Screening of a Novel Polysaccharide Lyase Family 10 Pectate Lyase from Paenibacillus polymyxa KF-1: Cloning, Expression and Characterization

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Abstract: Pectate lyase (EC 4.2.2.2) catalyzes the cleavage of α-1,4-glycosidic bonds of pectin polymers, and it has potential uses in the textile industry. In this study, a novel pectate lyase belonging to polysaccharide lyase family 10 was screened from the secreted enzyme extract of Paenibacillus polymyxa KF-1 and identified by liquid chromatography-MS/MS. The gene was cloned from P. polymyxa KF-1 genomic DNA and expressed in Escherichia coli. The recombinant enzyme PpPel10a had a predicted Mr of 45.2 kDa and pI of 9.41. Using polygalacturonic acid (PGA) as substrate, the optimal conditions for PpPel10a reaction were determined to be 50 °C and pH 9.0, respectively. The K_m, v_max and k_cat values of PpPel10a with PGA as substrate were 0.12 g/L, 289 µmol/min/mg, and 202.3 s⁻¹, respectively. Recombinant PpPel10a degraded citrus pectin, producing unsaturated mono- and oligogalacturonates. PpPel10a reduced the viscosity of PGA, and weight loss of ramie (Boehmeria nivea) fibers was observed after treatment with the enzyme alone (22.5%) or the enzyme in combination with alkali (26.3%). This enzyme has potential for use in plant fiber processing.

Keywords: Paenibacillus polymyxa; pectate lyase; cloning and expression; ramie degumming

1. Introduction

Pectin is a heteropolysaccharide mainly composed of α-1,4-linked galacturonic acids with different degrees of esterification [1,2]. Pectin is widely present in the cell walls of terrestrial plants and has a wide range of applications in industries such as food, medicine and fine chemicals [1,3]. Recent studies showed that the degradation products of pectin have good physicochemical properties, and wide application prospects as food additives, pharmaceutical molecules, and materials for cosmetics [4,5].

The complete degradation of pectin requires the synergistic work of a series of enzymes, including pectate lyase, pectin methyltransferase, and polygalacturonase [2,6]. Pectate lyases (EC 4.2.2.2), also called trans-eliminases, catalyze the cleavage of pectate via a β-elimination reaction to generate 4,5-unsaturated oligogalacturonates [7,8]. Recently, pectate lyase has received increasing attention because of its potential applications in various industries, such as in the extraction and clarification of fruit juices and wine, scouring of cotton fabric, retting of flax, degumming of ramie fibers, and treatment of pectic wastewater [7].

Ramie (Boehmeria nivea) fibers have wide application potential, and can be used to make clothing fabrics, twines and canvas [9]. The surface of ramie fiber is covered with gum-like
materials, which limit its application in the textile industry—The gum-like materials must be removed before application. The traditional degumming process is performed by alkaline treatment at high temperature, which damages the fibers and causes environmental pollution. However, the degumming could also be achieved by enzymatic catalysis performed by pectate lyase [10,11]. Enzymatic degumming has advantages compared with alkaline degumming because of its high efficiency, mild conditions and environmental compatibility.

According to the pH at which they show maximal activity, pectate lyases can be divided into acidic and alkaline groups. Alkaline pectate lyases are preferred for degumming ramie since pectin is more soluble in alkaline solution [12]. Alkaline pectate lyases have been isolated from various microbial sources, including bacteria and fungi [13]. The genus Bacillus is one of the main sources of pectic lyase [11,13–16]. In this study, Paenibacillus polymyxa KF-1 (also known as Bacillus polymyxa) was found to exhibit remarkable pectate lyase activity. A pectate lyase was identified from the enzyme extract of P. polymyxa KF-1 and expressed in Escherichia coli. The recombinant enzyme was purified, and its ability to remove the pectin from ramie was explored.

