BIOCHEMICAL CHARACTERIZATION OF A PECTATE LYASE FROM 
CALDICELLULOSIRUPTOR BESCII

by

PUJA CHANDRAYAN

(Under the Direction of Michael W.W Adams)

ABSTRACT

*Caldicellulosiruptor bescii* is a gram-positive thermophilic bacterium with the ability to grow on untreated plant biomass. Upon growth on switchgrass at 78°C, a set of genes encoding 94 proteins was up-regulated transcriptionally by more than 4-fold, in comparison to growth on glucose. Enzymes with activity towards pectin were among those that were most highly up-regulated. Of a total of four predicted pectinolytic enzymes, Cbes_1854, a multidomain protein of polysaccharide lyase family 3, was chosen for overexpression and purification in *E. coli*. The full-length protein (X-PL3) and just the catalytic domain (PL3) were purified with yields of approximately 10 mg/liter *E. coli* culture. Both purified recombinant proteins display alkaline pH optima and have a half-life of approximately one hour at 90°C. Under *in vitro* assay conditions, domain X does not play a significant role in catalysis. The catalytic domain, PL3, also shows a synergistic effect on sugar release from untreated switchgrass biomass during *in vitro* assays with a cellulase (Cel A) from *C. bescii*. In a collaborative study, the crystal structure of the PL3 domain was obtained. This provided insight into the catalytic mechanism of the enzyme.

INDEX WORDS: *Caldicellulosiruptor bescii*, Cbes_1854; Pectinolytic enzymes, Polysaccharide lyase family 3; Catalytic domain (PL3), Full-length protein (X-PL3); *E. coli*, Cellulase (Cel A)
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DEDICATION

This work dedicated to my parents:

Shri Laliteshwar Kumar Singh and Veena Kumari

For their blessings
ACKNOWLEDGEMENTS

I express my sincere gratitude to Dr. Adams for giving me an opportunity to work under his guidance. I would like to thank Dr. Lanzilotta and Dr. Maier for their advice on my research project. I am grateful to Dr. Irina Kataeva, who helped immensely in design and completion of the proposed research work. I am also thankful to Dr. Mirko Basen for giving me useful advice for the execution of my experiments. I would also like to thank my BESC team members Amanda Rhaesa and Israel Scott for reading drafts of my thesis and for many useful comments. I am also thankful to the lab manager, Farris Poole, for his continuous effort to make the workplace more efficient and productive. I also offer my sincere thanks to all other Adams lab members who helped me whenever it was needed.

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CHAPTER 1
INTRODUCTIONS AND LITERATURE REVIEW

Recalcitrance of plant biomass and role of pectin

The fossil fuel crisis has accelerated the search for renewable sources of energy [1-3]. Amongst the various available alternatives to fossil fuels, ethanol from cellulose has become a viable option [4-7]. As a feedstock, cellulose is the most abundant organic biopolymer, but exists in a complex form of lignocellulosic biomass [8,9]. Of the total lignocellulosic biomass, 70% is represented by plant cell walls but due to cell wall recalcitrance only 2% can be currently used for the production of biofuels [9,10]. It is a major challenge to develop a cost-effective bioconversion process [3,11,12]. The reported poor yield of this process is largely due to the natural resistance of the complex plant cell wall to deconstruction [8,13,14]. Cell wall resistance to enzymatic degradation is primarily due to the crystalline nature of cellulose, which is a major component of the plant cell wall, and its intimate association with cell wall polymers such as hemicellulose, pectin and lignin [9,15]. Ultrastructure studies show that the cell wall has three layers: (1) the middle lamella, which is outermost layer and has a significant amount of pectic polysaccharide and proteins; (2) the primary cell wall, which is the next layer after the middle lamella and largely consists of cellulose microfibrils cross-linked by hemicellulose and (3) the innermost layer of the plant cell wall termed the secondary cell wall, and this is mainly composed of cellulose microfibrils and lignin [9,15,16]. In general, there are primarily two types of cell walls depending on the type of crosslinking. Type-I plant cell walls are abundant in dicot plants and non-commelinoid monocot and contains xyloglucans (XyGs) as cross-linking
polysaccharides [9]. Type-II plant cell walls are prevalent in commelinoid grasses such as switchgrass and contain glucuronoarabinoxylans (GAXs) for crosslinking of the cell wall. Switchgrass is a widespread grass in eastern United States and has been proposed as an ideal plant for lignocellulose conversion [9,17,18]. Besides differences in hemicellulose content, cell walls also differ in the amounts of pectic polysaccharide and soluble protein that they contain [19].

Among the different polysaccharides within plant cell walls, pectin represents a very small amount, typically less than 5% of the total polysaccharide [19]. Nevertheless, pectin has an essential role in maintaining the structural integrity, mechanical strength and shape of the cell wall. Pectic polysaccharides together with a variety of proteins are present in the middle lamella and primary cell wall. There are three major types of pectic polysaccharide that have been found in plant cell walls and these are: (1) homogalacturonan, (HG) (2) rhamnogalacturonan I (RG-I) and (3) rhamnogalacturonan II (RG-II). HG and RG-I have a polygalacturonic acid backbone joined together by α-1-4 linkage while RG-II has a disaccharide backbone of rhamnose and polygalacturonic acid. Although RG-I and HG have a similar backbone, RG-I is heavily modified at the O-2 and O-3 positions by multiple hexose and pentose sugars while HG are mostly esterified by methyl group or by a simple xylose sugar [19].

By proportion in the primary cell wall, HG constitutes 65% of the total pectic polysaccharide, 20-25% is made up of RG-I and rest consists of RG-II and XGA. It is still not very clear how pectin and other polysaccharides are organized into the primary cell wall. To understand the organization in the primary cell wall, an in vitro attempt to extract pectin was carried out by mild salt and base that resulted into an insoluble pectin fraction with cellulose and hemicellulose. The insoluble pectin fraction with hemicellulose constitutes almost 10-50% of the
total pectin [20]. This observation demonstrates that the pectin lies in the insoluble network of polysaccharides within the plant cell wall rather than soluble phase. Recently, it has been shown that mutation of the homogalacturonan synthesis gene, GAUT1, of the model plant Arabidopsis thaliana, gives rise to a phenotype in which the cell wall contains less pectin [21]. Caldicellulosiruptor saccharolyticus, which lacks pectin-degrading capability, has been shown to grow better on transgenic plants with less pectin [21]. In a separate study, reduction of de-methyl-esterified homogalacturonan (HG) in A. thaliana increases the efficiency of enzymatic sugar release in bioconversion processes [21,22]. All of these observations suggest an important role for pectin in cell wall recalcitrance.

