Mechanistic Insights into Substrate Recognition and Catalysis of a New Ulvan Lyase of Polysaccharide Lyase Family 24

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ABSTRACT

Ulvan is an important marine polysaccharide. Bacterial ulvan lyases play important roles in ulvan degradation and marine carbon cycling. Until now, only a small number of ulvan lyases have been characterized. Here, a new ulvan lyase, Uly1, belonging to polysaccharide lyase family 24 (PL24) from the marine bacterium Catenovulum maritimum, is characterized. The optimal temperature and pH for Uly1 to degrade ulvan are 40°C and pH 9.0, respectively. Uly1 degrades ulvan polysaccharides in the endolytic manner, mainly producing D-Rha3S, consisting of an unsaturated 4-deoxy-L-threo-hex-4-enopyranosuronic acid and a 3-O-sulfated α-L-rhamnose. The structure of Uly1 was resolved at a 2.10-Å resolution. Uly1 adopts a seven-bladed β-propeller architecture. Structural and site-directed mutagenesis analyses indicate that four highly conserved residues, H128, H149, Y223, and R239, are essential for catalysis. H128 functions as both the catalytic acid and base, H149 and R239 function as the neutralizers, and Y223 plays a supporting role in catalysis. Structural comparison and sequence alignment suggest that Uly1 and many other PL24 enzymes may directly bind the substrate near the catalytic residues for catalysis, different from the PL24 ulvan lyase LOR_107, which adopts a two-stage substrate binding process. This study provides new insights into ulvan lyases and ulvan degradation.

IMPORTANCE

Ulvan is a major cell wall component of green algae of the genus Ulva. Many marine heterotrophic bacteria can produce extracellular ulvan lyases to degrade ulvan for a carbon nutrient. In addition, ulvan has a range of physiological bioactivities based on its specific chemical structure. Ulvan lyase thus plays an important role in marine carbon cycling and has great potential in biotechnological applications. However, only a small number of ulvan lyases have been characterized over the past 10 years. Here, based on biochemical and structural analyses, a new ulvan lyase of polysaccharide lyase family 24 (PL24) from the marine bacterium Catenovulum maritimum, is characterized. The optimal temperature and pH for Uly1 to degrade ulvan are 40°C and pH 9.0, respectively. Uly1 degrades ulvan polysaccharides in the endolytic manner, mainly producing D-Rha3S, consisting of an unsaturated 4-deoxy-L-threo-hex-4-enopyranosuronic acid and a 3-O-sulfated α-L-rhamnose. The structure of Uly1 was resolved at a 2.10-Å resolution. Uly1 adopts a seven-bladed β-propeller architecture. Structural and site-directed mutagenesis analyses indicate that four highly conserved residues, H128, H149, Y223, and R239, are essential for catalysis. H128 functions as both the catalytic acid and base, H149 and R239 function as the neutralizers, and Y223 plays a supporting role in catalysis. Structural comparison and sequence alignment suggest that Uly1 and many other PL24 enzymes may directly bind the substrate near the catalytic residues for catalysis, different from the PL24 ulvan lyase LOR_107, which adopts a two-stage substrate binding process. This study provides new insights into ulvan lyases and ulvan degradation.

KEYWORDS

ulvan, ulvan lyase, polysaccharide lyase family 24, marine bacterium, substrate recognition, catalytic mechanism
Marine macroalgae are major components of coastal ecosystems. Marine algal polysaccharides contain unique sugars, which are often heavily sulfated and differ substantially from terrestrial plant polysaccharides (1). Ulvan is the major cell wall polysaccharide of green algae of the genus *Ulva*, accounting for up to 29% of the dry weight (2, 3). Ulvan is a random polymer of 3-O-sulfated α-L-rhamnose (Rha3S), β-D-glucuronic acid (GlcA), α-L-iduronic acid (IdoA; the C-5 epimer of GlcA), and xylose (Xyl) (4). Based on the type of disaccharide unit, ulvan can be classified into one of three groups: A3S, B3S and U3S. A3S consists of Rha3S and GlcA, B3S consists of Rha3S and IdoA, and U3S is composed of Rha3S and Xyl (Fig. 1). A3S and B3S are the common disaccharide repetitive units within ulvan polysaccharides, and U3S occurs in smaller amounts. Because of its bioactivity and chemical versatility, ulvan has application potentials in many areas, including food, medicine, and biomaterials (5–7).

