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Pseudomonas aeruginosa AlgF is a protein-protein interaction mediator required for acetylation of the alginate exopolysaccharide

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Enzymatic modifications of bacterial exopolysaccharides enhance immune evasion and persistence during infection. In the Gram-negative opportunistic pathogen Pseudomonas aeruginosa, acetylation of alginate reduces opsonic killing by phagocytes and improves reactive oxygen species scavenging. Although it is well known that alginate acetylation in P. aeruginosa requires AlgI, AlgI, AlgF, and AlgX, how these proteins coordinate polymer modification at a molecular level remains unclear. Here, we describe the structural characterization of AlgF and its protein interaction network. We characterize direct interactions between AlgF and both AlgJ and AlgX in vitro and demonstrate an association between AlgF and AlgX, as well as AlgJ and AlgI, in P. aeruginosa. We determine that AlgF does not exhibit acetylesterase activity and is unable to bind to polymannuronate in vitro. Therefore, we propose that AlgF functions to mediate protein-protein interactions between alginate acetylation enzymes, forming the periplasmic AlgJFXK (AlgJ-AlgF-AlgX-AlgK) acetylation and export complex required for robust biofilm formation.

Biofilms are communities of bacterial cells surrounded and protected by a self-produced matrix containing lipids, exopolysaccharides, extracellular DNA, and proteins, and more complex structures such as membrane vesicles, bacteriophage, and amyloid fibers (1-3). The biofilm matrix promotes

adhesion and cohesion of bacterial cells and permits bacteria to adapt and thrive as a multicellular community despite environmental stresses (2, 4, 5). Bacterial biofilms can form in a variety of environments, including niches relevant to human health. They can grow on solid surfaces (e.g., medical devices), at an air-liquid interface (e.g., dental biofilms), or within semisolid media (e.g., sputum in the lungs of individuals with cystic fibrosis during chronic infection with Pseudomonas aeruginosa) (2).

Modification of exopolysaccharides within the biofilm confers protection to pathogenic bacteria during infection (6). Deacetylation of poly-N-acetyl-glucosamine is required for biofilm formation in Streptococcus epidermidis, Streptococcus aureus, Escherichia coli, and Yersinia pestis and provides resistance to neutrophil phagocytosis and enhances persistence in a mouse model of infection for S. epidermidis and S. aureus (6-10). Acetylation is another common modification. For example, acetylation of *Vibrio* polysaccharide is required for biofilm formation in Vibrio cholerae (11) and acetylation of cellulose is required for surface colonization in Pseudomonas fluorescens (12). Acetylation of alginate in P. aeruginosa is not only involved in forming the mature biofilm structure (13, 14) but also reduces opsonic killing by phagocytes, reduces susceptibility to enzymatic degradation, and aides in scavenging reactive oxygen species (6, 15–17).

Except for *algC*, the genes required for alginate biosynthesis in P. aeruginosa are clustered within a single operon (18). Alginate acetylation is known to require four proteins, AlgI, AlgJ, AlgF, and AlgX (19-23). After modification, alginate is exported from the cell by AlgX, AlgK, and AlgE (24). Belonging to the membrane-bound O-acetyltransferase family of proteins (19, 22), AlgI is hypothesized to receive an acetyl group from an unknown donor in the cytoplasm and transfer it to AlgJ in the periplasm (25). Although AlgJ exhibits acetylesterase activity in vitro, AlgJ is unable to bind or acetylate poly-mannuronate oligomers in vitro, suggesting that it is an intermediary in alginate acetylation (21). AlgX is a periplasmic acetyltransferase that can remove an acetyl group from an artificial donor and transfer it onto mannuronate polymer

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in vitro. In addition, chromosomal mutation of AlgX residues required for its acetyltransferase activity led to production of nonacetylated alginate (20, 21). Thus, it has been hypothesized that a relay takes place to transfer an acetyl group from the cytoplasm to the polysaccharide chain via AlgI, AlgI, and finally AlgX (19-21). AlgF is also localized to the periplasm and is required for alginate acetylation in vivo but little is known about its structure or function to date (19, 23).

Here, we describe the structure of AlgF determined by CS-Rosetta from chemical shifts data (26) and supported by interproton nuclear Overhauser effect (NOE) analysis. We demonstrate protein-protein interactions in vitro between AlgF and AlgJ, as well as AlgF and AlgX, by isothermal titration calorimetry (ITC) and an interaction between AlgF and AlgX identified by co-immunoprecipitation (co-IP) in P. aeruginosa. Based on these results, we propose that AlgF functions to mediate interactions between AlgJ and AlgX that are critical for the acetyl relay mechanism required for alginate acetylation, insights that allow us to propose the most detailed model yet of how alginate is modified prior to export.

Results

AlgF consists of two β -sandwich domains joined by a short linker region

AlgF is annotated as a periplasmic *O*-acetyltransferase in the Pseudomonas Genome Database for several species, including P. aeruginosa and Pseudomonas putida (27). This annotation is probably due to its demonstrated role in alginate acetylation

(19), but sequence alignments with the previously characterized O-acetyltransferases AlgJ (21) and AlgX (20, 21) reveal that AlgF lacks similarity to these and other known acetyltransferases. To gain insight into the role of AlgF in alginate acetylation, we first determined its structure. Despite exhaustive attempts, the protein proved to be recalcitrant to crystallization, and therefore CS-Rosetta models (28) were generated and validated using NMR spectroscopic techniques.

Uniformly 1H-,15N-, and 13C-labeled P. aeruginosa AlgF lacking its signal sequence, $AlgF_{Pa}^{30-216}$, was expressed and purified. Backbone, triple, and side-chain resonances were assigned by NMR (see Experimental procedures) and NOE distance restraints were collected for structural modeling (28–37). Alg F_{Pa} chemical shifts were assigned to 94.6% completeness. Analysis of the chemical shift data using the chemical shift index (CSI) as calculated by NMRView (28, 30) (Fig. S1) indicated that the N and C termini were not involved in any stable secondary structural elements as the CSI values were close to zero, as expected for regions with random coil characteristics. Sequence analysis revealed that the protein contains repeating homologous segments, suggesting the presence of a pair of tandem domains joined by a short interdomain region (Fig. 1A). The termini and the interdomain region (as well as many loop/turn regions) did not show any long-range NOE assignments (assignments more than four amino acid residues away in sequence), indicating that these regions are only in close contact with their sequential neighbors (Fig. S1). Analysis of the folded regions suggested that the structure was all β-strand as reflected by positive CSI scores

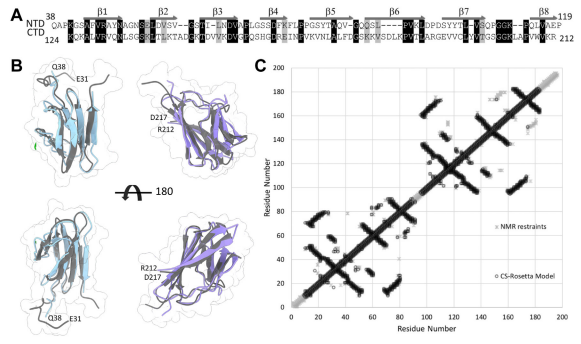


Figure 1. Comparison of CS-Rosetta and NOE-derived structures. A, sequence alignment of N-terminal (residues 38–119) and C-terminal (residues 124–212) domains of AlgF_{Pa} $^{30-216}$. Identical residues are shaded in *black*, similar residues are shaded in *gray*. Secondary structure elements are shown for each domain with secondary structure elements labeled across the top. B, superposition of the lowest energy CS-Rosetta (N-terminal, blue; C-terminal, purple) and NOE-based CYANA structures (lowest energy model in dark gray). The terminal residues are labeled. C, contact map analysis between the CS-Rosetta and NMR determined models. The inter-residue contacts observed in the NMR restraint-derived model and the CS-Rosetta-derived model are shown in gray asterisks and black circles, respectively. The inter-residue contacts are closely mirrored in structures determined by either method showing the similarity shared by the structures determined by independent methods. NOE, nuclear Overhauser effect.

and a lack of NOE patterns typically found in helical structures (i.e., backbone interproton NOEs seen between residues 3 or 4 amino acids apart, i-i+3 and i-i+4 in Fig. S1). There is good agreement with the secondary structure observed in the early atomic models calculated using NOE-derived distance restraints (Fig. S1, bottom row) and chemical shift data analysis. Despite the quality of the data there were an unacceptable number of clashes and NOE distance restraint violations in ensembles calculated using NOE-derived restraints in CYANA (29) (Table 1). This is probably the consequence of the 32% sequence identity between the N- and C-terminal domains (Fig. 1A), which results in a high degree of overlap in many chemical shifts resonances. Given the ~95% completeness of the chemical shift assignments, these data were therefore used in conjunction with the CS-Rosetta server to calculate structural models (28). As is typical for multidomain proteins, when the chemical shift data for the full-length protein was submitted to CS-Rosetta the algorithm failed to converge on a single structure. However, separating the shift data for the two domains provided well-converged structures that matched the NMR restraint-derived structures (Fig. 1, B and C) with backbone RMSD of 1.6 Å and 1.7 Å between the lowest energy CS-Rosetta and NMR-derived N-terminal and C-terminal domain models, respectively. All calculations using NOEderived distance restraints gave ensembles consisting of two tandem domains (Fig. 1). Four amino acids connect the two domains that we could not resolve structurally. Despite extensive searching, no unambiguous interdomain NOE

assignments could be found. These results are either because no interdomain NOEs exist or the structural and sequence similarity between the domains resulted in overlapping chemical shift assignments obscuring the interdomain NOE assignments.