**General features of the pectate lyases**

According to enzyme nomenclature, pectate lyase fits into enzyme class EC 4.2.2.2 and is a carbon-oxygen lyase. It uses uronic acid containing polysaccharide as a substrate. Uronic acid is a sugar acid with a carbonyl or hydroxyl group at position C1 or C6 of a monosaccharide that has undergone oxidation to form a carboxylic acid. In D-polygalacturonic acid the hydroxyl group at C6 position of D-galactose has been oxidized into a carboxylic acid [19,23,24]. Pectin is one of the complex polysaccharides made up from uronic acid carrying polysaccharides joined together by α1-4 glycosidic linkages. The substrate utilized by pectate lyase is a type of pectin that contains homogalacturonans and forms an integral part of plant cell wall. On the basis of the mechanism of catalysis and nature of substrate, the class of enzyme is categorized into the polysaccharide lyase family [24].

Pectate lyases from fungi and bacteria have been extensively characterized and have provided valuable insight regarding the general properties of this type of enzyme. It cleaves the carbon–oxygen bond of α1-4 glycosidic linkage between uronic acids, leaving an unsaturated
C4-C5 bond at the newly formed non-reducing end. Absorption at 235nm of the unsaturated product has been utilized for in vitro assays of pectate lyase. Pectate lyase activities are typically maximal in the alkaline pH range (8.5-10.5) and a metal ion is essential for catalysis. Ca$^{2+}$ is used as a metal cofactor in most enzymes, except for the few that use Co$^{2+}$, Mn$^{2+}$, and Ni$^{2+}$. The divalent cations participate like a cofactor in enzyme-substrate interaction by binding both the enzyme and the charged sugar substrate [23,24].

A systematic study of distribution of polysaccharide lyases among different domains of life has been made based on their protein sequence similarity and known catalytic features. On this basis, they have been organized into 22 families of polysaccharide lyase (PL) [25]. Among these only five families, PL1, PL2, PL3, PL9 and PL10, have been shown experimentally to be pectate lyases. Several structures have been determined for representatives of all five polysaccharide lyase families. By analyzing all available three-dimensional structures, it has been observed that there are three different kinds of topologies for the catalytic module: (1) a right-handed β-helix fold which is common with the families PL-1, PL-3, and PL-9, (2) an (α/α) 7 toroid in family PL2 and (3) an (α/α) 3 toroid in family PL-10. β-helix fold is more abundant among different pectate lyase and is structurally very unique [24].

Despite differences in substrate specificities and variation in structural topology all 22 families still have a common conserved mode of catalysis [26]. Catalysis involves a β-elimination reaction for the cleavage of the glycosidic bond. In contrast to glycosyl hydrolyase, polysaccharide lyase breaks the bond without the involvement of a water molecule and results in the formation of an unsaturated bond at the non-reducing end. The catalytic mechanism can be summarized into three steps: (1) abstraction of the C-5 proton on the sugar ring of a uronic acid or ester by a basic catalytic amino acid, (2) stabilization of the resulting anion by charge
delocalization into the C-6 carbonyl group and (3) lytic cleavage of the O-4: C-4 bond, facilitated by proton donation from a catalytic acid to yield a hexenuronic acid and hexuronic acid. The unsaturated product is one of the crucial pieces of evidence for the β-elimination [27].

**Caldicellulosiruptor bescii: the most thermophilic lignocellulolytic bacterium**

The crystalline nature of cellulose and presence of other heterogeneous polysaccharides has been reported as a major obstacle in development of a cost-effective process for making biofuels [12]. A strategy to develop an optimized and cost-effective pathway for production of biofuel or chemicals from insoluble complex cellulose is the subject of extensive research efforts and poses a major scientific and technical challenge. Current processes involve two major strategies. The first strategy is physio-chemical pretreatment of lignocellulosic biomass followed by enzymatic conversion into fermentable sugars, which are subsequently fermented to ethanol. The second strategy has been proposed on the basis of pioneering work done on *Clostridium thermocellum*. This process is known as consolidated bioprocessing (CBP), in which *C. thermocellum* carries out both saccharification and fermentation [4,10,28-30]. *C. thermocellum* has optimal growth temperature of 60°C and it is thought that high temperature increases the efficiency of plant biomass deconstruction [31]. High temperature also reduces the chances of microbial contamination, enhances the rate of catalysis, facilitates the simple purification of liquid biofuels from the growth medium and decreases the viscosity. Reduced viscosity helps in the fermentation of high loads of insoluble biomass [32]. Therefore, thermophilic cellulolytic microbes have been sought for efficient production of biofuels [31,33,34].

Among known thermophilic bacteria, species of the *Caldicellulosiruptor* genus have potential to be used in CBP due to their cellulolytic capability in the temperature range of 75-80°C [35]. All members of this genus are gram-positive, anaerobe, thermophilic and non-spore
forming bacteria. At present nine different species of *Caldicellulosiruptor* have been isolated and growth studies have been carried out [35]. Among them, *C. bescii* has the highest growth temperature and can grow up to 90°C with an optimum growth temperature of 78°C [36,37]. This strain was originally deposited in 1990 in the German Collection of Microorganisms and Cell Cultures (DSMZ) as DSM 6725 [38]. It was classified as a member of a new genus *Anaerocellum* and named as *Anaerocellum thermophilum* strain Z-1320. Subsequent growth studies on strain DSM 6725 with cellulose, hemicellulose and pectin showed marked differences compared to what was originally reported with strain Z-1320. For example, DSM 6725 was capable of growing on xylose and pectin while earlier it was shown that the strain Z-1320 could not utilize xylose and pectin [38]. The other notable difference was in the growth temperature of DSM 6725 (Figure 1.1 A) [38]. DSM 6725 strain could grow up to 90°C while the Z-1320 strain can grow up to 83°C only (Figure 1.1 A). These differences, together with 16S rRNA sequence analyses, showed that DSM 6725 was highly similar to *Caldicellulosiruptor saccharolyticus* suggesting that it is a new *Caldicellulosiruptor* genus. Subsequently, DSM 6725 was proposed as type strain of a new species named as *Caldicellulosiruptor bescii*. The rod-shaped morphology of *C. bescii* is shown in Figure 1.1 B [38].

