Catalytic Mechanism and Mode of Action of the Periplasmic Alginate Epimerase AlgG*

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Background: The alginate epimerase AlgG converts mannuronate to its C5 epimer guluronate at the polymer level.
Results: The structure of Pseudomonas syringae AlgG has been determined, and the protein has been functionally characterized.
Conclusion: His319 acts as the catalytic base, whereas Arg345 neutralizes the negative charge of the carboxylate group during catalysis.
Significance: This is the first structural characterization of a periplasmic alginate epimerase.

Pseudomonas aeruginosa is an opportunistic pathogen that forms chronic biofilm infections in the lungs of cystic fibrosis patients. A major component of the biofilm during these infections is the exopolysaccharide alginate, which is synthesized at the inner membrane as a homopolymer of 1→4-linked β-D-mannuronate. As the polymer passes through the periplasm, 22–44% of the mannuronate residues are converted to α-L-guluronate by the C5-epimerase AlgG to produce a polymer of alternating β-D-mannuronate and α-L-guluronate blocks and stretches of polymannurionate. To understand the molecular basis of alginate epimerization, the structure of Pseudomonas syringae AlgG has been determined at 2.1-Å resolution, and the protein has been functionally characterized. The structure reveals that AlgG is a long right-handed parallel β-helix with an elaborate lid structure. Functional analysis of AlgG mutants suggests that His319 acts as the catalytic base and that Arg345 neutralizes the acidic group during the epimerase reaction. Water is the likely catalytic acid. Electrostatic surface potential and residue conservation analyses in conjunction with activity and substrate docking studies suggest that a conserved electropositive groove facilitates polymannuronate binding and contains at least nine substrate binding subsites. These subsites likely align the polymer in the correct register for catalysis to occur. The presence of multiple subsites, the electropositive groove, and the non-random distribution of guluronate in the alginate polymer suggest that AlgG is a processive enzyme. Moreover, comparison of AlgG and the extracellular alginate epimerase AlgE4 of Azotobacter vinelandii provides a structural rationale for the differences in their Ca2+ dependence.

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The atomic coordinates and structure factors (codes 4NK6 and 4NK8) have been deposited in the Protein Data Bank (http://wwpdb.org/).
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Alginate is an unbranched anionic polysaccharide that is produced by brown algae (Phaeophyceae), Pseudomonas spp., and Azotobacter genera (1–3). Alginate is initially formed as a 1→4-linked poly-β-D-mannuronate polymer at the inner membrane and is subsequently selectively modified as it passes through the periplasm. These modifications alter the properties of the polymer and provide significant benefits to the organism. For example, in alginate-producing bacteria, mannuronate (M)2 residues can be selectively acetylated at the C2 and/or C3 positions (4), a modification that helps Pseudomonas aeruginosa evade host defense mechanisms (5). In both brown algae and alginate-producing bacteria, unacetylated mannuronate can be epimerized to its C5 epimer, α-L-guluronate (G) (6, 7). Brown algae and Azotobacter vinelandii express more than one epimerase and are capable of producing alginate rich in guluronate blocks, which in the presence of Ca2+ form gels that are important for structural integrity and cyst formation in brown algae and A. vinelandii, respectively (4, 6–10). In contrast, Pseudomonas spp. contain a single periplasmic epimerase. The alginate produced by these bacteria do not contain guluronate blocks but rather polymannuronic acid (poly(M)) blocks and blocks of alternating MG sequence (MG blocks) (4, 8, 11). The importance of epimerization in Pseudomonas spp. alginate is not clear, but guluronate incorporation, like acetylation, makes alginate more viscous, which could contribute to the ability of P. aeruginosa to evade host immune defenses (12).

Polymer level epimerization of sugar molecules is a rare modification that has only been found to date in three polysaccharides: alginate and the glycosaminoglycans heparin/heparan sulfate and dermatan sulfate (6). Heparin/heparan sulfate and dermatan sulfate are components of the extracellular matrix of animal tissue (13). Because of their negative charge, these polymers interact with a number of proteins to fulfill their roles in cell signaling, coagulation, and wound healing (14, 15).
Both glycosaminoglycans contain the uronic acid \(\alpha\)-iduronate, which is epimerized at its C5 position to \(\alpha\)-L-guluronate (6). Epimerization of heparan sulfate is essential for prenatal development as mice lacking the C5-epimerase die shortly after birth due to lung failure (16). Although alginate and heparin/heparan sulfate/dermatan sulfate are made by different organisms, they share some striking similarities. (i) All three polysaccharides are believed to be synthesized by a large multiprotein complex, the GAGosome in the case of mammalian systems. (ii) The differences in Ca\(^{2+}\) dependence observed between the two classes of enzymes.

**Experimental Procedures**

Cloning—The nucleotide sequence of \(algG\) from \(P. syringae\) pv. \(tomato\) strain DC3000 was obtained from the Pseudomonas Genome Database (32, 33) and used to design gene-specific primers. Residues 49–536 of \(algG\) were amplified from genomic DNA using the following forward and reverse primers: 5’-GCATCATATGTCAGGAGTTGCACCAAGG-3’ and 5’-GCAATACGTCTTCTCAGCTGGTCG-3’. (Italic font indicates that those nucleotides encode the protein.) The amplified PCR product was digested with NdeI and HindIII restriction endonucleases and cloned into a pET28a vector (Novagen) that had been linearized with the same enzymes. The resulting expression vector, pFW-AlgG49–536, encodes residues 49–536 of \(P. syringae\) \(algG\) fused to a cleavable N-terminal His\(_6\)-tag (His\(_6\)-AlgG49–536) for purification purposes.

The gene for \(Klebsiella\ pneumoniae\) \(\alpha\)-guluronate-specific lyase (\(alyA\)) was cloned in a similar fashion as described above using the following forward and reverse primers: 5’-GCATCATATGCTCAAGGAGTTGCACCAAGG-3’ and 5’-GCAATACGTCTTCTCAGCTGGTCG-3’. (Italic font indicates that those nucleotides encode the protein.) The amplified PCR product was digested with NdeI and HindIII restriction endonucleases and cloned into a pET28a vector (Novagen) that had been linearized with the same enzymes. The resulting expression vector, pFW-AlgG49–536, encodes residues 49–536 of \(P. syringae\) \(algG\) fused to a cleavable N-terminal His\(_6\)-tag (His\(_6\)-AlgG49–536) for purification purposes.