Ulvan lyases, first identified in 2011, can degrade ulvan, targeting the glycosidic 1,4 O-linkage between Rha3S and GlcA (A3S) or between Rha3S and IdoA (B3S), by a β-elimination mechanism with the formation of an unsaturated 4-deoxy-L-threo-hex-4-eno-pyranosiduronic acid (Δ) at the nonreducing end (Fig. 1) (8, 9). Until now, 17 ulvan lyases have been characterized (9), which are distributed into the polysaccharide lyase (PL) families 24, 25, 28, 37, and 40 in the Carbohydrate-Active enZYmes (CAZy) database (8–21). Of these, the structures of only three have been solved and characterized, namely, LOR_107 from *Alteromonas* sp. (the PL24 family) (18), PLSV_3936 from *Pseudoalteromonas* sp. (the PL25 family) (13), and NLR48 from *Nonlabens ulvanivorans* (the PL28 family) (19). The ulvan lyases LOR_107 from PL24 and PLSV_3936 from PL25 adopt the β-propeller scaffold, and the ulvan lyase NLR48 from PL28 adopts a β-jelly roll fold. The proposed catalytic mechanisms of these three lyases are significantly different. For LOR_107, the residue His146 is proposed to be both the catalytic base and acid aided by His167 and Tyr243 (18). PLSV_3936 adopts the typical His/Tyr mechanism, in which Tyr188 is most likely the catalytic base and His123 is most likely the catalytic acid (13). NLR48 uses different key catalytic residues, depending on the substrate at the +1 site. With GlcA at the +1 site, Tyr281 acts as the catalytic base and acid; with IdoA at the +1 site, Lys162 acts as the catalytic base and Tyr281 acts as the catalytic acid (19). Ulvan lyases play ecological roles in ulvan degradation in the ocean and have important applications in biotechnological fields, such as preparation of functional oligomers. Therefore, identification of more ulvan lyases and clarification of their characteristics, structures, and catalytic mechanisms will broaden our understanding of the degradation of ulvan polysaccharides.

The PL24 family currently contains four identified ulvan lyases in the CAZy database (LOR_61, LOR_107, PLSV_3875, and PLSV_3925). These are divided into two types: a short type (~59 kDa) and a long type (~110 kDa) (10). LOR_107 and PLSV_3875 belong to the short type, and LOR_61 and PLSV_3925 belong to the long type (10). The short-type ulvan lyases only contain the catalytic domain, whereas the long-type lyases comprise a catalytic domain and an unknown domain at the C terminus. Members of the PL24 family are known to specifically cleave the glycosidic bond between GlcA and Rha3S (A3S) by β-elimination, which differs from the other family ulvan lyases that can cleave both the glycosidic bonds of GlcA-Rha3S and IdoA-Rha3S (9). LOR_107 is the only enzyme with its structure reported in this family, adopting the β-propeller scaffold (18).

Of the 49 sequences of the PL24 family available in the CAZy database, 21 are from *Catenovulum* strains, indicating that *Catenovulum* is likely an important ulvan-degrading group. However, no ulvan lyases from *Catenovulum* have been characterized. *Catenovulum maritimum* Q1T is a marine bacterium isolated from the marine alga *Porphyra yezoensis* Ueda, collected from coastal areas of Weihai, China (22), which contains a putative PL24 ulvan lyase gene and a putative PL25 ulvan lyase gene. In this study, the putative PL24 ulvan lyase from *Catenovulum maritimum* Q1T, named Uly1, was characterized. Uly1 is a new ulvan lyase of the PL24 family, which is an endoenzyme with ΔRha3S as its main product. The three-dimensional structure of Uly1 was also solved at a 2.10-Å resolution, and the important residues in Uly1 for substrate
binding and catalysis were analyzed. A comparison of the conformation of the catalytic canyon and the residues involved in substrate binding between Uly1 and the PL24 ulvan lyase LOR_107 suggests that Uly1 may adopt a different substrate binding process from that of LOR_107. These results provide new insights into ulvan lyases and ulvan degradation in the ocean.