*C. bescii* grows well on cellulose, xylan and pectin and, remarkably, was shown to grow well on untreated plant biomass, including poplar and switchgrass [37]. Its capability of growing on untreated switchgrass considerably enhances the utility of *C. bescii* as a potential CBP microbe. A growth curve of *C. bescii* on wild type switchgrass is shown in Figure 1.2. A cell density of $1 \times 10^8$ cells/ml was reached on all complex substrates, which is similar to the cell density reported after growth on the simple sugar cellobiose (Figure 1.1) [37]. Quantitatively, after three successive growths on spent switchgrass, *C. bescii* can degrade up to 85% (w/w) of
Figure 1.1. Growth temperature and morphology of *C. bescii*. (A) Comparison of growth temperature of *C. bescii* with *A. thermophilum* Z-1320 [38]. (B) Scanning electron micrograph of *C. bescii* grown on 0.5% cellobiose. Taken from reference [38].
the initial switchgrass biomass (Figure 1.2) [39]. Considering its potential future application on an industrial scale, *C. bescii* exceeds all other cellulolytic microbes by showing uninhibited growth using up to 200 g/liter switchgrass as well as on similar concentration of crystalline cellulose (Figure 1.3) [40]. Interestingly, acid pretreated switchgrass (50 g/liter) inhibits growth of the *C. bescii*, showing the problems associated with the harsh pretreatment conditions [40].

Another crucial property of *C. bescii* is its ability to solubilize lignin, which could be very advantageous in enhancing the efficacy of CBP [39]. *C. bescii* can solubilize the cellulose microfibrils as well as the lignin components of cell walls at high temperature, thus making it a very promising candidate for CBP [36,37]. Currently, the bottleneck in using *C. bescii* on an industrial scale is its inability to produce ethanol, which is in contrast to members of the genus *Clostridium*, which generate ethanol when grown on cellulose [31]. In pursuit of a better technology and considering the recent development of genetic tools for *C. bescii*, it is possible to envision *C. bescii* as an efficient CBP-microbe in the not too distant future [41,42].

**Plant biomass and pectin deconstructing enzymes in *C. bescii***

The *C. bescii* genome encodes 2666 proteins among which 259 are categorized as carbohydrate active enzymes (CAZy) [25,43]. These CAZy genes are predicted to make 180 transcriptional units, 111 of which are multigene operons while 69 genes exist as single genes. CAZy enzymes often contain at least one cellulose-binding module (CBM), which factor greatly in their capacity to degrade cellulose and plant biomass [43,44]. For example, CBM3-containing enzymes attach to crystalline cellulose and CBM22-containing enzymes bind to xylan. *C. bescii* has enzymes carrying both of these CBM modules and moreover they also have other classes of CBM-containing enzymes. Importantly, some of the CBMs are specific to *C. bescii*. For example, CBM4_9 is present in all three extracellular pectinolytic enzymes [43,44].
Figure 1.2. Growth of *C. bescii* on untreated switchgrass. (A) Growth curve with hot water washed biomass (wSG) and successive growth curves on spent switchgrass SG1 and SG2, which is the insoluble biomass remaining at the end of growth that is used for next culture (taken from reference [37]). (B) Switchgrass degradation in successive growth experiments is plotted by comparing the residual switchgrass (%, w/w) in each step. The control involves incubating switchgrass without *C. bescii*. Taken from reference [39].
Figure 1.3. Growth of *C. bescii* on high concentrations of crystalline cellulose and switchgrass. 

(A) Green triangles, 5 g/liter; orange circles, 200 g/liter; and blue diamonds, 200 g/liter of crystalline cellulose or (B) unpretreated switchgrass. Taken from reference [40].
The potential role of CAZy enzymes in plant biomass degradation has been studied by transcriptional analyses and glycan immunoanalyses. Transcriptional analyses of the growth of *C. bescii* on unpretreated switchgrass versus growth on glucose revealed that there are 94 enzymes whose genes are up-regulated more than 4-fold when grown on untreated switchgrass and among these 18 have a functional role in carbohydrate metabolism. Of these 18 enzymes, 12 have the potential to bind and hydrolyze the cellulose, xylan, xyloglucans, pectin and mannans [39]. Up-regulated genes are shown in Table 1.1 [39].

Enzymes that hydrolyze cellulose, hemicellulose, xylan and xyloglucan are all involved in degradation of plant biomass [35,45]. However with the *C. bescii*, the presence of pectin-degrading enzymes among those that are up-regulated upon growth on unpretreated switchgrass is a novel and very interesting finding. Remarkably, two of the predicted pectin-degrading enzymes are up-regulated by more than 20-fold [39]. The importance of these pectin-degrading enzymes is also confirmed by immunoanalyyses where pectin-related epitopes showed increased extractability during switchgrass degradation [39]. This suggests that hydrolysis of specific pectin components facilitates biomass degradation by *C. bescii*.

Pectin-degrading enzymes fall into the category of polysaccharide lyase (PL) enzymes. The *C. bescii* genome contains 4 PLs. These are PL3 (Cbes_1854), PL9 (Cbes_1855), PL11 (Cbes_1853) and an unclassified PL (Cbes_2353), putatively active with polygalacturonate and other pectins, including rhamnogalacturonans. Gene clusters showing all the classified polysaccharide lyases has been shown in Figure 1.4. The three PL genes are located in a large CAZy cluster encoding several multidomain, multi-functional enzymes active against all major components of plant biomass. PL3 and PL9 have an N-terminal X domain of unknown function and and PL11 has a C-terminal CBM3 domain that is proposed to specifically bind with
Table 1.1: Up-regulated genes of *C. becii* after growth on switchgrass

<table>
<thead>
<tr>
<th>Gene</th>
<th>SP</th>
<th>CAZy modules</th>
<th>Catalytic activity</th>
<th>Binding</th>
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<tr>
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<td>Y</td>
<td>PL11-CBM3</td>
<td>rhamnogalacturonan lyase</td>
<td>cellulose</td>
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<td>PL (7)</td>
<td>unclassified pectate lyase</td>
<td>pectin</td>
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<tr>
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SP: signal peptide. Taken from reference [39].
Figure 1.4. Gene and domain organization of four predicted polysaccharide lyases of *C. bescii*.

(A) Domain organization for each gene is shown. The abbreviations are: SS, signal sequence; PL, polysaccharide lyase; CBM, cellulose binding modules and X, domain of unknown function.

(B) Gene organization of three extracellular pectate lyase genes as a cluster in the genome.
crystalline cellulose. The roles of these three PLs are to degrade untreated plant biomass as the three polysaccharide lyases are not highly up-regulated upon growth of *C. bescii* on cellulose or cellobiose. Two of them, PL3 and PL9, were part of the secretome at the beginning of growth on cellulose and cellobiose growth but they are downregulated as growth continues [46]. This result is not surprising because growth on cellulose does not release the same hydrolysis products as grow on switchgrass. Presumably, the initial hydrolysis of switchgrass biomass leads to exposure of pectin epitopes and release of soluble pectin-derived products that cause the up-regulation of genes encoding pectin-degrading enzymes.

**Goals of study**

Considering the importance of pectin-degrading enzymes in plant biomass degradation, the primary goal of this study was to carry out a biochemical study of three extracellular PL-type enzymes from *C. bescii* in order to provide insight their roles to plant biomass deconstruction. Ultimately, this information would determine which enzymes should be up-regulated to increase the efficiency of plant biomass degradation or be used to better design of enzyme mixtures for in *vitro* plant biomass degradation.

