Structure and stability of metagenome-derived glycoside hydrolase family 12 cellulase (LC-CelA) a homolog of Cel12A from Rhodothermus marinus

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Abstract

Ten genes encoding novel cellulases with putative signal peptides at the N-terminus, termed pre-LC-CelA-J, were isolated from a fosmid library of a leaf–branch compost metagenome by functional screening using agar plates containing carboxymethyl cellulose and trypan blue. All the cellulases except pre-LC-CelG have a 14–29 residue long flexible linker (FL) between the signal peptide and the catalytic domain. LC-CelA without a signal peptide (residues 20–261), which shows 76% amino acid sequence identity to Cel12A from Rhodothermus marinus (RmCel12A), was overproduced in Escherichia coli, purified and characterized. LC-CelA exhibited its highest activity across a broad pH range (pH 5–9) and at 90 °C, indicating that LC-CelA is a highly thermostable cellulase, like RmCel12A. The crystal structure of LC-CelA was determined at 1.85 Å resolution and is nearly identical to that of RmCel12A determined in a form without the FL. Both proteins contain two disulfide bonds. LC-CelA has a 16-residue FL (residues 20–35), most of which is not visible in the electron density map, probably due to structural disorder. However, Glu34 and Pro35 form hydrogen bonds with Ala mutation were therefore constructed and characterized. AFL-LC-CelA (residues 36–261) and E34A-LC-CelA with a single Glu34 → Ala mutation were therefore constructed and characterized. AFL-LC-CelA and E34A-LC-CelA had lower melting temperatures (Tm) than LC-CelA by 14.7 and 12.0 °C respectively. The Tm of LC-CelA was also decreased by 28.0 °C in the presence of dithiothreitol. These results suggest that Glu34-mediated hydrogen bonds and the two disulfide bonds contribute to the stabilization of LC-CelA.

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1. Introduction

A metagenomic approach is an efficient method to isolate novel enzymes useful for industrial purposes as well as for academic research [1–4]. In EXPO Park, Japan, leaves and branches cut from the trees are collected periodically, mixed with urea, and agitated for composting. The temperature increases up to ~70 °C inside this compost [leaf–branch compost (LC)] and then decreases to ~50 °C roughly one year later upon completion of composting. This compost is expected to be rich in various thermophilic microorganisms and is therefore a promising source of genes encoding novel thermostable enzymes. We have isolated a novel cutinase, LC-cutinase, with polyethylene terephthalate (PET) degrading activity [5], and 12 novel RNases H1, LC1 ~ LC12-RNases H1 [6], from this compost using a metagenomic approach. Structural and functional studies of these enzymes have shown that LC-cutinase is a kinetically robust protein with a slow unfolding rate [7]; LC9-RNase H1 is a bacterial RNase H1 with an atypical DEDN active site motif, evolutionarily distinct from those with a typical DEDD active site motif [8]; and LC11-RNase H1 is a prokaryotic RNase H1 with a unique substrate recognition mechanism, which interacts with four non-consecutive, instead of consecutive, 2′-OH groups of the RNA strand of RNA/DNA substrates [9,10]. However, no cellulases have been isolated from this compost.
Cellulases hydrolyze β-1,4-glycoside bonds in cellulose and have received much attention because of their potential application in the production of bioethanol from cellulose biomass [11–15]. Cellulases consist of three types of enzymes, endo-1,4-β-D-glucanase (endoglucanase; EC 3.2.1.4), exo-1,4-β-D-glucanase (cellobiohydrolase; EC 3.2.1.91 and 3.2.1.176), and β-glucosidase (EC 3.2.1.21). It has recently been shown that lytic polysaccharide monooxygenase enzyme A99, formerly known as glycoside hydrolase (GH) family 61, represent another class of cellulase degrading enzymes [16]. These enzymes synergistically hydrolyze cellulose to produce glucose [16,17]. According to the CAZy database (http://www.cazy.org/), endoglucanase, cellobiohydrolase and β-glucosidase are classified into twelve, five and six families respectively. GH family 12 is one of the most thermostable endoglucanases known [18]. RmCel12A has a putative signal peptide (SP, residues 1–17) and a flexible linker (FL, residues 18–37) at the N-terminus-free [19] and substrate-bound [20] forms. It has been proposed that dimerization, and a cluster of aromatic residues in the active site cleft, contribute to the adaptation of RmCel12A to a high temperature environment [20].

Thermostable enzymes are generally more useful than thermolabile ones in industry because of their higher temperature, chemical, and pH stability, lower production cost, and longer shelf life [21–23]. In addition, as the reaction temperature increases, the reaction rate increases, the solubility of the substrate and product increases, viscosity of the reaction mixture decreases, and the risk of microbial contamination decreases. Therefore, it would be informative to examine whether thermostable enzymes with cellulose-degrading activity can be isolated from a leaf–branch compost by a metagenomic approach.

In this report, we showed that ten novel cellulases were isolated from leaf–branch compost by a metagenomic approach. We also showed that one of them, LC-CelA, which has high amino acid sequence identity to RmCel12A, was highly thermostable. X-ray crystallographic studies and structure-based mutational studies indicate that hydrogen bonds formed between the C-terminal region of the FL and the central region of the protein, and the two disulfide bonds contribute to the stabilization of LC-CelA.

2. Materials and methods

2.1. Cells, plasmids, and enzymes

Escherichia coli BL21-CodonPlus(DE3)-RP was from Stratagene (La Jolla, CA, USA). Plasmid pET25b was from Novagen (Madison, WI, USA). E. coli BL21-CodonPlus(DE3)-RP transformants were grown in lysogeny broth (LB) medium (10 g Tryptone; 5 g Yeast extract; 10 g NaCl in 1 L H2O) supplemented with 50 mg L\(^{-1}\) ampicillin.

