Novel Molecular Insights into the Catalytic Mechanism of Marine Bacterial Alginate Lyase AlyGC from Polysaccharide Lyase Family 6*

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Alginate lyases that degrade alginate via a β-elimination reaction fall into seven polysaccharide lyase (PL) families. Although the structures and catalytic mechanisms of alginate lyases in the other PL families have been clarified, those in family PL6 have yet to be revealed. Here, the crystal structure of AlyGC, a PL6 alginate lyase from marine bacterium Glaciecola chathamensis S18K62, was solved, and its catalytic mechanism was illustrated. AlyGC is a homodimeric enzyme and adopts a structure distinct from other alginate lyases. Each monomer contains a catalytic N-terminal domain and a functionally unknown C-terminal domain. A combined structural and mutational analysis using the structures of AlyGC and of an inactive mutant R241A in complex with an alginate tetrasaccharide indicates that conformational changes occur in AlyGC when a substrate is bound and that the two active centers in AlyGC may not bind substrates simultaneously. The C-terminal domain is shown to be essential for the dimerization and the catalytic activity of AlyGC. Residues Tyr130, Arg187, His242, Arg265, and Tyr304 in the active center are also important for the activity of AlyGC. In catalysis, Lys220 and Arg241 function as the Brønsted base and acid, respectively, and a Ca2+ in the active center neutralizes the negative charge of the C5 carboxyl group of the substrate. Finally, based on our data, we propose a metal ion-assisted catalytic mechanism of AlyGC for alginate cleavage with a state change mode, which provides a better understanding for polysaccharide lyases and alginate degradation.

Brown algae are an important source of primary production in the marine ecosystem and represent a huge marine biomass (1). Alginates are the major polysaccharides produced by brown algae, which may reach 40% of the dry weight of algal biomass (2). Algin is a linear polysaccharide composed of α-L-guluronate (G)2 and its C5 epimer β-D-mannuronate (M), which are arranged in three ways, polyguluronate (PG), polymannuronate (PM), and alternating GM or random heteropolymorphic M/G stretches (P(MG)) (3). Alginates are widely used in the food, chemical, and pharmaceutical industries because of their ability to form gels and to chelate metal ions (4–6). In addition, alginates are also a major constituent of biofilm produced by some heterotrophic bacteria from the genera Pseudomonas and Azotobacter (7). Alginate lyases are synthesized by brown seaweeds, marine molluscs, and a variety of microbes (8). They play an important role in the marine carbon cycle and have important applications in biotechnological and chemotherapeutic fields, such as preparation of functional oligosaccharides (9) and protoplasts of algae (10) and treatment of cystic fibrosis (11, 12). With the discovery and characterization of novel enzymes, further applications of alginate lyases may be found. Alginate lyases degrade alginates through a β-elimination reaction, targeting the glycosidic 1→4 O-linkage between the monomers. A double bond is formed between C4 and C5, yielding a 4-deoxy-L-erythro-hex-4-enopyranosyluronic acid at the nonreducing end (13). According to their substrate specificities, alginate lyases are classified into three types, PM-specific lyases (EC 4.2.2.3), PG-specific lyases (EC 4.2.2.11), and bifunctional lyases that can degrade both PM and PG (EC 4.2.2.1). Most alginate lyases studied are endolytic enzymes that cleave glycosidic bonds inside polymers and release unsaturated oligosaccharides, and only a few exolytic alginate lyases that remove monomers or dimers from the ends of polymers are reported (14–17). In the Carbohydrate-Active enZymes database (CAZY database), alginate lyases are distributed in seven polysaccharide lyase (PL) families (PL5, -6, -7, -14, -15, -17, and -18) (18). Whereas those of alginate lyases from PL5, -7, -14, -15, -17, and -18 have been reported, the three-dimensional structures and catalytic mechanisms of the PL6 alginate lyases still remain unknown (7, 15, 19–22). Clarifying the structure and catalytic

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2 The abbreviations used are: G, α-L-guluronate; PL, polysaccharide lyase; M, β-D-mannuronate; NTD, N-terminal domain; CTD, C-terminal domain; PDB, Protein Data Bank; TBA, thiorbitaric acid; r.m.s.d., root mean square deviation; M4, α-tetramannuronate acid tetrasodium salt; DLS, dynamic light scattering; DS, dermatan sulfate; PG, polyguluronate; PM, polymannuronate.

