The Crystal Structure of Pectate Lyase Pell from Soft Rot Pathogen Erwinia chrysanthemi in Complex with Its Substrate*\(^{\text{S}}\)

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The crystallographic structure of the family 3 polysaccharide lyase (PL-3) Pell from Erwinia chrysanthemi has been solved to 1.45 Å resolution. It consists of an N-terminal domain harboring a fibronectin type III fold linked to a catalytic domain displaying a parallel β-helix topology. The N-terminal domain is located away from the active site and is not involved in the catalytic process. After secretion in planta, the two domains are separated by E. chrysanthemi proteases. This event turns on the hypersensitive response of the host. The structure of the single catalytic domain determined to 2.1 Å resolution shows that the domain separation unveils a “Velcro”-like motif of asparagines, which might be recognized by a plant receptor. The structure of Pell in complex with its substrate, a tetragalacturonate, has been solved to 2.3 Å resolution. The sugar binds from subsites −2 to +2 in one monomer of the asymmetric unit, although it lies on subsites −1 to +3 in the other. These two “Michaelis complexes” have never been observed simultaneously before and are consistent with the dual mode of bond cleavage in this substrate. The bound sugar adopts a mixed 2\(^1\) and 3\(^1\), helical conformation similar to that reported in inactive mutants from families PL-1 and PL-10. However, our study suggests that the catalytic base in Pell is not a conventional arginine but a lysine as proposed in family PL-9.

Polysaccharide lyases (EC 4.2.2.x) are polysaccharide-degrading enzymes that cleave glycosidic bonds of C\(_2\) uronic acid polymers. They play a central role in the recycling of plant material and are potent virulence factors of plant pathogenic bacteria and fungi. In contrast to the 111 sequence-derived families of glycoside hydrolases, polysaccharide lyases have been grouped into only 18 families in the CAZy data base (1). Five polysaccharide lyase families (PL-1, 2, 3, 9, and 10)\(^{4}\) contain pectate lyases (EC 4.2.2.2 and EC 4.2.2.9). These enzymes cleave polymeric α-1,4-linked galacturonic acid (GaLA) within the pectate component of the cell wall by β-elimination mechanism, leaving an unsaturated C\(_4\)–C\(_5\) bond at the newly formed non-reducing end. The activity is maximal in the pH range 8.5–10.5 and is metal ion-dependent; Ca\(^{2+}\) is the general cofactor except for the members of the family PL-2, which depend on Co\(^{2+}\), Mn\(^{2+}\), and Ni\(^{2+}\) (2, 3). The divalent cation mediates enzyme-substrate interaction by binding between the protein and the sugar (4). Various pectate lyases prefer either polygalacturonic acid or partially methylated pectin as substrate (5, 6).

Atomic structures have been determined for representatives of all polysaccharide lyase families: PL-1 (7–10), PL-2 (3), PL-3 (11), PL-9 (12), and PL-10 (13, 14). They reveal three topologies for the catalytic module: 1) a right-handed parallel β-helix fold common with the families PL-1, PL-3, and PL-9, first observed for family 1 pectate lyase PelC from Erwinia chrysanthemi (7); 2) an (α/α)\(_2\) toroid in family 2, recently described in pectate lyase YePel2A from Yersinia enterocolitica (3); 3) an (α/α)\(_3\) toroid in family PL-10, first reported for the catalytic module of the pectate lyase Pe110A from Cellulvibrio japonicus (14). The superhelix fold is also found in family PL-6 and in the glycoside hydrolase family GH-28, if one considers only enzymes cleaving glycosidic bonds (15–17). The α-barrel folds from families PL-2 and PL-10 are unique among polysaccharide lyases and have limited similarities to a few glycoside hydrolases (18). Interestingly, the structures of inactive mutants from family 1 PelC and family 10 Pel10Acm in complex with oligogalacturonate and Ca\(^{2+}\) show a similar pattern of enzyme-substrate interactions in their active sites (4, 14). To a certain extent, this observation could be extended to family 2 YePel2A complexed with trigalacturonate but without a metal cation (3). Such local structural equivalence suggests a common β-elimination mechanism among pectate lyases regardless of the topology of the protein. The catalytic reaction is initiated by an essential basic amino acid.  

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\(^{1}\) The atomic coordinates and structure factors (code 3B4N, 3B90, and 3B8Y) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

\(^{2}\) The on-line version of this article (available at http://www.jbc.org) contains two supplemental tables.

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\(^{4}\) The abbreviations used are: PL, polysaccharide lyase; GaLA, tetragalacturonic acid; Fn, fibronectin; GH, glycoside hydrolase; MES, 4-morpholinethanesulfonic acid; r.m.s.d., root mean square deviation.

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acid that abstracts the proton from the carbon C5. This residue appears to be an arginine in families PL-1 (7–10), PL-2 (3), and PL-10 (13, 14) and a lysine in family PL-9 (12). Whether a lysine or an arginine is the catalytic base for family PL-3 remains to be determined (11).