2.2. Construction of DNA library and screening

The 4-month-old leaf–branch compost made in EXPO Park, Japan was used to construct a metagenomic DNA library. The temperature and pH of this leaf–branch compost are 67 °C and pH 7.5. Extraction of DNA from this compost and construction of a DNA library for metagenomic study using CopyControl™ Fosmid Library Production kit (EPICENTRE Biotechnologies, Madison, WI, USA) were performed as described previously [5]. This DNA library was spread on LB-agar plates containing 12.5 μg mL\(^{-1}\) chloramphenicol, 0.01% L-arabinose, 0.5% CM-cellulose, and 0.1 mg mL\(^{-1}\) trypan blue. CM-cellulose and trypan blue have been used as a cellulose substrate and a chromogenic dye respectively for detection of cellulolytic activity [24]. The resultant plates were incubated at 37 °C for several days. Plasmids were extracted from colonies, which form halos around them due to hydrolysis of CM-cellulose. Genes encoding CM-cellulose degrading enzymes were identified by transposon mutagenesis using EZ-Tn5TM-<T7-KAN-2> Promoter Insertion kit (EPICENTRE Biotechnologies), according to the procedures recommended by the supplier. Nucleotide sequence of the gene was determined by an ABI Prism 3100 DNA sequencer (Applied Biosystems, Tokyo, Japan). Oligonucleotides for sequencing were synthesized by Hokkaido System Science (Sapporo, Hokkaido, Japan).

2.3. Construction of plasmids

For construction of plasmid pET-LC-CelA used to overproduce LC-CelA (residues 20–261 of pre-LC-CelA) in a form with Met-Asp at the N-terminus, the gene encoding LC-CelA was amplified by PCR using the fosmid vector harboring the pre-LC-CelA gene as a template. The sequences of the PCR primers were 5′-GATCCATGGATCTGTTCCCAGAG-3′ for 5′-primer and 5′-GAAATTCTACATTACGGGTGTCGTCG-3′ for 3′-primer, where underlines represent the Ncol site for 5′-primer and EcoRI site for 3′-primer.

The resultant DNA fragment was digested with Ncol and EcoRI, and ligated into the Ncol- EcoRI sites of pET25b to generate plasmid pET-pelB-LC-CelA, which harbors the gene encoding the pelB-LC-CelA fusion protein. The small Ndel and Ncol fragment encompassing the pelB leader sequence was then deleted from this plasmid by digestion with Ndel and Ncol, the 5′-overhang filled in by Klenow DNA polymerase, and re-ligation to generate pET-LC-CelA.

For constructions of plasmids pET-LC-CelA-His and pET-ΔFL-LC-CelA-His used to overproduce LC-CelA-His and ΔFL-LC-CelA-His (residues 36–261) in a form with Met at the N-terminus and a His-tag at the C-terminus, the genes encoding LC-CelA and ΔFL-LC-CelA were amplified by PCR using the fosmid vector harboring the pre-LC-CelA gene as a template. The sequences of the 5′-primers were 5′-GATGAGCATATGCGTTCACACACAG-3′ for LC-CelA and 5′-GAGATCATATGCGCTACAGTAGCGC-3′ for ΔFL-LC-CelA. The sequence of the 3′-primer was 5′-TTCGTCGACGGCTGCAGTGCAATGAC-3′ for both LC-CelA and ΔFL-LC-CelA. In these plasmids, underlines represent the Ndel site for 5′-primer and Sall site for 3′-primer. The resultant DNA fragments were digested with Ndel and Sall, and ligated into the Ndel- Sall sites of pET25b.

Plasmid pET-E34A-LC-CelA-His used to overproduce E34A-LC-CelA-His in a form with Met at the N-terminus and a His-tag at the C-terminus was constructed by PCR with the KOD-plus mutagenesis kit (Toyobo Co., Ltd., Osaka, Japan), according to the procedures recommended by the supplier. Plasmid pET-LC-CelA-His was used as a template. The mutagenic primers were designed in such a way that the GAG codon for Glu34 was changed to GCT for Ala.

PCR was performed in 25 cycles with Gene Amp PCR system 2400 (Applied Biosystems) using KOD DNA polymerase (Toyobo). PCR primers and mutagenic primers were synthesized by Hokkaido System Science. The nucleotide sequence was confirmed by an ABI Prism 3100 DNA sequencer (Applied Biosystems).

2.4. Overproduction and purification

E. coli BL21-CodonPlus(DE3) transformants with the pET25b derivatives were cultivated at 37 °C. When the absorbance of the culture at 600 nm reached around 0.5, isopropyl-β-D-thio galactopyranoside (IPTG) was added to the culture medium and cultivation was continued for an additional 4 h. Cells were then harvested by centrifugation at 6000 g for 10 min, suspended in

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**References:**

[11–15, 16, 17, 18, 19, 20, 21–23, 24]
10 mM Tris–HCl (pH 8.0) containing 1 mM EDTA (TE buffer), disrupted by sonication lysis, and centrifuged at 30,000g for 30 min. The supernatant was collected, dialyzed against TE buffer, incubated at 70 °C for 30 min for heat treatment, and centrifuged at 30,000g for 30 min. The subsequent purification procedures were carried out at 4 °C. For purification of LC-CelA–His and its derivatives with a C-terminal His-tag, the supernatant obtained after heat treatment was dialyzed against 20 mM Tris–HCl (pH 7.0) containing 10 mM imidazole and 0.3 M NaCl, and applied to a Ni Sepharose 6 Fast Flow column (GE Healthcare) equilibrated with the same buffer. The protein was eluted from the column by linearly increasing the imidazole concentration from 10 to 300 mM. The fractions containing the protein were collected and dialyzed against 10 mM Tris–HCl (pH 7.0). For purification of LC-CelA without a His-tag, the supernatant obtained after heat treatment was loaded onto a HiTrap Q HP column (GE Healthcare, Tokyo, Japan) equilibrated with TE buffer containing 1 mM DTT. The protein was eluted from the column by linearly increasing the NaCl concentration from 0 to 1 M. The fractions containing the protein were collected and applied to a Hi-Load 16/60 Superdex 200 pg column (GE Healthcare) equilibrated with TE buffer containing 50 mM NaCl for gel filtration chromatography. The fractions containing the protein were pooled.