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The atomic coordinates and structure factors (codes 5GKD and 5SGK) have been deposited in the Protein Data Bank (http://wwpdb.org/).
mechanism of the PL6 alginate lyases will broaden our understanding on alginate lyases.

In the PL6 family, only three alginate lyases have been characterized (23–25). AlyMG (475 amino acid residues) is a polyMG-specific alginate lyase from *Stenotrophomas maltophilia* KJ-2 (24). AlyP (398 amino acid residues) from *Pseudomonas* sp. OS-ALG-9 has a greater specificity to PM than to PG (23). OalS6 (named AlgS6 in the NCBI Protein Database, but changed to OalS6 when published) (770 amino acid residues) from *Shewanella* sp. Kz7 is an exo-type oligoalginate lyase that prefers to depolymerize the PG block (25). In addition to alginate lyases, the PL6 family also contains a chondroitinase B (ChonB, also named CslB) (506 amino acid residues) from *Pedobacter heparinus*. ChonB is a glycosaminoglycan (GAG) lyase with disaccharide polymer dermatan sulfate (DS) as its sole substrate. ChonB adopts a right-handed β-helix fold and has a calcium-dependent catalytic machinery (26, 27).

In this study, a PL6 alginate lyase, AlyGC, from marine bacterium *Glaciecola chathamensis* S18K6T (28, 29) was characterized, and the structures of wild-type (WT) AlyGC (2.2 Å resolution) and of an inactive mutant in complex with an alginate tetrasaccharide (2.6 Å resolution) were solved. Based on structural and mutational analyses, the molecular mechanism of AlyGC for substrate catalysis was explained. The results provide a better understanding of the PL6 alginate lyases.

**Results and Discussion**

Two-domain Structure of AlyGC Predicted by Sequence Analysis—The gene *alyGC* predicted to encode a PL6 alginate lyase (AlyGC) was cloned from the genome of *G. chathamensis* S18K6T. *alyGC* is 2265 bp in length and encodes a protein of 754 amino acid residues containing a predicted 28-residue signal peptide. According to the blast result against the NCBI non-redundant protein database, the putative protein AlyGC (unless otherwise stated, AlyGC discussed hereafter is 727 aa in length without the predicted signal peptide) shows the highest identity (58%) to OalS6, a characterized PL6 alginate lyase from *Shewanella* sp. Kz7 (25). Sequence analysis using Conserved

![FIGURE 1. Schematic domain diagram of PL6 enzymes.](image)

**FIGURE 1. Schematic domain diagram of PL6 enzymes.** The signal peptides were predicted by SignalP 4.1 Server. ChonB (GenBank™ ACU03011.1; PDB code 1DBG) is from *P. heparinus* DSM 2366; AlyP (GenBank™ BAA01182.1) is from *Pseudomonas* sp. OS-ALG-9; AlyMG (GenBank™ AFC88009.1) is from *S. maltophilia* KJ-2; OalS6 (GenBank™ AHC69713.1) is from *Shewanella* sp. Kz7; and AlyGC (GenBank™ BAEM00000000.1) is from *Glaciecola chathamensis* S18K6T. 
Domain Search suggests that AlyGC contains two domains, an N-terminal domain (NTD, Met1–Asn373) and a C-terminal domain (CTD, Gln374–Leu727). The NTD of AlyGC belongs to the PL6 family; however, the CTD of AlyGC does not display obvious similarity to any known functional protein sequences. Although not reported, sequence analysis indicates that the precursor of alginate lyase OalS6 also has a similar CTD with unidentified function (Fig. 1). Thus, these two CTDs may represent an uncharacterized protein domain. Taken together, sequence analysis indicates that AlyGC is a PL6 alginate lyase with a two-domain structure.

Characterization of AlyGC—AlyGC without the predicted signal peptide was overexpressed in *Escherichia coli*. Because AlyGC is a member of the PL6 family that contains alginate lyase and chondroitinase B, the activities of recombinant AlyGC toward alginate sodium and DS were measured. AlyGC showed activity toward alginate sodium but no detectable activity toward DS (Fig. 2A), indicating that AlyGC is an alginate lyase, rather than a GAG lyase. Among alginate sodium, PM and PG, AlyGC displayed the highest activity toward PG, indicating its preference to PG (Fig. 2A). Thin layer chromatography (TLC) analysis showed that AlyGC released monosaccha-
rides from PG (Fig. 2B), indicating that AlyGC is an exo-type lyase, consistent with that reported for OalS6 (25). With alginate sodium as substrate, AlyGC exhibited the highest activity at 30 °C and pH 7.0 (Fig. 2, C and D). Although from a marine bacterium, AlyGC showed very low level of salt tolerance (Fig. 2E).