The enterobacterium *E. chrysanthemi* is a causative agent of soft rot disease in a wide variety of plants. Among a set of pectin-depolymerizing enzymes produced by *Erwinia*, pectate lyases are the major pectinases and play a key role in plant tissue maceration. The pectate lyase Pell belongs to family PL-3 and is the only known example from the *E. chrysanthemi* pectinases that comprises two functional modules, termed the N-terminal domain (residues 20–116) and the catalytic domain (residues 117–344) (6). The close homologue Pell-3 from *Erwinia carotovora* (19) has a similar N-terminal extension, suggesting analogous protein architecture.

Pell possesses a 19-residue signal peptide that is cleaved during export by the Sec system, following which the protein is secreted into the external medium by the type 2 secretion system (20). The two structural modules are separated in planta by *E. chrysanthemi* proteases (6). The resulting catalytic domain (residues 117–344) possesses similar enzymatic properties *in vitro* to the full-length protein. However, in contrast to the full-length Pell, it elicits a necrotic reaction associated with an active defense by plants. This hypersensitive response is characteristic of a set of effector proteins secreted by many plant pathogenic bacteria by the type 3 secretion systems (21).

The catalytic domain of Pell shows a sequence identity of 26% with the unique structural representative of the family PL-3, the alkaline pectate lyase Pel-15 of *Bacillus sp.* strain KSM-P15 (11). This protein exhibits the classical superhelical fold, and its putative active site is located in a conserved cleft that stretches along the external core of the superhelix.

Here, we present the crystallographic structures of full-length Pell and that of the single catalytic domain termed Pellcat, in order to visualize the rearrangements that likely occur *in planta* after maturation. Furthermore, the associated structure of Pell in complex with a polygalacturonate fragment (a tetragalacturonate GalA₄) provides new insight into the catalytic mechanism.

**MATERIALS AND METHODS**

*Protein Purification*—PelI was expressed in *Escherichia coli* BL21 (DE3) (Stratagene) cells carrying a high expression plasmid with the *pelI* gene and purified to homogeneity as described before (22). The module Pellcat was obtained by treatment of pure full-length Pell with the *E. chrysanthemi* protease PrtC (6). Pellcat was then separated from the N-terminal domain by chromatography on Protein-Pak CM 8HR (Waters) column equilibrated with 20 mM sodium acetate, 0.1 mM EDTA, pH 5.0 buffer. The protein was eluted with a 0–0.5 M NaCl linear gradient in the same buffer and then concentrated to 15 mg/ml in a Centricon 10 (Amicon) in 10 mM Tris-HCl with 0.1 mM CaCl₂.

*Construction of Mutant Proteins and Enzyme Assay*—Single amino acid mutations were introduced in the Pell sequence by site-directed mutagenesis using the QuikChange kit (Stratagene). The primers used are PellK224R (5’-ggcgggaaacggagagtggctgcagcacattc-3’), PellK224R (5’-tcgggtaaacagcagagtttgctcctgtggcaggg-3’), PellIR252K (5’-ggtgaacagggaaatgaggtcctggcagctgcc-3’), and the corresponding reverse complementary primers (mutated bases are in bold). The nucleotide sequences of mutant genes were checked (Genome express). The mutant proteins were produced and extracted from *E. coli* BL21 cells as described above for the wild type Pell. Their amounts were estimated by immunoblotting with Pell antibodies, and then equivalent quantities of each mutant protein and the wild type Pell were used in enzyme assay. Pectate lyase activity was measured spectrophotometrically by monitoring the formation of unsaturated products from pectin. The assay mixture contained 50 mM Tris-HCl, pH 8.5, 1 or 3 mM CaCl₂, and 0.5 g-liter⁻¹ pectin with a degree of methylation of 45%. The appearance of unsaturated products was monitored at 37 °C over a period of 30 s with 6-s intervals and used to calculate enzyme activity as described (6).

*Crystallization*—Crystals of Pell were grown as reported previously (22). They belong to space group P2₁ with unit-cell parameters *a* = 61.6 Å, *b* = 70.7 Å, *c* = 73.4 Å, β = 112.8° and contain two molecules in the asymmetric unit. Crystals of Pellcat were obtained in the same conditions, i.e. by the vapor diffusion method from a solution containing 10 mM zinc sulfate, 100 mM MES, pH 6.5, 25% polyethylene glycol 550 (w/v) at a temperature of 292 K. A 1:1 ratio of protein to reservoir solution was used. Crystals belong to space group P2₁₂₁, and have cell dimensions *a* = 45.4 Å, *b* = 62.6 Å, *c* = 128.1 Å. The calculated *V*ₘᵢₙ = 1.9 Å³/Da for two molecules in the asymmetric unit.