Earlier efforts to make multidomain cellulase A (Cbes_1867) from *C. bescii* in *E. coli* were unsuccessful due to proteolysis in linker region that is present between two domains of cellulase A [47]. In this study, pectin-degrading enzymes were chosen to demonstrate overexpression in *E. coli* due to its proposed role in untreated switchgrass degradation. Pectin-degrading enzymes also have similar linker regions like Cbes_1867. Therefore, overexpression of pectin-degrading enzyme in *E. coli* is a challenge due to potential proteolytic susceptibility in the domain linker region of the protein [48].
Here the goal was the heterologous expression of pectin-degrading enzymes and the determination of their biochemical properties, including pH and temperature optima and temperature tolerance.
CHAPTER 2

CLONING, EXPRESSION AND PURIFICATION OF A PECTATE LYASE IN E. COLI

Introduction

The genes encoding four predicted pectin-degrading enzymes are up-regulated during growth of *C. bescii* on switchgrass in comparison with growth on glucose [39]. The corresponding genes are Cbes_1853, Cbes_1854, Cbes_1855 and Cbes_2353. Cbes_2353 is predicted to be an intracellular protein and belongs to an unclassified pectate lyase family [43]. The CAZY classification is unknown for this protein. The other three enzymes are predicted to be extracellular. All three extracellular enzymes share similar domain architecture with signal sequence at the N-terminus. Following the signal sequence, there are catalytic and substrate-binding domains. The domain arrangement is shown in Figure 1.4 (chapter 1). In all three proteins the catalytic and binding domains are separated by linkers, composed of repetitive amino acid sequences rich in proline, threonine and serine. Linkers promote proper interaction of the catalytic domains with insoluble substrate [49-52]. Presumably, proline residues provide conformational rigidity to binding and catalytic domains while serine and threonine provides sites for O-glycosylation that plays roles in enhancing the protein stability in the extracellular environment [53].

Cbes_1853 has a catalytic module of the PL11 family followed by a CBM3 domain at the C-terminus. The PL11 family has predicted rhammogalacturonan lyase activity and the CBM3 domain proposed to bind crystalline cellulose. In comparison to Cbes_1853, the arrangement of catalytic and substrate binding domains is reversed in Cbes_1854 and Cbes_1855. Both proteins
contain catalytic domains, PL3 and PL9, respectively. The PL3 and PL9 families have been characterized as pectate lyases. The substrate-binding domain of both proteins shares homology with CBM4 (subfamily 9). The family 4 CBM module is composed of approximately 150 residues and is mostly found in mostly in bacterial enzymes not in archaeal enzymes. The binding of this module to xylan, glucan and amorphous cellulose has been confirmed [54].

The goal here was to overexpress multidomain Cbes_1854 in *E. coli*. Besides the full-length protein, its two domains, ‘X’ and PL3, were also cloned and expressed separately in *E. coli* for biochemical and structural studies.

**Materials and Methods**

*Polymerase chain reaction (PCR):* PCR amplification of X, X-PL3 and PL3 was carried out with 20 ng *C. bescii* genomic DNA as template, in a final reaction volume of 25 µl, using AccuPrime™ Pfx DNA Polymerase from Invitrogen (Carlsbad, CA, USA) according to the manufacturer’s instructions. Primers are shown in Table 2.1.

*Restriction endonuclease digestion of DNA:* DNA samples were digested with restriction endonucleases, KpnI and XhoI, in their specific reaction buffers. Both restriction enzymes and the buffers were purchased from New England Biolabs (NEB, Ipswich, MA, USA). Digestions were carried out overnight (using 1U of enzyme/µg DNA) at 37°C in the presence of 0.1 mg/ml bovine serum albumin (BSA) as recommended by the supplier.

*Agarose gel electrophoresis:* DNA purification was performed using the StrataPrep gel extraction kit (Agilent Technologies, Santa Clara, CA, USA) and DNA was finally eluted in either elution buffer (10 mM Tris.Cl, pH 8.5) or in autoclaved distilled water.

*Ligation and transformation:* Digested and purified plasmids and inserts were set up for cohesive-end ligation using T4 DNA Ligase (NEB) in 1X buffer supplied. An insert: vector ratio
of 3:1 was used. Ligation reactions were incubated at 14-16°C for 18-20 h with 0.1 mg/ml BSA. For heat shock transformation, the ligation mix (10 μl containing ~ 20 ng DNA) was mixed with *E. coli* XL1 Blue-MRF’ (Agilent Technologies, Santa Clara, CA) and after heat shock at 42°C for 1 min, 1 ml Luria Broth medium was added into the transformed cells. This was followed by incubation at 37°C for 1 hr. Aliquots of the 50-100 μl cells was then plated on LB agar supplemented with an antibiotic ampicillin (100 μg/ml).

**Plasmid preparation for DNA sequencing:** Plasmid DNA for sequencing was purified using the StrataPrep Plasmid Miniprep Kit (Agilent). Purified DNA was sent for sequencing to the Macrogen sequencing facility (Macrogen, MD, USA) using T7 promoter primers.

**Design of the overexpression vector:** Three overexpression vectors for expression of full-length Cbes_1854(X-PL3), the catalytic domain only (PL3) and domain X was designed under the control of the T7 promoter using Novagen expression vector pET-45b. A plasmid map of the two constructs those were successfully expressed in *E. coli* is shown in Figure 2.1. Expression of recombinant proteins in *E. coli*: For expression of the proteins from the cloned genes in pET-45b-PL3 and pET-45b-X-PL3, the constructs were transformed into the *E. coli* strain BL21-Codon plus(DE3)-RIL strain. Cells were inoculated to a final concentration of 1 % in a fresh LB medium containing ampicillin (100 ug/ml) and were grown overnight to an OD600 of 0.6-0.7 at 37°C with shaking at 220 rpm. Production of recombinant proteins was induced with 1 mM IPTG.

**Heat treatment of cell free extract:** Cells expressing PL3 and X-PL3 domains were lysed by treatment with 1 mg/ml lysozyme and heated at 80°C for 30 minutes. Extracts were clarified by centrifuging at 14000 rpm for 30 min.