Overall Structure of AlyGC—A similarity search at Protein Data Bank (PDB) revealed that the closest homologue to AlyGC is ChonB (PDB code 1DBG), with 29% sequence identity (covering only 46% of the AlyGC sequence), which indicates that no suitable structure model can be used for AlyGC structure construction. Therefore, to solve the structure of AlyGC, the crystals of both WT AlyGC and SeMet-AlyGC were obtained. Then HKL2MAP was used for heavy atom searching, and Phenix.autosol was used for phasing and density modification. The values of the figure of merit (FOM) and BAYES-CC of the phasing result are 0.348 and 39.88, respectively. Finally, combined with the data of WT AlyGC, the structure of AlyGC was solved at 2.2 Å resolution. The AlyGC structure belongs to the P21 space group. Four AlyGC molecules (each contains 727 residues) are found in one asymmetric unit. Only two of the four have interactions (Fig. 3A), and the other two do not show the same symmetry-related interaction. In addition, both gel filtration and dynamic light scattering (DLS) analyses indicate that AlyGC presents as dimers in solution (Fig. 3, B and C). Therefore, AlyGC is a dimeric enzyme. The electron density of AlyGC
starting with the first PB1 strand (Glu$^{36}$–Lys$^{42}$) and finishing with the penultimate PB3 strand (Asn$^{370}$–Asn$^{373}$), which includes 12 PB1 strands, 14 PB2 strands, and 14 PB3 strands. The CTD consists of eight coils starting with the first PB1 strand (Leu$^{478}$–Leu$^{486}$) and finishing with the penultimate PB3 strand (Val$^{699}$–Glu$^{701}$), which includes 9 PB1 strands, 11 PB2 strands, and 9 PB3 strands. There are only two $\alpha$-helices in the structure of AlyGC, designated HA1 and HA2. The NTD is capped by HA1 (Pro$^{9}$–Lys$^{19}$) preceding the first PB1 strand and the CTD by HA2 (Thr$^{451}$–Ser$^{458}$) preceding the first PB1 strand. The linker between the NTD and the CTD consists of two $\beta$-strands followed by a long loop of 42 residues. The Ca$^{2+}$ is located in the T3-PB1-T1 region of coils 6 and 7 in the NTD.

A structure-based homology search for AlyGC was performed using the DALI server (35). The result indicates that ChonB is the closest structural homologue with a Z-score of 39.7 and a root mean square deviation (r.m.s.d.) of 2.3 on 395 aligned Ca residues. Other structural homologues include the mannuronan C5 epimerase AlgE4 from Azotobacter vinelandii (PDB code 2PYG; Z-score: 28.4, and r.m.s.d.: 2.8 on 310 residues) (36) and the K5 lyase from Escherichia virus k1–5 (PDB code 2X3H; Z-score: 26.6, and r.m.s.d.: 2.8 on 301 residues) (37). AlyGC exhibits similar topology with these proteins. However, these homologues adopt simple right-handed parallel $\beta$-helix folds, whereas AlyGC adopts a tandem $\beta$-helix fold. ChonB has an $\alpha$-helix (Leu$^{360}$–Arg$^{372}$) in the catalytic cleft, which, however, is lacking AlyGC. Because this $\alpha$-helix in ChonB is an important component of the catalytic cleft and residue Arg$^{362}$ on this $\alpha$-helix directly interacts with the substrate DS (26, 27), this structural difference between ChonB and AlyGC may be related to their different substrate specificity (Fig. 4A). In addition, the catalytic cleft of ChonB is more L-like compared with that of AlyGC (Fig. 4, B and C).