2. Results and Discussion

2.1. Screening of Pectate Lyase from P. polymyxa KF-1 by LC-MS/MS

Genes encoding pectate lyases have been cloned from Bacillus spp., such as for the pectate lyase from B. tequilensis SV11; PEL168, Apel and rPelB from B. subtilis; BliPelA from B. licheniformis; Bsp165PelA from Bacillus spp. N16-5; Pel-15E from Bacillus spp. strain KSM-P15; and Pel SWU from Bacillus spp. RN1 [13–18]. Here, P. polymyxa exhibited pectin-degradation activity. Using citrus pectin as the substrate, the pectate lyase activity accumulated with prolonged cultivation time, up to 36 h (Figure S1). These results suggested that P. polymyxa is a good candidate to screen for pectate lyase. The supernatant of P. polymyxa KF-1 culture (i.e., the fermentation broth) was purified by ion exchange chromatography performed using an SP Sepharose fast-flow column. Two peaks with pectate lyase activity were observed in the elution profile (Figure 1); the fraction eluted by 0.2 M NaCl (fraction 12) exhibited the highest pectate lyase activity and was sent for LC-MS/MS analysis. A total of 28 proteins were identified from the fraction, including four pectate lyases, one pectin esterase and 23 proteins which were not pectin-degrading enzymes. As shown in Table 1, ten peptides were retrieved and assigned to four pectate lyases, which belonged to PL families 1, 3, 9, and 10, respectively, predicted using the CAZy database (http://www.cazy.org/) [19]. The amino acid sequence of the protein with UniProt accession number E3EEN8 (NCBI protein ID WP_013370345.1) was consistent with a PL family 10 pectate lyase. The PL10 enzyme showed the highest intensity in LC-MS/MS analysis (Table 1), which suggested that the enzyme may be a major component of the pectin-degrading enzymes of P. polymyxa KF-1. Only five pectate lyases belonging to PL family 10 have been characterized, including PelA from B. alcalophilus NTT33 [20], Pel-15E from Bacillus spp. strain KSM-P15 [21], PelA from Azospirillum irakense [22], rPelA from Treponema pectinovorum ATCC 33768 [23], and r-PL_D from Xanthomonas campestris ACCC 10048 [24]. Reported PL10 enzymes had different structure from the reported members in PL family 1, 3, and 9, and were active at alkaline conditions, which may be potential in textile industry [20–24]. However, the properties especially the substrate specificity of PL10 enzymes are still unclear. The application of PL10 enzymes on ramie degumming have not been studied. Understanding of PL family 10 pectate lyases are needed for their application. Thus, the gene encoding the PL family 10 enzyme from P. polymyxa KF-1 was chosen for cloning and expression in the present study.
Figure 1. Elution profile of pectate lyase from P. polymyxa KF-1 culture broth by SP Sepharose fast-flow column chromatography. (■) Pectate lyase activity (A235); (○) protein concentration determined by absorbance at 280 nm (A280); (--) NaCl concentration. The arrow indicates the fraction used for LC-MS/MS analysis.

### Table 1. Pectate lyases identified from P. polymyxa KF-1 by LC-MS/MS analysis.

<table>
<thead>
<tr>
<th>Peptide Sequence</th>
<th>Intensity (×10^5)</th>
<th>Uniprot Accession No.</th>
<th>NCBI Accession No.</th>
<th>Signal Peptide</th>
<th>PL Family</th>
<th>Pfam Family</th>
<th>Predicted Mw (kDa)</th>
<th>Predicted pI</th>
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<tbody>
<tr>
<td>QPFDSDILDNTYR</td>
<td>1.69E7</td>
<td>E3E6N8</td>
<td>WP_013370345.1</td>
<td>1-33</td>
<td>10</td>
<td>PF09492</td>
<td>45.24</td>
<td>9.41</td>
</tr>
<tr>
<td>SKDGVELGTFDNEATTTEIR</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>EPGTVNITGGGAYHAYDK</td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>TVVADPDFTLDGDSQK</td>
<td>1.43E5</td>
<td>E3E6F5</td>
<td>WP_013369567.1</td>
<td>1-33</td>
<td>3</td>
<td>PF03211</td>
<td>24.619</td>
<td>9.19</td>
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<tr>
<td>VNMLEDNSINVK</td>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>GAADSIQLGDFLK</td>
<td>0.44E7</td>
<td>E3E7F9</td>
<td>WP_013373703.1</td>
<td>1-34</td>
<td>9</td>
<td>—</td>
<td>46.988</td>
<td>5.50</td>
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<tr>
<td>GLASADDVSVLPSPTR</td>
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<tr>
<td>GSDLGSCGPSSICNQASVIBMSNLDDLQWNIEPSAAK</td>
<td>0.05E7</td>
<td>E0RB75</td>
<td>WP_013308307.1</td>
<td>1-32</td>
<td>1</td>
<td>PF0544</td>
<td>72.777</td>
<td>6.18</td>
</tr>
</tbody>
</table>