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In this study, we present the first crystal structure of a periplasmic alginate epimerase. Our 2.1-Å structure of \(Pseudomonas syringae\) \(AlgG\) reveals that the protein, as expected, adopts a parallel \(\beta\)-helix fold with 11 complete coils and one incomplete coil. The \(\beta\)-helix is capped N-terminally by an unusually elaborate lid structure comprising a central helix flanked by two antiparallel \(\beta\)-sheets. Site-directed mutagenesis and functional analysis suggest that the catalytic mechanism utilizes His\(_{319}\) as the catalytic base, Arg\(_{345}\) neutralizes the charge on the uronic acid, and water acts as the catalytic acid.
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pRC5 plasmid (34). The resulting expression vector, pFW-G-Lyase21–307, encodes residues 21–307 fused to a cleavable N-terminal His$_6$ tag (His$_6$-G-Lyase21–307) for purification purposes.

Mutants of AlgG were constructed using the QuickChange® Lightning site-directed mutagenesis kit (Stratagene) according to the manufacturer’s instructions. The fidelity of all constructs was verified by DNA sequencing (ACGT DNA Technologies Corp.).

**Protein Expression and Purification—Escherichia coli** BL21(DE) competent cells (Stratagene) transformed with pFW-AlgG49–536 or mutant AlgG were grown in 500 ml of terrific broth supplemented with 0.05 mg/ml kanamycin at 310 K. After the A$_{600}$ of the cell culture reached 0.6, protein expression was induced with isopropyl β-D-1-thiogalactopyranoside to a final concentration of 1.0 mM. The induced cells were incubated for 4 h at 310 K and harvested via centrifugation at 6,000 × g for 10 min at 277 K. The resulting cell pellet was stored at 193 K until required. His$_6$-G-Lyase21–307 was expressed as described above for His$_6$-AlgG49–536. For selenomethionine (SeMet) incorporation, the pFW-AlgG49–536 expression vector was transformed into *E. coli* B834 Met– competent cells (Novagen) and grown in 500 ml of selenomethionine-containing medium with 0.05 mg/ml kanamycin at 310 K using the protocol described by Lee et al. (35).

His$_6$-AlgG49–536 was purified from the cell pellet of two 500-ml bacterial cultures after resuspension of the cell pellets in 40 ml of buffer A (50 mM MES, pH 6.4, 500 mM NaCl, 5% (v/v) glycerol, 2 mM MgSO$_4$, 10 mM imidazole) containing one tablet of SIGMAFAST protease inhibitor EDTA-free mixture (Sigma). The resuspended cells were lysed by sonication (Misonix Sonicator 3000), and cell debris was removed by centrifugation at 38,000 × g for 30 min at 277 K. The supernatant containing His$_6$-AlgG49–536 was then loaded onto a 5-ml Ni$^{2+}$-NTA Superflow cartridge (Qiagen) pre-equilibrated with buffer A. After washing the Ni$^{2+}$-NTA Superflow cartridge (Qiagen) pre-equilibrated with buffer A to separate untagged from tagged AlgG. The flow-through and the wash fraction (20 ml of buffer A) were pooled and concentrated by centrifugation (1240 × g at 277K) using a concentrator with a 30-kDa-molecular mass cutoff (Millipore) and run on a HiLoad 16/60 Superdex 200 prep grade gel filtration column (GE Healthcare) equilibrated with buffer C. Fractions corresponding to AlgG49–536 were pooled, methylated as described in Truebestein et al. (36), concentrated, and rerun on a HiLoad 16/60 Superdex 200 prep grade gel filtration column (GE Healthcare) equilibrated with buffer C. Fractions corresponding to methylated AlgG49–536 were pooled and concentrated to 4.4 mg/ml, and the protein was stored in buffer C at 277 K until required. SeMet-incorporated AlgG and the AlgG D317A mutant were purified as described above for the wild-type protein.

**Crystallization, Data Collection, Structure Determination, and Analysis—Methylated AlgG49–536** concentrated to 4.4 mg/ml was subjected to crystallization trials using commercially available sparse matrix screens (Emerald BioSystems, Hampton Research, and Qiagen). Methylated AlgG49–536 was crystallized using the hanging drop vapor diffusion method at 293 K with 2 µl of protein and crystallization solution in a 1:1 ratio over a reservoir containing 0.2 ml of crystallization solution. Initial crystals appeared after 7 days in condition 78 of JCSG+ suite (Qiagen; 20% (w/v) PEG 3350, 0.24 mM sodium malonate, pH 7.0) and diffraction to 9 Å. These crystals were harvested and used to make a 20× seed stock as described by Douglas Instruments Ltd. and stored at 253 K. Methylated AlgG49–536 was rescreened using the microseeding technique (1 µl of protein solution, 1 µl of crystallization solution, and 0.5 µl of 1× seed stock over 0.2 ml of crystallization solution). Crystals appeared after 2–3 months in 1.26 M (NH$_4$)$_2$SO$_4$, 0.1 M HEPES, pH 7.5 (condition 15 of Wizard2, Emerald Biosystems) and diffracted to 2.1 Å. The crystals grew to maximum dimensions of 300 × 150 × 150 µm. Methylated selenomethylion-incorporated AlgG and mutant AlgG were crystallized in the same way. Showers of methylated selenomethylion-incorporated AlgG crystals appeared after 5 days and grew to maximum dimensions of 75 × 40 × 40 µm. Mutant D317A AlgG crystals appeared after 2–3 months and grew to maximum dimensions of 300 × 100 × 100 µm.

In preparation for data collection, the crystals were cryopro- tection by soaking them for 10–30 s in crystallization solution supplemented with 20% (v/v) glycerol prior to vitrification by flash freezing and subsequently stored in liquid nitrogen. All data were collected at beam line X29 at the National Synchrotron Light Source (Brookhaven National Laboratory). For the native wild-type protein, 360 images of 1° oscillation on an Area Detector Systems Corp. Q315 charge-coupled device detector with a 250-mm crystal-to-detector distance with an exposure time of 0.4 s per image were collected. The data were processed, integrated, and scaled using the HKL-2000 program suite (37) (Table 1). SeMet single wavelength anomalous dispersion data consisting of 720 images of 0.5° oscillation on an Area Detector Systems Corp. Q315 charge-coupled device detector with a 350-mm crystal-to-detector distance with an exposure time of 0.4 s per image were also collected and pro-
cessed, integrated, and scaled using the HKL-2000 program suite (37) (Table 1). The SeMet single wavelength anomalous dispersion data in conjunction with HKL2MAP (38) were used to locate 13 of 15 selenium sites. Density-modified phases were calculated using SOLVE/RESOLVE (39). The electron density map was interpretable, and the model (residues 69–492) was manually built using Coot (40), briefly refined using PHENIX.REFINE (41), and subsequently used as a search model using the PHENIX AutoMR wizard to determine the structure of native AlgG(49–536) by molecular replacement. The AlgG D317A(49–536) structure was determined using the PHENIX AutoMR wizard using the native structure as a search model and refined using PHENIX.REFINE.