Conformational Change in AlyGC When Binding an Alginate Tetrasaccharide—To obtain an inactive AlyGC mutant for enzyme-substrate complex crystallization, site-directed mutations on AlyGC were conducted based on structural analysis. The result indicated that when Lys$^{220}$ or Arg$^{241}$ was mutated to alanine, the mutated AlyGC was almost inactive (<5% activity of WT AlyGC) toward PG. Thus, mutants K220A and R241A were crystallized with alginate oligosaccharides, respectively. Finally, the crystal structure of R241A in complex with tetramannuronic acid (M4) was solved to 2.6 Å resolution. There are two R241A molecules in one asymmetric unit of the crystal structure R241A-M4, each containing a Ca$^{2+}$ (Fig. 5A). Of the two active centers ($\alpha$ and $\beta$) in the dimeric R241A-M4, only the active center $\alpha$ binds an M4 beside the Ca$^{2+}$ (Fig. 5B). Structural comparison of R241A-M4 and WT AlyGC indicates that conformational changes occur in the R241A-M4 dimer (Fig. 5C). Compared with WT AlyGC, the entrance of the active center $\alpha$ that binds the M4 is enlarged and that of the active center $\beta$ without substrate is smaller. To characterize this, we compared the closest distances between two loops (1 and 2) at the entrances of the catalytic clefts in WT AlyGC and R241A-M4. Loop 1 (Ile$^{188}$–Leu$^{201}$) is from the NTD of one monomer, and loop 2 (Asp$^{520}$–Ile$^{523}$) is from the CTD of another monomer. The closest distance between loops 1 and 2 is that between His$^{192}$ and Asp$^{526}$ in AlyGC structures. In WT AlyGC, the dis-
tance between His192 and Asp526 near the active center (5.6 Å) is similar to that near the active center (4.4 Å). However, in R241A-M4, the distance between His192 and Asp526 near the active center (11.8 Å) and that near the active center (3.3 Å) (Fig. 5D), much smaller than the diameter of a carbohydrate chain. Therefore, based on the crystal structure of R241A-M4, it seems that the dimeric AlyGC can only accommodate one substrate molecule in one of the two active centers, although we cannot ensure that it is this case when AlyGC catalyzes substrates in solution.

Function of the CTD—The CTD in AlyGC has no conserved sequence or predicted function. To investigate the role of the CTD, a CTD-truncated mutation of AlyGC, ΔCTD (Met1–Asp434), was constructed. Gel filtration analysis showed that ΔCTD presents as monomers in solution (Fig. 6A), indicating that the CTD is essential for the dimerization of AlyGC. Compared with WT AlyGC, ΔCTD lost 94.3% activity toward PG (Fig. 6B), suggesting that, for AlyGC activity, the CTD is indispensable, and the dimerization of AlyGC is also necessary. Structural analysis shows that a loop from the CTD (Arg627–His638) stretches into the catalytic center. Mutations of the residues Asp631 and Ser633 on this loop to alanines led to significant decreases in the enzyme activity (Fig. 6B). Altogether, our data indicate that the CTD of AlyGC is crucial for its dimerization and catalytic activity.
AlyGC is essential for the dimerization and the catalytic activity of AlyGC.

Role of the Ca\(^{2+}\) in Catalysis—ChonB, the only enzyme with a solved structure in PL6 family, contains a Ca\(^{2+}\) coordinated by Asn\(^{213}\), Glu\(^{243}\), and Glu\(^{245}\) (26, 27). The Ca\(^{2+}\) in AlyGC is coordinated by Asn\(^{181}\), Glu\(^{213}\), Glu\(^{215}\), and Glu\(^{184}\), all of which are conserved in the characterized PL6 alginate lyases (Figs. 7 and 8A). Among these residues, Asn\(^{181}\), Glu\(^{213}\), and Glu\(^{215}\) correspond to the coordinative residues of Ca\(^{2+}\) in ChonB (Fig. 7). In the R241A-M4 complex structure, the Ca\(^{2+}\) also interacts with the carboxyl group of the substrate. These structural data suggest that the Ca\(^{2+}\) in AlyGC may be involved in catalysis, just as that in ChonB (26). To support this, we performed site-directed mutations on the residues coordinating the Ca\(^{2+}\). Substitution of any of the four residues to alanine resulted in severe loss in enzymatic activity (Fig. 8B). Furthermore, when metal chelator EDTA was added to the reaction mixture, the enzyme activity decreased with the increase of EDTA concentration, and 0.2 mM EDTA completely abolished the enzyme activity (Fig. 8C). However, the activity of apo-AlyGC in which the Ca\(^{2+}\) was completely depleted by EDTA could be recovered by the addition of Ca\(^{2+}\) and almost fully recovered by 0.2 mM Ca\(^{2+}\). In contrast, Mg\(^{2+}\) could only recover 33.4% of the enzyme activity, and Mn\(^{2+}\) had no effect on the recovery of apo-AlyGC activity (Fig. 8D). Taken together, these data indicate that the Ca\(^{2+}\) in AlyGC is the biological metal ion and is involved in catalysis.