1 Signal peptides were predicted using the SignalP 4.1 server [http://www.cbs.dtu.dk/services/SignalP/][25].

2 Molecular weight and pI were predicted using the ExPASy Compute pI/Mw tool [https://web.expasy.org/ compute_pi/][26].

2.2. Gene Cloning and Sequence Analysis

Analysis of the P. polymyxa genome revealed six genes which were predicted to encode pectate lyases belonging to PL family 1, 3, 9, and 10, respectively. The accession numbers in NCBI for the genes were WP_013372506.1, WP_014599567.1, WP_013369567.1, WP_013370345.1, WP_013373703.1, and WP_013308307.1. Only one PL10 enzyme was retrieved. The gene encoding pectate lyase from P. polymyxa KF-1 was cloned. The open reading frame of 1209 bp encoded a protein of 402 amino acids. The N-terminal 33 amino acids were predicted to be signal peptide by the SignalP 4.1 server. The predicted Mr/pI values were 45.24 kDa and pH 9.41. Generally, pectate lyases are classified into PL families 1, 2, 3, 9, and 10 according to the CAZy database [19]. The amino acid sequence of PpPel10a from P. polymyxa KF-1 showed similarity to the characterized PL family 10 pectate lyases (e.g., AF121904.1, 35%; JQ723690, 40%) (Figure 2). Using the family 10 polysaccharide lyase from Cellvibrio cellulosa (PDB ID: 1GXN, identity = 44.79%) as a template [27], the structure of PpPel10a was modelled; it displayed a predominantly α-helical structure with short β-strands and irregular coils (Figure S2). The structure of family 10 pectate lyases is different from that of the pectate lyases from PL families 1, 3 and 9, which have a parallel β-helix fold [28]. From the amino acid sequence alignment it is deduced that residues D138, N139, R273, E276, R355 and R370 of PpPel10a were responsible for catalysis.
2.3. Expression of Recombinant PpPel10a and Enzyme Purification

The gene encoding the mature pectate lyase PpPel10a was expressed in E. coli BL21 (DE3) cells. After induction with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 25 °C for 10 h, pectate lyase activity was detected mainly in the cell lysate, suggesting the soluble expression of recombinant PpPel10a. Recombinant PpPel10a was purified by Ni-NTA column chromatography, eluted by 50 mM imidazole. The specific activity of purified enzyme was determined to be 290 U/mg (Table 2), which was in the range of reported microbial pectate lyases (45–600 U/mg) [10–12]. Using polygalacturonic acid (PGA) as substrate, the K_m and v_max values for the enzyme were determined to be 0.12 g/L and 289 μmol/min/mg, respectively. The k_cat was calculated to be 202.3 s⁻¹.

Table 2. Purification of P. polymyxa KF-1 Pel10a and of recombinant PpPel10a from E. coli.

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Volume (mL)</th>
<th>Total Activity (U)</th>
<th>Total Protein (mg)</th>
<th>Specific Activity (U/mg)</th>
<th>Yield (%)</th>
<th>Purification</th>
</tr>
</thead>
<tbody>
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<td>Purification</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fermentation broth</td>
<td>200</td>
<td>2762.5</td>
<td>85</td>
<td>32.5</td>
<td>100.0</td>
<td>1.0</td>
</tr>
<tr>
<td>SP Sepharose FF column</td>
<td>9</td>
<td>1066.85</td>
<td>9.5</td>
<td>112.3</td>
<td>38.6</td>
<td>3.5</td>
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<tr>
<td>Heteroexpression</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crude enzyme extract</td>
<td>200</td>
<td>2,276.64</td>
<td>216</td>
<td>105.4</td>
<td>100</td>
<td>1.0</td>
</tr>
<tr>
<td>Ni-NTA column</td>
<td>10</td>
<td>9889</td>
<td>34.1</td>
<td>290</td>
<td>43.4</td>
<td>2.7</td>
</tr>
</tbody>
</table>
The molecular weight of purified recombinant PpPel10a was determined to be approximately 45.2 kDa by SDS-PAGE (Figure 3), which was similar to reported values for PL10 enzymes, such as PelA from *B. alcalophilus* NTT33 (~35 kDa), Pel-15E from *Bacillus* sp. strain KSM-P15 (~33 kDa), PelA from *A. irakense* (44.5 kDa), rPelA from *T. pectinovorum* ATCC 33768 (~49 kDa), and r-PL D from *X. campestris* ACCC 10048 (~38 kDa) [20–24].

