}$ retained higher activity in acidic range of the pH spectrum ( $\mathrm{pH} 2.0-4.5$ ) than the other deletion derivatives. The results of thermal activity and stability assay indicated the relatively low influence of the N-terminal CBM on enzymatic properties of PglA. However, deletion of the C-terminus in PglA , consisting of an Ig-like fold and a basic terminal domain, significantly increased the thermal tolerance of the enzyme, which was first observed with bacterial $\beta-1,3$-glucanase. These observations imply the negative impact of the C-terminal CBM on enzymatic acidic and thermal stabilities of catalytic domain. A similar negative effect has been identified for endo- $\beta$-1,4-glucanase from Bacillus subtilis JA18 in that truncation of the cellulose binding domain improved thermal stability of the enzyme [35]. Carbohydrate binding is not the sole function of the non-catalytic modules [36]. The impact of CBMs on enzymatic thermal stability of a multidomain xylanase has been discovered by Winterhalter et al. in 1995 [37]. Increasing literature showed controversial data on CBMs affecting glycoside hydrolase's thermostability. The comparison of catalytic module alone with catalytic module fused to CBM showed either improvement of thermostability [36-38], or on the contrary appeared to have a negative impact [35,39]. In this report, the negative impact of C-terminal CBM containing an Ig-like structure on thermostability as well as acidic stability of $\beta-1,3$-glucanase was revealed.
Proteins possessing the Ig-like fold are commonly found in bacteria, and are most often involved in cell-cell adhesion, or extracellular glycohydrolysis [40]. Recent years,
there is increasing structural information on the existence of these domains in glycoside hydrolase family proteins. Ig-like domain is generally considered being involved in carbohydrate binding and assist in hydrolysis reactions for insoluble substrates [33]. The influence of Ig-like domain on enzymatic stability and activity of GH family proteins has been mainly illustrated in cellulose hydrolases. Liu et al. revealed that deletion of the C-terminal Ig-like domains in GH5 family cellulose resulted in a significant loss of enzymatic activity toward soluble and insoluble cellulose substrates [41]. Many members of family 9 of glycoside hydrolases have an N -terminal Ig-like domain which is responsible for the hydrolytic activity and stability of the catalytic domain [33,42]. Tight structural interaction between the Ig-like and GH9 domains was evidenced by analyzing the crystal structure of the Ig-GH9 module pair [42]. It can be inferred that the C-terminal Ig-like structure also physically associate with the active site of the catalytic domain of PglA and hampered the thermal activity and stability of the enzyme. In contrast, the negligible impact of the N -terminal peptide on enzymatic stability and activity indicated it is very likely to fold as an independent structural module and show little interactions with the catalytic module of PglA.

## Conclusions

In the present study, a new $\beta$-1,3-glucanase gene PglA was cloned from Paenibacillus sp. S09. Truncation forms of PglA were successfully expressed in E.coli. Both the N -and C-terminal non-catalytic domains exerted carbohydratebinding functions, while the C-terminal region containing an Ig-like fold had negative impact on thermostability of catalytic domain. There is increasing documentation that the CBMs of bacterial glycoside hydrolases containing Ig-like structure. As the first report describing the characterization of a GH16 $\beta$-1,3-glucanase with an C-terminus Ig-like region, the present study improved our understanding on the module relationships between the enzyme's catalytic and non-catalytic domains and could provide information that may suggest an approach to rationally modify the characteristics of an enzyme for industrial application.

## Additional files

Additional file 1: Figure S1. Conserved domain sequence alignment of GH16 endo- $\beta-1,3$-glucanases. The organisms and accession numbers are as follows: Nocardiopsis sp.F96 (BAE54302.1), Cellulosimicrobium cellulans (AAC44371.1), Paenibacillus mucilaginosus KNP414 (YP_004643413.1), Bacillus clausii KSM-K16 (YP_174203.1), Bacillus circulans IAM1165 (BAA04469.1), Paenibacillus sp.CCRC 17245 (ABJ15796.1), Phaeodacty/um tricornutum CCAP 1055/1 (XP_002181321.1). The most conserved regions were used for degenerated primers. Identical amino acids are indicated by solid black and gray. The GH16 active site and two glutamate residues are indicated by frame and asterisks respectively.