Both the N- and C-terminal domains of $AlgF_{Pa}$ form an eight-stranded β-sandwich with backbone ensemble RMSDs for the CS-Rosetta determined models of 1.4 Å and 1.6 Å, respectively (Figs. 2A, S2 and Table 1). The AlgF_{Pa} β -sandwich has two distinct sides; one side is flatter with longer strands $(\beta 1, \beta 4, \beta 7, \text{ and } \beta 8)$, while the other side has a slight curve with shorter strands and longer loops (β2, β3, β5, and β6) (Fig. S2). This asymmetric β-sandwich with one flat face and one curved face was found in both the CS-Rosetta and NOE-derived models (Fig. 1). The N- and C-terminal domains are structurally similar to each other and superpose with an average ensemble backbone RMSD of 2.1 Å (Fig. 2A). Superimposition of the N- and C-terminal domains with the AlphaFold (38) model of AlgF_{Pa} which predicts the structure with high confidence (Fig. S3), reveals a backbone RMSD of 1.5 and 1.6 Å, respectively (Fig. 2B). Overall, comparison to the AlphaFold model suggests that the N- and C-terminal domains interact to form a compact structure.

To gain insight into the function of $AlgF_{Pa}$, the surface characteristics of each domain were further analyzed with respect to charge and sequence conservation. The theoretical pI values were calculated to be 4.60 and 9.61 for the N- and Cterminal domains, respectively, and these differences are

Table 1 Summary of model statistics for AlaF_{Pc}³⁰⁻²¹⁶

| | CS-Rosetta | | | |
|--------------------------------------|-------------------|-------------------|---|----------------------|
| | N-terminal domain | C-terminal domain | NOE derived | |
| Experimental restraints | | | | |
| CS-Rosetta input | | | CYANA input | |
| Residue range submitted | 30-119 | 120-222 | Assigned chemical shifts (%) | 2043 (94.6) |
| Folded Core ^a | 38-119 | 124-212 | Short range restraints ^b | 2267 |
| 13Ca shifts | 88 | 103 | Medium range restraints ^b | 438 |
| ¹³ C ^b shifts | 79 | 94 | Long range restraints ^b | 2145 |
| 13C' shifts | 84 | 100 | Angle restraints | 238 |
| ¹⁵ N shifts | 75 | 95 | H-Bond restraints | 130 |
| ¹ H ^N shifts | 75 | 95 | | |
| ¹ H ^a shifts | 98 | 111 | | |
| Assigned chemical shifts (%) | 914 (93) | 1148 (94) | | |
| Ensemble RMSD values | , , | , , | | |
| Residue range | 44-119 | 127-211 | $42-120^{\circ}$ | 125-211 ^c |
| Backbone atoms Å (# atoms) | 1.38 (304) | 1.59 (340) | 0.32 (316) | 0.24 (348) |
| Heavy atoms Å (# atoms) | 6.23 (1116) | 8.83 (1300) | 10.77 (1136) | 11.67 (1340) |
| Model Quality Measurements | , , | , | ` ' | ` ′ |
| Ramachandran statistics ^d | | | | |
| Most favoured regions (%) | 98 | 95 | | 77.6 |
| Additionally allowed regions (%) | 2 | 4 | | 21.9 |
| Generously allowed regions (%) | 0 | 0 | | 0.4 |
| Disallowed regions (%) | 0 | 1 | | 0.2 |
| Other Quality statistics | | | | |
| Bad contacts/1000 residues | 1 | 0 | | 33 |
| Violated restraints | N/A | N/A | $ m Distance > 0.5 ~ \AA \ Angle > 10^{ m o}$ | 91 0 |

a Resides that were determined by CS-Rosetta to be part of the folded core (regions with chemical shift values close to that of a random coil were truncated by the algorithm prior to fragment generation and model generation).

^b Definitions for restraint ranges: short range |i-j| < 1, medium range 1 < |i-j| < 5, long range |i-j| > 5.

The domains are aligned separately as the lack of inter domain contacts generates highly heterogeneous models with regard to relative domain positioning.

d Results for CS-Rosetta models from the PDB validation report and for the CYANA based models from the CYANA output. The statistics for the CYANA model are for the fulllength protein.



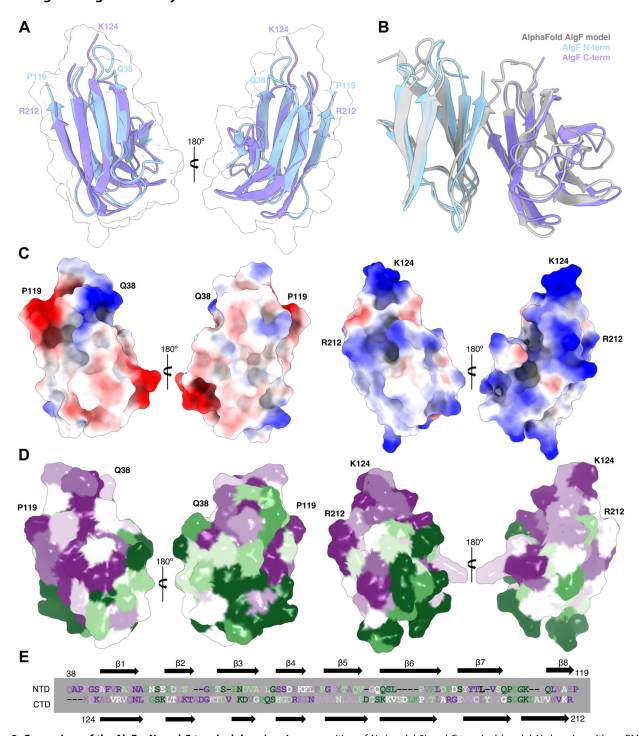


Figure 2. Comparison of the AlgF $_{Pa}$ N- and C-terminal domains. A, superposition of N- (model 9) and C-terminal (model 1) domains with an RMSD of 1.35 Å. The average backbone RMSD for the N- and C-terminal domain ensembles is 2.1 Å. B, structural superimposition of the N- and C-terminal domains of AlgF $_{Pa}$ with the AlphaFold model of AlgF $_{Pa}$. C, electrostatic potential surface representations of the N- and C-terminal domains (*left and right*, respectively). Shown in same orientation as *panel A*. The coulombic potential range from -10 (red) to 10 (blue) kcal/(mol·e). D, ConSurf analysis of conserved residues in the N- and C-terminal domains (*left and right*, respectively). Shown in same orientation as *panels A and C*. Surface representation colored by level of conservation ($green \ variable \ to \ purple$ highly conserved). E, sequence alignment (as in Fig. 1) colored by conservation level.

reflected in the calculated coulombic surface potential maps (Fig. 2C). The N-terminal domain of $AlgF_{Pa}$ has patches of negatively and positively charged residues, while the C-terminal domain is mostly positively charged. Using the ConSurf server (39), we identified highly conserved surface patches on both N- and C-terminal domains (Fig. 2, D and E). When

conservation was analyzed in the context of the compact AlphaFold2 AlgF $_{Pa}$ structure, highly conserved patches on both domains become buried, further suggesting that the two domains interact (Fig. S4). Specifically, the highly conserved residues Arg46, Ala50, Ser72, Ser73, and Val105 on the N-terminal domain and Asn132, Leu133, Val209, and Arg211 on

the C-terminal domain are buried in the AlphaFold AlgF_{Pa} model (Fig. S4A). Of these highly conserved residues, Arg46 on the N-terminal domain and Arg211 and Asn132 on the Cterminal domain are involved in electrostatic interactions (Fig. S4B). The most electropositive/electronegative regions of AlgF are not involved in mediating interactions between the two domains. The highly conserved residues Val105 on the Nterminal domain and Val209 and Leu133 on the C-terminal domain are involved in hydrophobic interactions (Fig. S4C). Less conserved residues, Tyr48 and Val115 on the N-terminal domain and Tyr196 and Val207 on the C-terminal domain, are also involved in hydrophobic interactions between the two domains (Fig. S4C). Analysis of the AlgF_{Pa} AlphaFold model by the Proteins, Interfaces, Surfaces, and Assemblies (PISA) server (40) also predicts that the N- and C-terminal domains interact, with an interaction interface buried surface of 752.4 Å². PISA indicates that the following residues become buried or are solvent inaccessible in the interaction interface: Arg46, Tyr48, Ala50, Ser72, Ser73, Val105, and Val115 on the N-terminal domain are buried, while Leu133, Tyr196, Val207, Val209, and Arg211 on the C-terminal domain are buried (Fig. S4D). These data support a compact AlgF structure where the N- and C-terminal domains interact.