The structure of the Michaelis complex was obtained by soaking a crystal of Pell for 24 h in the crystallization buffer depleted in zinc sulfate and with increasing concentrations of GalA₄ (1, 2, 3 mM). These successive soakings were performed in calcium-free solutions at pH 6.5 to avoid enzymatic reaction. The soaking experiment induces a significant modification of unit cell parameters up to *a* = 61.2 Å, *b* = 68.7 Å, *c* = 73.4 Å, β = 113.9°.

*X-ray Diffraction Experiments and Structure Determinations*—Before data collection, all crystals were rapidly transferred into a cryoprotectant solution consisting of the crystallization solution plus 15% (v/v) ethylene glycol. Synchrotron data were collected at 100 K at European Synchrotron Radiation Facility beamlines, Grenoble, France. The programs XDS and XSCALE were used for data reduction and scaling (statistics in Table 1). 5% of randomly selected reflections were kept apart for cross-validation, and the crystallographic refinement was performed with CNS using the method of slow cooling simulating annealing (23). After each cycle, the model was manually improved using the graphic program TURBO FRODO (24). Final statistics are presented in Table 1. The quality of the refined structures was assessed with PROCHECK (25).

*Pell*—The first model of Pell was obtained at 1.6 Å resolution by single wavelength anomalous dispersion phasing of a gold derivative (22). A native structure was subsequently determined to 1.45 Å resolution from synchrotron data collected at 100 K at a wavelength of 0.94 Å at beamline BM30A. The crystallographic refinement yields an *R*-factor of 16.8% (*R*_free 19.5%). The model contains two monomers, termed A and B. Residues 108–118 were not visible in electron density maps.
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TABLE 1
Data collection and refinement statistics

<table>
<thead>
<tr>
<th>Data collection</th>
<th>Pell</th>
<th>Pell-GalA4</th>
<th>Pell-GalA5</th>
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<td>ID14–3, ESRF</td>
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<td>P2₁</td>
<td>P2₁,2,2</td>
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<td>61.2</td>
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<td>b (Å)</td>
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<td>62.6</td>
</tr>
<tr>
<td>c (Å)</td>
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<td>73.4</td>
<td>128.1</td>
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<td>2.3 (2.4–2.3)</td>
<td>2.1 (2.5–2.1)</td>
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<td>72,076 (7,117)</td>
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<td>91.9 (77.6)</td>
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<td>R_cryst/R_free (%)</td>
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<td>10.8 (18.6)</td>
</tr>
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<td>10.6 (5.4)</td>
<td>18.4 (5.6)</td>
<td>8.0 (5.3)</td>
</tr>
</tbody>
</table>

Crystallographic refinement

Asymmetric unit content (2 monomers) (2 monomers) (2 monomers)
Number of protein atoms (non-hydrogen) 4,746 4,500 3,330
Number of water molecules 752 171 325
Cations 2 Ca²⁺, 6 Zn²⁺ 1 Ca²⁺, 1 Zn²⁺ 2 Ca²⁺, 4 Zn²⁺
Other non-hydrogen atoms 33 98 10
Overall B-factor (Å²) 17 19.8 21.0
Ramachandran plot (allowed) 533 (99.1%) 505 (99.4%) 371 (99.2%)
Ramachandran plot (generously allowed) 5 (0.9%) 3 (0.6%) 3 (0.8%)

According to geometric criteria and to the highest peaks in the 2Fo – Fc maps, five Zn²⁺ and two Ca²⁺ ions substitute the seven gold ions observed in the derivative structure. Zn²⁺ ions were further identified by collecting synchrotron data around the zinc K edge and calculating anomalous difference Fourier maps. An extra Zn²⁺ ion was positioned in the final structure as well as one SO₄²⁻, seven ethylene glycol, and 752 water molecules.

Pell_cata — Data were collected to 2.1 Å resolution at beamline ID14–3. The structure was solved by molecular replacement using the program AMoRe (26) and a two-body search taking as reference the structure of full-length PelI (residues 117–344). The first two residues of Pell_cata (residues 117–118) are not visible in electron density maps as in the Pell structure. The final R-factor is 20.4% (R_free 27.1%). Two Ca²⁺ and four Zn²⁺ ions are observed at positions similar to those of Pell. Two SO₄²⁻ and 325 water molecules have been placed in the asymmetric unit.