Figure 3. SDS-PAGE analysis of recombinant PpPel10a. M, molecular weight markers; lane 1, supernatant of lysed recombinant *E. coli* BL21 (DE3) cells carrying plasmid pET28a-pppel10a; lane 2, PpPel10a purified by Ni-NTA column chromatography.

2.4. Enzymatic Properties of Purified PpPel10a

The optimal pH for activity of PpPel10a was determined to be pH 9.0; the enzyme was very active over a wide pH range (≥70% of maximum activity at pH 7.0–11.0) (Figure 4a). Alkaline pectate lyases have mainly been cloned from *Streptomyces* and *Bacillus* spp. [24]. The optimal pH of PpPel10a was similar to that of Apel from *B. subtilis* (pH 9.0) and r-PL-STR from *Streptomyces* sp. S27 (pH 9.5), but higher than that of Pel from *B. subtilis* 168 (pH 8.0) [15,21,29]. PpPel10a showed stability over a wide pH range; the enzyme retained >50% of its initial activity after being held at pH 5.0–11.0 for 24 h (Figure 4b). The pH stability of PpPel10a was broader than those of the pectate lyases PEL168 from *B. subtilis* 168 (pH 7.0–9.5), Apel from *B. subtilis* (pH 7.0–10.0), and r-PL-STR from *Streptomyces* sp. S27 (pH 7.0–10.0) [13,15,29]. So far, only one PL10 enzyme r-PL D from *X. campestris* ACCC 10048 was characterized on its pH stability [24]. r-PL D was stable over a broad pH range, retaining more than 50% of its initial activity after incubation at pH 3.0–12.0. PpPel10a showed similar alkaline-tolerance property as r-PL D, which indicated that alkaline-tolerance may in common among PL10 enzymes. The optimal temperature for activity of PpPel10a was 50 °C (Figure 4c), which was consistent with that of Apel from *B. subtilis* (50 °C), but lower than those of PelA from *B. licheniformis* 14A (70 °C), and BacPelA from *B. clausii* (70 °C) [14,30]. PpPel10a was quite stable at <50 °C, but lost >50% of its initial activity after incubation at 60 °C for 30 min (Figure 4d). These remarkable alkaline-active, alkali-stable and thermostable properties make PpPel10a an excellent candidate for application in industries requiring alkaline reaction conditions.
PpPel10a exhibited remarkable tolerance to metal ions and chemical reagents. As Table 3 shows, Fe$^{2+}$, Co$^{2+}$, SDS and Triton X-100 were strong inhibitors of enzymatic activity at both 1 and 5 mM. Ca$^{2+}$ was reported to be essential for the activity of most pectate lyases (optimal Ca$^{2+}$ concentration in the range 0.1 to 1.0 mM). The enzymatic activity of PpPel10a was enhanced by the addition of Ca$^{2+}$; a dose-dependent effect was observed at 0.1–2.5 mM [31]. The presence of 2.5 mM Ca$^{2+}$ enhanced the enzyme activity by 1.98-fold compared with that in the absence of added Ca$^{2+}$ (Figure 5). Therefore, the addition of 2.5 mM Ca$^{2+}$ was employed in further experiments. Mg$^{2+}$, Zn$^{2+}$ and Al$^{3+}$ at 1 mM slightly enhanced the enzymatic activity, while other metal ions and chemicals tested showed no significant effect on the enzymatic activity at 1 or 5 mM.

**Table 3.** Effect of metal ions and chemical reagents on the enzyme activity of PpPel10a.

<table>
<thead>
<tr>
<th>Metal Ion or Chemical</th>
<th>1 mM</th>
<th>5 mM</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaCl</td>
<td>99.3 ± 1.9</td>
<td>86.7 ± 3.25</td>
</tr>
<tr>
<td>KCl</td>
<td>90.2 ± 2.0</td>
<td>79.0 ± 0.55</td>
</tr>
<tr>
<td>CaCl$_2$</td>
<td>195.5 ± 2.65</td>
<td>176.6 ± 3.2</td>
</tr>
<tr>
<td>MgCl$_2$</td>
<td>113.7 ± 0.55</td>
<td>94.5 ± 4.65</td>
</tr>
<tr>
<td>FeSO$_4$</td>
<td>65.1 ± 1.5</td>
<td>54.5 ± 0.65</td>
</tr>
<tr>
<td>EDTA</td>
<td>89.3 ± 0.15</td>
<td>63.1 ± 0.45</td>
</tr>
<tr>
<td>ZnCl$_2$</td>
<td>115.2 ± 0.95</td>
<td>88.0 ± 1.55</td>
</tr>
<tr>
<td>AlCl$_3$</td>
<td>119.6 ± 0.35</td>
<td>106.1 ± 3.1</td>
</tr>
<tr>
<td>CoCl$_2$</td>
<td>72.2 ± 3.4</td>
<td>51.4 ± 2.4</td>
</tr>
<tr>
<td>SDS</td>
<td>61.7 ± 1.95</td>
<td>37.7 ± 0.15</td>
</tr>
<tr>
<td>Tween-20</td>
<td>108.0 ± 1.0</td>
<td>122.5 ± 0.55</td>
</tr>
<tr>
<td>Triton X-100</td>
<td>8.5 ± 0.75</td>
<td>4.3 ± 0.15</td>
</tr>
</tbody>
</table>
2.5. Identification of Products from Citrus Pectin on Degradation by PpPel10a