Structural alignments were performed in PyMOL (Schrodinger, LLC) or using the Dali pairwise comparison server (42). Structure figures were generated using PyMOL (Schrodinger, LLC). Quantitative electrostatics were calculated using the PDB2PQR (43) and APBS (44) software. Surface residue conservation was calculated using the ConSurf program (45) using the T-Coffee (46) alignment of AlgG sequences from Pseudomonas spp. and A. vinelandii (accession numbers NP_252235 (P. aeruginosa PAO1), NP_791068 (P. syringae pv. tomato strain DC3000), YP_258150 (Pseudomonas protegens Pf-5), YP_273381 (P. syringae pv. phaseolicola 1448A), YP_610004 (Pseudomonas entomophila L48), YP_002870646 (Pseudomonas fluorescens SBW25), ZP_18874249 (Pseudomonas chlororaphis subsp. aureofaciens 30-84), ZP_16384839 (Pseudomonas avellanae BPIC 631), ZP_10143451 (Pseudomonas syringae algB), and CAA61231 (A. vinelandii)).

A β-d-mannurionate trisaccharide was modeled into the catalytic site of the wild-type AlgG structure using Coot (40). Subsequently, a dummy set of Fαβ of the model of AlgG with the β-d-mannurionate trisaccharide was created at a resolution of 3 Å using PHENIX.FMODEL (41). This model was then energy-minimized using PHENIX.REFINE (41). The β-d-mannurionate trisaccharide was taken from the structure of extracellular alginate epimerase complexed with a mannurionate trisaccharide (Protein Data Bank code 2PYH) (23).

Epimerase Activity Assay—Deacetylated poly(M) was prepared from P. aeruginosa FRD462 as described in Chitnis and Ohman (26), lyophilized, and dissolved in water at a concentration of 20 mg/ml. A fixed time coupled assay for AlgG, modified from Jerga et al. (31), was used to determine the activity of wild-type AlgG and its mutant variants. In the first step, 4 mg of poly(M) and 500 μg of AlgG were mixed in 300 μl of buffer C and incubated at 310 K for 40 h. AlgG and its mutant variants were subsequently inactivated by heating the mixture to 373 K for 10 min. The mixture was then centrifuged at 10,000 × g for 2 min to separate the alginate from the denatured epimerase. In the second step, the volume of the alginate solution was adjusted to 1 ml with 100 mM Tris-HCl, pH 7.5, and 2.5 μg of K. pneumoniae Aly 1-guluronate-specific lyase (His6-G-Lyase21–307) was added. The lyase reaction was terminated after 10 min at room temperature by heating the reaction mixture to 373 K for 10 min followed by centrifugation at 10,000 × g for 2 min. Unsaturated uronic acids, the product of the lyase reaction, were measured at 230 nm in a 1-cm-path length quartz cuvette in a Ultrospec 2100 UV/visible spectrometer (GE Healthcare) (47). The absorbance value of the K. pneumoniae Aly G-specific lyse on untreated poly(M) (control) was subtracted from the value of AlgG-treated poly(M). Relative activities for the AlgG mutants were calculated setting the value of the wild-type enzyme to 100%. All activity assays were performed in triplicate.

The folding of AlgG mutants that showed <5% of wild-type activity was assessed by circular dichroism spectroscopy. Circular dichroism spectra of purified AlgG (0.3 mg/ml) were recorded in 1 mM MES, pH 6.4, 10 mM NaCl at 293 K on a Jasco J-810 spectropolarimeter. The scans were done in triplicate.

Electrospray Ionization Mass Spectrometry (ESI-MS) Alginate Binding Assay—AlgG was dialyzed against aqueous 100 mM ammonium acetate, pH 7 using microconcentrators (Millipore Corp., Bedford, MA) with a molecular mass cutoff of 30 kDa and immediately used in the ESI-MS binding assay. A single chain Fv fragment (molecular mass, 26,539 Da) of the monoclonal antibody (mAb) Se155-4 was used as a reference protein to correct for nonspecific carbohydrate-protein binding during the ESI process.

Stock solutions of each of the individual alginate polymer ligands, tetramer through dodecamer (48), were prepared by dissolving the solid compounds in ultrafiltered water (Milli-Q, Millipore) at a concentration of 1 mM. Ligand solutions were stored at 253 K until needed.

The binding measurements were carried out on a Synapt G2 quadrupole-ion mobility separation-time-of-flight (Q-IMS-TOF) mass spectrometer (Waters, UK). A nanoflow ESI source was used. To perform nanoflow ESI, tips were produced from borosilicate tubes (1.0-mm outer diameter, 0.68-mm inner diameter) pulled to ~5-μm outer diameter at one end using a P-1000 micropipette puller (Sutter Instruments, Novato, CA).

A platinum wire was inserted into the nanoflow ESI tip, and a ~1.0-kV voltage was applied to carry out the ESI. Mass spectra were obtained in positive ion mode using cesium iodide (concentration, 30 ng/μl) for calibration. A cone voltage of 35 V was used, and the source block temperature was maintained at 343 K. I on transmission parameters (injection voltages) into the trap and transfer ion guides were maintained at 5 and 2 V, respectively. Argon was used in the trap and transfer ion guides at a pressure of 2.22 × 10−7 and 3.36 × 10−7 millibars, respectively. Data acquisition and processing were carried out using MassLynx (v4.1).

The determination of the association constant K_a, value for a given protein-ligand (P-L) interaction using the direct ESI-MS assay is based on the measured ratio (R) of the total abundance of the 1:1 P-L complex (Reaction 1), K_a is calculated using Equation 1.

\[ P + L = P-L \] (Reaction 1)

\[ K_a = \frac{R}{[L]_0 - \frac{R}{1 + [P]_0}} \] (Eq. 1)

where R is given by Equation 2.
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**RESULTS**

**AlgG Adopts a Right-handed Parallel β-Helix Fold**—To better understand how alginate is modified as it passes through the periplasm prior to export, we have undertaken structural studies of the C5-epimerase AlgG. As our initial attempts to crystallize *P. aeruginosa* AlgG were unsuccessful, we generated a number of constructs for various AlgG homologues. Ultimately, a construct lacking the predicted type I export signal yielded high resolution diffraction quality crystals after the protein had been methylated. The crystals took 2–3 months to grow. The structure of *P. syringae* AlgG was determined using selenomethionine incorporation and the single wavelength anomalous dispersion technique and was refined against a native data set to 2.1-Å resolution (Table 1). AlgG crystallized in space group *P*3*2*1 with unit cell dimensions of *a* = 126.6 Å, *c* = 97.7 Å, and *γ* = 120° with one molecule in the asymmetric unit. The final model was refined to an *R*~work~ of 18.8% and *R*~free~ of 23.6% (Table 1). Although the construct used produced protein that contains residues 49–536, the final refined model comprises residues 69–492 (Table 1). Analysis of the AlgG crystals by trypptic digest followed by LC/MS (data not shown) was able to detect a sequence starting at residue 128 and ending with residue 518, indicating that residues 493–518 were still part of the crystallized protein but could not be built due to the poor quality of the electron density. The available density suggests that this disordered C-terminal tail interacts