The production level of the protein in E. coli cells and the purity of the protein were analyzed by SDS–polyacrylamide gel electrophoresis [25] using a 12% polyacrylamide gel, followed by staining with Coomassie brilliant blue (CBB). The amount of the protein was estimated from the intensity of the band visualized by CBB staining using the Scion Image program. The N-terminal amino acid sequence of the protein was determined by a Procise automated sequencer model 491 (Applied Biosystems). The protein concentration was determined from the UV absorption on the basis that the absorbance of a 0.1% (1.0 mg mL\(^{-1}\)) solution at 280 nm is 3.37 for LC-CelA, 2.96 for LC-CelA–His, 3.15 for AFL-LC-CelA–His and 2.97 for E34A-LC-CelA–His. These values were calculated by using \(c = 1.526 \text{ M}^{-1} \text{ cm}^{-1}\) for tyrosine and 5225 M\(^{-1}\) cm\(^{-1}\) for tryptophan at 280 nm [26].

2.5. Sequence analysis

Blast searches of the amino acid sequences of metagenome-derived cellulases deduced from their nucleotide sequences were performed using the DDBJ blastp search tool ([http://blast.ddbj.nig.ac.jp/blastn?lang=en](http://blast.ddbj.nig.ac.jp/blastn?lang=en)). The flexible and hydrophilic regions of the protein were predicted by using the PROTSCALE tool ([http://web.expasy.org/protscale/](http://web.expasy.org/protscale/)). Calculation of pI corresponding amino acid region is performed by using the Compehl program ([http://compel.mbl.de/](http://compel.mbl.de/)).

2.6. Determination of enzymatic activity

The enzymatic activity was determined at the temperatures indicated by the dinitrosalicic acid (DNS) stopped method [27] using CM-cellulose as a substrate. The reaction mixture (100 µL) contained 100 mM sodium phosphate (pH 7.0) and 1% (w/v) CM-cellulose (low viscosity grade, Sigma–Aldrich Co., St. Louis, MO, USA). The enzymatic reaction was initiated by adding an appropriate amount of the enzyme and terminated by adding 10 µL of 10% SDS and boiling for 5 min. The reaction time was 10 min. The resultant solution was mixed with 300 µL of the DNS solution prepared as described previously [27], boiled for 5 min, and cooled on ice. After centrifugation at 17,000g for 5 min, an aliquot of the supernatant (100 µL) was withdrawn, diluted twice by distilled water, and measured for absorption at 500 nm (\(A_{500}\)). The amount of reducing sugars released from the substrate was estimated from the \(A_{500}\) value by using glucose as standard. One unit of enzymatic activity was defined as the amount of enzyme that produced 1 µmol of reducing sugars per min.

For analysis of the temperature dependence of activity, the activity was determined at pH 7.0 and various temperatures ranging from 40 to 100 °C. For the analysis of pH dependence of activity, the activity was determined at 90 °C and various pHs ranging from pH 4.0 to 10.5. The buffers used for this analysis were 100 mM sodium citrate (pH 4.0–6.0), 100 mM sodium phosphate (pH 6.0–8.0), and 100 mM glycine–NaOH (pH 8.0–10.5).

2.7. Measurement of circular dichroism (CD) spectra

The far-UV (200–260 nm) CD spectrum of the protein was measured at 25 °C on a J-725 spectropolarimeter (Japan Spectroscopic, Tokyo, Japan). The protein was dissolved in 10 mM Tris–HCl (pH 7.0). The protein concentration was 0.1 mg mL\(^{-1}\) and a cell with an optical path length of 2 mm was used. The mean residual ellipticity (°, deg cm\(^2\) dmol\(^{-1}\)) was calculated using an average amino acid molecular mass of 110 Da.

2.8. Thermal denaturation

The thermal denaturation curve of the protein was obtained by monitoring the change in CD values at 222 nm as the temperature was increased. The protein was dissolved in 10 mM Tris–HCl (pH 7.0) containing 3.0 M GdmHCl. The protein concentration and optical path length were 0.1 mg mL\(^{-1}\) and 2 mm, respectively. The rate of temperature increase was 3.0 °C min\(^{-1}\). The temperature of the midpoint of the transition, \(T_m\), was calculated by curve fitting of the resultant CD values versus temperature data on the basis of a least-square analysis.

2.9. Crystallization

For crystallization, LC-CelA was dialyzed against 10 mM Tris–HCl (pH 8.0) and concentrated to 10 mg mL\(^{-1}\) using an ultrafiltration system Centricron (Millipore, Billerica, MA, USA). The crystallization conditions were initially screened using crystallization kits from Hampton Research (Aliso Viejo, CA, USA) (Crystal Screens I and II) and Emerald BioStructures, Inc & Emerald BioSystems (Bainbridge Island, WA, USA) (Wizard I, II, III and IV). The conditions were surveyed using sitting-drop vapor-diffusion method at 4 °C and 20 °C. Drops were prepared by mixing 1 µL each of the protein and reservoir solutions, and were vapor-equilibrated against a 100 µL reservoir solution. Crystals of LC-CelA appeared after a few weeks in Wizard I No. 28 [0.1 M 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) (pH 7.5), 0.2 M NaCl and 20% (w/v) polyethylene glycol (PEG) 3000]. To improve the crystal quality, the crystallization conditions were further optimized. Diffraction-quality crystals were obtained when the protein concentration was increased to 11.7 mg mL\(^{-1}\) and the reservoir solution was changed to 0.1 M HEPES (pH 7.5) containing 0.2 M NaCl and 23% (w/v) PEG 3350.