Additional file 2: The detailed procedures of SEFA-PCR method for amplification of $\beta-1,3$-glucanase gene.
Additional file 3: Figure S2. Protein structure prediction of PgIA. (A) Tertiary structure homology modeling. The PDB IDs of the templates used for modeling were shown in parentheses; ( $B$ ) Secondary structure prediction by PSIPRED 3.0 with a-helixes shown in orange and $\beta$-sheets in blue.

## Abbreviations

GH16: Glycoside hydrolase family 16; I-PCR: Inverse PCR; SEFA-PCR: Self-formed adaptor PCR; CMC: Carboxymethyl cellulose; DNS: Dinitrosalycilic acid; CBM: Carbohydrate-binding module; PglA: Full-length Paenibacillus sp. S09 $\beta$-1,3-glucanase; PgIA $\triangle C$ : C-terminus truncated PglA; PgIA $\triangle \mathrm{N}$ : N -terminus truncated PglA; Pg|A-CD: Catalytic domain of PglA; PglA-N: N-terminal domain of PglA; PglA-C: C-terminal domain of PglA.

## Competing interests

All of authors declare no competing interests.

## Authors' contributions

RC performed the cloning, expression and purification experiments, analyzed the data and was involved in drafting the manuscript. JPC and XHY carried out the enzyme activity assays. SMW participated in the sequence analysis and truncation design. YW assisted with sequence analysis and enzyme characterization. JFZ contributed to study conception and design, analysis and interpretation of data, as well as drafting and revision of the manuscript, and acquisition of funding. All authors read and approved the final manuscript.