Structurally similar proteins are involved in protein-protein and/or protein-ligand interactions

The AlgF $_{Pa}$ CS-Rosetta and AlphaFold2 models were submitted individually to the DALI server (41, 42) to identify structurally similar proteins. Based on the nature of ligands bound, we identified four different classes that represent most of the hits identified: 1, protein binding (e.g., PepT2 solute carrier family 15 extracellular domains, BcpA); 2, cholesterol binding (e.g., pneumolysin); 3, carbohydrate binding (e.g., rhamnogalacturonase); and 4, hormone/vitamin binding (e.g., transthyretin) (Fig. S5). Although all four classes contain structurally similar β -sandwich domains, each class is capable of binding chemically distinct ligands.

Class 1 proteins have the most structural homology to $AlgF_{Pa}$. Within class 1, solute carrier family 15 proteins (PepT1 and PepT2 (43)) are involved in oligopeptide transport across mammalian cell membranes and the extracellular domains of these proteins bind trypsin and recruit it to the site of dietary peptide uptake. PepT1 and PepT2 extracellular regions consist of two tandem β-sandwich domains joined by a short linker and most closely resemble the β-sandwich domain structure of $AlgF_{Pa}$ (Fig. S6A). However, only PepT2's domains appear to interact to form a compact particle, similar to $AlgF_{Pa}$ (Fig. S6A). Despite the similar structures, PepT1 and PepT2 share 19% and 14% amino acid sequence identity, respectively, to the AlgF N-terminal domain. Sequence comparisons to the $AlgF_{Pa}$ C-terminal domain are similar; 14% and 16% for PepT1 and PepT2, respectively. The bacterial pilin adhesion protein BcpA is critical for maintaining cell-cell contacts through intramolecular amide bonds, resulting in pili fiber formation on the cell surface (44). While BcpA has three tandem

β-sandwich domains which are structurally similar to the Nterminal and C-terminal domains of AlgF_{Pa}, the amino acid sequence identity between any two of the aligned domains is less than 6%. Class 2 consists of cholesterol-binding proteins including pneumolysin, a toxin found in pathogenic Streptococcus pneumoniae that forms a pore in eukaryotic membranes (45). Only the cholesterol-binding domain of the toxin forms a β-sandwich that is structurally similar to the N-terminal and C-terminal domains of $AlgF_{Pa}$. Domain II of rhamnogalacturonase in class 3 is suggested to be involved in oligosaccharide binding and resembles a single domain of AlgF (22, 23). The class 4 protein transthyretin is named for its role in transporting thyroxine and retinol in serum (46). While transthyretin consists of β-sandwich structures, the domains are oriented differently compared to $AlgF_{Pa}$ (Fig. S6B) (46). Most of the remaining unclassified hits (i.e., not belonging to classes 1-4) also have a role in binding small molecules and have little similarity to the structure of AlgF. In the case of multidomain proteins identified by the DALI search, the AlgFlike β-sandwich domain functions to bind small molecules, while the rest of the respective protein carries out the biological function. Since AlgF lacks any ancillary domains, we hypothesize that it most likely functions solely to bind either a small molecule or protein ligand.

In vitro binding analyses reveal direct interactions between AlgF, AlgJ, and AlgX

Since class 1 proteins with a β-sandwich structure are involved in binding proteins and AlgF has been proposed to interact with the acetylation machinery (47), we next investigated whether AlgF is involved in protein-protein interactions. Binary interactions between P. putida homolog constructs $\mathrm{Alg}\mathrm{F}_{Pp}^{30\text{-}215}$, $\mathrm{Alg}\mathrm{J}_{Pp}^{75\text{-}370}$, and $\mathrm{Alg}\mathrm{X}_{Pp}$ were probed in vitro using ITC. These proteins could be obtained in higher yields and were more stable than their P. aeruginosa counterparts thus enabling the ITC experiments (Fig. 3). We found that $AlgJ_{pp}$ and $AlgX_{pp}$ bind to AlgF_{Pp} with K_d values of 86 \pm 12 μ M and 178 \pm 5 μ M, respectively. Titration of $AlgX_{Pp}$ into $AlgJ_{Pp}$ demonstrated a decreasing heat of enthalpy as the titration proceeded, however, the data could not be reliably fit as the heats of emission were small and the titration did not reach saturation. The data suggest a weak interaction with a dissociation constant in the mM range.

AlgF is not an acetyltransferase and does not bind alginate

In $\Delta algF$ strains of *P. aeruginosa*, previous studies found that alginate exopolysaccharide is produced but not acetylated (19). The fold of AlgF did not suggest a particular enzymatic role and the 3D structure alone was not sufficient to provide insight into the function of AlgF. Unlike AlgJ and AlgX, AlgF shared no structural similarities with acetyltransferase enzymes. As AlgF is required for alginate acetylation, we investigated if this requirement was the result of a previously uncharacterized enzymatic activity.



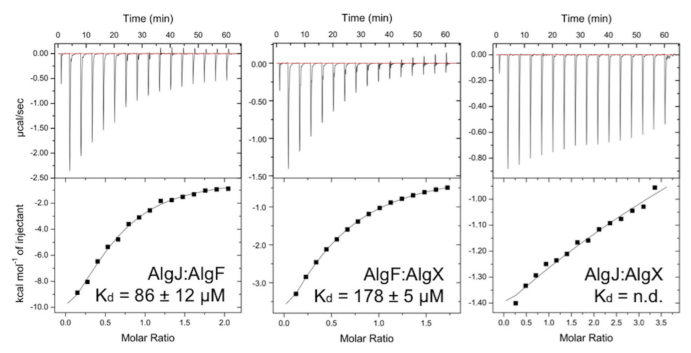


Figure 3. ITC analyses of $AlgF_{p_p}^{29-215}$ protein–protein interactions with $AlgJ_{p_p}^{75-370}$ and $AlgX_{p_p}$. Protein–protein interactions were investigated by binary titrations of AlgJ:AlgF (left), AlgF:AlgX (center), and AlgJ:AlgX (right) using ITC. Data were fit and analyzed with the Origin software package as shown, and K_d values were determined. The AlgJ:AlgX interaction (right) was too weak to be analyzed.isothermal titration calorimetry.

An assay was carried out to determine whether $AlgF_{pp}$ had acetylesterase activity, the first step in the acetyltransferase reaction. Removal of an acetate group from the pseudosubstrate 3-carboxyumbelliferyl acetate results in release of a fluorescent product, and reaction progress can be monitored through fluorescence spectroscopy. The acetylesterase activity of P. aeruginosa $AlgF_{Pp}$, $AlgX_{Pp}$, and $AlgJ_{Pp}$ was measured independently, in combination, and in the presence of an acetyl group acceptor (a nonacetylated mannuronic acid decasaccharide; ManA₁₀). No acetylesterase activity was observed for $AlgF_{pp}$ (Fig. 4). Addition of $AlgF_{pp}$ to either $AlgJ_{pp}$ or $AlgX_{Pp}$ did not increase observed acetylesterase activity, with or without presence of ManA₁₀ (Fig. 4). Similarly, addition of $AlgF_{Pp}$ to a combination of $AlgJ_{Pp}$ and $AlgX_{Pp}$ did not result in an increase acetyltransferase activity, with or without presence of ManA₁₀, suggesting that the presence of AlgF does not influence overall acetylesterase activity. Therefore, the proposed formation of an acetylation machinery complex, although required for alginate acetylation in vivo, does not affect the acetylesterase activity of AlgJ or AlgX in vitro.