2.10. X-ray diffraction data collection and structure determination

X-ray diffraction data set of LC-CelA was collected at −173 °C without cryoprotectant at a wavelength of 0.9 Å with the beam line BL44XU at SPring-8 (Hyogo, Japan). Diffraction data set was indexed, integrated and scaled using the HKL2000 program suite [28]. The structure was solved by the molecular replacement method using MOLREP [29] in the CCP4 program suite [30]. The 1.85 Å structure of rnbCel12A (PDB: 1H0B) was used as a starting model. Automated model building was done by using ArpWarp
3. Results and discussion

3.1. Cloning of cellulase genes from metagenomic DNA library

The size of the metagenomic DNA library prepared from the leaf–branch compost was approximately 2.1 × 10^8 colony forming units (CFU). The restriction fragment length polymorphisms of 10 randomly selected clones using restriction enzyme BamH1 showed non-redundant patterns and an average insert size of 35 kb. Screening of the library for genes encoding cellulase degrading enzymes was performed using CT-agar plates (agar plates containing CM-cellulose and trypan blue). Of approximately 6000 clones screened, 24 clones gave a halo on CT-agar plates when they were incubated at 37 °C for 3 days. Ten of them, which gave a halo on CT-gellan gum plates (gellan gum plates containing CM-cellulose and trypan blue) at 50 °C, were chosen to determine the nucleotide sequences of the genes responsible for halo formation. Determination of the nucleotide sequences of these genes by transposon mutagenesis indicated that all these genes harbored genes encoding cellulases, that were different from one another. Because these cellulases have a putative signal peptide (SP) at their N-terminus (as described below), they are termed pre-LC-CelA–J. Clones harboring the genes encoding pre-LC-CelA, pre-LC-CelD, pre-LC-CelF, pre-LC-CelG, and pre-LC-Cell produced a halo on CT-gellan gum plates even at 80 °C, suggesting that these cellulases are highly thermostable.

3.2. Amino acid sequences of pre-LC-CelA–J

Blast searches of the amino acid sequences of pre-LC-CelA–J deduced from their nucleotide sequences indicated that all the cellulases are novel, as summarized in Table 1. The number of constituent amino acid residues in these cellulases varies from 261 to 782. The highest identity between the amino acid sequence of either one of these cellulases and that available from the database varies from 42% to 76%. This result indicates that a metagenomic approach is useful to increase our knowledge on sequence space of cellulase. The nucleotide sequences of the genes encoding pre-LC-CelA–J are deposited in GenBank under accession numbers KF626648–KF626657.

The amino acid sequences of pre-LC-CelA–J are schematically shown in Fig. 1. Domain searches using the SMART tool [34] allow us to predict a SP, a catalytic module of cellulase, and a cellulose binding domain/module. All ten cellulases have a 16–36 residue long SP at their N-terminus, suggesting that they are secretory proteins. They also have a catalytic module of cellulase. Three (pre-LC-CelA, pre-LC-CelD and pre-LC-CelE) have a catalytic module of GH family 12 cellulase, two (pre-LC-CelF and pre-LC-CelG) have a catalytic module of GH family 9 cellulase, and two (pre-LC-CelB and pre-LC-CelC) have a catalytic module of GH family 6 cellulase. Pre-LC-Cell, pre-LC-CelH, and pre-LC-CelC have catalytic modules of GH family 3, 4, and 51 cellulases respectively. In addition, pre-LC-CelD and pre-LC-CelF have a cellulose binding domain II (CBII) and pre-LC-CelG has a cellulose binding module 3 (CBM3). Analyses of the hydrophilic and flexible regions using the PROTSacle tool allow us to predict a hydrophilic flexible linker (FL). All the cellulases, except for pre-LC-CelG, have a 14–29 residue long FL between the putative SP and the catalytic domain. Pre-LC-CelG has a 28 residue long FL within a catalytic module of cellulase. These FLs are acidic, because their pl values range from 3.28 to 5.52.

Pre-LC-CelA–J are probably secreted in microorganisms as LC-CelA–J, because SPs are removed by signal peptidase upon secretion. Of them, LC-CelA is expected to be the most stable cellulase, because it shows high amino acid sequence identity to GH family 12 cellulase from R. marinus (RmCel12A) (Table 1). RmCel12A has a 20 residue long FL (residues 12–31) [35]. Removal of this FL decreases the Tm value of RmCel12A by 8.4 °C and its half-life at 90 °C from 5 h to 2 h [35], suggesting that the FL contributes to the stabilization of RmCel12A. However, this stabilization mechanism remains to be understood, because the crystal structure of RmCel12A has been determined only in a form without the FL [19,20]. Therefore, in order to examine whether LC-CelA is, like RmCel12A, a highly thermostable enzyme, and to analyze the role of the FL, we decided to overproduce LC-CelA in E. coli, purify and characterize it, and determine its crystal structure.

3.3. Comparison of amino acid sequences of pre-LC-CelA and pre-RmCel12A

The amino acid sequence of pre-LC-CelA is compared with that of pre-RmCel12A in Fig. 2. Both proteins consist of 261 residues. However, the regions predicted as SP (residues 1–19) and FL (residues 20–35) in pre-LC-CelA are slightly different from those predicted as SP (residues 1–17) and FL (residues 18–37) in pre-RmCel12A. According to the crystal structure of AFL-RmCel12A (RmCel12A without FL, residues 38–261) in complex with the substrate [20], Trp9, Trp26, Trp69, Trp159, Trp161, Tyr163, and Trp209

| Table 1 | List of cellulases isolated from leaf–branch compost and proteins with the highest amino acid sequence identities. |
|-----------------------------------------------|-------------------------------------------------------------------------------------------------|
| Cellulases | No. of residues | Protein with the highest sequence identity | Source organism | Accession No. | Identity (%) | |
| Pre-LC-CelA | 261 | Glycoside hydrolase family 12 | Rhodothermus marinus (65 °C) | G25J29 | 76 |
| Pre-LC-CelB | 596 | Cellulase | Plesiotricystis pacifica SIR-1 (18 °C) | A663J7 | 50 |
| Pre-LC-CelC | 515 | α-L-Arabinofuranosidase-like protein | Opitutus terrae | B1ZZS5 | 45 |
| Pre-LC-CelD | 400 | Glycosyl hydrolase family 12 | Thermosipora spira (50 °C) | D6Y3Q3 | 45 |
| Pre-LC-CelE | 286 | Cellulase 12A | Streptomyces sp. 11AG8 (35 °C) | OQK9H1 | 43 |
| Pre-LC-CelF | 782 | Uncharacterized protein | Streptosporangium roseum (26 °C) | DB8B08 | 62 |
| Pre-LC-CelG | 577 | Glycosyl hydrolase family 9 | Micrococcus sp. PCC 7113 (25 °C) | K9W9M6 | 42 |
| Pre-LC-CelH | 742 | Cellulase | Candidatus methylomoribidis oxyfera (25 °C) | D3M1F3 | 64 |
| Pre-LC-CelI | 592 | Glucosidase-like glycosyl hydrolase | Xenococcus sp. PCC 7305 (25 °C) | L8M232 | 57 |
| Pre-LC-CelJ | 326 | Glycoside hydrolase family 6 | Frankia sp. EUN11 (30 °C) | D3D8M1 | 53 |