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## References

1. Zverlov W, Volkov IY, Velikodvorskaya TV, Schwarz WH: Highly thermostable endo-1,3- $\beta$-glucanase (laminarinase) LamA from Thermotoga neapolitana: nucleotide sequence of the gene and characterization of the recombinant gene product. Microbiology 1997, 143:1701-1708.
2. Henrissat B: A classification of glycosyl hydrolases based on amino acid sequence similarities. Biochem J 1991, 280:309-316
3. De la Cruz J, Pintor-Toro JA, Benítez T, Llobell A, Romero LC: A novel endo- $\beta-1,3$-glucanase, BGN13.1, involved in the mycoparasitism of Trichoderma harzianum. J Bacteriol 1995, 177:6937-6945.
4. Liu B, Xue X, Cui S, Zhang X, Han Q, Zhu L, Liang X, Wang X, Huang L, Chen $X$, Kang $Z$ : Cloning and characterization of a wheat $\beta-1,3$-glucanase gene induced by the stripe rust pathogen Puccinia striiformis f. sp. tritici. Mol Biol Rep 2010, 37:1045-1052.
5. Bielecki S, Galas E: Microbial $\beta$-glucanases different from cellulases. Crit Rev Biotechnol 1991, 10:275-304.
6. Gacto M, Vicente-Soler J, Cansado J, Villa TG: Characterization of an extracellular enzyme system produced by Micromonospora chalcea with lytic activity on yeast cells. J Appl Microbiol 2000, 88:961-967.
7. Shi P, Yao G, Yang P, Li N, Luo H, Bai Y, Wang Y, Yao B: Cloning, characterization, and antifungal activity of an endo-1,3- $\beta$-D-glucanase from Streptomyces sp. S27. Appl Microbiol Biotechnol 2010, 85:1483-1490.
8. Blättel V, Larisika M, Pfeiffer P, Nowak C, Eich A, Eckelt J, König H: $\beta-1,3$-glucanase from Delftia tsuruhatensis strain MV01 and its potential application in vinification. Appl Environ Microbiol 2011, 77:983-990
9. Miyanishi N, Iwamoto Y, Watanabe E, Odaz T: Induction of TNF-a production from human peripheral blood monocytes with $\beta-1,3$-glucan oligomer prepared from laminarin with $\beta-1,3$-glucanase from Bacillus clausii NM-1. J Biosci Bioeng 2003, 95:192-195
10. O'Connell E, Piggott C, Tuohy M: Purification of exo-1,3- $\beta$-glucanase, a new extracellular glucanolytic enzyme from Talaromyces emersonii. Appl Microbiol Biotechnol 2011, 89:685-696.
11. Yamamoto M, Aono R, Horikoshi K: Structure of the $87-\mathrm{kDa} \beta-1,3$-glucanase gene of Bacillus circulans IAM1165 and properties of the enzyme
accumulated in the periplasm of Escherichia coli carrying the gene. Biosci Biotechnol Biochem 1993, 57:1518-1525.
12. Hong TY, Meng M: Biochemical characterization and antifungal activity of an endo-1,3- $\beta$-glucanase of Paenibacillus sp . isolated from garden soil. Appl Microbiol Biotechnol 2003, 61:472-478.
13. Cheng YM, Hong TY, Liu CC, Meng M: Cloning and functional characterization of a complex endo- $\beta-1,3$-glucanase from Paenibacillus sp. Appl Microbiol Biotechnol 2009, 81:1051-1061.
14. Tanabe $Y$, Oda M: Molecular characterization of endo- $1,3-\beta$-glucanase from Cellulosimicrobium cellulans: effects of carbohydrate-binding module on enzymatic function and stability. Biochim Biophys Acta 1814, 2011:1713-1719.
15. Palumbo JD, Sullivan RF, Kobayashi DY: Molecular characterization and expression in Escherichia coli of three $\beta-1,3$-glucanase genes from Lysobacter enzymogenes strain N4-7. J Bacteriol 2003, 185:4362-4370.
16. Xiu A, Kong Y, Zhou M, Zhu B, Wang S, Zhang J: The chemical and digestive properties of a soluble glucan from Agrobacterium sp. ZX09. Carbohyd Polym 2010, 82:623-628.
17. Sambrook J, Russell DW: Molecular cloning: a laboratory manual. New York: Cold Spring Harbor Laboratory Press, Cold Spring Har; 2001.
18. Wang S, He J, Cui Z, Li S: Self-formed adaptor PCR: a simple and efficient method for chromosome walking. Appl Environ Microbiol 2007 73:5048-5051.
19. Petersen TN, Brunak S, Von Heijne G, Nielsen H: SignalP 4.0: discriminating signal peptides from transmembrane regions. Nat Methods 2011, 8:785-786.
20. Geer RC, Gonzales NR, Gwadz M, Hurwitz DI, Lanczycki CJ, Lu F, Lu S, Marchler GH, Song JS, Thanki N, Yamashita RA, Zhang D, Bryant SH: CDD: conserved domains and protein three-dimensional structure. Nucleic Acids Res 2013, 41:348-352.
21. Buchan DW, Ward SM, Lobley AE, Nugent TC, Bryson K, Jones DT: Protein annotation and modelling servers at University College London. Nucleic Acids Res 2010, 38:W563-W568.
22. Arnold K, Bordoli L, Kopp J, Schwede T: The SWISS-MODEL workspace: a web-based environment for protein structure homology modelling Bioinformatics 2006, 22:195-201.
23. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, Thompson JD, Gibson TJ, Higgins DG: Clustal W and Clustal X version 2.0. Bioinformatics 2007 23:2947-2948.
24. Kimura M : A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. J Mol Evol 1980, 16:111-120.
25. Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S: MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol Biol Evol 2011, 28:2731-2739.
26. Miller GL: Use of Dinitrosalicylic acid reagent for determination of reducing sugar. Anal Chem 1959, 31:426-428.
27. Yamamoto M, Ezure T, Watanabe T, Tanaka H, Aono R: C-terminal domain of $\beta-1,3$-glucanase H in Bacillus circulans IAM1165 has a role in binding to insoluble $\beta$-1,3-glucan. FEBS Lett 1998, 433:41-43.
28. Cheng YM, Hsieh FC, Meng M: Functional analysis of conserved aromatic amino acids in the discoidin domain of Paenibacillus $\beta-1,3$-glucanase. Microb Cell Fact 2009, 8:62.
29. Van Bueren AL, Morland C, Gilbert HJ, Boraston AB: Family 6 carbohydrate binding modules recognize the non-reducing end of $\beta-1,3$-linked glucans by presenting a unique ligand binding surface. J Biol Chem 2005, 280:530-537.
30. Asano T, Taki J, Yamamoto M, Aono R: Cloning and structural analysis of bglM gene coding for the fungal cell wall-lytic $\beta$-1,3-glucan-hydrolase BgIM of Bacillus circulans IAM1165. Biosci Biotechnol Biochem 2002, 66:1246-1255
31. Dvortsov IA, Lunina NA, Chekanovskaya LA, Schwarz WH, Zverlov W, Velikodvorskaya GA: Carbohydrate-binding properties of a separately folding protein module from $\beta$-1,3-glucanase Lic16A of Clostridium thermocellum. Microbiology 2009, 155:2442-2449.
32. Koizumi N, Masuda S, Maeda K, Isoda Y, Yatsunami R, Fukui T, Nakamura S: Additional carbohydrate-binding modules enhance the insoluble substrate-hydrolytic activity of $\beta-1,3$-glucanase from alkaliphilic Nocardiopsis sp. F96. Biosci Biotechnol Biochem 2009, 73:1078-1082.
33. Liu H, Pereira JH, Adams PD, Sapra R, Simmons BA, Sale KL: Molecular simulations provide new insights into the role of the accessory immunoglobulin-like domain of Cel9A. FEBS Lett 2010, 584:3431-3435.
34. Araki R, Ali MK, Sakka M, Kimura T, Sakka K, Ohmiya K: Essential role of the family-22 carbohydrate-binding modules for $\beta-1,3-1,4$-glucanase activity of Clostridium stercorarium Xyn10B. FEBS Lett 2004, 561:155-158.
35. Wang Y, Yuan H, Wang J, Yu Z: Truncation of the cellulose binding domain improved thermal stability of endo- $\beta-1,4$-glucanase from Bacillus subtilis JA18. Bioresour Technol 2009, 100:345-349.
36. Verjans P, Dornez E, Segers M, Van Campenhout S, Bernaerts K, Beliën T, Delcour JA, Courtin CM: Truncated derivatives of a multidomain thermophilic glycosyl hydrolase family 10 xylanase from Thermotoga maritima reveal structure related activity profiles and substrate hydrolysis patterns. J Biotechnol 2010, 145:160-167.
37. Winterhalter C, Heinrich P, Candussio A, Wich G, Liebl W: Identification of a novel cellulose-binding domain within the multidomain 120 kDa xylanase XynA of the hyperthermophilic bacterium Thermotoga maritima. Mol Microbiol 1995, 15:431-444.
38. Couturier M, Feliu J, Haon M, Navarro D, Lesage-Meessen L, Coutinho PM, Berrin JG: A thermostable GH45 endoglucanase from yeast: impact of its atypical multimodularity on activity. Microb Cell Fact 2011, 10:103.
39. Selvaraj T, Kim SK, Kim YH, Jeong YS, Kim YJ, Phuong ND, Jung KH, Kim J, Yun HD, Kim H: The role of carbohydrate-binding module (CBM) repeat of a multimodular xylanase (XynX) from Clostridium thermocellum in cellulose and xylan binding. J Microbiol 2010, 48:856-861.
40. Fraser JS, Yu Z, Maxwell KL, Davidson AR: Ig-like domains on bacteriophages: a tale of promiscuity and deceit. J Mol Biol 2006, 359:496-507.
41. Liu G, Qin Y, Hu Y, Gao M, Peng S, Qu Y: An endo-1,4- $\beta$-glucanase PdCel5C from cellulolytic fungus Penicillium decumbens with distinctive domain composition and hydrolysis product profile. Enzyme Microb Technol 2013, 52:190-195.
42. Kataeva IA, Uversky VN, Brewer JM, Schubot F, Rose JP, Wang BC, Ljungdahl LG: Interactions between immunoglobulin-like and catalytic modules in Clostridium thermocellum cellulosomal cellobiohydrolase CbhA. Protein Eng Des Sel 2004, 17:759-769.
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[^1]:    ${ }^{\text {a }}$ The restriction sites included in the oligonucleotides are underlined