Even though $AlgF_{Pp}$ demonstrated a lack of acetylesterase activity, some of the AlgF-related \mathcal{B} -sandwich proteins bind sugars/small molecules. Thus, the ability of $AlgF_{Pa}$ to bind carbohydrate polymer was investigated using an electrospray ionization mass spectrometry (ESI-MS) binding assay. Nine mannuronic acid oligomers ranging from 4 to 12 sugars in length (ManA₄ to ManA₁₂) were tested. $AlgF_{Pa}$ did not display any affinity for the oligomers tested. The approximate K_a values measured were less than 500 M^{-1} for all oligomers tested (Table S1). This is similar to what was observed previously for AlgJ (21). In contrast, AlgX binds to mannuronic acid oligomers in a length-dependent manner (21). As the data

suggest that AlgF does not bind alginate in isolation, it most likely functions as a protein–protein interaction mediator between AlgJ and AlgX.

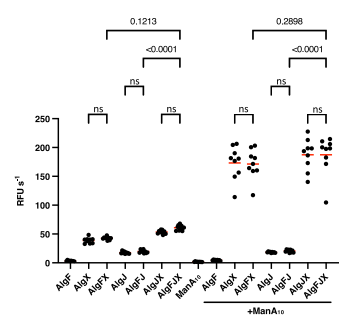
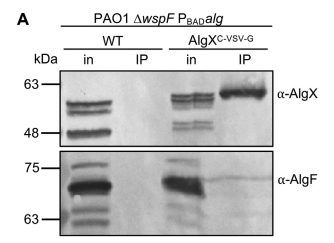


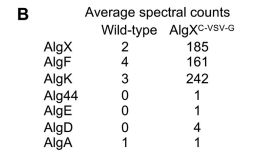
Figure 4. Acetylesterase activity of AlgF and the putative acetylation complex. Enzyme activity was measured by hydrolysis of acetate from the pseudosubstrate ACC. Release of the 7-hydroxycoumarin-3-carboxylic acid fluorescent product was measured ($\lambda_{\rm ex}=386$ nm and $\lambda_{\rm em}=446$ nm). Values represent three technical replicates for three biological replicates. Red lines represent the mean. The reactions were performed with the addition of 2 mM ACC to 5 μ M of each protein. Buffer contained 50 mM sodium Hepes pH 7.6 and 75 mM NaCl at 25 °C. ManA $_{10}$ denotes the addition of 1 mg/ml chemically synthesized polymannuronate decasaccharide. Statistical significance was determined using a one-way ANOVA test.

AlgIJFX form a complex in P. aeruginosa

The in vitro binding data obtained by ITC suggests that AlgF interacts with AlgX and AlgJ and that AlgF may mediate the formation of a complex when all three proteins are present. To examine whether these findings also hold true in P. aeruginosa, we performed co-IP experiments from lysed P. aeruginosa cells expressing vesicular stomatitis virus glycoprotein (VSV-G)-tagged AlgX, AlgJ, or AlgF. The VSV-G sequence was introduced at the C terminus of each gene directly on the chromosome of PAO1 ΔwspF P_{BAD}alg (48). In this strain, the native algD promoter has been replaced by araC-PBAD, allowing for high level inducible expression of the algD operon in the presence of arabinose. Clarified lysates were applied to agarose resin conjugated to anti-VSV-G mAbs, and the elution from the resin after washing was analyzed by Western blot using protein-specific polyclonal antibodies. The corresponding untagged protein was used as a negative binding control. When $AlgX_{Pa}^{C-VSV-G}$ was supplied as the bait, $AlgF_{Pa}$ was identified as an interaction partner (Figs. 5A and S10). Alg F_{Pa} was not identified in the elution from the negative control, indicating that the observed interaction is not due to nonspecific binding with the co-IP resin (Fig. 5A). To confirm this finding, co-IP eluates from six independent co-IP experiments using AlgX $_{Pa}^{C-VSV-G}$ as the bait were analyzed by ESI-MS. Alg F_{Pa} , as well as the previously identified interaction partner $AlgK_{Pa}$ (24, 49), were significantly enriched in the $AlgX_{Pa}^{C-VSV-G}$ eluate *versus* the untagged negative control (Fig. 5B), confirming the interaction between $AlgX_{Pa}$ and AlgF_{Pa}. When AlgF_{Pa} $^{\text{C-VSV-G}}$ was supplied as the bait, no coeluting proteins were identified, suggesting that addition of the VSV-G tag to AlgF_{Pa} may have disrupted the stability or function of $AlgF_{Pa}$.

Co-IP could not be performed successfully with $AlgI_{Pa}$ as the bait due to the instability and aggregation of $AlgI_{Pa}$ after solubilization from P. aeruginosa membranes. Attempts to optimize extraction using various detergents were unsuccessful, precluding the ability to generate an AlgI_{Pa}-specific polyclonal antibody. While we were able to generate a VSV-G-tagged construct of AlgI_{Pa} that could complement acetylation in an algI deletion mutant, AlgIPa also aggregates in Laemmli buffer under all conditions tested, preventing its detection by Western blot. To determine whether the destabilization of $AlgI_{Pa}$ during the detergent extraction step of co-IP experiments would have an effect on the stability of other acetylation proteins, a mutual stability analysis was performed using a $\triangle algI$ variant to mimic the loss of AlgI_{Pa} due to detergent extraction-mediated aggregation. When expression of AlgJ_{Pa} was analyzed by Western blot, a significant reduction in steady-state protein levels was observed in the $\triangle algI$ background versus WTWT (Figs. 5C and S11). Complementation of algI at the neutral attTn7 site on the P. aeruginosa chromosome restored $AlgJ_{Pa}$ to WT levels (Fig. 5C), suggesting that the reduction in whole-cell $Alg J_{Pa}$ levels was due specifically to the deletion of algI. Indeed, when co-IP experiments were performed with $AlgJ_{Pa}^{C-VSV-G}$ as the bait, no interaction partners were identified, likely due





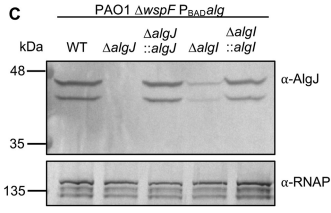


Figure 5. AlgX-AlgF and AlgI-AlgJ interact in *Pseudomonas aeruginosa. A*, co-IP from whole-cell lysates of *P. aeruginosa* expressing VSV-G-tagged AlgX as the bait. Proteins applied to the anti-VSV-G co-IP resin (input, in) and the elution from the resin after washing (immunoprecipitate) were analyzed by Western blot using AlgX- and AlgF-specific antibodies. A strain expressing untagged AlgX was used as a negative binding control. *B*, analysis of co-IP eluates from the experiment described in *panel A* by ESI-MS. Spectral counts were the average of six independent co-IP experiments using AlgX^{C-VSV-G} as the bait. Only alginateiosynthetic proteins identified with a minimum of one spectral count in this analysis are listed. *C*, Western blot analysis of whole-cell lysates of the indicated *P. aeruginosa* strains using AlgJ-specific antibodies. Antisera recognizing the β-subunit of RNA polymerase was used as a loading control. Co-IP, coimmunoprecipitation; ESI-MS, electrospray ionization mass spectrometry; VSV-G, vesicular stomatitis virus glycoprotein.

to $AlgI_{Pa}$ aggregation after solubilization of the membranes and the resultant effects on the stability of $AlgJ_{Pa}$. Overall, these data support the presence of a physiological interaction between $AlgX_{Pa}$ and $AlgF_{Pa}$ and imply also that $AlgJ_{Pa}$ and $AlgI_{Pa}$ interact based on the observed stability requirement of $AlgI_{Pa}$ for $AlgI_{Pa}$.