**Rapid screening of small expression cultures:** Expression of proteins PL3 and X-L3 in BL21-
Figure 2.1. Plasmid map for overexpression constructs. (A) Overexpression vector for full-length protein (X-PL3) and (B) overexpression vector for only catalytic domain (PL3).
Table 2.1: List of primers used in this study

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence (5’ – 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>X (Forward)</td>
<td>CGGGAATTCGGTAATGAAGTTGCAAGCGAGCG</td>
</tr>
<tr>
<td>X (Reverse)</td>
<td>CCGCTCGAGTTATGCTGACCGCTCACACTTTGAAATTAC</td>
</tr>
<tr>
<td>PL3 (Forward)</td>
<td>CGGGAATTCGACGGGTTTTAGTTATACAGAT</td>
</tr>
<tr>
<td>PL3 (Reverse)</td>
<td>CCGCTCGAGTTATGATTTGATGTCTGTGGATGAGCGG</td>
</tr>
</tbody>
</table>
Codon plus(DE3)-RIL cells (10 ml culture) were confirmed by affinity chromatography of heat-treated cell free extracts using the N-terminal 6x-His tag. Cells were induced with 1 mM IPTG for 16 h at 15°C for overnight. The cell pellet was collected and suspended in 20 mM sodium phosphate pH 7.0 containing 0.3 M NaCl (buffer A) and the cells were disrupted by the addition of 1 mg/ml lysozyme. Cell debris was removed by centrifugation at 10,000 rpm for 30 min. The Supernatant was heated at 80°C for 20 min, and, after removing denatured protein by centrifugation at 10,000 rpm for 30 min, the supernatant was applied onto a minispin HisTrap FF affinity column (GE Healthcare, USA) equilibrated with buffer A. After washing the column, the protein was eluted using imidazole (0.5 M) in buffer A.

**Protein expression and purification:** BL21-Codon plus(DE3)-RIL cells from a 1 liter culture were induced with 1 mM IPTG for 16 h at 15°C. The cell pellet was collected and suspended in buffer A and the cells were disrupted using a French press (800 psi). Cell debris was removed by centrifugation at 10000 rpm for 30 min. The supernatant was heated at 80°C for 20 min, and after removing denatured protein by centrifugation at 10 000g for 30 min, the supernatant was applied onto a His Trap FF affinity column (5 ml, GE Healthcare) equilibrated with buffer A. After washing the column, the PL3 module was eluted using an imidazole gradient (0–0.5 M) in buffer A. Based on SDS–PAGE analysis, fractions containing PL3 were combined, concentrated using ultrafiltration (Amicon, 10 kDa cutoff) and dialyzed against 20 mM phosphate of pH 7.0.

**Results and Discussion**

**Characterization of PL3 and X-PL3:** The full-length protein contains a PL3 family catalytic domain, which has pectate lyase activity, while the X domain shows some similarity with carbohydrate-binding domains present in several CAZy proteins [25]. Sequence alignment analysis of the X domain (39-222) predicts it to be a member of concanavalin A-like lectin superfamily, which binds with sugars, glycoproteins and glycolipids. Association of this X
domain with pectate lyase suggests that this is a putative pectin-binding domain. The X domain has also been proposed to be a member of cellulose binding domain family 4 CBM4_9 [39].

**Purification of PL3 and X-PL3**: X-PL3 was cloned from a thermophilic bacterium, therefore, it was expected that the X and PL3 domains, as well as full-size X-PL3, would tolerate high temperature. We found that the X domain was insoluble upon induction in *E. coli* at either 37°C or 18°C. We were unable to recover the X domain from the insoluble fraction. In contrast, full-length X-PL3 was soluble. This indicates that the presence of the PL3 domain, which is separated from the X domain by a long linker, mediates proper folding of the X domain as well as maintaining its stability. We speculate that solubilization of the attached X domain occurs due to specific domain interactions with the PL3 domain as shown for other CAZy enzymes [55]. The presence of a 44 amino acid long flexible linker (Figure 2.2) might play an important role in mediating domain interactions. In many cases, the presence of a linker sequence separating domains in a recombinant protein expressed in *E. coli* results in proteolysis of the linker region [47,56]. This can arise because of lack of glycosylation or phosphorylation of the linkers region by host *E. coli*. Consequently, in recombinant proteins linkers are very susceptible to intracellular proteolysis. It is important to note that we did not observe proteolysis of X-PL3.

The PL3 and X-PL3 proteins were not degraded after incubation at 70°C for 30 min as the intensities of their bands on the SDS gel were similar to those seen in unheated cell extracts (Figure 2.3 A). The proteins also retained 100% of their activity after heating of the cell extract, again confirmed that they were unaffected by heat treatment. These conditions were therefore used as a first heat-treatment step during the large-scale purification of each enzyme. Both enzymes contain an N-terminal 6xHis tag, which was used for one-step purification by affinity chromatography using a 5ml GE-HisFF column. Each protein bound to the column and eluted
MIKSKNKKEEVWVMSNRKILAIIVSIVSLFTGIGLRENVEAKAATLTDFFDGDNGRDI
WTSNGSWSVVDGSKVLSQASTGSEARAYTGGSSWSDYTVEAVKVLNVDSSS
GAVIVRYKNSGNYALVLRGSKIEIGKKNLNSWSTLAFSFTLDQDTWYNVKLE
NGSKLVYGYNNSQVLSADLSITTGKAGLIADRCVAEFDDVVVNSVSSTGAPTPPTTP
TSSVTPPTSTPTPTPTKTPPTPTSTPVTPQAPVPTPTPTPNTGGVLVTDTITIVKSGQTYDGKG
KIIAQQMGDGSQSENQKPIFKLEKGANLKNVIIGAGCDGIHYGVNVDNVVVWED
VGEDALTIVKSEGVEVIGGSKEAADKVFDQNLACPFTKVKNFTATNIGKLVRQNG
NTTFKVVIYLEDVTLNVKSCVAKSDSPVSELWYHNBLNVNNCKTLFEPSQSIQHY
Y
atgataaaactaagataaaagaggaggttttgtgtgtgtgatactgtagtcgtttgtagcagatgcggaagaggatggtgccattgtgctgagaattttctttctttacttttagttgtagtggtgtttcattgttacagggattgggttgccgaagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgtagtttgagttttatgtgtgtgtgtgtgtgtgtgagttttatg
Figure 2.2. The protein (upper) and DNA sequence (lower) of Cbes_1854 (X-PL3). The indicated residues are: residues 1-38, signal sequence (italics); Residues 39-222, X domain (bold); Residues 223-267, linker sequence (italics) and Residues 268-460, PL3 domain (bold). The primer sequences are underlined in the DNA sequence.
Figure 2.3. SDS PAGE gel of expressed proteins. (A) 1, protein marker; 2, heated cell-free extract of PL3; 3, unheated cell-free extract of PL3; Lane 4, unheated cell free extract of X-PL3 and Lane 5, heated cell-free extract of X-PL3. Lanes 1-4 depict the purification of PL3 domain and lanes 5-8 depict the purification of X-PL3. The lanes are (B) 1, flowthrough; 2, wash; 3-4, elution fractions; 5, flowthrough; 6, wash; 7-8, elution fractions and 9, protein marker.
Figure 2.4. Diagram of the three-dimensional structure of the *C. bescii* PL3 domain. Diagram was made using PyMol software using structure of PL3 domain (PDB: 3T9G) [59]. The calcium ion is in magenta.
with increasing concentration of imidazole. This indicates that the tags were readily accessible. After two steps of purification, X-PL3 and PL3 were homogeneous, as indicated by SDS gel electrophoresis (Figure 2.3). The molecular masses estimated from the SDS gel were similar to those calculated based on the deduced amino acid sequences (46 and 21 kDa for X-PL3 and PL3, respectively) suggesting that no proteolysis occurred. After purification from two liters of E. coli cultures, approximately 20 mg of each protein (PL3 and X-PL3) was obtained. The purified proteins were used for crystallization.