Pell-GalA₄ — Data were collected to 2.3 Å resolution at a wavelength of 0.93 Å at beamline ID14–3. Attempts to solve the structure with the complex by a crystallographic rigid body refinement were unsuccessful due to non-isomorphous crystals (shift in β angle). The structure was solved by molecular replacement using the program AMoRe (26) and a two-body search based on the substrate-free structure. Fo – Fc maps were inspected after crystallographic refinement using CNS. The tetrasaccharide bound to molecule A is unambiguously defined in electron density, whereas the one bound to molecule B is weakly defined at its non-reducing end. The four sugar rings of the two ligands were built and oriented in the electron density. No electron density was found for segments 148–150 and 216–222 from monomer A and segments 77–81, 145–150, and 215–222 from monomer B. These amino acids were deleted from the refined structure as well as one Ca²⁺ and five Zn²⁺ ions. Further cycles of crystallographic refinement yield an R-factor of 20.0% (R_free 27.9%). The final model contains two GalA₄, one Ca²⁺ ion, one Zn²⁺ ion, and 171 water molecules.

Structure Analysis — Secondary structure elements were assigned according to main-chain hydrogen bonds and ψ/φ angles calculated with STRIDE (27). Pair-wise structure comparison was accomplished with Dalili (28). Images of sequence alignments were prepared using ESPript/ENDscript (29). Interactions between tetragalacturonate and Pell were examined with LIGPLOT (30). Other images were generated with Molscript/Bobscript (31, 32). Atomic coordinates and structure factors of Pell, Pell_cata, and Pell-GalA₄ have been deposited in the Protein Data Bank (PDB) under the codes 3B4N, 3B9O, and 3B8Y, respectively.

RESULTS AND DISCUSSION

Overall Fold of Pell — The structure of full-length Pell has been determined to 1.45 Å resolution. The protein crystallizes with two monomers, A and B, in the asymmetric unit but is monomeric in solution according to gel filtration experiments (not shown). The two monomers A and B are very similar, and their Cα traces can be superimposed with a root mean square deviation (r.m.s.d.) of 0.4 Å.

Pell is composed of a fibronectin type III domain (Fn3D) at its N terminus (residues 20–107) fused to a catalytic domain that displays a parallel β-helix fold (residues 119–344). The connecting segment where cleavage by E. chrysanthemi proteases occurs (residues 108–118) (6) was not observed in the density and is not modeled. This linker is also predicted as natively disordered by the Regional Order Neural Network server (33), and the crystallographic packing was carefully examined to identify the biological monomer.

The more plausible arrangement of entire protein is presented in Fig. 1A. It possesses the largest buried interface between the two domains, 750 Å², as well as the shortest dis-
FIGURE 1. Structural alignment of the catalytic domain of Pell. A, superimposition of the catalytic domain of Pell (blue) with Pell15 (red) (Protein Data Bank code 1EE6) and PellC (green) (Protein Data Bank code 2EWE). Bound calcium ions are shown in Pell, Pell15, and PellC with blue, red, and green spheres, respectively. The bound sugar in PellC is shown in yellow. The disordered linker in Pell is shown by a dashed line. B, structure-based sequence alignment with secondary structure elements. Top, red and blue triangles indicate the putative catalytic residue of Pell and PellC, respectively. Bottom, green numbers, red stars, and blue circles indicate Pell disulfide bridges, residues at the interdomain interface (contact distances < 3.2 Å), and residues coordinated to a Ca^{2+} ion, respectively.
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tance between residues 107 and 119, 23 Å. Such spacing agrees with the 11 missing residues. The occluded face of the catalytic domain is also partially covered in numerous polysaccharide lyases by N-terminal or C-terminal extensions. Moreover, this arrangement between domains evokes the one recently described in a modular bacterial glycoside hydrolase YeGH28 (Protein Data Bank code 2UVE) (16). The Fn3 domain is bound to the opposite of the active site in both proteins but is oriented perpendicularly to the β-helix axis in Pell and parallel in YeGH28.

Fn3 Domain—The N-terminal domain of Pell exhibits a conventional seven-stranded Fn3 fold with a Greek key motif composed of four β-strands termed βC, βC’, βF, and βG (Fig. 1). As reported in some other animal and bacterial Fn3Ds, the last strand βG is split into two short strands (34). The Fn3D does not contain any disulfide bonds and interacts with the catalytic domain via its flat βA, βB, βE, façade (Fig. 1A). Two hydrophobic residues, Met22 and Tyr23, give extensive contacts at this interface, whereas polar residues mediate hydrogen bonds (Ser27, Ser38, and Thr72, Fig. 1B).

A structural search performed with the DALI server shows that this N-terminal domain superimposes with the best match on an Fn3 module of a neural cell adhesion molecule (r.m.s.d. of 2 Å for 85 aligned Cα with an amino acid identity of 13%) (Protein Data Bank code 1CFB) (35). It superimposes also extremely well on the Fn3D of the modular YeGH28 polygalacturonase (16), which includes an additional helix-loop-helix motif that further stabilizes the interdomain interface (r.m.s.d. of 2 Å for 79 aligned Cα and a sequence identity of 11%).