Important Residues in the NTD for Substrate Catalysis—The oligosaccharide is bound in the active center of AlyGC as shown in Fig. 5 and mainly interacts with the T1-PB1-T3 region of the NTD. One end of the catalytic groove of AlyGC is nearly blocked, suggesting that AlyGC is an exo-type lyase, corresponding to the TLC result (Fig. 2B). We adopt the nomenclature proposed by Davies et al. (38) by convention. \(-n\) and \(+n\) represent the nonreducing terminus and the reducing terminus, respectively, and subsites are labeled from \(-n\) to \(+n\). Therefore, the tetrasaccharide binding to AlyGC is positioned at subsites \(-1\), \(-1\), \(-2\), and \(-3\), and the constituent mannuronate residues are named from M\(^{-}\) to M\(^{+}\) (Fig. 7). Mutation of Lys\(^{220}\) or Arg\(^{241}\) to alanine led to complete loss of the enzyme activity (Fig. 8B). Therefore, according to these data and the mechanism of \(\beta\)-elimination reaction, Lys\(^{220}\) is the Brønsted base, and Arg\(^{241}\) is the Brønsted acid in the cleavage reaction.
of AlyGC on alginate, just as the corresponding residues in ChonB (26).

According to the WT AlyGC and R241A-M4 structures, the hydrophilic residues Tyr130, Arg187, His242, Arg265, and Tyr304 in the active center (Fig. 8A) may interact with the substrate. Residues Tyr130, Arg187, Arg265, and Tyr304 are conserved in all characterized PL6 alginate lyases, and His242 is conserved in all characterized PL6 enzymes, including ChonB (Fig. 7). Site-directed mutations of these residues to alanine decreased the activity of AlyGC (Fig. 8B). These data indicate that these hydrophilic residues are important for the activity of AlyGC.

Circular dichroism (CD) spectra show that the curves of all the variants are similar to that of WT AlyGC, suggesting that WT AlyGC and the variants have similar secondary structures. Therefore, the activity loss in the variants is caused by amino acid replacement, rather than by structural change (Fig. 8E). These data indicate that these hydrophilic residues are important for the activity of AlyGC.

Catalytic Mechanism of AlyGC—Based on our structural and biochemical results on AlyGC, we propose a metal ion-assisted mechanism of this PL6 alginate lyase for alginate cleavage with a possible state change. In the absence of substrate, AlyGC is in the resting state, in which the sizes of the entrances of the two active centers are similar (Fig. 9A). When alginate or an oligosaccharide enters one of the catalytic cavities, the active center binding the substrate is enlarged and the other one is smaller. The enzyme is in the active state. The enzyme molecule returns to the resting state.

FIGURE 9. Catalytic mechanism of AlyGC. A, model of the catalytic mode of AlyGC. Step 1, the resting state. In the absence of substrate, the sizes of the entrances of the two active centers of AlyGC are similar. The whole enzyme is presented as surface view with one monomer colored in pale cyan and another in wheat. Loop 1 is colored in dark blue, and the loop 2 is colored in pink. A schematic diagram is presented on the right. Step 2, the active state. A substrate enters one of the catalytic cavities. The active center binding the substrate is enlarged, and the other one is smaller. The enzyme is in the active state. The substrate is shown as a red sphere in the right schematic diagram. Step 3, return to the resting state. The product is released and the enzyme molecule returns to the resting state. B, catalytic mechanism of AlyGC on alginate degradation. The Ca²⁺ forms interactions with the carboxyl group of the A₁+ and activates the Cα hydrogen of A₁+ and Lys²⁰⁵ functions as a nucleophilic base to attack the Cα of A₁+, and Arg²⁴¹ functions as the Brønsted acid to accept an electron. Electron transfer is presented with red arrows.

of AlyGC on alginate, just as the corresponding residues in ChonB (26).