* The optimum growth temperature of each organism is shown in parenthesis.
Trp45, Trp62, Trp104, Trp195, Trp197, Tyr199, and Trp245 respectively in pre-RmCel12A) form an extensive aromatic network in the active site cleft, to which the substrate binds. Asn24, His67, Arg100, Met136, Pro137, and Gly138 (Asn60, His103, Arg136, Met172, Pro173, and Gly174 respectively in pre-RmCel12A) also form this cleft. Two catalytic residues, Glu124 and Glu207 (Glu160 and Glu243 respectively in pre-RmCel12A) are located at this cleft. All of these residues are fully conserved in pre-LC-CelA.

3.4. Overproduction and purification of LC-CelA and LC-CelA-His

Because the SP of pre-RmCel12A is cytotoxic to E. coli cells, and the production level of pre-RmCel12A greatly increases on removal of the SP [35], LC-CelA (residues 20–261) was overproduced in E. coli without the SP, either in a non-His-tagged or a His-tagged form. LC-CelA in a non-His-tagged form with Met-Asp at its N-terminus is simply designated as LC-CelA, whereas LC-CelA in a His-
tagged form with Met at the N-terminus and a His-tag at the C-terminus is designated as LC-CelA-His. LC-CelA and LC-CelA-His were used for structural analysis and biochemical characterization respectively. Upon induction for overproduction, LC-CelA and LC-CelA-His equally accumulated in E. coli cells, in both soluble and insoluble forms. The fraction of the protein in soluble form was 20–30%. This protein was purified to give a single band on SDS-PAGE by heat treatment, followed by nickel affinity chromatography (LC-CelA-His) or ion-exchange and gel filtration chromatographies (LC-CelA) (data not shown). The amount of protein purified from 1 L culture was typically 3 mg for both LC-CelA and LC-CelA-His. The N-terminal amino acid sequence of LC-CelA was calculated mass (26.9 kDa), suggesting that LC-CelA exists as a monomer.

The molecular mass of LC-CelA was estimated to be 29 kDa by gel filtration chromatography. This value is comparable to the calculated mass (26.9 kDa), suggesting that LC-CelA exists as a monomer.

AFL-RmCel12A has been reported to exist as a dimer [20]. The interface of the two monomers is stabilized by two salt bridges formed between Glu4 (Glu40 in pre-RmCel12A) of one monomer and Arg47 (Arg83 in pre-RmCel12A) of the other. Arg47 is conserved as Arg81 in LC-CelA, whereas Glu4 is replaced by Thr38 in LC-CelA. LC-CelA does not form a dimer, probably because the salt bridges that stabilize the interface of two monomers are not formed.

3.5. Activity of LC-CelA-His

The temperature dependence of the enzymatic activity of LC-CelA-His was analyzed by measuring the activity at various temperatures ranging from 40 to 100 °C using CM-cellulose as a substrate. As shown in Fig. 3A, LC-CelA-His exhibited the highest activity at 90 °C. This temperature was slightly lower than, but comparable to, that of RmCel12A in a form without the SP, which has been reported to be ≥100 °C [35]. However, the specific activity of LC-CelA-His determined at 60 °C (4.2 units mg⁻¹), was slightly higher than that of RmCel12A without the SP determined at 65 °C (3.1 units mg⁻¹) [35]. These results indicate that LC-CelA-His is a highly thermostable enzyme, like RmCel12A, and is slightly more active than RmCel12A.

The pH dependence of the enzymatic activity was analyzed by measuring the activity at various pHs ranging from pH 4.0 to 10.5. As shown in Fig. 3B, LC-CelA-His exhibited nearly maximal activity across a broad pH range (pH 5.0–9.0). RmCel12A also exhibits nearly maximal activity over a broad pH range (pH 5.0–8.0), but shows only ~50% of the maximal activity at pH 9.0 [18]. This result indicates that the pH range suitable for activity of LC-CelA-His is larger than that of RmCel12A.

3.6. Stability of LC-CelA-His against heat inactivation

To analyze the stability of LC-CelA-His against irreversible heat inactivation, the enzyme (0.05 mg mL⁻¹) was incubated in 100 mM sodium phosphate (pH 7.0) at various temperatures ranging from 60 to 100 °C for 30 min, and the residual activity was determined at pH 7.0 and 60 °C. As shown in Fig. 4, LC-CelA-His almost fully retained activity after incubation at 90 °C for 30 min, whereas it almost totally lost activity after incubation at 95 °C for 30 min. This result is consistent with the observation that the optimum temperature for LC-CelA-His activity is 90 °C (Fig. 3A). LC-CelA-His almost completely lost activity after incubation at 100 °C for 30 min (Fig. 4), but it exhibited approximately 80% of the maximal activity in activity assays at 100 °C (Fig. 3A), probably because the enzyme hydrolyzed the substrate before it was fully denatured in those assay conditions.