Fn3Ds are found in many eukaryotic proteins, where they are frequently implicated in cell adhesion and signaling processes (36). They display a variety of binding modes with other proteins, especially via their BC, CE, and FG loops (37, 38). Fn3Ds have also been reported in bacterial carbohydrate active enzymes, where they may act as spacers between sugar-binding and catalytic domains (34). It has also been suggested that they can help binding of sugar groups or promote hydrolysis (39, 40). However, the distinctive features of carbohydrate binding modules such as a large aromatic platform or a bound Ca2+ (41) are not detected on the Fn3D surface of Pell. Moreover, Pell displays the same enzyme activity on pectin regardless of the presence of its N-terminal domain (6). In addition, following crystal-soaking experiments, a pectin fragment, tetragalacturonate GalA5, has been detected only within the catalytic domain, but not in the Fn3D of Pell (see below).

Thus, there is no evidence of direct involvement of this Fn3D (or any other modular carbohydrate active enzymes) in polysaccharide recognition and degradation. On the other hand, eukaryotic Fn3Ds are known for their ability to interact with cell receptors, and this bacterial module may have been a potential pathogen elicitor. However, the hypersensitive response of the plant following the proteolytic cleavage of Pell seems to be triggered by the sole catalytic domain freed from the Fn3D (6).

Catalytic Domain—The catalytic domain is made of eight coils stacked on top of one another (Fig. 1). As in pectate lyases of families 1, 3, and 9, each coil contains three consecutive strand-turn motifs termed (PBn-Tn)n=1,3 (7, 8, 12). The characteristic ladders of inward-facing asparagines observed in family PL-1 and PL-9 within regular stacked turns (7, 12) are absent in Pell as in Pel-15 of the same family PL-3 (11). The stability of the cylindrical core is ensured by inner stacks of aliphatic residues and by five disulfide bonds (highlighted in Fig. 1B).

The structures of Pell and Pel-15 superimpose well on each other with a r.m.s.d. of 1.7 Å on 177 Cα atoms of 226 (Fig. 1A). The main differences lie in the two extremities of the β-helix. The catalytic domain of Pell begins with a portion of random coil whereas Pel-15 harbors a β-strand. This feature might be related to the proximity of the cleavage site in Pell and increases accessibility to proteases. The unique disulfide bond of Pel-15 is conserved in Pell in positions Cys177 and Cys182. Finally, Pell displays in its C-terminal a distinctive β-hairpin made of strands PB1.8 and PB1.8 (Fig. 1). The structure of Pell is more distantly related to that of PelC of family PL-1. 176 Cα atoms of 226 overlap with a r.m.s.d. of 2.9 Å, and the sequence identity is only 16%. Two disulfide bonds of PelC maintaining extended loops have no structural equivalents in Pell.

Turns (Tn)n=1,3 of the β-superhelix generally start by an amino acid in the conformation of a left-handed α helix (αL). The secondary structure assignment of Fig. 1B shows that several turns T of Pell and Pel-15 are made of a single amino acid in αL conformation, which sharply bends the solenoid fold. Of particular interest, the T2 amino acids from coils 2 to 5 line up at the surface of the superhelix (Asn163, Asn186, Gly218, Asn240) and form a highly accessible “Velcro”-like motif of asparagines that anchors the Fn3D. The 2.1 Å structure of the catalytic domain alone, Pellcat, has been determined. This structure is quasi-equivalent to that of Pell and demonstrates that this region retains its rigidity when the Fn3D is removed. Hence, this T2 pattern of outward-facing asparagines (NNGN) located in the interdomain interface of full-length Pell becomes accessible after proteolytic maturation. It then might be recognized by an unknown receptor of the infected plant, leading to the defense response.

Ca2+ and Zn2+ Ions—Calcium ions are essential for the catalytic mechanism of all pectate lyases except the family PL-2, which employs other divalent metal cations (2, 3). They may neutralize the charges of the acidic substrate as observed in the inactive mutant R218K of PelC and pentagalacturonate GalA5 that possesses four Ca2+ ions in the binding site (4). Pell was purified in the presence of calcium and was crystallized at pH 6.5 with zinc sulfate (22). One Ca2+ ion is present in an equivalent position in each molecule A and B of Pell and Pellcat. It binds to an external region of the β-superhelix and coordinates the main-chain carbonyl O of Ile192 and Tyr218 and the side-chain Oδ of Asp191 and Asn213.