According to the WT AlyGC and R241A-M4 structures, the hydrophilic residues Tyr¹³⁰, Arg¹⁸⁷, His²⁴², Arg²⁶⁵, and Tyr³⁰⁴ in the active center (Fig. 8A) may interact with the substrate. Residues Tyr¹³⁰, Arg¹⁸⁷, Arg²⁶⁵, and Tyr³⁰⁴ are conserved in all characterized PL6 alginate lyases, and His²⁴² is conserved in all characterized PL6 enzymes, including ChonB (Fig. 7). Site-directed mutations of these residues to alanine decreased the activity of AlyGC (Fig. 8B). These data indicate that these hydrophilic residues are important for the activity of AlyGC.

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Structure and Catalytic Mechanism Comparison of Alginate Lyases—Alginate lyases fall into seven PL families, including PL5-7, -14, -15, -17, and -18 (18). With the three-dimensional structures of WT AlyGC and its complex being solved, at least one alginate lyase structure has been revealed in each family.
A phylogenetic tree was constructed for the characterized alginate lyases in all the seven PL families, in which each family contains at least one alginate lyase with reported structure. Although alginate lyases are classified into seven PL families according to their primary structures, their three-dimensional structures can be grouped into 3-fold types. The alginate lyases from PL5, -15, and -17 adopt a (α/α)n toroid fold; those from PL7, -14, and -18 adopt a β-jelly roll fold, and AlyGC from PL6 adopts a parallel β-helix fold.

**FIGURE 10.** Comparison of structures and catalytic mechanisms of alginate lyases. A, overall structures of the alginate lyases in seven PL families. PL6 alginate lyase AlyGC (5GKD) is in yellow; PL5 alginate lyase AI-II (1QAZ) is in gray; PL7 alginate lyase AI-II (2CWS) is in wheat; PL14 alginate lyase vAL-1 (3A0N) is in green; PL15 alginate lyase Atu3025 (3AFL) is in blue; PL17 alginate lyase Alg17c (4NEI) is in pink; and PL18 alginate lyase aly-SJ02 (4Q8K) is in cyan. Brønsted acids and bases are shown as red sticks, and residues for neutralization are shown as yellow sticks. B, maximum likelihood tree of the characterized alginate lyases. PL14 alginate lyases circled with dotted line are from viruses or eukaryotes. Other lyases are from prokaryotes. One or two lyases with solved structures are shown in red as representatives in each family. C, schematic diagram of the histidine/tyrosine catalytic mechanisms of alginate lyases except for PL6 alginate lyases. Asx (Asp or Asn), Glx (Glu or Gln), His, or Arg neutralizes the acidic carboxyl group. Tyr functions as a Brønsted acid, and Tyr or His functions as a Brønsted base.
PL6 adopts a tandem parallel β-helix fold (Fig. 10, A and B). Thus, AlyGC of the PL6 family represents a new alginate lyase fold.

In addition, the catalytic mechanism of AlyGC is also significantly different from those of other alginate lyases. Previous studies showed that alginate lyases from the other families adopt the Tyr/His (Tyr) mechanism to catalyze a typical alginate lyase reaction. In the reaction, residues Asx (Asp/Asn), Glx (Glu/Gln), His, or Arg neutralize the negative charge of C5 carboxyl group to lower the pKa of the C5 proton; a His (sometimes a Tyr) functions as the general base to attract the C5 proton, and a Tyr functions as the general acid to donate a proton to the O4 atom, thus forming a double bond between C4 and C5 (Fig. 10C) (8, 39). However, for AlyGC, a Ca²⁺, rather than a residue, neutralizes the negative charge of C5 carboxyl group; a Lys acts as the base, and an Arg acts as the acid. Thus, AlyGC of the PL6 family adopts a catalytic mechanism different from that of the other families.

**Experimental Procedures**

**Materials and Strains**—Chondroitin sulfate B sodium salt from porcine intestinal mucosa and sodium alginate (viscosity, 15–20 cps) were purchased from Sigma. PolyM, polyG (6–8 kDa), and oligosaccharide substrates (purity ≥97%) were purchased from Zzstandard (China). Strain *G. chathamensis* S18K6 was ordered from China General Microbiological Culture Collection Center (CGMCC) and grown in marine broth 2216 (BD Biosciences-Difco, America) at 20 °C. *E. coli* was cultured in Luria-Bertani (LB) medium at 37 °C.