High-performance gel permeation chromatography (HPGPC) analysis showed that citrus pectin (~406,922 Da) was degraded to smaller molecular weight species (~231,899 Da and ~2085 Da) by PpPel10a (Figure 6). The oligomers released from citrus pectin by PpPel10a were observed on high-performance anion exchange chromatography (HPAEC) (Figure 7).

Figure 5. Effect of Ca\(^{2+}\) addition on the activity of PpPel10a.

Figure 6. High-performance gel permeation chromatography analysis of the lytic products of citrus pectin produced by PpPel10a. (—) Citrus pectin; (-----) lytic products of citrus pectin.

Figure 7. High-performance anion exchange chromatography analysis of the lytic products of citrus pectin produced by PpPel10a. Black line, citrus pectin; red line, lytic products of citrus pectin.
The oligosaccharides were identified by ESI-MS. Negative ESI-MS gave strong peaks at \( m/z \) 175 (unsaturated galacturonic acid, uG1), \( m/z \) 351 (unsaturated bigalacturonide, uG2), \( m/z \) 527 (unsaturated trigalacturonide, uG3), \( m/z \) 703 (unsaturated tetragalacturonide, uG4), and \( m/z \) 879 (unsaturated pentagalacturonide, uG5) (Figure 8) [16]. The abundance was in the order uG2 > uG3 > uG4 > uG1 > uG5. These results indicate that citrus pectin was degraded by PpPel10a, producing a mixture of 4,5-unsaturated monogalacturonic acid and oligogalacturonic acid, confirming the trans-elimination reaction mechanism of PpPel10a. Generally, exo-acting pectate lyase produces only monomer or dimer, while endo-acting pectate lyase releases larger oligomers [16]. The HPAEC and ESI-MS results showed that uG2 and uG3 were the main products released from citrus pectin by PpPel10a, indicating PpPel10a is an endo-acting enzyme. The catalytic behavior of PpPel10a was similar to that of Pel SWU, the PL family 1 endo-acting enzyme from Bacillus spp. RNI, but different from some other reported endo-acting pectate lyases, such as PelA from *B. licheniformis* 14A and PL from *B. subtilis*, which produced unsaturated oligogalacturonides uG2 and uG3, but not uG1 [14–16]. Only two PL10 enzymes have been reported on their catalytic behavior. PelA from *A. irakense* was an endo-acting pectate lyase, producing multiple unsaturated oligogalacturonates (uG2 to uG9) without the accumulation of uG1 [22]. r-PL D from *X. campestris* ACCC 10048 was an exo-type pectate lyase, the major product was determined to be uG2 [24]. Thus, to our knowledge, this is the first report of a PL10 enzyme that can accumulate uG1 from pectin. It is deduced that PL10 enzymes may possess both endo- and exo-acting activities. The catalytic mechanism of PL10 enzymes need to be further studied.

![Figure 8.](image)

**Figure 8.** Electrospray ionization-mass spectrometry analysis of the lytic products of citrus pectin produced by PpPel10a.

2.6. Application of PpPel10a in Viscosity Reduction and Ramie Degumming

The viscosity of a PGA solution decreased gradually during incubation at 50 °C for 1 h on addition of PpPel10a; the relative viscosity was reduced by 33.2% after 1 h of incubation (Figure 9). The reducing sugars released from PGA indicated that <3% of the galacturonosidic linkages were cleaved, which indicated that PpPel10a acts as an endo-pectate lyase toward PGA.