RESULTS AND DISCUSSION

Sequence analysis of Uly1. The *uly1* gene of the marine bacterium *Catenovulum maritimum* Q1T is 1,539 bp in length and was predicted to encode a putative PL24 ulvan lyase (Uly1) of 512 amino acid residues. Uly1 contains a 20-residue signal peptide predicted by the SignalP 4.1 server and a catalytic domain (A21 to Q512). Phylogenetic analysis of Uly1 and other reported ulvan lyases suggests that Uly1 is a PL24 ulvan lyase (Fig. 2). Only 4 ulvan lyases have been characterized in the PL24 family. Of these, Uly1 has the highest sequence identity (50%) with the ulvan lyase LOR_107 (covering 94% of the Uly1 sequence) (18). Nevertheless, differences in the remaining 50% of the sequence leave open the possibility that Uly1 may be a new PL24 ulvan lyase with some different properties from the other characterized ulvan lyases. Therefore, we further characterized Uly1.

Expression and characterization of Uly1. The gene of Uly1, without the signal peptide sequence, was overexpressed in *Escherichia coli* BL21(DE3). SDS-PAGE analysis showed that the recombinant Uly1 was approximately 55 kDa (Fig. 3A), consistent with its theoretical molecular mass of 55.8 kDa. Gel filtration analysis indicated that Uly1 is present as a monomer in solution (Fig. 3B). Because all the enzymes characterized in the PL24 family are ulvan lyases, the activity of Uly1 toward ulvan was measured along with three other polysaccharides, chondroitin sulfate, heparan sulfate, and alginate, to investigate its substrate specificity. Of these polysaccharides, the recombinant Uly1 was active only toward ulvan, with a specific activity of 20.16 ± 1.40 U/mg at 40°C, demonstrating that Uly1 is an ulvan lyase. The optimal temperature of Uly1 toward ulvan was 40°C (Fig. 3C). The thermostability of Uly1 was further investigated. Approximately 50% of the activity was lost after incubation at 40°C for 1 h, and approximately 70% of its activity was lost after incubation at 50°C for 15 min (Fig. 3D).

Uly1 exhibited maximal activity at pH 9.0 (Fig. 3E) and retained more than 80% of its activity after incubation at pH 6.0 to 11.0 for 1 h (Fig. 3F). The optimum pH of a majority of the characterized ulvan lyases ranged from 7.5 to 9.0 (9), which reflects their adaptation to the alkaline marine environment. In saline solutions containing 0 to 4 M NaCl, Uly1 exhibited its highest activity at 0.5 M NaCl, which was 1.5-fold higher than

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**FIG 1** The disaccharide repetition units encountered in ulvan and the degradation product of ulvan lyases on ulvan. Three disaccharide repetition units, A₃S, B₃S, and U₃S, exist in ulvan. Only A₃S and B₃S can be decomposed by ulvan lyases, leading to a reducing end on one fragment and an unsaturated ring (Δ, 4-deoxy-L-threo-hex-4-enopyranosiduronic acid) on the nonreducing end of the other fragment.
that at 0 M NaCl (Fig. 3G), indicating the adaptation of Uly1 to the salt concentration of the seawater. While no metal ions among Mg$^{2+}$, Li$^+$, K$^+$, Ca$^{2+}$, and Cu$^{2+}$ had an enhancing effect on its activity, 1 mM Co$^{2+}$ and Mn$^{2+}$ reduced Uly1 activity by 53.40 and 91.44%, respectively (Table 1).