3.7. Thermal denaturation of LC-CelA-His

To analyze the stability of LC-CelA-His more quantitatively, thermal denaturation of this protein was analyzed at pH 7.0 in the presence of 3 M guanidine hydrochloride (GdnHCl) by monitoring the change in CD values at 222 nm. Thermal denaturation of the protein was reversible in this condition. It was necessary to use 3 M GdnHCl because the protein was not fully denatured even at 100 °C in the presence of <2 M GdnHCl. Fig. 5 shows the thermal denaturation curve of LC-CelA-His measured in the presence of 3 M GdnHCl. The midpoint of the transition of this curve, Tm, is 86.8 °C (Table 2). The Tm value of RmCel12A has been reported to be 102.9 °C in the absence of denaturant [35]. Because LC-CelA-His is not fully denatured even at 100 °C in the absence of denaturant, LC-CelA-His may be nearly as stable as RmCel12A.

3.8. Activity and stability of LC-CelA

To examine the effects of attachment of a C-terminal His-tag, the activity and stability of LC-CelA were determined and compared with those of LC-CelA-His. The activity of LC-CelA was determined at 60 °C using CM-cellulose as a substrate. The specific activity of LC-CelA determined at this temperature was 4.3 units mg⁻¹, which was comparable to that of LC-CelA-His (4.2 units mg⁻¹). The stability of LC-CelA was analyzed at pH 7.0 in the presence of 3 M GdnHCl by CD spectroscopy. Thermal dena-

**Fig. 3.** Optimum temperature and pH for activity of LC-CelA-His. The temperature (A) and pH (B) dependencies of the enzymatic activity of LC-CelA-His are shown. The activity was determined at pH 7.0 and the temperatures indicated (A) or at 90 °C and the pHs indicated (B) using 1% (w/v) carboxymethyl-cellulose (CM-cellulose) as a substrate, as described in Experimental procedures. The buffers used to analyze the pH dependence of the activity were 100 mM sodium citrate (pH 4.0–6.0), 100 mM sodium phosphate (pH 6.0–8.0) and 100 mM Glycine–NaOH (pH 8.0–10.5). The experiment was carried out at least twice, and errors from the average values are indicated by vertical lines.
Denaturation of LC-CelA was reversible in this condition. The $T_m$ value of this protein was determined to be 86.0 °C, which was comparable to that of LC-CelA-His (86.8 °C). These results indicate that attachment of a C-terminal His-tag does not significantly affect the activity or stability of LC-CelA.

### 3.9. Crystal structure of LC-CelA

To examine whether the FL region of LC-CelA is folded into a defined structure, the crystal structure of LC-CelA was determined at 1.85 Å resolution. The asymmetric unit of the crystal structure consists of two LC-CelA molecules (A and B). Both molecules consist of residues 33–261. The N-terminal Met-Asp and most of the FL region (residues 20–32) are not visible in the electron density map probably due to structural disorder. The structures of these two molecules are nearly identical with a root-mean-square deviation (RMSD) value of 0.08 Å for 229 Ca atoms. We used the structure of molecule A in this study. Details of the data-collection statistics and refinement are summarized in Table 3.

The overall structure of LC-CelA (molecule A) is shown in Fig. 6A. This structure belongs to the typical $\beta$-jelly roll fold and consists of two $\beta$-sheets that pack with each other and one $\alpha$-helix. The structure of LC-CelA excluding the FL region is nearly identical to that of AFL-RmCel12A (PDB: 1H0B) with a RMSD value of 0.33 Å for 226 Ca atoms. Superimposition of the structure of LC-CelA around the active site onto that of the AFL-RmCel12A-substrate (cellopentose) complex determined by the soaking method [20] shows that the steric configurations of all aromatic and non-aromatic residues forming the active site cleft of LC-CelA, including two acidic catalytic residues (Glu158 and Glu241), are nearly identical to those of RmCel12A (Fig. 6B). In the active site cleft of LC-CelA, a band of aromatic residues, mostly tryptophan residues, probably provides a series of glucose-binding subsites through stacking interactions with glucose residues, and charged and polar residues provide specificity through hydrogen bonding interactions with sugar molecules, as reported for RmCel12A [20]. In the crystal structure of AFL-RmCel12A, a HEPES molecule binds to the active site [19]. However, electron density of this molecule was not clearly detected in the LC-CelA structure, suggesting that no HEPES molecule binds strongly to the active site of LC-CelA. This may be due to slight differences in the active site geometry. The LC-CelA structure shows that two disulfide bonds are formed between Cys40 and Cys67 and between Cys100 and Cys105. The Cys40–Cys67 bond apparently anchors the N-terminal region of LC-CelA, and the Cys100–Cys105 bond is conserved only in GH family 12 cellulases from various organisms, whereas the Cys100–Cys105 bond is conserved only in GH family 12 cellulases from R. marinus and Streptomyces species.

As noted above, most of the FL region is structurally disordered. However, the C-terminal region of the FL (Asp33-Pro35) is visible in the electron density map (Fig. 6C), indicating that this region is not included in the FL. Interestingly, this region interacts with the central region (loop between $\beta$A1 and $\beta$B1 strands) to the central region ($\beta$A2 strand). The Cys100–Cys105 bond forms a short loop (residues 101–104). The Cys40–Cys67 bond is well conserved in GH family 12 cellulases from various organisms, whereas the Cys100–Cys105 bond is conserved only in GH family 12 cellulases from R. marinus and Streptomyces species.

![Fig. 4](image1.png)

Fig. 4. Stability of LC-CelA-His and its derivatives against heat inactivation. LC-CelA-His was incubated at pH 7.0 and the temperatures indicated for 30 min in the absence of DTT (filled circle) or in the presence of 1 mM DTT (open circle). AFL-LC-CelA-His (filled square) and E34A-LC-CelA-His (filled triangle) were also incubated at the same condition in the absence of DTT. The residual activities were determined at pH 7.0 and 60 °C using CM-cellulose as a substrate, as described in Experimental procedures. The experiment was carried out at least twice, and errors from the average values are indicated by vertical lines.

