

Membrane Topology of Outer Membrane Protein AlgE, Which Is Required for Alginate Production in *Pseudomonas aeruginosa*[∇]

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Received 6 December 2009/Accepted 13 January 2010

The ubiquitous opportunistic human pathogen *Pseudomonas aeruginosa* secretes a viscous extracellular polysaccharide, called alginate, as a virulence factor during chronic infection of patients with cystic fibrosis. In the present study, it was demonstrated that the outer membrane protein AlgE is required for the production of alginate in *P. aeruginosa*. An isogenic marker-free *algE* deletion mutant was constructed. This strain was incapable of producing alginate but did secrete alginate degradation products, indicating that polymerization occurs but that the alginate chain is subsequently degraded during transit through the periplasm. Alginate production was restored by introducing the *algE* gene. The membrane topology of the outer membrane protein AlgE was assessed by site-specific insertions of FLAG epitopes into predicted extracellular loop regions.

Pseudomonas aeruginosa is an ubiquitous opportunistic human pathogen responsible for chronic infections of the lungs of patients with cystic fibrosis (CF), in whom it is the leading cause of mortality and morbidity (9). The establishment of a chronic infection in the lungs of patients with CF coincides with the switch of *P. aeruginosa* to a stable mucoid variant, producing copious amounts of the exopolysaccharide alginate; this is typically a poor prognostic indicator for these patients (24, 31). Alginate is a linear unbranched exopolysaccharide consisting of 1,4-linked monomers of β -D-mannuronic acid and its C-5 epimer, α -L-guluronic acid, which is known to be produced by only two bacterial genera, *Pseudomonas* and *Azotobacter* (34). The switch to a mucoid phenotype coincides with the appearance of a 54-kDa protein in the outer membrane; this protein has been identified and has been designated AlgE (13, 31).

The genes encoding the alginate biosynthesis machinery are located within a 12-gene operon (*algD-alg8-alg44-algK-algE-algG-algX-algL-algI-algJ-algF-algA*). AlgA and AlgD, along with AlgC (not encoded in the operon), are involved in precursor synthesis (34). Alg8 is the catalytic subunit of the alginate polymerase located at the inner membrane (35). AlgG is a C-5 mannuronan epimerase (19). AlgK contains four putative Sell-like repeats, similar to the tetratricopeptide repeat motif often found in adaptor proteins involved in the assembly of multiprotein complexes (3, 10). AlgX shows little homology to any known protein, and its role is unclear (14). Knockout mutants of AlgK, AlgG, and AlgX have nonmucoid phenotypes, although they produce short alginate fragments, due to the activity of the alginate lyase (AlgL), which degrades the nascent alginate (1, 14, 19–21, 36). AlgF, AlgI, and AlgJ are involved in acetylation of alginate, but they are not ultimately

required for its production (12). The membrane-anchored protein, Alg44, is required for polymerization and has a PilZ domain for the binding of c-di-GMP, a secondary messenger essential for alginate production (16, 25, 33). The periplasmic C terminus of Alg44 shares homology with the membrane fusion proteins involved in the bridging of the periplasm in multidrug efflux pumps (11, 43). The periplasmic alginate lyase, AlgL, appears to be required for the translocation of intact alginate across the periplasm (1, 26). AlgE is an outer membrane, anion-selective channel protein through which alginate is presumably secreted (30). A protein complex or scaffold through which the alginate chain can pass and be modified and which spans the periplasm bridging the polymerase located (Alg8) at the outer membrane pore (AlgE) has been proposed (21). Indeed, it has been demonstrated that both the inner and the outer membranes are required for the *in vitro* polymerization of alginate (35).

The requirement of AlgE for the biosynthesis of alginate in *P. aeruginosa* was first observed by complementation of an alginate-negative mutant derived by chemical mutagenesis with a DNA fragment containing *algE* (8). Secondary structure predictions suggested that AlgE forms an 18-stranded β barrel with extended extracellular loops. Several of these loops show high densities of charged amino acids, suggesting a functional role in the translocation of the anionic alginate polymer (29, 30). Preliminary analysis of AlgE crystals has been reported (48).

In this study, the role of AlgE in alginate biosynthesis was investigated and the membrane topology of AlgE was assessed by site-directed insertion mutagenesis.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The bacterial strains, plasmids, and oligonucleotides used in the present study are listed in Table 1. The *Escherichia coli* strains were grown in LB medium at 37°C. *E. coli* S17-1 was used for conjugative transfer of suicide plasmid pEX100T: Δ algEGm and flipase-encoding plasmid pFLP2. When they were required, the following antibiotics were used at the indicated concentrations: ampicillin, 100 μ g/ml; gentamicin, 10 μ g/ml; and streptomycin, 30 μ g/ml. The *P. aeruginosa* strains were grown in LB medium or *Pseudomonas* isolation agar (PIA) medium (20 g of peptone, 10 g of K₂SO₄, 1.4 g

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[∇] Published ahead of print on 22 January 2010.