To determine the polysaccharide degradation pattern, the enzymatic products of Uly1 over a time course of 0 to 12 h were separated on a Superdex peptide 10/300 GL.
column. The products formed five large peaks and some small ones. These peaks displayed changes in amount but consistency in location with the increase in degradation time from 15 min to 12 h (Fig. 4A), indicating that Uly1 has several specific cleavage sites on ulvan and thus is an endolytic lyase. The peaks separated by gel filtration chromatography were further identified by negative electrospray ionization mass spectrometry (ESI-MS). Only the highest peak was successfully identified, with the others failing, probably due to the small amount and/or the impurity of these peaks. (One peak contains several products, and the amount of each is quite low.) The highest peak had an m/z value of 401, which is consistent with the molecular weight of the disaccharide D-Rha3S (Fig. 4B). This indicates that the main product released from ulvan by Uly1 is D-Rha3S.

Kopel et al. reported that the PL24 ulvan lyases LOR_107, LOR_61, PLSV_3875, and PLSV_3925, with different molecular masses or sources of strains, all preferentially cleaved GlcA residues (10). Therefore, as a PL24 ulvan lyase, Uly1 may also preferentially cleave GlcA; however, this needs further confirmation.

Overall structure of Uly1. At the time the structure of Uly1 was solved, a similarity search of the Protein Data Bank (PDB) database showed that no suitable structure model could be used for Uly1 structure construction. Therefore, the crystal structure of the wild-type (WT) Uly1 was solved by the single-wavelength anomalous dispersion method using a selenomethionine (SeMet) derivative. Uly1 crystals belong to the P1211 space group, and the structure was solved at a resolution of 2.10 Å. The statistics for refinement are summarized in Table 2. Structure data show that each asymmetric unit contains two Uly1 molecules. Gel filtration analysis indicated that Uly1 is present as a monomer in solution (Fig. 3B), suggesting that the intermolecular contacts observed in the crystal originate from crystal packing. Uly1 adopts the fold of a seven-bladed b-propeller. Each propeller consists of four antiparallel b-strands (Fig. 5A). The catalytic center is located at the canyon on the top of the propeller (Fig. 5A). After the structure of Uly1 was solved, three ulvan lyase structures were successively reported, including those of PLSV_3936 from PL25 (13), LOR_107 from PL24 (18), and NLR48 from PL28 (19). The b-propeller structure of Uly1 is similar to those of PLSV_3936 and LOR_107. The root mean square deviation (RMSD) value (based on all amino acid residues in their structures) between Uly1 and PLSV_3936 is 3.37 Å, and that between Uly1 and LOR_107 is 1.32 Å. As shown in Fig. 5B, PLSV_3936 and Uly1 are significantly different in overall structure, and LOR_107 is the closest structural homologue. It has been reported that two Ca2+ ions are present in each independent molecule of the two LOR_107 molecules in the asymmetric unit at an identical position distant from the conserved catalytic residues, and they appear to play a strictly structural role (18). In the Uly1 structure, the electron density map indicates that there are many metal ions that are identified as Ca2+ based on their coordination geometry. However, these Ca2+ ions do not exist in exactly the same positions in the two molecules of an asymmetric unit. Inductively coupled plasma-optical emission spectrometry (ICP-OES) results showed that each Uly1 monomer contains only approximately 0.1 Ca2+. All these data

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<tr>
<th>Compound</th>
<th>Relative activity (%)</th>
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<tr>
<td></td>
<td>0.25 mM</td>
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<tr>
<td>Mg2+</td>
<td>102.71 ± 5.91</td>
</tr>
<tr>
<td>Li+</td>
<td>103.35 ± 8.53</td>
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<tr>
<td>K+</td>
<td>106.09 ± 9.57</td>
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<tr>
<td>Mn2+</td>
<td>60.68 ± 6.01</td>
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<tr>
<td>Ca2+</td>
<td>106.46 ± 2.16</td>
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<tr>
<td>Cu2+</td>
<td>105.21 ± 14.37</td>
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<tr>
<td>Co2+</td>
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*aThe enzyme activity of Uly1 at 40°C and pH 9.0 without the addition of metal ion was taken as 100%. All experiments were repeated three times.
suggest that the bound Ca$^{2+}$ ions in the Uly1 crystal structure are most likely obtained from calcium acetate in the crystallization solution.