In molecule A of Pell, the Ca2+ ion also coordinates two water molecules and displays a classical octahedral geometry. In molecule B, the Ca2+ ion binds three water molecules, giving rise to pentagonal bipyramidal coordination geometry. In both sites, coordination distances are in the range 2.2–2.5 Å. The bound Ca2+ displays a high affinity for the site. Indeed, it has been observed in crystals of Pell grown from protein purified in the presence of the chelating agent EDTA (data not shown). This Ca2+ ion is not located in the active site like those identified in PelC but stabilizes a long turn T3 comprising Tyr218.
This loop is rich in glycine and forms a lid over the bound Ca$^{2+}$ (Fig. 1A). A structural Ca$^{2+}$ is found at an equivalent position in Pel-15, crystallized at pH 6.5 (11). However, in both PelI and PelICata, the two equivalent aspartic residues Asp$^{173}$ and Asp$^{195}$ already coordinate two Zn$^{2+}$ ions provided by the crystallization buffer. These interactions may interfere with the binding of catalytic calcium ions at this site even at alkaline pH. Interestingly, Zn$^{2+}$ ions have a strong inhibitory effect on PelI activity (data not shown). The other Zn$^{2+}$ ions mediate contacts between domains in the crystal. All Zn$^{2+}$ ions display tetrahedral coordination geometry with an average distance of 2.0 Å.

The Conformation of Bound Substrate—The structure of PelI in complex with pectin fragment GalA$_4$ has been determined at a resolution of 2.3 Å after crystal soaking in the absence of calcium. One substrate molecule binds in each molecule A and B to an elongated groove conserved in families PL-1, PL-3, and PL-9. The binding site of PelI stretches from coils 3 to 6, and its base is formed from amino acids of strands PB1. The reducing end of the substrate, termed Ada1, is oriented toward coil 6. The sugar subsites have been identified after superimposition between PelI-GalA$_4$ and inactive PelC-GalA$_5$ (4). One galacturonate unit was not visible in PelC, and the substrate occupies subsites −1 to +3.

Of particular interest in PelI, the substrate binds from subsites −2 to +2 in molecule A, whereas it lies on subsites −1 to +3 in molecule B (Figs. 2 and 3). These two sugar binding modes are consistent with the dual mode of bond cleavage in this substrate. PelI cleaves GalA$_4$ with a frequency of 62% between moieties Ada3 and Ada4 and with a frequency of 38% between moieties Ada2 and Ada3 (43). Thus, the two observed Michaelis complexes perfectly confirm the positions of the two scissile glycosidic bonds in GalA$_4$ acted...
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FIGURE 4. Superimposition between bound sugars in PelI and PelC. The putative catalytic base Lys$^{224}$ of PelI is colored in shades of blue and red in monomers A and B, respectively. Bound sugars are colored using the same scheme and are numbered from the reducing end to the non-reducing end (A1 to A4 and B1 to B4). Bond angles (θ) and torsional rotations (φ, ω) about glycosidic bonds are given in supplemental data. Catalytic base Arg$^{218}$ bound sugar, and catalytic Ca$^{2+}$ ions of PelC are shown in green. The essential Arg$^{215}$ has been re-introduced as in the active enzyme.

upon by Pell. In both molecules, the sugar folds so as to form a
2_1 helix (2 units/helical turn) between subsites −1 and +1 and a
3_1 helix between subsites +1 and +2 (Fig. 4). Such distorted
helical topology between the essential subsites −1 and +2 has
also been reported in PelC and in Pel10Acm complexes with
substrate and Ca$^{2+}$ (4, 14). The latter protein exhibits an α-barrel fold, (α/α)$_n$ instead of the classical β-superhelix fold. There-
fore, the conserved conformation of the substrate at this central
zone −1 to +2 appears to be a requisite for the lyase activity and
can be induced even in the absence of Ca$^{2+}$.

Rearrangements in the Active Site—The structure of Pell-
GalA$_4$, indicates that the active site does not overlap with the
Fm3D interface on the surface of the catalytic domain. However,
in Pell crystals, positive subsites of monomer A and negative
subsites of monomer B are partially buried in non-crystallo-
graphic interfaces. Monomers A and B shift by ~2 Å with
respect to each other during the soaking of Pell crystals with
tetrasaccharide. As a consequence, the substrate has gained
access to the active site in Pell-GalA$_4$ and loops 144–150 and
215–222 are disordered at these interfaces. These structural
constraints at the A/B interfaces are very likely to be responsi-
ble for the two sugar binding modes observed in the crystal.
They also have an impact on adjacent Asp$^{223}$ and Lys$^{224}$ located
at subsite +1 whose side chains are disordered. Beside these
structural rearrangements due to the crystal packing, it can be
noted that strand PB1’8, which is poorly conserved in the PL-3
family, has moved by 2 Å in Pell-GalA$_4$ to accommodate the
substrate. Hence, Lys$^{317}$ and Thr$^{322}$ come close to the tetrasac-
charide to bind it at subsites −1 and +2, respectively (Fig. 3).

Interestingly, the two Zn$^{2+}$ ions coordinated to the acidic cluster in Pell (Asp$^{173}$, Asp$^{195}$) disappear in Pell-GalA$_4$ (the soaking experiment was performed in zinc-depleted solutions) and the side chain of Asp$^{195}$ has rotated by 106° around χ$_{1}$. It
now has a suitable orientation to bind the sugar at subsite −1
(Fig. 2). Finally, a sulfate molecule has been substituted by the
substrate at subsite +2 of monomer B (Fig. 2). This sulfate
molecule was stabilized by the amine N$_{ε}$ of Lys$^{224}$ and by the
guanidinium of Arg$^{252}$. The latter residue is the unique arginine
of the active site and raises the possibility of it catalyzing the
β-elimination of the proton at subsite +1.