TABLE 1. Bacterial strains, plasmids and oligonucleotide used in this study

Strain or plasmid	Characteristics	Reference or source
Strains		
<i>P. aeruginosa</i>		
PDO300	<i>mucA22</i> isogenic mutant derived from PAO1	18
PDO300 Δ <i>algE</i>	Isogenic <i>algE</i> deletion mutant derived from PDO300	This study
<i>E. coli</i>		
TOP10	<i>E. coli</i> cloning strain	Invitrogen
S17-1	<i>thi-1 proA hsdR17</i> ($r_K^- m_K^+$) <i>recA1</i> ; <i>tra</i> gene of plasmid RP4 integrated in chromosome	39
Plasmids		
pGEM-T Easy	Ap^r , P_{lac}	Invitrogen
pBBR1MCS-5	Gm^r ; broad-host-range vector; P_{lac}	23
pBBR1MCS-5: <i>algE</i>	HindIII-BamHI fragment comprising <i>algE</i> inserted into vector pBBR1MCS-5	This study
pBBR1MCS-5: <i>algE</i> DelL7	<i>algE</i> fragment with 57-bp deletion in region encoding putative surface exposed loop 7 inserted into pBBR1MCS-5	This study
pBBRMCS-5: <i>algEL1</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 189th bp of the ORF	This study
pBBRMCS-5: <i>algEL2</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 336th bp of the ORF	This study
pBBRMCS-5: <i>algEL3</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 468th bp of the ORF	This study
pBBRMCS-5: <i>algEL4</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 582nd bp of the ORF	This study
pBBRMCS-5: <i>algEL5</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 711th bp of the ORF	This study
pBBRMCS-5: <i>algEL6</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 882nd bp of the ORF	This study
pBBRMCS-5: <i>algEL7</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 1035th bp of the ORF	This study
pBBRMCS-5: <i>algEL8</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 1269th bp of the ORF	This study
pBBRMCS-5: <i>algEL9</i> FLAG	<i>algE</i> fragment with a 24-bp insertion (encoding the FLAG epitope) inserted after the 1428th bp of the ORF	This study
pEX100T	Ap^r Cb^r ; gene replacement vector containing <i>sacB</i> gene for counterselection	17
pEX100T: Δ <i>algEGm</i>	Ap^r Cb^r Gm^r ; vector pEX100T with SmaI-inserted <i>algE</i> deletion construct	This study
pPS865	Ap^r Gm^r ; source of 1,100-bp BamHI fragment comprising <i>aacC1</i> gene flanked by Flp recombinase target site signal sequences	17
pPFLP2	Ap^r Cb^r ; broad-host-range vector encoding Flp recombinase	17

MgCl₂, 0.025 g of Triclosan, 20 ml of glycerol per liter) at 37°C, and when they were required, antibiotics were added at the appropriate concentrations. The antibiotic concentrations used for the *P. aeruginosa* strains were as follows: gentamicin, 300 µg/ml, and carbenicillin, 300 µg/ml. All chemicals were purchased from Sigma-Aldrich (St. Louis, MO).

Isolation, analysis, and manipulation of DNA. General cloning procedures were performed as described previously (37). All pBBR1MCS-5-derived plasmids were transferred to *P. aeruginosa* strains via electroporation, as described previously (7). DNA primers, deoxynucleoside triphosphates, and *Taq* and platinum *Pfx* polymerases were purchased from Invitrogen. The DNA sequences of the plasmid constructs were confirmed by DNA sequencing.

Construction and confirmation of *algE* deletion mutants. Two regions of the *algE* gene were amplified by using *Taq* polymerase and primers algE1N-Ec5, algE1C-Ba, algE2N-Ba, and algE2C-Ec5. Region algEN (469 bp) comprised bases 136 to 585 and region algEC (445 bp) comprised bases 1047 to 1472 relative to the designated *algE*-coding region (41). Both PCR products were hydrolyzed by using BamHI and were inserted into the pGEM-T Easy vector (Promega). Vector pPS865 (17) was hydrolyzed with BamHI. The fragment containing the *aacC1* gene (which encodes gentamicin acetyltransferase) flanked by two Flp recombinase target sites was inserted into the BamHI site of plasmid pGEM-TEasy:*algENC*, resulting in plasmid pGEM-TEasy: Δ *algEGm*. The DNA of the 1,989-bp *algEGm* fragment was amplified by using *Pfx* polymerase and primers algE1N-Ec5 and algE2C-Ec5, and the corresponding PCR product was inserted into SmaI site of vector pEX100T (17), resulting in plasmid pEX100T: Δ *algE* Ω Gm.