![Fig. 5](image2.png)

Fig. 5. Thermal denaturation curves of LC-CelA-His and its derivatives. Thermal denaturation curves of LC-CelA-His (thin dashed line), AFL-LC-CelA-His (thick solid line) and E34A-LC-CelA-His (thick dashed line) obtained in the absence of DTT and that of LC-CelA-His obtained in the presence of 1 mM DTT (thin solid line) are shown. These curves were obtained at pH 7.0 in the presence of 3M GdnHCl by monitoring the change in CD values at 222 nm as described in Experimental procedures.

### Table 2

| Protein                  | [DTT] (mM) | Specific activity$^a$ (U/mg) | Relative activity (%) | $T_m$ (°C) | $\Delta T_m$ (°C) |
|-------------------------|------------|-----------------------------|----------------------|------------|------------------|
| LC-CelA-His             | 0          | 4.2                         | 100                  | 86.8       | –                |
| AFL-LC-CelA-His         | 1          | 4.2                         | 93                   | 58.8       | –28.0            |
| E34A-LC-CelA-His        | 0          | 3.9                         | 96                   | 72.1       | –14.7            |

$^a$ The specific activity was determined at 60 °C either in the presence or absence of DTT using CM-cellulose as a substrate, as described in Experimental procedures. Each experiment was carried out at least twice and the average value is shown. Errors are within 10% of the values reported.

$^b$ The melting temperature ($T_m$), which is the temperature of the midpoint of the transition, was determined from the thermal denaturation curves shown in Fig. 5.

$^c$ $\Delta T_m = T_m$ determined – 86.8 °C.
Table 3
Data collection and refinement statistics of LC-CelA.

| Crystal          | LC-CelA       |
|------------------|---------------|
| Wavelength (Å)   | 0.900         |
| Space group      | C121          |
| Cell parameters  |               |
| a, b, c (Å)      | 130.78, 59.58, 74.95 |
| \(\pi, \gamma, \beta\) (°) | 90.00, 122.89 |
| Molecules/asymmetric unit | 2               |
| Resolution range (Å) | 50.00–1.85 (1.88–1.85) |
| Reflections measured | 297,834       |
| Unique reflections | 41,281         |
| Redundancy       | 7.2 (7.4)     |
| Completeness (%) | 99.4 (99.6)   |
| \(R_{merge}\) (%) | 12.0 (37.1)   |
| Average \(I/\sigma(I)\) | 21.8 (5.6)    |
| Refinement statistics |          |
| Resolution limits (Å) | 62.9–1.9       |
| No. of atoms     | 1798/507      |
| Protein/water    | 16/20.5       |
| \(R_{merge}\) (%) | 12.0 (37.1)   |
| \(R_m\) (%)       | 15.2          |
| Water            | 27.2          |
| Ramachandran plot statistics |        |
| Preferred regions (%) | 96.9          |
| Allowed regions (%) | 3.1           |
| Molprobity score | 1.35          |

\(a\) Values in parentheses are for the highest-resolution shell.

\(b\) \(R_{merge} = \frac{\sum |I_{hkil} - \langle I_{hkil} \rangle|}{\sum I_{hkil}}\), where \(I_{hkil}\) is an intensity measurement for reflection with indices \(hkil\) and \(\langle I_{hkil} \rangle\) is the mean intensity for multiply recorded reflections.

\(c\) Free \(R\)-value was calculated using 5% of the total reflections randomly and omitted from refinement.

\(d\) MolProbity score combines the clashscore, rotamer, and Ramachandran evaluations into a single score, normalized to be on the same scale as X-ray resolution [32].

3.10. Stability of AFL-LC-CelA-His and E34A-LC-CelA-His

To examine whether hydrogen bonding interactions between the FL and central regions contribute to the stabilization of LC-CelA, AFL-LC-CelA-His and E34A-LC-CelA-His were constructed. AFL-LC-CelA-His and E34A-LC-CelA-His are the LC-CelA-His derivatives without the FL and with a single Glu34→Ala mutation, respectively. These proteins were purified to give a single band on SDS-PAGE as was LC-CelA-His by heat treatment, followed by the nickel affinity chromatography. The amounts of these proteins purified from 1 L culture were similar to that of LC-CelA or LC-CelA-His (3 mg). The far-UV CD spectra of AFL-LC-CelA-His and E34A-LC-CelA-His were similar to that of LC-CelA-His (Fig. 7), suggesting that neither removal of the FL region nor a single mutation in the FL significantly affects the gross structure of LC-CelA.

The stability of AFL-LC-CelA-His and E34A-LC-CelA-His with respect to heat inactivation was analyzed at pH 7.0 by incubating these proteins at various temperatures for 30 min and determining their residual activities at 60 °C. The results are shown in Fig. 4. Unlike LC-CelA-His, which retains maximal activity after incubation at 90 °C, AFL-LC-CelA-His lost approximately 60% and 100% of its activity after incubation at 85 °C and 90 °C respectively. Similarly, E34A-LC-CelA-His lost approximately 20 and 60% of its activity after incubation at 85 °C and 90 °C respectively. These results indicate that the thermal stability of the proteins decreases in the order LC-CelA-His > E34A-LC-CelA-His > AFL-LC-CelA-His.

The stability of AFL-LC-CelA-His and E34A-LC-CelA-His was also analyzed at pH 7.0 in the presence of 3 M GdnHCl using CD spectroscopy as described above for LC-CelA-His. Thermal denaturation of the proteins was reversible in this condition. The thermal denaturation curves of AFL-LC-CelA-His and E34A-LC-CelA-His are shown in Fig. 5. The midpoints of the transitions of these curves, \(T_m\), are summarized in Table 2. The \(T_m\) values of AFL-LC-CelA-His and E34A-LC-CelA-His are lower than that of LC-CelA-His by 14.7 °C and 12.0 °C, respectively, indicating that the FL region of LC-CelA contributes to protein stability (as does the FL of RmCel12A). The difference in \(T_m\) values between LC-CelA-His and E34A-LC-CelA-His accounts for 82% of that between LC-CelA-His and AFL-LC-CelA-His. This result suggests that Glu34-mediated hydrogen bonding interactions are the major forces responsible for the FL-dependent stabilization of LC-CelA. Hydrogen bonds have been reported to make large contributions to protein stability [36]. The FL region (Leu20-Glu32) may not significantly contribute to the stabilization of LC-CelA.