The Brønsted Base—In the sugar binding site of Pell, the pos-
sible catalytic bases are Lys$^{224}$, Lys$^{249}$, and Arg$^{252}$. It is notewor-
thy that these three amino acids are invariant in Pel-15 where
they display a similar orientation of their side chains.

In Pell-GalA$_4$, the guanidinium group of Arg$^{252}$ firmly binds
the carboxylic group of the sugar at subsite +2 (Fig. 3), which
prevents any displacement toward subsite +1. The side chain of
the neighboring Lys$^{249}$ is also involved in a hydrogen bond with
the sugar at subsite +2. This orientation is surprising because
Lys$^{249}$ is equivalent in sequence to the catalytic base Arg$^{218}$ of
PelC. Thus, a similar extended conformation of its side chain
toward subsite +1 (leaving group) was expected. However,
Arg$^{218}$ belongs to a turn T3 in PelC and adopts an αL confor-
lation, whereas the isostructural Lys$^{249}$ has a β conformation in Pell. Such divergence in the hydrogen bond network of the
two proteins explains the opposite direction taken by the side
chains of these two equivalent residues.

The last candidate, Lys$^{224}$, is the only invariant basic amino
acid of the active site in family PL-3 and is well conserved in
family PL-1 (Fig. 1B). It is close to the flexible segment 215–222,
and its side chain is disordered in Pell-GalA$_4$. Nevertheless, its
amine function N$_{ε}$ can point next to the C$_{5}$ atom of the sugar at
substrate +1 if the orientation of the side chain in substrate-free
monomer B is taken as a model (its side chain has another
orientation in substrate-free monomer A because it is hydro-
gen-bonded to Oδ of Asp$^{195}$). Hence, Lys$^{224}$ seems the best
candidate for a catalytic base within Pell. To better address this
question, conservative point mutations K224R and K249R, as
well as R252K, have been introduced and analyzed (Table 2).

The mutant K224R is completely defective in lyase activity,
unlike K249R and R252K, keeping ~40–60% and 6–9% of wild
type activity, respectively (Table 2). Interestingly, at a higher
Ca$^{2+}$ concentration in the substrate medium, enzymatic activi-
ties of K249R and R252K mutants are less affected. Hence, it
seems likely that this cation can partially compensate for these
mutations either by changing the pK$_{a}$ of the residues or by
altering the substrate conformation. These three mutations
maintain the electrostatic character of the binding site, and
Lys$^{224}$ appears again as the best candidate catalytic base. A
lysine has also been proposed as the catalytic base for Pel9A
from family PL-9, although without the support of a liganded
structure (12). The putative catalytic bases of Pell and Pel9A are
not equivalent in sequence, but their amine functions N$_{ε}$ are
located in similar positions in the active site.

TABLE 2
Pectate lyase activity detected with the wild-type and mutant proteins
Initial activity rate, in % to wild type, measured in the presence of either 1 mM or 3
mM CaCl$_{2}$

<table>
<thead>
<tr>
<th></th>
<th>1 mM CaCl$_{2}$</th>
<th>3 mM CaCl$_{2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>R252K</td>
<td>6.3</td>
<td>9.1</td>
</tr>
<tr>
<td>K249R</td>
<td>38</td>
<td>64</td>
</tr>
<tr>
<td>K224R</td>
<td>0.7</td>
<td>0.9</td>
</tr>
</tbody>
</table>
CONCLUSION

Our work demonstrates that modular Pell, despite its 11-amino acid-long loop sequence, is a compact protein with two interacting domains. Notably, the interdomain interface comprises a remarkable pattern of stacked asparagines from the catalytic domain. The hypersensitive response of the plant might be elicited by exposure of this motif (NNGN) following elimination of the Fn3 domain. Hence, the response would be independent of lyase activity as observed in HpW from Erwinia amylovora (44).

Our work also answers the question raised by the structure of Pel-15 (11) by showing that the catalytic base in family PL-3 is not an arginine as in family PL-1, PL-2, and PL-10 but a lysine. The use of a lysine, whose pK_a is 10.5, as a base in enzymes active at pH ⁰ 9.0 seems more appropriate than the use of an arginine, whose pK_a is 12.5. Ca²⁺ ions close to the arginine in the Michaelis complex could, however, lower its pK_a to a value that would make the enzyme active (4). The putative catalytic base Lys²²⁴ of Pell is located one coil below the catalytic base Arg²¹⁸ found in the paradigm PelC from family PL-1 (Fig. 1B). The specific activity of Pell on polygalacturonate is lower than that of PelC, and K249R substitution in Pell may have resulted in a PL-1-like efficient enzyme (Lys²⁴⁹ in Pell is equivalent to Arg²¹⁸ in PelC), yet this mutant completely lacks activity. Structural comparisons suggest that the key feature is the main-chain conformation of this amino acid, αl in family PL-1 against β in family PL-3.