[^2]:    ATGAATCGACACGTCCTGCTTCATCCGTATCTCCACCGGAAGCGTTGCCTCTGCTCCTGGCCTTGACGCTGCTGACGGGCATCGCCCTGTTCCCGGCCTCCACCGCGCAGGCGGCGACGACCGTGACGTCGATGACGTACTTCTCTGCC
    

    301 GACAGCGTTGGCGGCTATGTGTACAACCAGAACTGGGCCATTGGAACGACAGCGGCACCTATGGCTACTGGTTCACCCTCTCCGCCACGACCGAGCTGCAGCTCTACTCCAACGCGAACAGCAGCGTCACACTCAACTACACGCTCGTC TTCCAGAATGTCAATGAAACGACCATTACCTCGATGACACCGACCCAGGGCCCGCAATTGACCGCAGGGTATACCGGCGGCGCAGGCTTCACCTATCCGGTCTTCAACAACGATCCCTCCATCCCGTATGCAGCCGTAGCCGGCGATCTG
    
    
    
    1051 TACTGGGCCGACTACATCTACGGCCTCTGGTTCCGGCCGATCCAGGAGGATATGCAGATCCGCATCGGCTATCCGCTGAATGGCCAGTCCGGCGGCAGCGTCGGCAGCAACTTCGTCACCTATACGCTGATCGGCAACCCGAACGCGCCG
    201 CGACCCGATGTGAGCGACCAGGGCGACGTCGAGATCGGCACGCCCACCGATCCGGCCATCGCAGGATGGAATCTGTATTGGCAGGATGAATTCGCCGGCAGCGCGCTCGATCTGAACAAGTGGAACTACGAGACCGGCTACTACATCGGC
    1351 AACGACCCCAATCTGTGGGGCTGGGGCAACGCCGAGATGCAGCACTATACGACGAGCACGCAAAATGTCTTCGTCGCTGACGGCAAACTCAACATCCGAGCGCTCCACGATTACCAATCGTTCCCGCAGGACCCGAACCGCTACGCGACC
    
    1501 TACTCCTCCGGCAAGATCAACACCAAGGACAACATGTCGCTGCAGTACGGCCGCGTCGATATCCGCGCCAAGCTGCCGACTGGCGATGGCGTCTGGCCGGCACTGTGGATGCTGCCGGAGGACTCCGTCTACGGCGCATGGGCGGCATCA
    
    
    1951 GCGGTCGGCGGCCACTTCGACGGCGGGCGTACGCCCGACCCGTCCGACATCCCGGCGACGATGCAGATCGACTACGTGCGGGTGTACAAAGAGGGCGCGGGCGGCGGTCCGGGCAACCCGGGCGGCAACGTCGCGGTGACCGGCGTTAGC
    
    2251 CTCGCTGCTGGCACGACGACCGTAACCGCCACGACCGCAGACGGCAACCGCACCGCCTCGGCGACGATCACCGTCGTGCCGCCACCGACGACGACCGTCATCATCGGCGATAGCGTGCGCGGCATCCGAAACACCGGCGACAACCTGCCTC
    2401 TTCTACGTCAACGGCGCAACCTACGCCGACCTGCACTACAAGGTGAACGGCGCGGTCAGCCTAATGTCGCGATGACGCACACAGGAGGCGGCAACTACACCTACCCGGTGCATGGCCTCCAACAACGCGATACCGTCGAATACTTCTCC
    2551 ACCTACAACCCCGGCAACGGCGCGCTAGACACGCCTTGGCAGACTTATGTGCATGGGGTAACACAAGGTGTTGTTGAGTAA

[^3]:    Data are shown as mean $\pm S D(n=3)$.