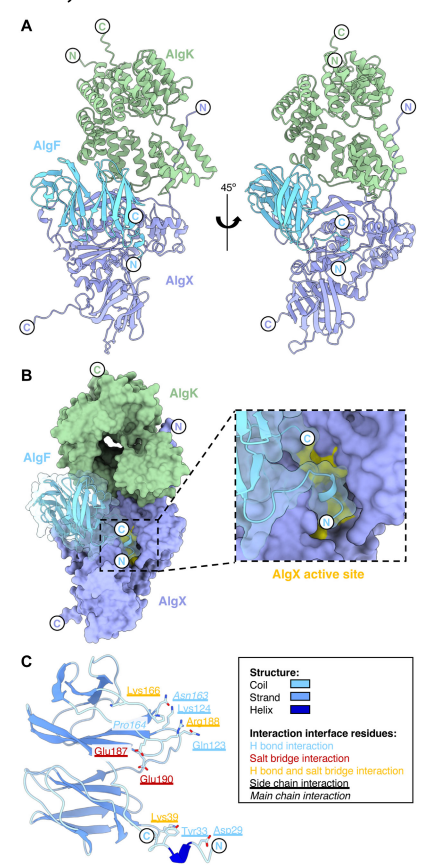


Figure 6. Model of the AlgKXF_{Pp} complex involved in alginate acetylation and export. A, AlphaFold2 model of the Pseudomonas putida AlgKXF complex (AlgK_{Pp}, green; AlgX_{Pp}, periwinkle; AlgF_{Pp}, light blue). B, surface representation of the AlgKXF_{Pp} complex with the AlgX_{Pp} active site highlighted in yellow. C, AlgF_{Pp} residues involved in the interaction interface with AlgX_{Pp}. The model of AlgF_{Pp} is colored by secondary structure (coil, light blue; strand; blue; helix, dark blue). Residues involved in hydrogen bonding interactions, salt bridge interactions, or hydrogen bonding and salt bridge interactions are represented by blue, red, and yellow text, respectively. Residues that are underlined represent side chain interactions and residues that are italicized represent main chain interactions. In both panels the N terminiand C terminiare represented by N and C, respectively.

Predictive modeling of the AlgKXF and AlgIJ complexes

The structure of the $AlgKX_{Pp}$ complex has been previously determined experimentally (PDB: 7ULA) and AlphaFold2 was shown to accurately predict the structure/interaction interface of the AlgKX_{Pp} complex across different *Pseudomonas* species (24). Building on the success of the AlphaFold2 model of $AlgKX_{Pp}$, we next sought to determine whether the program could predict how $AlgF_{pp}$ may interact with $AlgX_{pp}$ and $AlgK_{Pp}$. Using this approach, we were able to generate a highconfidence model of the $AlgKXF_{Pp}$ complex involved in alginate modification and export (Figs. 6, A and B and S7). The predicted AlgKXF_{Pp} complex shows that AlgK_{Pp} and AlgX_{Pp} maintain the same interaction interface as previously reported in the experimentally determined structure (24). Analysis of the AlgKXF_{Pp} model by the PISA server (40) suggests that $AlgF_{Pp}$ only interacts with $AlgX_{Pp}$. The modeled $AlgXF_{Pp}$ interaction interface was calculated to have a buried surface area of 1545 Å² mediated by 12 hydrogen bonds and six salt bridges (Fig. 6C). Most notably, ten out of eleven predicted interacting residues on $AlgF_{Pp}$ interact with $AlgX_{Pp}$ using their side chain atoms and all eleven residues are present on coil regions of $AlgF_{Pp}$. Three $AlgF_{Pp}$ interaction interface residues are present on the N-terminal domain (Asp29 and Tyr33 involved in hydrogen bonding and Lys39 involved in both hydrogen bonding and salt bridge interactions), while the remaining nine are present on the C-terminal domain (Gln123, Lys124, Asn163, Pro164, Lys166, Glu187, Arg187, and Glu190) (Fig. 6C). Thus, both the N- and C-terminal domains are predicted to be required for the interaction with $AlgX_{Pp}$. In $AlgF_{Pa}$, these interaction interface residues correspond to Asp29, Tyr34, Lys40, Gln124, Lys125, Asn164, Pro165, Lys167, Ala188, Arg189, and Glu191. Of the nine residues that are present in the $AlgF_{Pa}$ experimentally determined structure, four of these residues are highly conserved, four are conserved, and one is less conserved (Fig. 2E). Building on this model, we next attempted to model the $AlgKXFJ_{pp}$ complex using AlphaFold2. To expand this prediction to include $AlgJ_{Pp}$, due to the 1400 residues limitation in AlphaFold2, we only included the C-terminal region of Alg K_{Pp} that binds to Alg X_{Pp} when predicting the AlgKXFJ_{Pp} complex. Although the predicted models revealed a consistent $AlgKXF_{Pp}$ interaction interface (as shown in Fig. 6), the predicted alignment error plot revealed the interaction of $AlgJ_{pp}$ with $AlgKFX_{pp}$ could not be accurately predicted as the predicted alignment error values for AlgJ_{Pp} were estimated to be >25 Å (Fig. S8). Thus, we were unable to generate a high-confidence model to illustrate how $AlgF_{Pp}$ and $AlgJ_{Pp}$ may interact.

Given that our mutual stability data from P. aeruginosa suggests an interaction between $AlgI_{Pa}$ and $AlgJ_{Pa}$, we also used AlphaFold to model the AlgIJ $_{Pp}$ complex. A highconfidence model of the $AlgIJ_{pp}$ complex was generated for this part of the predicted acetylation complex (Fig. S9). This complex suggests that it is predominantly the transmembrane domain of AlgJ that interacts with AlgI. Analysis by PISA reveals that the interaction interface area between AlgI and AlgJ is 2122 Å², mediated by ten hydrogen bonds and three salt

bridges. The N-terminal helix of AlgJ is inserted into the inner membrane and is predicted to pack between the first and last helices of AlgI (Fig. S9).

Discussion

In this study, we present the structure of AlgF and, using in vitro studies coupled with analyses from P. aeruginosa, establish that the proteins involved in alginate acetylation (AlgI, J, F, and X) interact to form an acetylation complex that is linked to the outer membrane export machinery via an interaction between AlgX and AlgK. We found that AlgF consists of two β-sandwich domains joined by a linker, and our functional characterizations suggest that AlgF is unlikely to function as an alginate acetyltransferase as it lacks acetylesterase activity and is unable to bind alginate in vitro. Considering that most of its structural neighbors are involved in protein-ligand interactions, we propose that AlgF functions as a protein-protein interaction mediator within the alginate biosynthetic system to coordinate an AlgJFX periplasmic acetylation complex.

A search for AlgF structural homologs using the DALI server revealed proteins involved in ligand binding. The tandem two-domain architecture of AlgF resembles the binding module of the eukaryotic peptide transporter PepT2, which mediates an interaction between trypsin and the membrane transporter domain (43). This functionality could be mirrored with AlgF localizing and/or strengthening interactions between the acetyltransferase proteins and the rest of the alginate biosynthetic complex. As evidence of this role for AlgF, we have been able to establish that AlgF binds to both AlgX and AlgJ in vitro. We originally hypothesized that one domain of AlgF binds AlgJ at the inner membrane while the other binds to AlgX, however, structural prediction of the AlgKXF_{Pa} model suggests that both domains are required for its interaction with AlgX. Thus, further binding studies in conjugation with targeted mutagenesis of AlgF are required to ascertain which regions of AlgF are involved in interacting with either AlgX or AlgJ.

Previous studies have demonstrated that alginate is produced but not acetylated when either AlgJ or AlgX enzymatic activity is compromised (20, 21). Taking into consideration all the data presented on AlgJFX thus far, this suggests that transfer of an acetyl group from AlgJ to AlgX occurs and is required for polymer acetylation. As no direct interaction between AlgJ and AlgX was observed in this study, we hypothesize that AlgF is necessary to bring AlgJ and AlgX together in close enough proximity for acetyl relay and that lack of AlgF would decouple the acetyltransferase process. AlgI has also been found to be critical for acetylation of the nascent polymer (19, 22, 25). Our mutual stability studies demonstrate that AlgI may be part of the acetylation complex. This hypothesis is further strengthened by our AlphaFold2 model that suggests that AlgI is linked to the acetylation machinery primarily through the transmembrane domain of AlgJ (47). While we have been unable to model the entire AlgIJFX complex, we propose that AlgIJFX serve as an acetyl relay to transfer an



acetate from AlgI, across the inner membrane first to AlgI, then AlgX and finally to the polymer (Fig. 7). Comparably, acetylation of cellulose in *P. fluorescens* requires the proteins WssF/H/I/G which are homologous to the alginate proteins, AlgX/I/J/F (12). The AlgF-like protein, WssG, in acetylated cellulose biosynthesis is poorly characterized and its role remains unknown (12). Future experiments along the same lines as those presented here could help establish whether WssG is also a protein-protein interaction mediator and critical for the formation of a WssH/I/G/F complex.