The ramie degumming effect of PpPel10a was studied. After treatment with 1 U/mL PpPel10a, 22.5 ± 0.68% of the weight of the ramie fibers was lost, whereas 10.1 ± 0.57% weight loss was obtained on treatment with 50 mM sodium glycine buffer, pH 9.0. In combined enzyme and chemical (sodium hydroxide) treatment, the weight loss was 26.3 ± 0.68%, which was higher than on chemical treatment alone (0.5% w/v NaOH, 19.9 ± 0.87%; 1% w/v NaOH, 25.9 ± 0.65%). Although the weight losses showed no significant difference between chemical-enzyme combination and 1% sodium hydroxide, the amount of sodium hydroxide used in the chemical-enzyme combination was 50% lower, which is better for the environment.
was higher than those of these pectate lyases. It is reported that the total gum content of ramie fibers
whiter compared with that treated with 50 mM sodium glycine buffer (pH 9.0) or by sodium hydroxide
alone, which is favorable for the textile industry. The textiles were observed by scanning electron microscopy (SEM). Figure 10 shows that the enzyme-treated and enzyme-chemical treated samples showed a smoother surface than the control group, which suggested that the gum-like material, i.e., the pectin, was removed.

Recently, ramie degumming by some Bacillus spp. alkaline pectate lyases was reported [11,32]. Treatment with pectate lyases from B. pumilus DKS1, B. subtilis 7-3-3, and B. pumilus ATCC7061 resulted in 17%, 13.5% and 23.1% weight loss, respectively. The degumming efficiency of PpPel10a was higher than those of these pectate lyases. It is reported that the total gum content of ramie fibers is approximately 30–34% [14]. The reason of the incomplete degumming by PpPel10a may derive from the pH stability of PpPel10a at extreme alkaline environment. The pH of 0.5% NaOH in 20 mL of 50 mM sodium glycine buffer (pH 9.0) was determined to be ~12.5. The activity and stability of PpPel10a in this solution was studied. PpPel10a showed moderate activity at this pH, 62.5% of its activity detected. Therefore, the degumming efficiency needs to be further improved by improving the stability of PpPel10a or by using PpPel10a combined with other enzymes.

Figure 9. Reduction of PGA viscosity by PpPel10a (–) and galacturonoside bond cleavage (–).

The ramie textile treated by enzyme and enzyme plus chemical became much softer, smoother and
whiter compared with that treated with 50 mM sodium glycine buffer (pH 9.0) or by sodium hydroxide
alone, which is favorable for the textile industry. The textiles were observed by scanning electron microscopy images of single fibers (5000×).

Figure 10. Ramie fibers treated with buffer (50 mM sodium glycine buffer, pH 9.0, negative control) (A,F); chemically (0.5% w/v sodium hydroxide (B,G), or 1% w/v sodium hydroxide (C,H)); enzymatically (1 U/mL PpPel10a) (D,I); or with enzyme-chemical combination (1 U/mL PpPel10a and 0.5% w/v sodium hydroxide) (E,J). Images A–E: extrinsic features of ramie fibers; images F–J: scanning electron microscopy images of single fibers (5000×).
Degumming of ramie fibers needs robust pectate lyases that are active and stable at alkaline pH and moderate temperature [10–12]. PpPel10a may have potential for use in this way because of its alkaline-tolerance and thermostability. In addition, PpPel10a was easily prepared since purification of the enzyme required only one step. Altogether, the enzyme-chemical combination degumming was more environmentally friendly than the traditional chemical degumming process. In the future, the degumming process will be further optimized.

3. Materials and Methods

3.1. Bacterial Strains, Plasmid, and Substrates

*P. polymyxa* KF-1 was collected by our laboratory and deposited in the China Center for Type Culture Collection (CCTCC AB 2018146). *Escherichia coli* DH5α was used for gene cloning procedures. pET-28a(+) and *E. coli* BL21 (DE3) cells (Novagen, Madison, MI, USA) were used for recombinant enzyme production. Citrus pectin (P8030) and PGA (P8510) were purchased from Solarbio (Beijing, China).

3.2. Screening of Pectate Lyase from *P. polymyxa* KF-1 by LC-MS/MS

Strain *P. polymyxa* KF-1 was inoculated into 100 mL Luria-Bertani (LB) broth and shaken at 200 rpm at 30 °C for 36 h. After that, the supernatant of culture broth was obtained by centrifugation at 5000 × g and 4 °C for 10 min. The supernatant was loaded onto a SP Sepharose fast-flow column (1.5 × 5 cm; GE Healthcare, Little Chalfont, Buckinghamshire, UK). The column was eluted by a stepwise gradient of 0–0.5 M sodium chloride (72 mL) in 25 mM Tris-HCl, pH 7.5. The pectate lyase activity of each fraction was measured with 0.2% PGA as substrate. The reaction was carried out in 50 mM sodium glycine buffer (pH 9.0) at 50 °C for 15 min. The absorbance increase at 235 nm was recorded. The extinction coefficient of unsaturated galacturonic acid at 235 nm is 4600 M⁻¹ cm⁻¹ [33]. One unit (U) of activity was defined as the amount of enzyme needed to produce 1 µmol/min unsaturated oligogalacturonide.