**Analysis of the important residues involved in catalysis.** To investigate the crucial amino acid residues in Uly1, we attempted to crystallize different Uly1 mutants with their substrate or products, but this failed. Instead, we made a structural alignment of Uly1 and LOR_107 in complex with the ulvan tetrasaccharide (PDB code 6BYT) (Fig. 6A). The Uly1 residues R239, H149, H128, and Y223 are all close to the tetrasaccharide and are highly conserved in the PL24 ulvan lyases (Fig. 6A and B). The catalytic mechanism of LOR_107 has recently been revealed (18). In LOR_107, H146 plays the roles of both catalytic acid and base, R259 and H167 act as neutralizers of the charge of the GlcA acidic group, and Y243 plays a supporting role in properly orienting the imidazole ring of H146 (18). R239, H149, H128, and Y223 of Uly1 correspond to R259, H167, H146, and Y243 of LOR_107, respectively. To confirm the role of these residues in Uly1 catalysis, these four residues were mutated to alanine, and the activities of the mutants were measured. Based on the structural alignment, the basic amino acid residues H149 and R239 extend toward the carboxylic group of GlcA at the +1 position and can interact with it (Fig. 6A). Mutations on H149 and R239 decreased the activity of Uly1 by approximately 50% (Fig. 6C), indicating that H149 and R239 are likely to act as neutralizers to the negative charge of the +1 carboxyl group. H128 is close to the C-5 atom at subsite +1, and the glucosidic bond (Fig. 6A) and the mutation H128 to alanine almost completely abolished the enzymatic activity of Uly1 (Fig. 6C), indicating

**FIG 4** The enzymatic products of ulvan generated by Uly1 over a time course of 0 to 12 h. (A) Gel filtration analysis of the enzymatic products over a time course of 0 to 12 h. Gel filtration chromatography analysis was performed using a Superdex peptide 10/300 GL column monitored at a wavelength of 210 nm. Ulvan treated with Uly1 that was preactivated by boiling was taken as the control. (B) ESI-MS analysis of the major peak indicated by an arrow in panel A. The major peak has an m/z value of 401, consistent with the molecular weight of the disaccharide ΔRha3S.
that H128 plays a crucial role in catalysis. As with H128, Y223 is also close to the glucosidic bond (Fig. 6A). However, when Y223 was mutated to alanine, the enzyme activity decreased only approximately 40% (Fig. 6C), suggesting that Y223 plays a supporting role in catalysis. Y223 may act as an alternative catalytic acid or help properly orient the imidazole ring of H128, as Y243 does in LOR_107 (Fig. 6A) (18). Therefore, in the catalysis of ulvan, H128 of Uly1 is most likely to be both the catalytic acid and base, H149 and R239 neutralize the charge of the GlcA acidic group, and Y223 plays a supporting role in catalysis.