**Gene Cloning and Mutagenesis**—The genome DNA of *G. chathamensis* S18K6 was previously shotgun-sequenced and submitted to NCBI (GenBank™ BAEM00000000.1). Gene (WP_007984897.1) (22) in this genome was deduced to encode a PL6 alginate lyase. This gene was named *alyGC* in this study. The *alyGC* gene was amplified from the genomic DNA of *G. chathamensis* S18K6 via PCR and cloned into the vector pET-22b-alyGC as the template by using a QuickChange kit (Agilent Technologies).

**Protein Expression and Purification**—Recombinant proteins of WT AlyGC and its mutants were overexpressed in *E. coli* BL21 (DE3) and cultured at 20 °C for 16 h in LB broth containing 100 μg/ml ampicillin under the induction of 0.3 mM isopropyl 1-thiogalactopyranoside (IPTG). Selenomethionine (SeMet)-labeled AlyGC was expressed by inhibiting endogenous methionine biosynthesis in *E. coli* BL21 (DE3) in defined media (40). Cells grown overnight in LB medium were harvested and inoculated in the M9 medium containing 100 mg/liter lysine, phenylalanine, and threonine, 50 mg/liter isoleucine, leucine, and valine, 5.2% (w/v) glucose, and 0.65% (w/v) yeast nitrogen base (YNB), which was then cultured at 37 °C. When the A₆₀₀ reached 0.6, the culture was cooled to 6 °C and then protein expression was induced by 0.3 mM IPTG for 8 h at 26 °C. Recombinant proteins were purified by using an Ni-NTA affinity column (Qiagen) and an Ion- Exchange chromatography column (HiPrep 16/60 Sephacryl S-300, GE Healthcare) (41). The recombinant protein was dialyzed against 20 mM Tris-Cl (pH 8.0) and concentrated to 10 mg/ml.

**Diffraction data and refinement statistics of WT-AlyGC, SeMet-AlyGC, and R241A-M4**

| Parameters                   | WT-AlyGC | SeMet-AlyGC | R241A-M4 |
|------------------------------|----------|-------------|----------|
| Data collection              | P 2 1 1  | P 2 1 1     | P 2 1 1  |
| Space group                  |          |             |          |
| Unit cell                    | a, b, c  | 116.6       | 116.3    |
| α, β, γ (°)                  |          | 90.0        | 90.0     |
| Wavelength (Å)               |          | 0.9785      | 0.9785   |
| Resolution (Å)               |          | 50.2–19.21  | 50.2–19.21|
| Redundancy                   |          | 4.6 (4.7)   | 6.9 (6.8)|
| Completeness (%)             |          | 98.6 (99.2) | 98.1 (98.4)|
| Rmerge (%)                   |          | 0.124 (0.450) | 0.141 (0.592) |

**Phasing**

| Figure of merit              | 0.348    |
| BAYES-CC                     | 39.88    |
| Sites                        | 45       |

**Refinement statistics**

| Resolution (Å)               | 37.03–2.19 (2.27–2.19) | 49.30–2.56 (2.66–2.56) |
| Rwork (%)                    | 16.6       | 19.1       |
| Rfree (%)                    | 19.7       | 22.6       |
| Figure of merit              | 38.66      | 79.85      |
| Sites                        | 45         | 45         |

* Numbers in parentheses refer to data in the highest resolution shell.
* Rmerge = Σhkl|I(hkl) − ⟨I(hkl)⟩|/ΣhklI(hkl),
* Probability-weighted average of the cosine of the phase error before and after density modification.
* The Ramachandran plot was calculated by PROCHECK program in CCP4i program package.
15 °C and 50 mg/liter l-SeMet was added. Fifteen min later, the culture was incubated at 15 °C for 14 h under the induction of 0.4 mM IPTG.

The recombinant proteins were first purified by nickel-nitritoltriacetic acid resin (Qiagen, Germany) and then fractionated by anion exchange on a Source 15Q column (GE Healthcare) and gel filtration on a Superdex G-200 column (GE Healthcare). Aldolase (158 kDa), conalbumin (75 kDa), and ovalbumin (43 kDa) from GE Healthcare were used as protein size standards.