3.11. Stability of LC-CelA-His in the presence of DTT

Proteins are usually [7,37–40], but not always [41,42], destabilized by removal of a native disulfide bond. To examine whether the two disulfide bonds of LC-CelA contribute to protein stability, thermal denaturation of LC-CelA-His was analyzed as described above, at pH 7.0 in the presence of 1 mM dithiothreitol (DTT), which reduces disulfide bonds to thiol groups, and 3 M GdnHCl. Thermal denaturation of LC-CelA-His was reversible in this condition. The thermal denaturation curve of LC-CelA-His measured in the presence of 1 mM DTT and 3 M GdnHCl is shown in Fig. 5. The \(T_m\) value of LC-CelA-His determined in the presence of 1 mM DTT is lower than that determined in the absence of DTT by 28.0 °C (Table 2). This result indicates that disulfide bonds contribute significantly to the stabilization of LC-CelA-His. The \(T_m\) value of LC-CelA-His (59.8 °C) determined in the presence of 10 mM DTT and 3 M GdnHCl was comparable to that (58.8 °C) determined in the presence of 1 mM DTT and 3 M GdnHCl, suggesting that the disulfide bonds of LC-CelA-His are fully reduced by 1 mM DTT.

It remains to be determined which disulfide bond contributes more to the stabilization of LC-CelA. However, it has been reported that the contribution of disulfide bonds to protein stability increases as the number of the residues in the loop formed by the disulfide bond increases [37]. That number is 26 for Cys40→Cys67 and four for Cys100→Cys105. Therefore, the Cys40→Cys67 bond may contribute significantly more to the stabilization of LC-CelA than the Cys100→Cys105 bond.

Disulfide bonds do not account for the differences in stability between LC-CelA or RmCel12A and a GH family 12 cellulase from Streptomyces species, because the two disulfide bonds of LC-CelA are conserved in these proteins. The \(T_m\) values of RmCel12A and Cel12A from Streptomyces sp. 11AG8 (StCel12A), which represents GH family 12 cellulases from Streptomyces species, have been reported to be 102.9 °C [35] and 65.7 °C [43] respectively in the absence of denaturant, indicating that RmCel12A is more stable than StCel12A by 37.2 °C. Because LC-CelA is nearly as stable as RmCel12A, LC-CelA is also more stable than StCel12A. Proteins are stabilized relative to each other by a combination of factors, including increased number of ion pairs, increased number of hydrogen bonds, increased number of disulfide bonds, increased number of proline residues in loop regions, and increased interior hydrophobicity [36]. Therefore, we compared these structural factors in LC-CelA, RmCel12A and StCel12A (PDB: 10A4). The ratio of...
interior apolar residues to total interior residues in LC-CelA (66.7%) is comparable to that in RmCel12A (65.2%), but significantly higher than that in StCel12A (60.8%). Similarly, the number of ion pairs in LC-CelA (10) is comparable to that in RmCel12A (8), but higher than that in StCel12A (5). However, the number of hydrogen bonds and proline residues in loop regions of LC-CelA and RmCel12A are not consistently higher than in StCel12A. The number of hydrogen bonds is 202 in LC-CelA, 177 in RmCel12A, and 197 in StCel12A. The number of proline residues in loop regions is 8 in LC-CelA, 7 in RmCel12A, and 13 in StCel12A. These results suggest that differences in interior hydrophobicity and in the number of ion pairs at least partially account for the difference in stability between LC-CelA or RmCel12A and StCel12A.

3.12. Activities of ΔFL-LC-CelA-His and E34A-LC-CelA-His

The enzymatic activities of ΔFL-LC-CelA-His and E34A-LC-CelA-His were determined at 60 °C in the absence of DTT using CM-celullose as a substrate. The activities of these proteins were not determined at the optimum temperature for activity of LC-CelA-His (90 °C), because they are thermally denatured at this temperature (Fig. 4). The specific activities of ΔFL-LC-CelA-His and E34A-LC-CelA-His determined in the absence of DTT were slightly lower than, but comparable to, that of LC-CelA-His (Table 2). These data indicate that the enzymatic activity of LC-CelA-His is not significantly affected by removal of, or mutation in, the FL. The specific
activity of LC-CelA determined in the presence of 1 mM DTT was identical to that determined in the absence of DTT, indicating that the enzymatic activity of LC-CelA-His is not significantly affected by reduction of the disulfide bonds.

4. Conclusion

LC-CelA, a homolog of the highly thermostable GH family 12 cellulase from *R. marinus* (*RmCel12A*), was isolated from leaf–branch compost by a metagenomic approach and its crystal structure was determined. This structure is the first of a GH family 12 cellulase with an acidic flexible linker (FL) at the N-terminus. The structure shows that most of the regions predicted as FL is disordered. This result supports the hypothesis that the FL is necessary to separate a hydrophobic signal peptide that anchors the enzyme to the cell from the catalytic core [35]. However, Glu34 and Pro35 located at the C-terminus of the FL assume a defined structure and form hydrogen bonds with the central region of the protein. The structure also shows that LC-CelA contains two disulfide bonds (Cys40–Cys67 and Cys100–Cys105). LC-CelA is a highly thermostable enzyme with an optimum temperature for activity of 90°C (Cys40–Cys67 and Cys100–Cys105). LC-CelA is a highly thermostable enzyme with an optimum temperature for activity of 90°C (Cys40–Cys67 and Cys100–Cys105). LC-CelA contains two disulfide bonds (Cys40–Cys67 and Cys100–Cys105). It has been reported for this enzyme that mutations in the N-terminal region [45] and engineering of an N-terminal disulfide bond [46–49] stabilize the protein.

Cellulases with improved activity and stability have been engineered [50,51]. However, studies of protein engineering of GH family 12 cellulases are still limited. The results presented in this work provide valuable information that serves as a structural toolbox for the engineering of GH family 12 cellulases with improved function.

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