**Comparison of the residues involved in the substrate binding of Uly1 with those of LOR_107.** LOR_107 adopts a two-stage substrate binding process containing a substrate sliding event (18). During this process, N263 of LOR_107 plays a crucial role by moving its side chain, making the catalytic canyon narrower and pinning the substrate to the active site (Fig. 7A) (18). However, it was found that in Uly1, the side chain of S243, which spatially corresponds to N263 of LOR_107, is too short to fulfill the same function as N263 of LOR_107 (Fig. 7B). It was also found that the substrate-binding canyon of Uly1 is much narrower than that of LOR_107, mainly caused by three
loops on one side of the active center. These loops (L1, residues 300 to 314; L2, residues 361 to 369; and L3, residues 389 to 392) are closer to the other side of the canyon in Uly1 than those in LOR_107 and show low flexibility, with relatively low B factors (Fig. 7C). R308 on the L1 of Uly1 occupies the position of the modeled carboxyl group of the −2 site GlcA in the structural alignment of Uly1 with the LOR_107 mutant (R259N) with tetrasaccharide (PDB code 6BYX) and LOR_107 mutant (R320N) with tetrasaccharide (PDB code 6BYT) are colored in yellow, gray, purple, and green, respectively. The bound tetrasaccharides in the complex structures of R259N and R320N occupying the −2 to +2 subsites are shown as salmon and red sticks, respectively. The residues of Uly1 and LOR_107 are labeled in orange and gray letters, respectively. (B) Sequence alignment of Uly1 and characterized PL24 ulvan lyases from the CAZy database. Identical and similar amino acid residues are shaded. Red dots indicate the residues involved in catalysis. The noncatalytic domains of the long ulvan lyases (ulvan lyases LOR_61 and PLSV_3925) are not included. (C) Enzymatic activities of WT Uly1 and its mutants toward ulvan. The activity of WT Uly1 was taken as 100%.

FIG 6 Analysis of the important residues in the active site of Uly1. (A) Structural alignment of the important residues of Uly1 and the ulvan lyase LOR_107. The amino acid residues in the structure of Uly1, wild-type LOR_107 structure, and the complex structures of the LOR_107 mutant (R259N) with tetrasaccharide (PDB code 6BYX) and LOR_107 mutant (R320N) with tetrasaccharide (PDB code 6BYT) are colored in yellow, gray, purple, and green, respectively. The bound tetrasaccharides in the complex structures of R259N and R320N occupying the −2 to +2 subsites are shown as salmon and red sticks, respectively. The residues of Uly1 and LOR_107 are labeled in orange and gray letters, respectively. (B) Sequence alignment of Uly1 and characterized PL24 ulvan lyases from the CAZy database. Identical and similar amino acid residues are shaded. Red dots indicate the residues involved in catalysis. The noncatalytic domains of the long ulvan lyases (ulvan lyases LOR_61 and PLSV_3925) are not included. (C) Enzymatic activities of WT Uly1 and its mutants toward ulvan. The activity of WT Uly1 was taken as 100%.
In LOR_107, R320 coordinates the initial binding of the substrate (Fig. 7A) (18). When R320 of LOR_107 was mutated to alanine, the mutation resulted in a complete loss of the enzyme activity (18). However, when R300 in Uly1, corresponding to R320 in LOR_107, was mutated to alanine, the activity of the mutant was clearly detectable (remained ~20%) (Fig. 6C), suggesting that R300 plays an important role in the catalysis of Uly1 but may not play a similar role as R320 does in LOR_107.

Altogether, based on these analyses, it is likely that Uly1 directly binds the substrate near the catalytic residues for catalysis, unlike LOR_107, which undergoes a two-stage substrate binding event in the catalytic process (18).

The proposed substrate binding and catalytic mechanisms of Uly1. Based on the above structural and biochemical results, the possible substrate binding and catalytic mechanisms of Uly1 for ulvan depolymerization are proposed (Fig. 8). Uly1 possesses a narrower catalytic canyon than that reported for the PL24 ulvan lyase LOR_107. When the substrate approaches the catalytic canyon, the residues involved in substrate binding make the substrate directly bind near the catalytic residues. The key residues H128, H149, R239, and Y223 then mediate the catalytic reaction. H149 and R239 neutralize the negative charge on the carboxylic group of the +1 subsite, H128 acts as both the catalytic acid and base, and Y223 plays a supporting role in catalysis. Although the molecular mechanism of Uly1 catalysis is similar to that of LOR_107, the substrate binding process of Uly1 is likely significantly different from the reported two-stage process of LOR_107 due to differences in the width of the catalytic canyons and the arrangement of amino acids in the catalytic canyons of these two enzymes.