**Crystallization of PL3 domain:** The purified proteins were supplied to Dr. M. Himmel of National Renewable Energy Laboratory, CO, for structure studies. A 1.5-Å resolution X-ray structure of the catalytic module of C. bescii family 3 pectate lyase was obtained (PDB: 3T9G) [57]. The structure is similar to the previously solved structure of a family 3 pectate lyase from Bacillus sp. strain KSM-P15 (Pel15; PDB: 1EE6) [58]. The overall structure has β-helix fold, which is common for pectate lyases from families 1 and 3. The structural model for the *C. bescii* PL3 domain is shown in Figure 2.4. One calcium ion is bound with each chain in the crystal. Structural comparison with Pel15 suggests a strong structural similarity with a root-mean-square deviation of 0.93 Å. This structural similarity is significant considering that *C. bescii* is a thermophile and *Bacillus* sp. is a mesophile and that the sequence identity between PL3 and Pel 15 is only 53%. Notably, Pel15 from *Bacillus sp.* has an optimum temperature of 30°C while PL3 of *C. bescii* is expected to be optimally active and stable at the growth temperature of *C. bescii* (78°C), see chapter 3.

However our collaborators were unable to crystallize X-PL3 under any of the conditions used. This was not surprising, as the majority of multi-domain CAZy proteins with domains separated by even short linkers have not been crystallized. The only approach at present to
determine the structure of multi-domain proteins is to use protein modeling using structures of the individual domains [60].
CHAPTER 3
BIOCHEMICAL CHARACTERIZATION OF TWO DOMAINS OF CBES_1854, A FAMILY 3 PECTATE LYASE

Introduction

Cbes_1854 of the thermophilic bacterium C. bescii is a family 3 pectate lyase and belongs to subfamily 1, as predicted by amino acid sequence analysis. In addition to the C. bescii enzyme, there are three other pectate lyases in subfamily 1, from the mesophilic bacteria Bacillus sp. KSM 15, Bacillus subtilis subsp. and Penibacillus barcinonensis BP 23 (Table 3.1). Among these, the structures of the enzyme from Bacillus sp. KSM P15 and Bacillus subtilis subsp. are available. It is assumed that family 3 pectate lyase have the same catalytic mechanism as that of family 1 pectate lyase but this has not been confirmed structurally [58]. This assumption is based on the fact that both families share the same β-helix fold. The goal of this work was the biochemical characterization of C. bescii X-PL3 and PL3 including determining their pH optima. It was expected that they would be similar to those of family 1 pectate lyases, which prefer alkaline pH. The other goals were to (A) determine the temperature optimum of both enzymes; (B) evaluate the substrate specificity of X-PL3; (C) reveal the catalytic mechanism through a collaborative effort by structure determination with bound substrate and (D) determine if this enzyme works synergistically with another cellulolytic enzyme of C. bescii, cellulase A. This is the most abundant cellulolytic enzyme in extracellular fraction of C. bescii [46].

Materials and Methods

Protein sample preparation: The PL3 and X-PL3 proteins were obtained by expression and one
**Table 3.1: Comparison of different family 3 pectate lyases**

<table>
<thead>
<tr>
<th>Protein Name</th>
<th>Organism</th>
<th>Temp. opt.(°C)</th>
<th>pH opt.</th>
<th>Subfamily</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pectin lyase (Ply; PlyAI4)</td>
<td><em>Bacillus sp. I4</em></td>
<td>50</td>
<td>10.5</td>
<td>3</td>
<td>[61]</td>
</tr>
<tr>
<td>Pectate lyase (Pel15)</td>
<td><em>Bacillus sp. KSM15</em></td>
<td>50-55</td>
<td>10.5</td>
<td>1</td>
<td>[62]</td>
</tr>
<tr>
<td>Pectate lyase C (PelC)</td>
<td><em>Bacillus subtilis subsp.</em></td>
<td>65</td>
<td>10</td>
<td>1</td>
<td>[63]</td>
</tr>
<tr>
<td>Pectate lyase I (PelI)</td>
<td><em>Dickeya dadantii 3937</em></td>
<td>37</td>
<td>9</td>
<td>5</td>
<td>[64]</td>
</tr>
<tr>
<td>Pectate lyase A (PelA)</td>
<td><em>Penibacillus barcinonensis BP-23</em></td>
<td>50</td>
<td>10</td>
<td>1</td>
<td>[65]</td>
</tr>
<tr>
<td>Pectate lyase B (PelB)</td>
<td><em>Pectobacterium carotovorum SCC3193</em></td>
<td>37</td>
<td>9.5</td>
<td>5</td>
<td>[66]</td>
</tr>
<tr>
<td>Pectate lyase (Pel-3)</td>
<td><em>Pectobacterium carotovorum subsp.cart .71</em></td>
<td>37</td>
<td>ND</td>
<td>5</td>
<td>[67]</td>
</tr>
<tr>
<td>Endo Pectate lyase I (PL 1)</td>
<td><em>Streptomyces thermocarboxydus B-1</em></td>
<td>50</td>
<td>9</td>
<td>4</td>
<td>[68]</td>
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<tr>
<td>Pectate lyase (X-PL3)</td>
<td><em>C. bescii</em></td>
<td>85</td>
<td>9</td>
<td>1</td>
<td>This work</td>
</tr>
</tbody>
</table>

ND: not determined
Pel15 and PelC have three-dimensional structure.
step purification as described in chapter 2. Samples were prepared by dialysis of protein from the Ni-NTA elution fractions against 20mM Tris buffer pH 8.0 containing 100 mM KCl. The protein was quantified by absorption at 280 nm, using the calculated extinction coefficient of the PL3 and X-PL3 shown in Table 3.2. Some other properties based on amino acid sequence was predicted by Vector NTI software are also shown in Table 3.2 [69].