In addition to the AlgI/J/F/X interactions outlined here, previous studies have demonstrated that AlgX and AlgK form a robust complex that couples alginate acetylation and export (24). Furthermore, given that AlgE localization is dependent on the presence of AlgK (50), formation of an AlgEKX outer membrane secretion complex has also been proposed (24). Within the inner membrane, it has been suggested that Alg8 and Alg44 form a synthase copolymerase complex (51). Alg44's periplasmic domain has been proposed to interact with AlgX and AlgK (52). These previously established interaction networks highlight the mechanisms required for alginate synthesis. Our ability to observe the well-established AlgX-AlgK interaction using co-IPs reinforces the results obtained here that link AlgX to AlgF and the rest of the acetylation machinery. The interactions between the periplasmic acetylation proteins have been found to be relatively weak by ITC with µM affinities. Thus, it may be possible that this weak binding is an artifact of *in vitro* protein studies of binary rather than native multiprotein interactions. The acetylation proteins may interact with a higher affinity once the rest of the biosynthetic complex and alginate substrate is present, allowing for a more stable complex in vivo. This hypothesis is not without precedent in large protein complexes (53–55). Specifically, the VirB type IV secretion system complex functions as a minimal set of VirB7 to VirB10 proteins, while the addition of VirB1 to VirB4 increases the activity of the transport complex dramatically, as well as increases the abundance of macrocomplex protein interactions (54). Weak µM interactions between proteins may also indicate the presence of a transient overall biosynthetic complex (55), where a number of subcomplexes may be present in the alginate biosynthesis pathway with variable intrasubcomplex and intersubcomplex affinities. Neither high-affinity interactions nor stable complex formation may be needed (or desirable) in order to fine-tune alginate modification, as the degree of acetylation is seen to vary depending on bacterial strain and species or on growth conditions and may even vary over the course of biofilm development, maintenance, and dispersal (56-59). If this is the case, then structural determination of subcomplexes with the alginate biosynthetic system may be more reasonable and realistic compared to determination of the complex in its entirety.

The structural determination of AlgF, structural homology searches, and characterization of protein function have

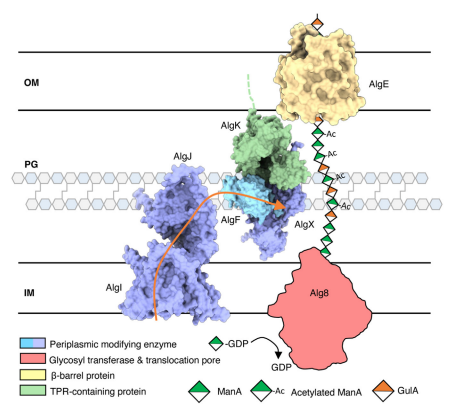


Figure 7. Model of alginate O-acetylation machinery. Algl receives an acetyl group from an as-yet unknown donor in the cytoplasm and the acetate group is passed through the inner membrane and transferred to AlqJ. AlqF mediates interactions in the periplasm allowing for the acetate to be passed from AlaJ to AlaX, where AlaX O-acetylates the newly polymerized mannuronic acid. OM, PG, IM denote the outer membrane, peptidoglycan, and inner membrane, respectively. Orange arrow indicates how the acetyl group is transferred between Algl, AlgJ, and AlgX.

enabled us to identify that AlgF mediates protein-protein interactions in the alginate acetylation machinery. The formation of an AlgIJFX acetylation complex supports the previously proposed relay mechanism for alginate modification, where AlgF mediates interactions between AlgJ and AlgX for acetylation of the polymer (19-21). For the first time, we are able to build upon established protein networks described above and show links from the inner membrane complex AlgI/J to AlgX/ K at the outer membrane. These complexes are linked via AlgF and join the modification subcomplex in the inner membrane/ periplasm with the outer membrane export subcomplex. These data are suggestive of a large protein interaction network and support the hypothesis of a trans-envelope macrocomplex, encompassing all of the exopolysaccharide biosynthetic proteins (19, 60). Furthermore, recent studies on the homologous cellulose biosynthetic machinery have shown the presence of an inner membrane subcomplex formed by cytoplasmic, transmembrane, and periplasmic proteins (61-64). Future work with alginate proteins may uncover similar interactions with cytoplasmic components including sugar precursor enzymes, thus expanding our knowledge of how alginate polymerization and precursor biosynthesis may be coupled. Understanding the biosynthesis of exopolysaccharides, including the alginate system, will require further studies of the protein interaction networks not only at a protein-protein level but also at a cellular level. The results presented here provide the first insight into a membrane-spanning polysaccharide secretion complex with significant etiological consequences for patient outcomes in cystic fibrosis.

Experimental procedures

Bacterial strains, plasmids, and growth conditions

A detailed list of the bacterial strains and plasmids used in this study can be found in Table S2. All P. aeruginosa mutant and complemented strains were derived from PAO1 (65) and were constructed using allelic exchange and miniTn7 mutagenesis (66, 67), as described below. Unless otherwise stated, lysogeny broth (LB) was used for growth of all strains. LB contained, per liter of ultrapure water, 10 g tryptone, 5 g NaCl, and 10 g yeast extract. Vogel-Bonner minimal medium (VBMM) was prepared as a 10 × concentrate containing, per liter of ultrapure water, 2 g MgSO4•7H2O, 20 g citric acid, 100 g K2HPO4, and 35 g NH4HPO4 and was diluted to $1 \times$ as needed. To prepare solid media, 1.5% (w/v) agar were added to LB or VBMM. Where appropriate, antibiotics were added to growth media. For E. coli, 10 µg/ml gentamicin, 100 µg/ml carbenicillin, or 50 µg/ml kanamycin was used. For P. aeruginosa, 30 or 60 μg/ml gentamicin was used depending on the application, as described below.

P. aeruginosa and P. putida gene expression in E. coli

The nucleotide sequences of AlgF from *P. aeruginosa* PAO1 and *P. putida* KT2440 and of AlgX from *P. putida* KT2440 were obtained from the *Pseudomonas* Genome Database (27). AlgF $_{Pa}^{30-216}$ and AlgF $_{Pp}^{29-215}$ were PCR amplified from genomic DNA. Primer sequences are as indicated in Table S3.

Primers account for entire full-length proteins without the respective N-terminal signal sequence as predicted by SignalP (68) and introduce NdeI and XhoI restriction sites. The codonoptimized gene for expression in E. coli of full-length $AlgX_{Pp}$ included flanking NdeI and XhoI restriction sites at the 5' and 3' ends, respectively, was synthesized by BioBasic. Alg ${\sf F}_{\it Pa}^{-30\text{-}216}$ and codon-optimized $AlgX_{Pp}$ were incorporated into the pET24b vector for C-terminal His6-tag protein expression. $AlgF_{Pp}^{29-215}$ was incorporated into the pET28a vector with a 3' stop codon for N-terminal His6-tag protein expression. For each protein construct, E. coli BL21 CodonPlus (λDE3) cells (Stratagene) were transformed with expression vector and grown in LB broth containing 50 μg/ml kanamycin at 37 °C. Once the A_{600} of the culture reached 0.6, protein expression was induced by the addition of IPTG to a final concentration of 1 mM. The cell culture was incubated at 18 °C for 16 h prior to being harvested by centrifugation at 6700g for 25 min at 4 $^{\circ}$ C. Cell pellets were stored until needed at –20 $^{\circ}$ C. AlgJ $_{Pa}^{79-379}$ and AlgJ_{Pp} $^{75-370}$ were expressed as described previously (21).

Purification of His6-tagged protein

The cell pellet from 1 L of bacterial culture was thawed and resuspended in 50 ml of lysis buffer (50 mM Tris pH 8.0, 500 mM NaCl, 0.5 M EDTA, 1 mM DTT, 1 mM PMSF, 2% (v/ v) glycerol, 1 mg/ml lysozyme) with one Roche Complete protease-inhibitor cocktail (EDTA-free) tablet. The resuspended pellet was incubated at 4 °C for 30 min. Cells were homogenized at 15,000 psi using an Emulsiflex C3 (Avestin Inc.) for three passes or until fully lysed. Cell lysate was centrifuged at 20,100g for 20 min at 4 °C to remove cellular debris. The resultant lysate supernatant was loaded onto Ni-NTA agarose resin and washed with 30 column volumes of 50 mM Tris pH 8, 500 mM NaCl, 2% (v/v) glycerol, and 30 mM imidazole. Protein was eluted using similar buffer with 300 mM imidazole and concentrated by centrifugation with 4 kDa (AlgF) or 30 kDa (AlgJ and AlgX) cut-off Vivaspin Turbo centrifugal concentrators (Sartorius). His₆-tagged protein was further purified using a HiLoad 16/60 Superdex 200 prep-grade size-exclusion column (GE Healthcare) in 50 mM Tris pH 7.5, 500 mM NaCl, and 2% (v/v) glycerol. Protein purification was monitored throughout by SDS-PAGE. Alg J_{Pp} and AlgX $_{Pp}$ proteins were purified as described previously (21, 69).