**Biochemical Characterization of AlyGC**—Protein concentration was determined with bovine serum albumin as the standard by using a BCA protein assay kit (Thermo Fisher Scientific). The activities of WT AlyGC and its mutants toward alginate and PG were measured by the ultraviolet absorption spectrometry method (7, 41). Briefly, a 200-μl mixture containing 25 μg/ml enzyme and 2 mg/ml substrate in 50 mM Tris-HCl (pH 7.5) was incubated at 30 °C for 30 min. After that, the reaction mixture was boiled for 10 min to terminate the reaction. Then an increase in the absorbance at 235 nm (A235) caused by the production of unsaturated uronic in the mixture was monitored. One unit of enzyme activity was defined as the amount of enzyme needed to produce an A235 increase of 0.1 per min. The enzyme assays toward DS were performed with the method described by Michel et al. (26). The action mode of AlyGC was measured by TLC using PG as the substrate. Monoguluronic acid standard and the product were separated using a solvent system of 1-butanol/acetic acid/water (4:6:1, v/v) and visualized by heating TLC plates at 90 °C for 15 min after spraying with 10% (v/v) sulfuric acid in ethanol.

The ion in AlyGC molecule was investigated by using ICP-OES (42). To determine the effect of EDTA on the activity of AlyGC, EDTA at different concentrations was added to the reaction mixtures containing 3.5 μg/ml enzyme and 2 mg/ml PG in 50 mM Tris-HCl (pH 7.5) before the enzyme activity was determined. Apo-AlyGC was prepared by the addition of 2 mM EDTA and subsequent desalination. The recovered activity of apo-AlyGC was measured after Ca2+, Mg2+, or Mn2+ was added. Ca2+ was added in the final concentrations of 0.05, 0.1, 0.15, and 0.2 mM. Mg2+ or Mn2+ was added in the final concentration of 0.2 mM. Enzyme activity was determined by measuring the absorbance of the mixture at 548 nm with the thio-barbituric acid (TBA) method (43). One unit of enzyme activity was defined as the amount of enzyme required to liberate 1 nmol of β-formylpyruvic acid per min at 30 °C.

**Crystallization and Data Collection**—WT AlyGC (15 mg/ml) was crystallized at 20 °C by the sitting drop method in the buffer containing 25% (w/v) polyethylene glycol (PEG) 1500 and 100 mM SPG (succinic acid, sodium dihydrogen phosphate, and gly- cine) (pH 8.5). Crystals of SeMet-AlyGC (10 mg/ml) were grown in the buffer containing 200 mM malonate sodium (pH 7.0) and 18% (w/v) PEG 3350. The inactive mutant R241A (10 mg/ml) mixed with M4 at a molar ratio of 1:15 was crystallized at 20 °C by the hanging drop method in the buffer containing 100 mM HEPES-NaOH (pH 7.3), 8% ethylene glycol, and 11% PEG 8000. X-ray diffraction data were collected on BL17U1 beam line at the Shanghai Synchrotron Radiation Facility using detector ADSC Quantum 315r (44). The initial diffraction data sets were processed by HKL2000 (45). Data collection statistics are shown in Table 1.

**Structure Determination and Refinement**—Heavy atoms were searched by SHELXD (46). The phase problems were solved by single-wavelength anomalous diffraction (SAD) method using Phenix program Autosol (47). Initial model building was finished by Phenix program AutoBuild (47). Refinement of the AlyGC structure was done by Phenix program Refine (47) and Coot (48) alternately. The quality of the final model is summarized in Table 1. All the structure figures were processed using the program PyMOL.

**Dynamic Light Scattering and Circular Dichroism Spectra**—The DLS experiment was performed on Dynaprot Titan TC (Wyatt Technology) at 4 °C using 5.7 mg/ml AlyGC in a buffer containing 10 mM Tris-HCl (pH 8.0) and 100 mM NaCl, and analyses were done by Dynamics 7.1.0 software. CD spectra were collected from 250 to 200 nm at a scan speed of 200 nm/min with a path length of 0.1 cm on a J-810 spectropolarimeter (Jasco, Japan) at 25 °C. The final concentration of the proteins for CD spectra was 2.5 μM in 10 mM Tris-HCl (pH 7.5).

**Author Contributions**—F. X. and F. D. performed all experiments. X. C. directed the experiments. F. X. and X. C. wrote the manuscript. P. W. and H. C. solved the structures. C. L. and P. L. analyzed the data. Y. Z. and X. C. designed the research. X. P. edited the manuscript.

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