Pectate lyase assay: A substrate stock of 0.25% polygalacturonic acid (PGA) was prepared in 100 ml water and the pH adjusted 9.0 by the addition of NaOH pellets. For the assay, 100 mM N-cyclohexyl-3-aminopropanesulfonic acid (CAPS) buffer containing 1mM CaCl₂ at 80°C was used, unless otherwise indicated. The amount of product was measured by the absorption at 235 nm, using a molar extinction coefficient of 4600 M⁻¹cm⁻¹. One unit of enzyme activity is defined as the amount of enzyme that produces 1 µmol of product /min [70].

To determine the pH dependence of the enzyme the same amount were added to different buffers, ranging from pH 4.0 to 11.0, and were incubated with 0.1% PGA for 3 min at 80°C. Data was collected at one-minute intervals to calculate the slope, which was used to calculate the specific activity. To determine the temperature profile, the same assay protocol was used at pH 9.0 but the temperature range used was between 40 and 90°C. For the comparative studies the results are expressed as relative activity (in %).

The optimum concentration of calcium for activity was determined using the same protocol. The substrate specificity was determined using 0.1% of all of the substrates listed in Table 3.3. The thermal stability of the protein was determined by incubating it at different temperatures for up to 48 hours. The thermal stability was assessed at pH 9.0 and pH 7.0. At different time intervals samples were removed and assayed at 80°C.
Table: 3.2. Predicted biophysical properties of the catalytic domain (PL3) and of the holoenzyme form (X-PL3) of Cbes_1854 from *C. bescii*

<table>
<thead>
<tr>
<th>Properties</th>
<th>X-PL3</th>
<th>PL3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molecular weight (kDa)</td>
<td>46,178</td>
<td>20,023</td>
</tr>
<tr>
<td>Isoelectric point (pI)</td>
<td>6.32</td>
<td>6.29</td>
</tr>
<tr>
<td>Molar extinction coefficient (A$_{280}$)</td>
<td>55,265</td>
<td>18,500</td>
</tr>
</tbody>
</table>
**Results and Discussion**

The pH, temperature and calcium dependence of the pectate lyase activity of PL3 and X-PL3 were determined using polygalacturonic acid as the substrate. The assay is based on measurement of the unsaturated reaction product, hexenuronic acid, formed after cleavage of the α 1-4 glycosidic linkage of PGA. Cleavage follows β elimination reaction and results in the unsaturated product, hexenuronic acid (Figure 3.1).

As shown in Figure 3.2 calcium ions were indispensable for the activity of both X-PL3 and PL3 with an optimum concentration of 1mM Ca++. Both enzymes also display pH optima of 9.0. At pH 9.0, the optimum temperature for the activity is between 85 and 90°C. As shown in Figure 3.3, the half-life for both, PL3 and X-PL3, at 90°C is approximately 1 hr but at pH 7.5 this increases to 5 hr and 2 hr for PL3 and X-PL3, respectively. These data suggests that X-PL3 is less thermostable than PL3. To determine the specificity of X-PL3, different types of esterified forms of citrus pectin were used as substrates (Table 3.3). It was found that X-PL3 shows the highest activity with 90% esterified citrus pectin, as shown in Table 3.4. No significant activity was observed when rhamnogalacturonan was used as substrate, although X-PL3 still shows comparable amount of activity using non-esterified polygalacturonan as a substrate.

The biochemical properties of enzymes from the PL3 family (Table 3.1) suggest that these enzymes have the same catalytic mechanism as that of family 1 pectate lyases (PL1). The catalytic mechanism of *Bacillus subtilis* pectate lyase (BsPel), the closest homolog to X-PL3 and a member of PL1, has been studied in detail. BsPel has two basic residues, lysine and arginine, as part of the active site where arginine acts as a catalytic base and abstracts a proton. The role of the arginine residue has been established by mutation [27]. As shown in Figure 3.1, the proposed
The detail descriptions of the substrates are present in following references [70-72].
Figure 3.1. Schematic diagram of the typical β-elimination cleavage of polygalacturonic acid. P⁺, Ca⁺²; B, catalytic base that abstracts the proton from C-5; and A-H, does protonation to the glycosidic oxygen. The unsaturated product is Hexenuronic acid and the diagram is taken from reference [26].
Figure 3.2. Temperature, pH and calcium (Ca^{++}) dependence of pectate lyase activity of PL3 and X-PL3. (A) Specific activity profile at pH 9.0; (B) Calcium requirement for pectate lyase activity; (C) pH profile and (D) Temperature profile. All assays were carried out using 0.1% polygalaturonic as the substrate.
Figure 3.3. Thermostability of PL3 and X-PL3. These plots depict the activity and thermal stability relationship between X-PL3 (A, C) and PL3 (B, D) at pH 9.0 (A, B) and 7.5 (C, D). The temperature range for measurement of thermal stability is between 60-90°C, as shown in D.
catalytic mechanism of BsPel is an anti-β elimination reaction that requires a basic residue to abstract the proton from the C5 atom of the substrate that has been charge stabilized by the two catalytically important Ca$^{2+}$ ions, together with the concerted action of an acidic residue that donates a proton to the carbonyl group, creating an enol intermediate. This is followed by elimination of the leaving group together with protonation of the O4 atom by another acidic residue or a water molecule acidified by the two catalytically important Ca$^{2+}$ ions, while the Ca$^{+2}$ atoms stabilize the enol intermediate [27,73].

In our collaborative effort, the three-dimensional structure of the catalytic domain (PL3) of *C. bescii* pectate lyase was obtained with the substrate trigalacturonic acid bound to the enzyme [74]. A comparison of catalytic residues in PL3 and BsPel, from family PL1, revealed that there is Lys 130 in the catalytic domain of PL3 in place of the catalytic residue Arg 279 of BsPel. The evidence for this comes through the substrate-bound structure of the PL3 domain (PDB: 4EW9) at 1.6-Angstrom resolution [74]. Figure 3.4 shows a diagram of the catalytic domain of PL3 with trigalacturonic acid bound. In this three-dimensional structure, Lys 130 is not correctly positioned to abstract the proton from C-5, rather Lys 108 is better positioned for proton abstraction in the active site (Figure 3.4). Therefore, it is likely that *C. bescii* PL3 uses this Lys 108 as a catalytic residue rather than Lys 130. This observation supports a different catalytic mechanism for the family 3 pectate lyase. Based on this, our collaborators have proposed a novel reaction mechanism based on Lys 108 as a catalytic base for the family 3 pectate lyase [74].