NMR structure determination

NMR studies on 1 H, 15 N, and 13 C AlgF $_{Pa}^{30-216}$ were carried out at a protein concentration of 1 mM in 50 mM phosphate buffer pH 6.8, 10 mM DTT, 2% (v/v) glycerol, 10% D $_{2}$ O. To produce uniformly labeled protein, cells were grown in minimal media supplemented with 1 g of 15 N-NH $_{4}$ Cl and 2 g of 13 C-glucose per liter, and the protein was expressed and purified as described above. The NMR spectra were collected at QANUC on either a Varian INOVA 500 MHz or 800 MHz NMR spectrometer with triple resonance cryoprobes. Backbone resonances were assigned using HNCACB, CBCA(CO) NH, HNCA, and HNCO triple resonance experiments and



side-chain resonances were assigned using CCC-TOCSY, HCC-TOCSY, and CT-heteronuclear single quantum coherence experiments (33). N- and C-NOESY-heteronuclear single quantum coherence (both aliphatic and aromatic) were used to obtain NOE distance restraints for structural determination purposes. Data were processed with NMRPipe and visualized with NMRDraw (https://www.niddk.nih.gov/researchfunding/at-niddk/labs-branches/laboratory-chemical-physics/ biophysical-nuclear-magnetic-resonance-spectroscopy-section/ software?from=NMRPipe/) (31). Spectral analysis was performed with either NMRView (30, 32) or analysis by CcpNmr (34, 35). Structures were calculated using CYANA (29) and with the CS-Rosetta server (28), and the models were visualized using PyMOL (The PyMOL Molecular Graphics System, Version 1.2, Schrödinger, LLC, https://pymol.org/2/). NOEderived models were compared to restraints using the Protein Structure Validation Suite (37) and the Analysis integrated RPF protocol (PyRPF) (34, 35). The CS-Rosetta determined ensembles of the N- and C-terminal domains have been deposited in the PDB with accession codes 6CZT and 6D10, respectively. Relevant NMR data has been deposited in the Biological Magnetic Resonance Bank, accession code 30450.

Structure analysis tools

Inter-residue contacts were determined by CMview (36) and visualized in Excel. Conservation analysis was performed using the ConSurf server with their automatically generated homolog search and multiple sequence alignment (39). Coulombic surface potentials were calculated in ChimeraX (70). Structures were visualized in PyMol and ChimeraX. A tertiary structure comparison search was conducted using the DALI server for the CYANA-determined AlgF N- and C-terminal domains (41).

Acetylesterase activity assay

Reactions were carried out as previously described (20, 21). Briefly, reactions contained 5 μ M of each protein (AlgJ_{Pa}⁷⁹⁻³⁷⁹, $AlgF_{Pa}^{30-216}$, $AlgX_{Pa}$) in 50 mM sodium Hepes pH 7.6 and 75 mM NaCl at 25 °C and were initiated with the addition of 3-carboxyumbelliferyl acetate (ACC) to 2 mM (dissolved in dimethylsulfoxide at stock concentrations of 20 mM). Deacetylated poly-mannuronate was prepared from P. aeruginosa FRD462 as previously described (71) and added at a concentration of 1 mg/ml. The final concentration of dimethylsulfoxide in each reaction was 2% (v/v). Hydrolysis was measured by fluorescence for 20 min with an excitation and emission wavelength of 386 and 446 nm, respectively (72). Reaction rates were calculated using a calibration curve for 7hydroxycoumarin-3-carboxylic acid, the fluorescent hydrolysis product of ACC. Background hydrolysis rates were measured and subtracted from reaction rates. Assays were carried out in triplicate in 96-well microtiter plates and measured using a SpectraMax M2 microplate reader (Molecular Devices). Data analysis was carried out in Prism 7 (GraphPad, https://www.graphpad.com), and statistical analyses were performed using an ordinary one-way ANOVA.

Alginate binding assay

Assays were performed as previously described (21) with a range of synthesized oligomannuronic acid oligomers (73).

Isothermal titration calorimetry

P. putida protein constructs were used for ITC analyses. ITC experiments were performed with a MicroCal Auto-(https://www.malvernpanalytical.com/en) in a buffer consisting of 50 mM Tris pH 7.8, 150 mM NaCl, 2% (v/v) glycerol, 50 mM arginine, and 50 mM glutamic acid at 20 °C. The Arg/Glu buffer components were used to reach the higher protein concentrations required for ITC experiments (74). ITC mixtures included: 1.86 mM AlgJ (syringe) titrated into 188 µM AlgF (cell) for AlgJ:AlgF; 4.17 mM AlgF (syringe) titrated into 500 µM AlgX (cell) for AlgF:AlgX; 2.97 mM AlgJ (syringe) titrated into 170 μM AlgX (cell) for AlgJ:AlgX. AlgX was placed in the cell due to a lower yield of protein. Runs were performed using 16 to 20 2.5 µl injections at an interval of 240 s. Data were analyzed using MicroCal Origin ITC Analysis software (https://www. malvernpanalytical.com/en).

P. aeruginosa strain construction

In-frame, unmarked algI and algI gene deletions in $P.\ aeruginosa\ PAO1\ \Delta wspF\ P_{BAD}alg\ (48)$ were generated using an established allelic replacement protocol (66). Construction of the gene deletion alleles was performed by amplifying flanking regions of the algI or algI ORFs and joining these flanking regions by splicing-by-overlap extension PCR (primers are listed in Table S3). The upstream forward and downstream reverse primers were tailed with EcoRI and HindIII restriction sequences, respectively, and the assembled $\Delta algI$ and $\Delta algI$ alleles were cloned into pEX18Gm.: $\Delta algI$ and pEX18Gm:: $\Delta algI$, were selected on LB agar containing 10 $\mu g/ml$ gentamicin and verified by Sanger sequencing using the M13F and M13R primers (Table S3).

The deletion alleles encoded by pEX18Gm:: $\Delta algI$ and pEX18Gm:: $\Delta algI$ were introduced into *P. aeruginosa* PAO1 $\Delta wspF$ P_{BAD}alg via biparental mating with the donor strain *E. coli* SM10 (75). Merodiploids were selected on VBMM agar containing 60 µg/ml gentamicin. SacB-mediated counter selection was carried out to select for double cross-over mutations on no-salt LB agar containing 15% (w/v) sucrose. Unmarked gene deletions were identified by colony PCR using primers that targeted the outside, flanking regions of *algI* or *algI* (Table S3). These PCR products were Sanger sequenced using the same primers to confirm the correct deletion.

Construction of strains encoding C terminally VSV-G—tagged AlgJ, AlgF, and AlgX was performed as above, with the following modifications. The upstream and downstream regions flanking the stop codon of *algX*, *algJ*, and *algF* were amplified using primer pairs whose upstream reverse and downstream forward primers were tailed with complementary sequence encoding the VSV-G peptide sequence (Table S3). The VSV-G sequence was encoded upstream of the stop

codon for each gene. The flanking upstream and downstream PCR products were then assembled by splicing-by-overlap extension PCR and cloned into pEX18Gm using SacI and HindIII restriction sites for AlgX and AlgJ and EcoRI and HindIII restriction sites for AlgF.

miniTn7 complementation

For gene complementation in P. aeruginosa, pUC18TminiTn7T-Gm, which allows for single-copy chromosomal insertion of genes (76), was modified to allow for arabinosedependent expression of complemented genes. The araC-P_{BAD} promoter from pJJH187 (77) was amplified using the primer pair miniTn7 pBAD F and miniTn7 pBAD R, the latter of which contains flanking sequence encoding Smal, Notl, PstI, and NcoI sites to generate a multiple cloning site downstream of the araC-PBAD promoter (Table S3). The resulting PCR product was cloned into the SacI and HindIII sites of pUC18T-miniTn7T-Gm to generate pUC18TminiTn7T-Gm-pBAD (Table S2).

The ORF corresponding to algJ and algI was amplified using the primer pairs AlgJ miniTn7 F + AlgJ miniTn7 R and AlgI miniTn7 F + AlgI miniTn7 R, respectively, which encode a synthetic ribosome binding site upstream of the start codon (Table S3). The resultant PCR products were cloned into pUC18T-miniTn7T-Gm-pBAD using the NcoI + SacI and NotI + NcoI sites, respectively, selected on LB agar containing 10 μg/ml gentamicin and 100 μg/ml carbenicillin, and confirmed by Sanger sequencing using the miniTn7 SeqF and miniTn7 SeqR primers (Table S3).