Our structure of the complex Pell-GalA₄ shows that the sugar should bind in pectate lyases with a characteristic distorted helical conformation. It indicates that this topology is not calcium-dependent, although this cation is required for pectate lyase activity. In Pell, the presence of a structural Ca²⁺ ion near the active site may influence the efficiency of the catalytic mechanism. This cation stabilizes the flexible loop 215–222 adjacent to Asp²²³ and the putative catalytic base Lys²²⁴. Without this structural Ca²⁺, the essential Lys²²⁴ is disordered in Pell-Gal₄. Catalytic Ca²⁺ ions certainly bind to the active site of Pell at alkaline pH. In any case, pectate lyases are secreted by bacteria in the plant tissue, where they act on cell wall pectin. Ca²⁺ ions are frequently coordinated to free carboxyl groups of pectin, so catalytic cations can be naturally found complexed with the substrate rather than bound to the enzyme.

The structures of inactive PelC and GalA₄ have previously revealed how these catalytic Ca²⁺ ions can mediate interactions between the protein and the sugar at alkaline pH (4). The three Ca²⁺ ions that are unambiguously defined in the electron density maps of PelC (the fourth Ca²⁺ has half occupancy) can be modeled in our structure of Pell in complex with GalA₄ with limited structural rearrangements. These ions are of particular interest because, in this configuration, they may connect together deprotonated carboxyl groups of Pell and of the sugar: one Ca²⁺ between Asp¹⁷³, Asp¹⁹⁵, and subsite sugar -1, a second between Asp¹⁹⁵ and subsite sugar +1, and a third between Asp²²³, Asp²⁵⁶, and subsite sugar +1. Asp²²³ is invariant in family PL-3 like the adjacent essential Lys²²⁴, and a sequence signature -DK- of the active site can be proposed for this family.

A detailed catalytic mechanism has been proposed for Pel9A, with a lysine as a Bronsted base and the support of a More O’Ferrall diagram (12). The interaction of the carboxyl group from the +1 sugar with positive charges can be an important factor for proton abstraction and stabilization of an anionic intermediate. In our high pH model of Pell, the two putative Ca²⁺ ions that are coordinated to the conserved Asp¹⁹⁵ and Asp²²³ may play this role by connecting the sugar carboxyl. A reasonable assumption is that the transiently protonated base Lys²²⁴ may then act as the catalytic acid to eliminate the substituent at O₄ and give a C₄–C₅ unsaturated product. Interaction of polar residues with the -1 and +1 sugar oxygen atoms can facilitate the whole reaction (Gln²²⁷, Asn²²⁹, and Lys³¹⁷ according to Fig. 3).

Finally, the comparison between Pell and Pell-GalA₄ sheds new light on the importance of the conserved acidic cluster Asp¹⁷³, Glu¹⁹⁴, and Asp¹⁹⁵ at the negative subsite. This acidic cluster can favorably complex with Zn²⁺ ions, which results in the obstruction of several Ca²⁺ binding sites and prevents the formation of an active enzyme-substrate complex. Moreover, the side chain of the putative catalytic base Lys²²⁴ needs to be released from its interaction with Asp¹⁹⁵ to reach an appropriate orientation at subsite +1 to perform β-elimination. In either PelC or in the complex PelC-GalA₄ from family PL-1, the equivalent lysine Lys¹⁹⁰ is not released from its contact with a similar acidic cluster and the catalytic base is Arg²¹⁸.

This head-to-head interaction between an acidic and a basic amino acid at the essential -1/+1 subsites evokes the acid-base mechanism observed in family GH28 at acidic pH between the catalytic aspartate and histidine residues (isostructural to Glu¹⁹⁴/Asp¹⁹⁵ and the putative catalytic base Lys²²⁴ in Pell). Thus, not only do the overall architectures of the modular GH28 and PL-3 resemble one another but the positions of their catalytic residues are also similar. The divergent evolution of carbohydrate-degrading enzymes bearing a β-superhelix fold may have given rise first to the families GH28 and PL-3 and then to family PL-1 with a catalytic base at a distinct position.

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REFERENCES

Structure of Pectate Lyase Pell with and without Substrate


The Crystal Structure of Pectate Lyase Pell from Soft Rot Pathogen *Erwinia chrysanthemi* in Complex with Its Substrate
Christophe Creze, Sandra Castang, Emmanuel Derivery, Richard Haser, Nicole Hugouvieux-Cotte-Pattat, Vladimir E. Shevchik and Patrice Gouet

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