Different substrate binding processes in the PL24 family. Among the ulvan lyase genes, genes encoding the PL24 family ulvan lyases are the most abundant. The PL24 family includes 50 ulvan lyase sequences, but only 5 (including Uly1) have been characterized. Results presented here suggest that Uly1 likely adopts a different substrate binding process from LOR_107. To investigate the universality of the two substrate binding processes in PL24 ulvan lyases, a sequence alignment of PL24 ulvan lyase sequences was performed. The results show that S243 and R308 of Uly1 are conserved.
in many PL24 ulvan lyases, and N263 and K329 of LOR_107 are conserved in many other PL24 lyases (Fig. 9). This implies that there are likely two types of substrate binding processes in the PL24 lyases. One type, represented by LOR_107, adopts a two-stage substrate binding process, while the other type, represented by Uly1, may directly bind the substrate near the catalytic amino acid residues for catalysis. The results in this study will provide a better understanding of the PL24 ulvan lyases and ulvan degradation and will be helpful for developing enzymatic tools for the utilization of green alga polysaccharides.

**MATERIALS AND METHODS**

**Gene synthesis, cloning, and mutagenesis.** The 1,539-bp full-length sequence of the Uly1 gene (GenBank accession no. KMT65093.1) from the marine bacterium *Catenovulum maritimum* Q17, which is
available in the CAZy database, was synthesized by the Beijing Genomics Institute (China). The gene without the signal peptide was then subcloned into the pET-22b vector (TaKaRa, Japan) with a C-terminal His tag. Site-specific mutagenesis was carried out by a QuikChange site-directed mutagenesis kit (Agilent, USA) using the plasmid pET-22b-Uly1 as the template. All the recombinant plasmids were verified by sequencing.

**Protein expression and purification.** Recombinant proteins of WT Uly1 and its mutants were overexpressed in *E. coli* BL21(DE3) (Vazyme, China) and were induced by the addition of 0.3 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 15°C for 16 h. Cells were collected and disrupted by sonication in buffer containing 50 mM Tris-HCl buffer (pH 8.0) and 100 mM NaCl. The SeMet derivative of Uly1 was overexpressed in *E. coli* BL21(DE3) in M9 minimal medium containing 100 mg/liter of lysine, phenylalanine, and threonine, 50 mg/liter of isoleucine, leucine, and valine, 5.2% (wt/vol) glucose, and 0.65% (wt/vol) yeast nitrogen base (YNB), which was then grown at 37°C. When the optical density at 600 nm (OD600) reached 0.6, the culture was cooled to 15°C and 50 mg/liter L-SeMet was added. Fifteen minutes later, the culture was incubated at 15°C at 100 rpm for 16 h under the induction of 0.4 mM IPTG. The recombinant proteins were first purified by nickel-nitrilotriacetic acid resin (Qiagen, Germany) and then by gel filtration chromatography on a Superdex 200 column (GE Healthcare, USA) in buffer containing 10 mM Tris-HCl (pH 8.0) and 100 mM NaCl. Carbonic anhydrase (29 kDa) and conalbumin (75 kDa) as the protein size standards from GE Healthcare were calibrated in the same buffer in the column.

**Biochemical characterization of Uly1.** The concentrations of Uly1 and its mutants were determined by the bicinchoninic acid (BCA) protein assay kit (Thermo, USA), with bovine serum albumin (BSA) as the standard. The metal ions in Uly1 were quantified using ICP-OES (23). The activities of WT Uly1 and its mutants toward ulvan (purchased from Elicityl, France) were measured by the dinitrosalicylic acid (DNS) method (11). The reaction system was composed of 50 mM Tris-HCl buffer (pH 9.0), 10 mg/ml substrate, and 0.5 mg/ml enzyme. The reaction was carried out at 40°C for 30 min at a final volume of 200 μl and terminated by the addition of 100 μl DNS. The reaction mixture was boiled at 100°C for 10 min and immediately cooled on ice for 2 min. Then, the absorbance values were determined at 540 nm. The amount of reducing sugars released into the mixture was determined with rhamnose as the standard. One unit of enzyme activity is defined as the amount of enzyme required to release 1 nmol reducing sugars per min (24).