**TABLE 1**

Data collection and refinement statistics

|                | AlgG SeMet | AlgG wild type | AlgG D317A |
|----------------|------------|----------------|------------|
| Data collection|            |                |            |
| Wavelength (Å) | 0.97920    | 1.075          | 1.075      |
| Temperature (K) | 100        | 100            | 100        |
| Space group    | *P*3*2*1   | *P*3*2*1       | *P*3*2*1  |
| Unit cell parameters (Å) | *a* = 126.4, *b* = 97.3; | *a* = 126.6, *b* = 97.7; | *a* = 125.6, *b* = 98.5; |
| Resolution (Å) | 50.0-2.5   | 50.0-2.1       | 50.0-2.3   |
| Total no. of reflections | 699,311    | 1,242,960      | 958,215    |
| No. of unique reflections | 31,584     | 53,161         | 40,540     |
| Redundancy     | 22.1 (22.3)| 23.4 (21.5)    | 23.7 (19.0)|
| Completeness (%) | 100 (100)  | 100 (99.7)     | 100 (99.7)|
| Average I/σ(I) | 28.3 (5.3) | 43.3 (6.8)     | 31.9 (4.0)|
| *R*merge (%)   | 12.4 (64.0)| 8.8 (58.7)     | 10.9 (72.4)|

An underlying assumption is that P-L and P have similar ionization and detection efficiencies (*i.e.* similar ESI-MS response factors) such that the gas-phase abundance ratio is equal to the concentration ratio in solution. This assumption has been shown to be valid in cases where L is small compared with P such that P and P-L are similar in size and surface properties (49, 50). Nonspecific binding of free ligand to the free and bound protein can occur during the ESI process, leading to changes in their measured relative abundances. To account for nonspecific binding, a non-binding protein (single chain Fv fragment) was added to the ESI-MS solution containing P and L. A corrected *R* value was obtained by the reference protein method using the following expression (51).

\[
R = \frac{Ab(P-L)}{Ab(P)} = \frac{[P-L]}{[P]} = \left(\frac{Ab_{app}(P-L)}{Ab_{app}(P)}\right) \left(\frac{Ab(P_{ref-L})}{Ab(P_{ref})}\right)
\]

where *Abapp*(*P*) and *Abapp*(*P-L*) are apparent measured abundances of ions corresponding to P and P-L ions and *Ab(P_{ref})* and *Ab(P_{ref-L})* are the measured abundances of ions corresponding to free P_{ref} and P_{ref} bound nonspecifically to L.

Mass Spectrometry—Approximately 15 AlgG crystals were harvested, washed in well solution, dissolved in 50 μl of H2O, and analyzed by the in-solution trypptic digest LC/MS method (Advanced Protein Technology Centre, The Hospital for Sick Children).
the predictions of Douthit et al. (53); several internal stacks of aliphatic residues located on the PB1, PB2, and PB3 sheets; and an N-terminal tail lacking regular secondary structure that is packed against the PB2 face (Fig. 2B). The hydrophobic core of the protein is capped N-terminally by a short helix, called the cap helix, comprising residues 142–148. This short cap helix is flanked by two antiparallel β-sheets, each of which consists of two β-strands that together form an elaborate lid over the N-terminal central core region (Fig. 2A). At the C terminus, the hydrophobic core of AlgG is packed against a symmetry-related molecule in a tail-to-tail arrangement. AlgG contains an additional 44 residues at the C terminus that could not be modeled in the current structure. The missing C terminus is predicted to contain a short β-strand, presumably completing the 12th and final coil, followed by a helix, which could potentially cap the hydrophobic core at the C terminus (56). Hence, it is likely that this tail-to-tail protein interaction is an artifact of crystal packing.

In addition to the N-terminal cap helix, there are four other helices present in the structure that we have termed the H1–H4 helices (Fig. 2B). Helix H1 (residues 83–89) is found within the mostly random coil N-terminal tail and packs against the PB2 surface (coils 2 and 3 of the β-helix). Helix H2 (residues 107–113) interacts with residues 266–272, helix H4, a part of the T1 loop of coil 4. Helix H3a/b (residues 116–126) belongs to the same extended loop as helix H2 and consists of a 310 helix (residues 116–118; helix H3a) and a short two-turn α-helix (residues 119–126; helix H3b).

**His<sup>319</sup> Is the Catalytic Base**—The parallel β-helix fold found in AlgG is most commonly associated with polysaccharide lyases (PLs) belonging to families PL1, PL3, PL6, and PL9 (57, 58). Polysaccharide lyases and epimerases not only share the same fold but also have been suggested to have similar reaction mechanisms (Fig. 1) (20). As polysaccharide lyases have been extensively characterized, it is beneficial to use this knowledge to understand the catalytic mechanism of alginate C5-epimerases (57). Members of the PL1, PL3, PL6, and PL9 families utilize either Ca<sup>2+</sup> or arginine to neutralize the negative charge on the carboxylate group of the uronic acid (Fig. 1) (57, 59, 60). Polysaccharide lyases with a β-helix fold typically have either an arginine or a lysine as the catalytic base, whereas the catalytic acid has been identified to be either arginine or a water molecule (57). Polysaccharide lyases of other folds use tyrosine or histidine as the catalytic base and tyrosine as the catalytic acid (57).