Based on the biochemical results presented here, the unknown domain ‘X’ does not provide any advantage to the catalytic PL3 domain in either degrading polygalacturonic acid or providing protein stability. From the NCBI-BLAST of just the ‘X’ domain against the non-
Table 3.4: Activity of X-PL3 with different substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Sp. Activity (U/mg)</th>
<th>Activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polygalacturonic acid</td>
<td>225 ± 23</td>
<td>45 ± 10</td>
</tr>
<tr>
<td>Apple pectin</td>
<td>394 ±13</td>
<td>78 ± 3</td>
</tr>
<tr>
<td>Citrus pectin (mixed)</td>
<td>385 ±16</td>
<td>76 ± 4</td>
</tr>
<tr>
<td>Rhamnogalactomannan</td>
<td>6 ± 1</td>
<td>1 ± 0.5</td>
</tr>
<tr>
<td>Pectic galactan</td>
<td>10 ± 2</td>
<td>2 ± 20</td>
</tr>
<tr>
<td>Citrus pectin (30%)</td>
<td>274 ±34</td>
<td>54 ± 12</td>
</tr>
<tr>
<td>Citrus pectin (60%)</td>
<td>362 ±14</td>
<td>72 ± 4</td>
</tr>
<tr>
<td>Citrus pectin (90%)</td>
<td>506 ±15</td>
<td>100 ± 3</td>
</tr>
</tbody>
</table>

One unit is 1 μmol of unsaturated galacturonic acid product /min.
redundant protein database, this domain was not found in archaea and is mainly found associated with the catalytic domain of the pectate lyases in different *Caldicellulosiruptor* species. In some cases it was found with a domain of glycosyl hydrolase family 28. GH28 is also proposed to function as a pectin-degrading enzyme. The association of domain X with pectin-degrading enzymes suggests that domain ‘X’ might functions as a pectin-binding domain.

During these studies it was found that the recombinant form X-PL3 and PL3 tended to aggregate at higher temperature (85°C) if the protein concentrations were above ~0.2mg/ml. This was not expected for a protein of thermophilic origin and suggests that glycosylation of the linker region of X-PL3 may have a positive role in bridging the interaction the domain ‘X’ and its catalytic domain. The linker region provides sites for O-glycosylation but this does not occur in recombinant proteins expressed in *E. coli*. These data suggest that glycosylation plays an important role in the protein stability at higher temperature.

Another objective of the present study was to provide purified pectate lyase in order to study of enzymatic degradation of biomass. The essential role of cellulases in biomass degradation has been well established but it is not known if other extracellular enzymes, such as pectate lyases, could enhance the biomass degradation. In collaboration studies with Dr. M. Himmel of National Renewable Energy Laboratory, CO, it was shown that the presence of *C. bescii* PL3 increases the amount of sugar released from untreated switchgrass biomass in comparison to using *C. bescii* cellulase A alone. In fact, the presence of PL3 addition reduces the load of cellulase A by ~33% [74]. Therefore, adding *C. bescii* pectate lyase to cellulases appears to be a promising approach to increase the efficiency of enzymatic biomass degradation.
Figure 3.4. Diagram of the catalytic domain PL3 with the reaction product trigalacturonic acid. Reaction products are marked and labeled (PDB: 4EW9) [74]. The catalytic residue, Lys 108, is in bold and labeled. The diagram was prepared using PyMol software [59].
SUMMARY

Caldicellulosiruptor bescii is the most thermophilic bacterium known with potential for plant biomass degradation. Remarkably, it grows on switchgrass biomass without any chemical treatment at 78°C [37,43]. C. bescii possesses a variety of enzymes with carbohydrate active domains (CAZy enzymes) that are potentially involved in the conversion of insoluble switchgrass into soluble fermentable sugars [43]. Transcriptional analyses after growth of C. bescii on glucose and switchgrass revealed that the genes encoding 94 enzymes were up-regulated and therefore apparently involved in switchgrass degradation. Among these 94 enzymes, 18 have a proposed functional role in carbohydrate metabolism. These include four potential pectinolytic enzymes, three of which are predicted to be extracellular (Cbes_1853, Cbes_1854 and Cbes_1855) [39]. Interestingly, Cbes_1854 is the most highly up-regulated enzyme during growth on switchgrass versus glucose. This was unexpected because pectin represents less than 5% of the total polysaccharides in plant biomass. Therefore, Cbes_1854 was chosen as a model multidomain enzyme for overexpression, purification and biochemical studies.

Cbes_1854 belongs to family 3 pectate lyases, of which there is no structure available for the catalytic domain with a bound substrate. In addition to biochemical studies, we also sought to provide the pure enzyme for the structural analysis so that information could be obtained on the mechanism of family 3 pectate lyases. Cbes_1854 encodes two domains. The X domain is of unknown function while PL3 is the catalytic domain. The role of the X domain is probably to bind the insoluble pectin-rich component of the complex biomass. Three constructs were cloned
and expressed separately in *E. coli*: domain X, catalytic domain (PL3) and full-length protein having both domains (X-PL3). Domain X was found in the insoluble fraction after expression in *E. coli* and attempts to recover the protein were not successful. In contrast, the other two proteins remained in the soluble fraction and could be purified using the 6X-His tag present at the N-terminus by affinity chromatography. The yields of each of these two proteins were approximately 10 mg/liter of *E. coli* culture.

Biochemical studies of PL3 and X-PL3 using polygalacturonic acid (PGA) as a substrate showed that both enzymes have similar alkaline pH optima (pH 8-9) and temperature optima (85-90°C). This observation also highlights the fact that under *in vitro* assay conditions domain X does not influence the catalytic properties of the PL3 domain. Interestingly, it was observed that the catalytic domain is more active against 90% esterified citrus pectin than it is with unesterified PGA. PL3 and X-PL3 are thermostable and each has a half-life of 1 h at 90°C at pH 9.0. Although the protein stability increases at physiological pH (pH 7.0), they retain only 20-30% of total activity at this pH value.

The three-dimensional structure of catalytic domain PL3 was obtained in a collaborative study and this has been deposited (PDB: 1EE6). It consists of a parallel β-helix fold. The structure with product of trigalacturonic acid, 4, 5-unsaturated digalacturonic acid and D-galacturonic acid, was also obtained and deposited (PDB: 4EW9). Surprisingly, unlike the previously described structure of a pectate lyase, that of BsPel, a homolog with same structural fold belonging to PL1 family, the PL3 catalytic module has only one basic residue, lysine, rather than two conserved basic residues lysine and arginine, present in active site. This suggests that PL3 has different catalytic mechanism from what was observed with BsPel.

To probe the role of Cbes_1854 in biomass degradation, untreated switchgrass biomass
was incubated with the most abundant cellulolytic enzyme, Cel A, from *C. bescii* supplemented with recombinant PL3. The presence of PL3 reduced the load of Cel A by ~33% while maintaining the same amount of sugar released from biomass. This demonstrates the synergistic effect of pectinolytic enzymes in biomass degradation. In the future, recombinant PL3 and X-PL3 have the potential to be used in enzyme mixtures to increase the efficiency of biomass degradation.
REFERENCES