The chromatographic fraction with maximum pectate lyase activity was analyzed by Orbitrap Fusion Lumos LC-MS/MS (Thermo Fisher Scientific, Waltham, MA, USA) by Beijing Bio-Tech Pack Technology Company Ltd. (Beijing, China). Briefly, the proteins were in-solution digested by 100 ng/mL trypsin, the peptides were extracted by a mixture of trifluoroacetic acid/acetonitrile/distilled water (5/50/45, v/v/v), then separated using Acclaim PepMap RSLC C18 column (Thermo Fisher Scientific, Waltham, MA, USA). The separated peptides were analyzed by nanoLC-MS/MS [34]. The data obtained was searched against the UniProt nonredundant protein database for *P. polymyxa* using Proteome Discoverer 1.4 (Thermo Fisher Scientific, Waltham, MA, USA) [35].

3.3. Sequence Analysis of PL Family 10 Pectate Lyase

The peptides identified by LC-MS/MS analysis were analyzed by Protein BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi). Amino acid sequence alignment was performed using MEGA version 6.06 (https://www.megasoftware.net/) [36]. Signal peptides were predicted using the SignalP 4.1 Server [25]. A structural model of the pectate lyase PpPel10a was generated using SWISS-MODEL with the family 10 polysaccharide lyase from *C. cellulosa* (PDB ID: 1GXN, identity = 44.79%) as the template [28,37].

3.4. Gene Cloning and Protein Expression

The pectate lyase gene PpPel10a (NCBI accession number WP_013370345.1) was amplified from genomic DNA of *P. polymyxa* KF-1 using primers 5′-CGGGATCCGAGCAGAATTTGACTGATGC-3′ and 5′-CCGCTCGAGTTATTGTGGCAACGGCTTGGACAG-3′. The PCR product was digested with BamHI and XhoI, and ligated into BamHI/XhoI-digested pET-28a(+). The ligation product was transformed into *E. coli* DH5α cells and the recombinant plasmid pET-28a-PpPel10a obtained was used to transform *E. coli* BL21 (DE3) cells [38]. The transformant was cultured in 200 mL LB medium.
supplemented with 30 µg/mL kanamycin [13]. Protein induction was initiated by the addition of 0.5 mM IPTG when the optical density of the culture at 600 nm reached 0.6. The culture was grown for another 10 h at 25 °C with shaking at 200 rpm [13].

Recombinant E. coli BL21 (DE3) cells were disrupted by sonication, and centrifuged at 12,000 × g at 4 °C for 30 min to remove cell debris. The supernatant obtained (200 mL) was loaded onto a Ni-NTA agarose column (1 × 5 cm). The column was eluted with a linear gradient of imidazole (10–200 mM) in 25 mM Tris-HCl buffer, pH 7.5 [4]. Fractions with pectate lyase activity were collected and dialyzed into 25 mM Tris-HCl buffer (pH 8.0). Recombinant enzyme PpPel10a was analyzed by 10% SDS-PAGE [39]. The protein concentration was determined using BCA reagent [40].

3.5. Enzymatic Characterization of Pectate Lyase

The effect of pH on the activity of PpPel10a was determined at pH 2.0–11.0 using 0.2% PGA as the substrate. The reactions were carried out at 50 °C for 15 min. The effect of pH on stability was determined by incubating purified recombinant PpPel10a at 4 °C for 24 h at different pH, then the enzyme activity was assayed in standard conditions (50 °C, 15 min). The buffers used in this study were: pH 2.0–6.0, 50 mM sodium acetate buffer; pH 6.0–8.0, 50 mM phosphate buffer; pH 8.0–11.0, 50 mM sodium glycine buffer. The effect of temperature on enzyme activity was studied at 20–80 °C at pH 9.0 for 15 min. Thermostability of PpPel10a was studied by measuring residual pectate lyase activity after preincubating the enzyme at different temperatures for up to 1 h [41]. Then, the residual activities were determined at pH 9.0, 50 °C, over 15 min.