**In vivo Studies of P. aeruginosa AlgG** Revealed that mutagenesis of each of the residues in the conserved DPHD motif (residues 317–320 in P. syringae) abolished epimerase activity (53). This motif is part of the carbohydrate-binding/sugar hydrolysis domain and is proposed to be part of the active site (22, 53, 55). In our AlgG structure, the DPHD motif is composed of Asp<sup>317</sup>, Pro<sup>318</sup>, and His<sup>319</sup>, which are located on the concave PB1 surface, and Asp<sup>320</sup>, which is part of the adjacent T1 loop of coil 6 (Fig. 3, A and B). To determine whether these residues are important for epimerase activity in P. syringae, alanine point mutants of residues Asp<sup>317</sup>, His<sup>319</sup>, and Asp<sup>320</sup> were made, and their ability to epimerize polymannuronate in vitro was tested. Pro<sup>318</sup> was not mutated as its side chain is not surface-accessible but part of the β-helix core. Using an end point assay, mutation of His<sup>319</sup> and Asp<sup>320</sup> was found to completely ablate epi-
structure reveals that the side chain of His$^{319}$ is in an unusual rotamer conformation ($\chi_1 = -118^\circ$). This high energy conformation and its presence in the DPHD motif suggests that His$^{319}$ plays a role in catalysis. His$^{319}$ is the only rotamer outlier present in the AlgG structure. This conformation appears to be due to residues Tyr$^{294}$, Asp$^{317}$, and Asp$^{320}$, which surround His$^{319}$ and prevent it from adopting a preferred rotamer conformation. Asp$^{320}$ O$^{\delta1}$ is in hydrogen bond distance to His$^{319}$ N$^{\delta1}$ (Fig. 3B), and mutating Asp$^{320}$ to alanine completely abolishes epimerase activity (Fig. 4). The His$^{319}$-Asp$^{320}$ interaction suggests that His$^{319}$ may be deprotonated over the pH range observed for optimal AlgG activity (pH 6–7.5) (31), a hypothesis supported by calculations using the heuristic pK\(_a\) prediction software PROPKA (61), which suggests that His$^{319}$ has a pK\(_a\) of 5.48. Because the formation of the His$^{319}$-Asp$^{320}$ interaction appears to be crucial for epimerase activity and a deprotonated His$^{319}$ would be capable of abstracting a proton from the C5 of mannanurionate, it would appear that His$^{319}$ likely acts as the catalytic base in the epimerase reaction. Asp$^{317}$ is also in close proximity to His$^{319}$ but too far for a hydrogen bond interaction as the distance between O$^{\delta1}$ of Asp$^{317}$ and N$^{\epsilon2}$ of His$^{319}$ is 3.7 Å. As we had found that the D317A AlgG mutant retains only ~5% wild-type epimerase activity, to further probe whether Asp$^{317}$ could play a role in the positioning of His$^{319}$, we determined the structure of the D317A mutant to 2.3-Å resolution (Table 1). The D317A mutant structure is very similar to the wild-type AlgG structure with a root mean square deviation of 0.2 Å over 424 Ca atoms (42). The only residue that changed in the active site region of the AlgG D317A mutant structure is His$^{319}$, which is no longer in a high energy conformation but adopts a more favorable side chain conformation, thus supporting our hypothesis that Asp$^{317}$ helps to position His$^{319}$ by forcing His$^{319}$ into a high energy conformation to avoid a steric clash between Asp$^{317}$ and His$^{319}$ (Fig. 3B). The His$^{319}$-Asp$^{320}$ interaction is not formed in the AlgG D317A structure as His$^{319}$ adopts the only possible low energy conformation that does not result in a steric clash. The loss of the His$^{319}$-Asp$^{320}$ interaction potentially explains why the D317A mutant only shows ~5% epimerase activity compared with wild-type levels. The side chain of the third residue likely to be important for positioning His$^{319}$ in wild-type AlgG, Tyr$^{294}$, sits above and approximately perpendicular to His$^{319}$ within van der Waals distance (shortest distance, ~3.7 Å). Mutating Tyr$^{294}$ to an alanine (Y294A) reduced AlgG epimerase activity to 24.5% of wild-type activity, whereas the Y294F mutant retained 66% activity (Fig. 4). These results suggest that Tyr$^{294}$ may also contribute to the proper orientation of His$^{319}$ and that the bulky side chain of Tyr$^{294}$ limits the potential side chain conformations that His$^{319}$ can adopt.

Further examination of the wild-type structure reveals that Tyr$^{294}$ is one of five conserved aromatic residues that lie directly above the active site (Fig. 3B). Mutation of the conserved aromatic residue Tyr$^{296}$ to alanine reduced the epimerase activity to 66% of wild-type levels (Fig. 4). Tyr$^{296}$ is within van der Waals distance of Asp$^{320}$ and could potentially help position Asp$^{320}$.

As it had previously been suggested for the calcium-dependant extracellular alginate epimerase AlgE4 from A. vinelandii (23) that Tyr$^{149}$ (Tyr$^{314}$ in AlgG P. syringae) was the catalytic base, we mutated Tyr$^{314}$ to phenylalanine and tested this AlgG mutant for epimerase function. This single point mutation reduced the epimerase activity to ~5% of wild-type levels (Fig. 4). CD spectroscopy confirmed that the Y314F mutant was folded (data not shown). The predicted pK\(_a\) of Tyr$^{314}$ is 13, suggesting that Tyr$^{314}$ would be protonated between pH 6 and 7.5 and therefore unlikely to function as the catalytic base in the AlgG epimerase reaction (61), making it more likely that Tyr$^{314}$ is involved in substrate recognition.

AlgG Uses Arg$^{245}$ to Neutralize the Negative Charge on the Uronic Acid during the Epimerase Reaction—Parallel β-helix polysaccharide lyases use either Ca$^{2+}$ or arginine to neutralize the negative charge of the uronic acid during catalysis (57). If
alginate epimerases use a similar reaction mechanism and because the AlgG epimerization reaction was found to be Ca^{2+}-independent, then it is likely that AlgG uses an arginine to neutralize the negative charge of the uronic acid (31). Examination of the AlgG structure suggests that the most likely candidate for this role is Arg^{345} (Fig. 3B). This arginine is in close proximity to the His^{319}.Asp^{320} diad and is held in position by three salt bridges. The N\textsubscript{1} of Arg^{345} interacts with the O\textsubscript{61} of Asp^{320}, and Ne and N\textsubscript{2} of Arg^{345} interact with O\textsubscript{61} and O\textsubscript{62} of Asp^{452}, respectively (Fig. 3B). To probe the role that Arg^{345} may play in catalysis, site-directed mutants that replaced the arginine with alanine, glutamine, glutamate, or lysine were generated. The R345K mutant reduced the epimerase activity to 36% of the wild-type level, whereas both R345A and R345Q mutants exhibited less than 10% of wild-type activity (Fig. 4). No epimerase activity was detected in the R345E mutant (Fig. 4). CD spectroscopy confirmed that the Arg^{345} mutants, which showed less than 10% epimerase activity, were folded (data not shown). The retention of 36% activity of the R345K mutant suggests that the positive charge of lysine may be able, in part, to perform the same role as Arg^{345}. In contrast, replacing Arg^{345} with the negatively charged glutamate completely abrogated epimerase activity, suggesting that the negative charge of glutamate might repel the negatively charged carboxylate group of mannuronate. The mutational results are consistent with the proposal that Arg^{345} is involved in neutralizing the negative charge of saccharide during catalysis.