The optimum temperature for Uly1 activity was determined at a range of 10 to 60°C at pH 9.0. The optimum pH for Uly1 activity was determined at 40°C in Britton-Robinson (B-R) buffer ranging from pH 5.0 to 12.0. B-R buffer was prepared with boric acid, acetic acid, and phosphoric acid, all at a final concentration of 0.04 M in an aqueous solution and was adjusted to different pHs with 0.2 M NaOH. The effect of NaCl on Uly1 activity was determined at NaCl concentrations ranging from 0 to 4.0 M.

The action mode and degradation products of Uly1 were analyzed by using ulvan as the substrate. The degradation reaction was carried out at 40°C for different times, from 0 to 12 h. The concentrations of the enzyme and substrate used were both 1 mg/ml. The ulvan degradation products were analyzed by gel filtration chromatography on a Superdex Peptide 10/300 GL column (GE Healthcare, USA) at a flow rate of 0.3 ml/min using 0.2 M ammonium hydrogen carbonate as the running buffer (25, 26). Elution was monitored at 210 nm using a UV detector. LabSolutions software was used for online monitoring and data analysis. The major peak was further identified by ESI-MS on an ion trap time of flight (TOF) hybrid mass spectrometer (LCMS-IT-TOF; Shimadzu, Japan). ESI-MS analysis was set in the

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**FIG 9** Sequence alignment of Uly1 and the other PL24 lyases. S243 and R308 of Uly1 and the corresponding residues of the other PL24 lyases are indicated with black triangles.
negative-ion mode and with the following parameters: source voltage at 3.6 kV, nebulizer nitrogen gas flow rate at 1.5 liter/min, heat block, curve desolvation line temperature at 200°C, and detector voltage at 1.8 kV. The mass acquisition range was set at 200 to 800.

Crystallization and data collection. The purified Uly1 protein was concentrated to ~10 mg/ml in the buffer containing 10 mM Tris-HCl (pH 8.0) and 100 mM NaCl. Crystals of WT Uly1 were obtained at 18°C using the sitting drop method in the buffer containing 200 mM calcium acetate hydrate (pH 6.5), 18% (wt/vol) polyethylene glycol 8000 (PEG 8000), and 100 mM sodium cacodylate trihydrate. Crystals of SeMet Uly1 were grown in the buffer containing 200 mM calcium acetate, 100 mM MES (morpholineethanesulfonic acid)-NaOH (pH 6.0), and 20% (wt/vol) PEG 8000. All the X-ray diffraction data were collected on the BL17U1 beamline at the Shanghai Synchrotron Radiation Facility using Area Detector Systems Corporation Quantum 315i. The initial diffraction data sets were processed by the HKL2000 program. Relevant data collection statistics are shown in Table 2.

Structure determination and refinement. Heavy atoms were searched by SHELXD. The phase problems were solved by the single-wavelength anomalous diffraction (SAD) method using Phenix program Autosol (27). Initial model building was finished by Phenix program AutoBuild (27). Refinement of these structures was performed using Coot and Phenix alternately (27, 28). The quality of the final model is summarized in Table 2. All structure figures were generated using PyMOL software. The RMSD values were calculated using a tool on the PDB database (https://www.rcsb.org/alignment).

Data availability. The atomic coordinates and structure factors of WT Uly1 (PDB code 7CZH) and SeMet Uly1 (PDB code 7DRQ) have been deposited in the Protein Data Bank.

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F.X., F.D. and X.-H.S. performed all experiments. P.W., Y.-Z.Z., X.-L.C., H.-H.F., and X.-Y.Z. directed the experiments. F.X. and F.D. wrote the manuscript. H.-Y.C., P.W., and C.-Y.L. solved the structures and analyzed the data. Y.-Z.Z. and X.-L.C. designed the research. X.-L.C. and A.M. edited the manuscript.

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