Complemented P. aeruginosa strains were generated through incorporation of miniTn7 vectors at the neutral attTn7 site on the P. aeruginosa chromosome via electroporation of miniTn7 vectors, along with the helper plasmid pTNS2, as previously described (67). Transposon mutants were selected on LB agar containing 30 µg/ml gentamicin.

Coimmunoprecipitation

One liter of LB, containing 0.5% (w/v) L-arabinose and 30 µg/ml gentamicin, was inoculated with a P. aeruginosa strain carrying a VSV-G-tagged alginate protein and allowed to grow overnight at 37 °C with shaking. The next day, cells were collected at 5000g for 20 min at 4 °C, resuspended in 50 ml of lysis buffer (20 mM Tris pH 8, 100 mM NaCl, 1 mM EDTA, 1 mg/ml lysozyme, 100 μg/ml DNase I, 2% (w/v) Triton X-100, 1 SIGMAFAST EDTA-free protease inhibitor cocktail tablet), and rocked for 2 h at 4 °C to allow for cell lysis. The cell lysate was subsequently clarified by centrifugation at 30,000g for 30 min at 4 °C. A sample of the clarified whole-cell lysate was collected before application to the immunoprecipitate (IP) resin as a representative example of the input into the experiment. The IP resin (Sigma anti-VSV-G mAb-agarose) was prepared by mixing 60 µl of slurry with 10 ml of wash buffer (20 mM Tris pH 8, 100 mM NaCl, 2% (w/v) Triton X-100), followed by collection of the IP resin by centrifugation at 100g for 2 min at 4 °C and removal of the supernatant. The clarified cell lysate was applied to the washed IP resin and incubated at 4 °C for 1 h with agitation. The IP resin was then collected by centrifugation at 100g for 2 min at 4 °C, and the supernatant was discarded. The resin was washed four times with 10 ml of wash buffer as above, followed by one wash with 10 ml of detergent-free wash buffer (20 mM Tris pH 8, 100 mM NaCl) to remove nonspecifically bound protein. Protein was then eluted from the resin by incubation in 110 µl of 0.2 M glycine, pH 2.2, for 15 min at room temperature, followed by collection of the resin by centrifugation at 100g for 2 min at 4 °C and removal of the supernatant containing eluted protein. The eluate was then neutralized by the addition of 40 µl of 1 M K₂HPO₄. Samples of the eluate were analyzed by ESI-MS by the SPARC BioCentre (The Hospital for Sick Children). For Western blot analysis, equal volumes of eluate and 2 × Laemmli buffer were mixed, heated at 95 °C for 10 min, and separated by SDS-PAGE, followed by Western blot as described below. As a negative control, an IP experiment was also performed using a P. aeruginosa strain expressing the corresponding untagged alginate protein.

Western blot sample preparation and analysis

To analyze protein levels from alginate-overproducing P. aeruginosa strains, 5 ml of LB containing 0.5% (w/v) Larabinose was inoculated with the appropriate strain and allowed to grow overnight at 37 °C with shaking. The next day, culture density was normalized to an $A_{600} = 1$, and 1 ml of the resulting culture was centrifuged at 5000g for 5 min to pellet cells. The cell pellet was resuspended in 100 µl of 2× Laemmli buffer, heated for 10 min at 95 °C, and analyzed by SDS-PAGE followed by Western blot.

For Western blot analysis, a 0.2 µm polyvinylidene difluoride (PVDF) membrane was wetted in methanol and soaked for 5 min in Western transfer buffer (25 mM Tris-HCl, 150 mM glycine, 20% (v/v) methanol) along with the SDS-PAGE gel to be analyzed. Protein was transferred from the SDS-PAGE gel to the PVDF membrane by wet blotting (25 mV, 2 h). The membrane was briefly rinsed in Tris-buffered saline (10 mM Tris-HCl pH 7.5, 150 mM NaCl) containing 0.5% (v/v) Tween-20 (TBS-T) before blocking in 5% (w/v) skim milk powder in TBS-T for 2 h at room temperature with gentle agitation. The membrane was briefly washed again in TBS-T before incubation overnight with primary antibody (1:1000 α -AlgJ, 1:250 α -AlgF, 1:1000 α -AlgX, 1:3000 α -PilP; described below) in TBS-T with 1% (w/v) skim milk powder at 4 °C. The next day, the membrane was washed four times in TBS-T for 5 min each before incubation for 1 h with secondary antibody (1:2000 dilution of Bio-Rad affinity purified goat α-rabbit immunoglogulin G conjugated to alkaline phosphatase) in TBS-T with 1% (w/v) skim milk powder. The membrane was then washed three times with TBS-T for 5 min each before development with 5-bromo-4-chloro-3-indolyl phosphate/ nitro blue tetrazolium chloride (BioShop ready-to-use BCIP/ NBT solution). Developed blots were imaged using a Bio-Rad ChemiDoc imaging system.



Antibody production and purification

PilP and AlgX antisera were generated and purified as described previously (20, 78). $AlgF_{Pa}^{30-216}$ and $AlgJ_{Pa}^{79-379}$ were purified as described above and used to generate antiserum from rabbits via a standard 70 day-protocol (Cedarlane Laboratories). The α -AlgF and α -AlgJ antibodies were further purified using a protocol adapted from Salamitou et al. (79). Briefly, 200 µg of purified $AlgF_{Pa}^{30-216}$ or $AlgJ_{Pa}^{79-379}$ were loaded on a 16% or 12% Tris-HCl polyacrylamide gel, respectively and transferred to a PVDF membrane. The membrane was stained with Ponceau S and the band corresponding to $\mathrm{AlgF}_{Pa}^{30\text{-}216}$ or $\mathrm{AlgJ}_{Pa}^{79\text{-}379}$ was cut out and blocked using PBS pH 7 with 0.1% (w/v) Tween 20 and 5% (w/ v) skim milk powder for 1 h. The membrane was then incubated with α-AlgF or α-AlgJ antisera overnight at 4 °C, followed by incubation at room temperature for 2 h. After washing in PBS, α-AlgF or α-AlgJ antibodies were eluted from the membrane by incubation in 700 µl of 0.2 M glycine pH 2.2 for 15 min, followed by neutralization with 300 µl of 1 M K₂HPO₄. The purified antibodies were dialyzed overnight against PBS, mixed 1:1 with glycerol, and stored at -20 °C. α -AlgF antibodies were used at a dilution of 1:250, α-AlgJ antibodies were used at a dilution of 1:1,000, α-AlgX antibodies were used at a dilution of 1:1,000, and anti-PilP antibodies were used at a dilution of 1:3000.

AlphaFold modeling

AlphaFold predictions were run using ColabFold v 1.5.2 with default parameters (*i.e.*, AlphaFold2_multimer_v3). No template information was used and the multiple sequence alignment options chosen were mmseq2_uniref_env and unpaired_paired. Five structures for each complex were predicted and used without relaxation using Amber. The sequences for *P. putida* were retrieve from UniProt accession numbers (AlgI: Q88ND2 AlgJ: Q88ND3; AlgF: Q88ND4; AlgX: Q88ND0; AlgK: Q88NC7) were used either as is (AlgI, AlgJ) or with their signal sequences (as determined by SignalP(69)) removed (AlgF, AlgX, and AlgK).

Data availability

The CS-Rosetta determined ensembles of the N- and C-terminal domains have been deposited in the PDB with accession codes 6CZT and 6D10, respectively. Relevant NMR data has been deposited to the BMRB under accession code 30450. The majority of software programs used in this report were configured and supported by the SBGrid consortium (80).

Supporting information—This article contains supporting information (21, 48, 75, 76, 81).

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Abbreviations—The abbreviations used are: Co-IP, coimmunoprecipitation; CSI, chemical shift index; ESI-MS, electrospray ionization mass spectrometry; IP, immunoprecipitate; ITC, isothermal titration calorimetry; LB, lysogeny broth; NOE, nuclear Overhauser effect; PISA, Proteins, Interfaces, Surfaces, and Assemblies; PVDF, polyvinylidene difluoride; TBS-T, Tris-buffered saline containing 0.5% (v/v) Tween-20; VBMM, Vogel-Bonner minimal medium; VSV-G, vesicular stomatitis virus glycoprotein.

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