His^{319}, Asp^{320}, and Arg^{345} are part of an extensive hydrogen bond and electrostatic interaction network of conserved residues that are arranged in a ringlike manner: His^{319}.Asp^{320}.Arg^{345}.Asp^{452}.Arg^{459}.Asp^{368}.Ser^{344} (Fig. 3B). The hydrogen bond-salt bridge ring is broken between His^{319} and Ser^{344} located just underneath His^{319}. Two of these residues, Asp^{368} and Ser^{344} in P. syringae, were found to be important for epimerase activity in vitro in P. fluorescens (mutants in P. fluorescens are D361N and S337F) (62). To test whether these residues were important for epimerase activity in vitro and to probe the importance of this network, we mutated Asp^{368} to asparagine and Ser^{344} to alanine. CD spectroscopy confirmed both mutants to be folded (data not shown). AlgG D368N showed 5% of wild-type activity, confirming the in vitro observations and suggesting that Asp^{368} is important for epimerase function (Fig. 4). The S344A mutant retained 54% of wild-type activity (Fig. 4). The reason for the disparity between in vitro and in vivo results is explained by that fact that phenylalanine replaced serine in vivo, probably causing AlgG to become unstable, whereas the smaller alanine does not perturb folding. The D452A mutant has a less severe impact on AlgG epimerase activity, retaining 63% of wild-type levels (Fig. 4). The AlgG R369A mutant shows 23% epimerase activity compared with wild-type activity, indicating that it is important for proper epimerase function and might be involved in substrate binding (Fig. 4).

Our analysis of the active site hydrogen bond network shows that several residues play a crucial role in epimerase activity.

A Conserved, Electropositive Groove on the Concave Face of AlgG Facilitates Alginate Binding—As the polymer is negatively charged, the alginate binding site is likely to have an overall positive electrostatic surface potential. Examination of the electrostatic surface properties of AlgG (Fig. 5A) indicates only one positively charged region on the enzyme located just below the putative catalytic site. This region is also highly conserved across all periplasmic AlgG alginate epimerases (Fig. 5B). We propose that this conserved region, which is in the extended substrate binding site. Given that the lengths of the mannuronate trisaccharides found in the alginate lyase A1-III (63) and the alginate epimerase AlgE4 (23) structures are 15.5 and 14.7Å, respectively, the substrate binding site of AlgG could accommodate at least nine monomers. In vitro characterization of AlgG has found that the enzyme requires a minimum of nine mannuronate residues for catalysis to occur, suggesting that the enzyme contains several substrate binding subsites (31). Ligand binding studies of wild-type AlgG with mannuronate oligomers ranging in length from four to 12 residues show that the ligand affinity does not significantly increase in oligomers longer than a nonamer (K\textsubscript{d} of 5.0 \times 10^{-4} M^{-1}) (Table 2). These data suggest that the AlgG substrate binding site contains at least nine subsites.

As our current attempts to co-crystallize AlgG with mannuronate oligomers have been unsuccessful, we modeled a mannuronate trisaccharide into subsites +1 to −2 of the putative binding site (Fig. 5, C and D). The orientation of mannuronate in subsite +1 (site of catalysis) was based on our proposed catalytic mechanism. Fixing the position of the mannuronate in subsite +1 limits the possible orientations of mannuronates for subsites −1 and −2, which were placed using Coot. The resulting model was subsequently energy-minimized.

According to our substrate-bound model, His^{339} could hydrogen bond to the carboxylate group of the mannuronate residue in subsite −1. The conserved Lys^{338} does not interact with the modeled trisaccharide but is in the vicinity of the mannuronate residue in subsite −1. To examine the influence of His^{339} and Lys^{338} on the epimerase activity of AlgG, we mutated Lys^{338} and His^{339} to alanine. The K338A mutant has an epimerase activity of 87% of wild-type levels, whereas the H339A mutant retains only 49% epimerase activity. This suggests that His^{339} plays a bigger role in substrate binding than Lys^{338} and further validates our substrate-bound model.

Our model suggests that Arg^{415} likely interacts with the carboxylate group of mannuronate in subsite −2 (Fig. 5D). Interestingly, the corresponding residue in P. fluorescens AlgG, Arg^{408}, is crucial for epimerase function in vitro as its mutation to cysteine results in a strain that only produces polymannuronate (62). To probe the role of Arg^{415}, we introduced the equivalent mutation in P. syringae AlgG (R415C), confirmed it to be folded (data not shown), and found that it completely abolishes epimerization activity in our in vitro assay (Fig. 4). We did not anticipate that the mutation of a single residue within the alginate binding site −13Å from the site of catalysis (His^{319}) would cause a complete loss of function in AlgG. The loss of epimerase activity in the R145C mutant supports the in vivo data.

Tyr^{392} is also located in close proximity to the mannuronate in subsite −2 although not within hydrogen or \pi bonding distance in our model. Because aromatic residues are frequently implicated in protein-carbohydrate interactions, we mutated Tyr^{392} to phenylalanine and alanine and tested each mutant for epimerase function. Our analysis shows that both the Y392F

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mutant and the Y392A mutant have ~65% of wild-type activity (Fig. 4). These results suggest that it is not the aromatic ring of the tyrosine residue but the hydroxyl group that may be important for AlgG epimerase function. The hydroxyl group of Tyr392 hydrogen bonds with N of Arg413. Hence, Tyr314 could be important for the correct positioning of Arg413.

Conservation of Active Site Residues of Periplasmic and Extracellular Alginate Epimerases Does Not Extend to the Polymer Binding Site—Although both periplasmic AlgG and the extracellular epimerase AlgE4 (Protein Data Bank code 2PYG) adopt the same parallel $\beta$-helix fold (23), these enzymes have