The effects of metal ions or chemical compounds (1 and 5 mM) on pectate lyase activity were measured by preincubation of PpPel10a in different metal ion solutions or chemicals at 4 °C for 24 h. Then, the residual activities were measured using 0.2% PGA as the substrate at pH 9.0 and 50 °C for 15 min [13]. Enzyme without preincubation was defined as 100% active. The effect of Ca²⁺ on pectate lyase activity was determined from 0.1 to 5 mM Ca²⁺. The enzymatic reaction without the addition of Ca²⁺ was set as 100% activity. The kinetic parameters of PpPel10a were measured using 0.1–5.0 mg/mL PGA in standard assay conditions (pH 9.0, 50 °C, 15 min). Kinetic values \( K_m \), \( v_{max} \) and \( k_{cat} \) were calculated from Lineweaver-Burk plots [42].

3.6. Analysis of Lytic Products from Citrus Pectin on Degradation by PpPel10a

The change of molecular weight of citrus pectin after treatment with PpPel10a was analyzed by HPGPC using a TSK-gel G-3000PWXL column (7.8 × 300 mm; Tosoh, Tokyo, Japan) [43]. The column was calibrated with standard dextrans (1, 5, 12, 25 and 50 kDa) (Sigma-Aldrich, St. Louis, MO, USA). The unsaturated galacturonic acids released from citrus pectin by PpPel10a were analyzed by HPAEC with a pulsed amperometric detector [44]. The unsaturated galacturonic acids were identified by ESI-MS with the amaZon speed ion ETD trap (Bruker, Bremen, Germany) using negative electrospray as the ionization process [45].

3.7. Application of PpPel10a in Viscosity Reduction and Ramie Degumming

Purified PpPel10a (1.0 U) and PGA (0.2% w/v) were incubated in 10 mL of 50 mM sodium glycine buffer (pH 9.0) at 50 °C for up to 1 h. Viscosity was determined using a glass capillary viscometer (1833, Shenyi Glass, Shanghai, China) at timed intervals [24,46]. The viscosity of PGA without enzymatic treatment was considered as 100%. The extent of PGA cleavage was determined by measuring the released reducing sugars using the DNS method [46,47].

For ramie degumming, 2 g of ramie fibers were boiled in distilled water for 10 min, then treated by chemical (0.5% or 1% NaOH), enzyme (1 U/mL purified recombinant PpPel10a), or a chemical-enzyme combination (1 U/mL purified recombinant PpPel10a in 0.5% sodium hydroxide) in 20 mL of 50 mM sodium glycine buffer (pH 9.0). The mixtures were shaken at 50 °C and 100 rpm for 4 h. Then, the ramie fibers were washed twice and dried to constant weight. The weight loss was recorded, and the surface of the ramie fibers was observed by SEM [10,11,13]. Ramie fibers treated with 50 mM sodium glycine...
buffer (pH 9.0) were used as the negative control. The experiments were performed with five parallels. For ramie degumming, the activity and stability of PpPel10a in the chemical-enzyme combination treatment (0.5% NaOH in 20 mL of 50 mM sodium glycine buffer, pH 9.0) was studied using 0.2% PGA as the substrate.

4. Conclusions

An alkaline pectate lyase was screened from *P. polymyxa* KF-1 and identified as belonging to PL family 10. The enzyme showed catalytic activity toward citrus pectin, producing unsaturated mono- and oligogalacturonates. The significant weight loss of ramie fibers treated with PpPel10a indicated the potential of this enzyme for application in ramie degumming. PpPel10a, which has good thermostability and pH stability, might be suitable for use in the textile industry.

**Supplementary Materials:** The following are available online, Figure S1: Measurement of *P. polymyxa* KF-1 pectate lyase activity during liquid culture, Figure S2: Structural model of PpPel10a generated by SWISS-MODEL using pectate lyase from *C. cellulosa* (PDB ID: 1GXN, identity = 44.79%) as the template.

**Author Contributions:** Conceptualization, J.G.; Data curation, Y.Z. and Y.Y.; Formal analysis, X.Z. and Y.L.; Funding acquisition, Q.L.; Methodology, Y.Z. and Y.Y.; Software, Y.Z., Y.Y. and J.G.; Validation, Y.Z.; Visualization, X.Z.; Writing—Original draft, Y.Z. and J.G.; Writing—Review & editing, J.G.

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Sample Availability: Not available.

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