![Structure-Function Analysis of P. syringae AlgG](image)
distinctly different metal dependences. To gain insight into why AlgE4 requires Ca\(^{2+}\) for its enzymatic activity but AlgG does not, we have compared the two enzymes. Superimposition of the maximum number of coils of both structures resulted in their respective active sites being misaligned. This was probably due to the fact that their respective active sites are located on different coils. The active site of AlgG, Tyr\(^{314}\) and \(^{317}\)DPH, is located on coil 6, whereas the active site of AlgE4, Tyr\(^{149}\) and \(^{152}\)DPH, is located on coil 4. As initial inspection suggested that the active sites are very similar, a more meaningful comparison was obtained by aligning both structures using their active site residues Tyr\(^{314}\)-Tyr\(^{149}\), Asp\(^{317}\)-Asp\(^{152}\), Pro\(^{318}\)-Pro\(^{153}\), His\(^{319}\)-His\(^{154}\), and Asp\(^{320}\)-Glu\(^{155}\). When AlgG and AlgE4 are aligned in this manner, residues 294–482 of AlgG and 122–290 of AlgE4, the \(\beta\)-helix fold, superimpose well with a root mean square deviation of 0.753 Å over 543 backbone atoms (Fig. 6A). Both active sites have a similar organization (Fig. 6B). The Tyr\(^{294}\) of AlgG that we propose aids in orientation of AlgG His\(^{319}\) is equivalent to Phe\(^{122}\) in the AlgE4 structure (Fig. 6B). However, Asp\(^{320}\) in AlgG does not superimpose well with Glu\(^{155}\) of AlgE4. The Glu\(^{155}\) of AlgE4 was found to be critical for its epimerase activity (23). Interestingly, there is no AlgE4 residue equivalent to Arg\(^{345}\) of AlgG. In addition, examination of the active site region of AlgE4 shows a strong negatively charged surface potential as this active site lacks positively charged residues (Fig. 7, A and C). One of the negatively charged residues located in the active site region, Asp\(^{178}\), is crucial for the epimerase function of AlgE4 (23). This residue superimposes with Ser\(^{344}\) of AlgG (Fig. 6B). A closer look at the superimposition of the active sites of AlgG and AlgE4 reveals that Glu\(^{155}\) in AlgE4 partly overlaps with Arg\(^{345}\) in AlgG (Fig. 6B). Indeed, AlgE4 Glu\(^{155}\) and Asp\(^{178}\) surround the guanidinium group of Arg\(^{345}\) of AlgG in the superimposition. Because AlgE4 is Ca\(^{2+}\)-dependent and the coordination of Ca\(^{2+}\) is achieved by negatively charged amino acids, it is feasible that residues Glu\(^{155}\) and Asp\(^{178}\) of AlgE4 are part of a putative Ca\(^{2+}\) binding site.

The N-terminal cap regions of AlgG and AlgE4 do not superimpose as AlgG is capped by an elaborate lid, whereas AlgE4 is capped by a single N-terminal \(\alpha\)-helix (Fig. 6A). Furthermore, the N-terminal tail of AlgG does not have an equivalent in

**FIGURE 6. AlgG and AlgE4 superimposition.** A, schematic representation of AlgG (gray) and AlgE4 (purple). AlgG and AlgE4 were structurally aligned according to their respective active sites. B, a detailed view of the superimposed active site residues of AlgG (gray) and AlgE4 (purple) are shown in stick representation.

**FIGURE 7. AlgE4 surface analysis.** A, electrostatic potential surface representation of AlgE4. The electrostatic properties of AlgE4 were determined with the APBS server. Positive surface potential is shown in blue, and negative surface potential is shown in red and contoured from \(-5\) to \(+5\) \(kT/e\). B, surface representation of residue conservation of AlgE4 (dark magenta, high residue conservation) to dark cyan (variable residue conservation). Residue conservation was produced with the ConSurf server using a T-Coffee alignment comprising AlgE1–7 sequences from A. vinelandii. C, surface representation of AlgE4. Residues Asp\(^{152}\), His\(^{154}\), and Glu\(^{155}\) are highlighted in white.
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AlgE4 (Fig. 6A). Interestingly, the packing of the molecules in the two structures is comparable as the C terminus of AlgE4 is capped by the C terminus of the second molecule in the asymmetric unit in a tail-to-tail fashion. This arrangement again appears to be due to crystal packing as full-length AlgE4 contains an 18-kDa regulatory domain at the C terminus, and this is not present in the construct used to determine the structure. Although the backbone atoms of the extended substrate binding site of both epimerases superimpose well, only a few residues in this region are conserved between the two proteins. One exception is Arg415 of AlgG that superimposes with AlgE4 Lys255, a residue that is 100% conserved within *A. vinelandii* AlgE proteins. This residue is also critical for AlgE4 function as mutating Lys255 to an alanine reduced AlgE4 epimerase activity to less than 10% of the wild-type level (23). Interestingly, the sequence conservation of the substrate binding site across the extracellular AlgE proteins is low, and the conserved area is smaller than the conserved substrate binding site of the periplasmic epimerases (Fig. 7, B and C).

**DISCUSSION**

To better understand how alginate is post-translationally modified as it passages through the periplasm, we determined the structure and functionally characterized the C5-epimerase AlgG. The protein adopts a parallel β-helix fold establishing that the periplasmic epimerase shares the same fold as the extracellular epimerase AlgE4 (23). This is in contrast to the glycosaminoglycan polysaccharide epimerases, heparin/heparan sulfate and dermatan sulfate epimerases, the only other known polymer level carbohydrate epimerases, that are predicted to adopt a α/α toroid fold (64, 65). Like the β-helix fold, the α/α toroid fold is also common in polysaccharide lyases, further emphasizing the strong connections between polysaccharide epimerases and lyases (57).

Because of their lack of sequence similarity and different characteristics, extracellular and periplasmic alginate epimerases were thought to have different mechanisms (6, 21, 31). Our data reveal that the active site architecture of both types of enzymes is very similar (Fig. 6B). The main difference in the active sites is that AlgE4 does not contain a residue equivalent to Arg345 found in AlgG but instead contains several acidic residues that we propose form the AlgE4 Ca2+ binding site. As found in most β-helix fold polysaccharide lyases, this Ca2+ would be in a position to neutralize the carboxylate group of the uronic acid during the reaction, thereby performing a role equivalent to that of Arg345. The coordination of calcium also helps explain why the active site region of AlgE4, which has a negative surface potential, is able to catalyze the epimerase reaction of the negatively charged polyanunnuronate. One reason why no Ca2+ was modeled in the active site of the AlgE4 structure is that calcium coordination may require the substrate, which is not present at this subsite in the AlgE4 structure. The *A. vinelandii* extracellular alginate epimerases also appear to use the histidine of their 100% conserved DPHE motif as the catalytic base. Although His154 does not interact with Glu155 in the structure of the extracellular alginate epimerase AlgE4, in a manner comparable with His139 and Asp320 in AlgG, the Nδ1 of His154 hydrogen bonds with the Oδ1 of Asp178, His154 is also predicted to have a pK_a of 3.77 and therefore would be able to act as the catalytic base over the pH range of optimal AlgE4 activity (pH 6.5–7.0) (61, 66). The lack of sequence conservation in the substrate binding sites of the AlgG and AlgE4 enzymes and between periplasmic and extracellular alginate epimerases in general could reflect the fact that their substrates are slightly different and/or that the epimerases have adapted to their respective environments. Periplasmic AlgG catalyzes the epimerization of polymannuronate, whereas the extracellular epimerases only encounter the secreted alginate chain, which has already been modified by the periplasmic alginate epimerase and the acetylation machinery of *A. vinelandii*.

Our structure suggests that AlgG adheres to the first steps of the catalytic mechanism found in β-helix polysaccharide lyases and, although less commonly found in these polysaccharide lyases, appears to use arginine to neutralize the negative charge of the carboxylate group of the uronic acid (59, 60). β-Helix polysaccharide lyases, such as the extracellular pectate lyases, more frequently use Ca2+ to neutralize the negative charge (57). As calcium is used as a signaling molecule in bacteria, its concentration is tightly regulated in the bacterial cell (67). As reduced periplasmic Ca2+ concentration was found to promote biofilm formation in *Pseudomonas* (68), it is possible that periplasmic epimerases have adapted to this environment. Ca2+-dependent *A. vinelandii* AlgE alginate epimerases are secreted to the extracellular space where calcium is found in abundance (24, 25). To complete the epimerization reaction, a proton from a catalytic acid is required, and water is the most likely catalytic acid in the AlgG epimerase reaction. An *A. vinelandii* extracellular alginate epimerase has been found to incorporate tritium into guluronate residues during the epimerization reaction (69). Given the similarity of the active site architectures and proposed similarity of the reaction mechanisms, it is likely that in AlgG the proton, which is added to the opposite face of the sugar to complete the epimerization reaction, comes from the solvent. Water is also frequently used as the catalytic acid in the β-elimination reaction mechanism of β-helix polysaccharide lyases (57). Examination of the AlgG structure reveals that Tyr314 coordinates a water molecule, which is in the vicinity of the C5 position of the mannnuronate in the +1 subsite and could be a potential candidate for the catalytic acid. The readdition of the proton to the opposite face of the sugar also explains why the reaction is irreversible (21). The C5 proton of the newly created guluronate points away from the active site, making it impossible for AlgG to convert it back to mannnuronate. Flipping the alginate polymer so that the C5 proton of guluronate is pointing toward the catalytic base would also not allow the reverse reaction to take place as the epimerization reaction causes a substantial conformational change in the sugar with mannnuronate adopting a C1C4 configuration, whereas guluronate typically adopts a C1C4 conformation. Hence, it is very unlikely that the +1 subsite could accommodate guluronate for the reverse epimerase reaction to occur. The main difference between the alginate epimerases reaction mechanism, including that of AlgG, and the reaction mechanism of β-helix polysaccharide lyases is the use of histidine rather than arginine or lysine as the catalytic base. This differ-
ence is reflected in the different pH optima of the enzymes with alginate epimerases and β-helix polysaccharide lyases having neutral and basic pH optima, respectively (66, 70–75). Taken together, our data suggest that AlgG has a slight variation of the otherwise conserved catalytic mechanism found in β-helix polysaccharide lyases. It is interesting to note that the polysaccharide epimerases, heparin/heparan sulfate and dermatan sulfate epimerases, follow the same catalytic mechanism as AlgG but are able to perform the reverse reaction as they contain two catalytic bases (76, 77). The fact that polysaccharide epimerases adopt folds that are prevalent in polysaccharide lyases and their similar reaction mechanisms suggest that polysaccharide epimerases are derived from polysaccharide lyases.

Our model of the mannuronate trisaccharide bound to AlgG suggests that for the epimerase reaction to occur the reducing end of the sugar needs to point toward the active site of AlgG. The same orientation is seen in the AlgE4-mannuronate structure (23). Alginate, cellulose, and poly-β-N-acetylglucosamine are synthase-dependent extracellular polysaccharides (78). The recent cellulose synthase study (79) modeled the nascent polymer with the reducing end as the first portion of the polymer chain to enter the periplasm. This modeling of the nascent chain was based on the finding that cellulose is polymerized at its non-reducing end (80). If the same polymerization mechanism takes place in the alginate system, the substrate binding site of AlgG, located below the active site where the trisaccharide is modeled, coordinates the nascent and unepimerized polymannuronate chain before the epimerase reaction occurs. Although we only modeled a trisaccharide into the substrate binding site, AlgG has been shown to require longer substrates for catalysis to occur. Data from Jerga et al. (31) suggest that a minimum of nine mannuronate residues are required, which suggests that the substrate binding site of AlgG contains several subsites. The occurrence of subsites is a characteristic of processive enzymes, and many processive enzymes contain a groove to which the substrate binds (81). The catalytic and substrate binding sites of AlgG lie on a shallow concave surface of the parallel β-helix and form such a groove. AlgG has a low affinity for its substrate with a $K_a$ of $5.0 \times 10^3 M^{-1}$ for the nonamer (Table 2), additionally supporting the processivity hypothesis as tight binding would be detrimental to the sliding of the polymer. The sliding movement of the polymer could be a consequence of polymannuronate synthesis. The loss of epimerase activity in the AlgG R415C mutant suggests that the coordination of the carboxylate group of mannuronate in subsite −2 and hence the binding of the substrate in the correct register is crucial for the epimerase reaction to occur and that several subsites are indeed present on AlgG. The negatively charged carboxylate moieties of polymannuronate are offset by ~180° in successive monomers due to the nature of the β-1→4 linkage. Because the alginate binding site of AlgG is lined with positively charged residues (Fig. 5, A and D), each mannuronate could potentially bind to any subsite, allowing polymannuronate to glide through the binding site. But maximally only every other mannuronate would be in the correct orientation to be converted to guluronate. This processive mode of action model would explain why there are only alternating sequences of M and G but no guluronate doublets found in Pseudomonad alginites (4, 8, 11, 26). In vitro AlgG was found to produce G blocks probably because the polymer and enzyme are free to dissociate in the in vitro assay (31). Another explanation for the mode of action of AlgG in vivo could be that a polymer where a mannuronate has been converted to a guluronate is a better substrate for the epimerase (preferred attack mode of action). In this case, AlgG should show a higher affinity for substrates containing guluronate residues than for pure mannuronate substrates if this preferred attack mode of action is followed. However, AlgG shows no preference for MG over MM substrates (21), thereby excluding a preferred attack mode of action. Thus, only a processive mode of action can explain the non-random distribution of guluronate in alginate (4, 82). The extracellular alginate epimerases AlgE4 and AlgE6 have been found to be processive, adding support to the suggestion that AlgG follows this mechanism (66, 82–84).

In this study, we have determined the first structure of a periplasmic alginate epimerase and have shown that it adopts a β-helix fold with an elaborate lid structure. Subsequent bioinformatics analysis, site-directed mutagenesis, enzymatic analysis, and substrate modeling have allowed us to propose a catalytic mechanism and mode of action that is able to explain the composition of mannuronate and guluronate found in alginate from Pseudomonas spp. Furthermore, comparing our structure of AlgG with the structure of the extracellular epimerase AlgE4 has enabled us to identify a probable Ca$^{2+}$ binding site in AlgE4. Calcium is essential for catalysis in this enzyme and is believed to be involved in neutralizing the negative charge of mannuronate during the epimerase reaction of extracellular alginate epimerases.

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