E. coli S17-1 was used as the donor for the transfer of plasmid pEX100T: Δ *algE* Ω Gm into the *P. aeruginosa* strains, and transconjugants were selected on mineral salt medium (38) containing gentamicin and 5% (wt/vol) sucrose. Cells

growing on this selective medium should have emerged from double-crossover events. Gene replacement was confirmed after subculture of the cells on PIA medium containing gentamicin and by PCR with primers algEup and algEDnA.

E. coli S17-1 was used to transfer the Flp recombinase-encoding vector pPFLP2 (17) into *P. aeruginosa* PDO300 Δ *algE* Ω Gm strains, and after 24 h of cultivation on PIA medium containing 5% (wt/vol) sucrose, the gentamicin- and carbenicillin-sensitive cells were analyzed by PCR for the loss of the gentamicin resistance-conferring cassette.

Complementation of isogenic *algE* deletion mutants. For complementation of the *algE* deletion mutants, the *algE* open reading frame (ORF) of *P. aeruginosa* PAO1 was amplified by using *Pfx* polymerase and primers algEN(HiSDNd) and algEC(Ba). This product was then inserted into the pGEM-T Easy vector, resulting in plasmid pGEMTEasy:*algE*. The *algE* fragment was released by hydrolysis with HindIII and BamHI and was inserted into the HindIII and BamHI sites of broad-host-range vector pBBR1MCS-5 (23), resulting in plasmid pBBR1MCS-5:*algE*. In addition, a plasmid encoding AlgE with a 9-amino-acid deletion in the highly charged putative surface-exposed loop 7 region was constructed. This was constructed by amplifying the region upstream of the deletion point by using *Pfx* polymerase and primers algEN(HiSDNd) and algEDel7N(Ba) and the region downstream of the deletion point by using primers algEDel7C(Ba) and algEC(SaI). The two products were hydrolyzed with HindIII and BamHI or BamHI and SacI, respectively, and inserted into the HindIII and SacI sites of pBBR1MCS-5, resulting in plasmid pBBR1MCS-5:*algE*DelL7.

In order to probe the membrane topology of AlgE, insertions of the FLAG (DYKDDDDK) epitope were introduced into all of the nine putative surface-exposed loops. The insertions were constructed by site-directed, ligase-independent mutagenesis (SLIM) (6). Briefly, two PCRs were completed for each construct by using plasmid pGEMTEasy:*algE* as the template: one with the

"F_{FLAG}" and the corresponding "R_S" primer (algEL1F_{FLAG} and algEL1R_S, algEL2F_{FLAG} and algEL2R_S, algEL3F_{FLAG} and algEL3R_S, algEL4F_{FLAG} and algEL4R_S, algEL5F_{FLAG} and algEL5R_S, algEL6F_{FLAG} and algEL6R_S, algEL7F_{FLAG} and algEL7R_S, algEL8F_{FLAG} and algEL8R_S, and algEL9F_{FLAG} and algEL9R_S) and one with the F_S and the corresponding R_{FLAG} primer (algEL1F_S and algEL1R_{FLAG}, algEL2F_S and algEL2R_{FLAG}, algEL3F_S and algEL3R_{FLAG}, algEL4F_S and algEL4R_{FLAG}, algEL5F_S and algEL5R_{FLAG}, algEL6F_S and algEL6R_{FLAG}, algEL7F_S and algEL7R_{FLAG}, algEL8F_S and algEL8R_{FLAG}, and algEL9F_S and algEL9R_{FLAG}). The plasmid template was removed by hydrolysis with DpnI. The two PCR products were mixed in equimolar amounts, and hybridization was achieved by incubation in H buffer (150 mM NaCl, 25 mM Tris, 20 mM EDTA, pH 8.0) at 99°C for 3 min, followed by three cycles of 65°C for 5 min and 30°C for 40 min. The resulting mixture was used to transform competent *E. coli* TOP 10 cells. Selection for cells coating the new plasmid was performed on ampicillin-containing medium, and the plasmids were extracted. Insertion of the 24-bp FLAG-encoding region was confirmed by DNA sequencing of the ORF, resulting in plasmids pGEMTEasy:algEL1FLAG to pGEMTEasy:algEL9FLAG. The *algE* (FLAG)-containing fragments were hydrolyzed and inserted into pBBR1MCS-5, as described above, resulting in plasmids pBBR1MCS-5:algEL1FLAG, pBBR1MCS-5:algEL2FLAG, pBBR1MCS-5:algEL3FLAG, pBBR1MCS-5:algEL4FLAG, pBBR1MCS-5:algEL5FLAG, pBBR1MCS-5:algEL6FLAG, pBBR1MCS-5:algEL7FLAG, pBBR1MCS-5:algEL8FLAG, and pBBR1MCS-5:algEL9FLAG.

Alginate production assays. Two milliliters of an overnight culture was harvested at 4°C and washed twice with saline. Then, 200 µl of the cell suspension was plated onto PIA medium and incubated 72 h at 37°C. Cells were scraped off of two agar plates by using a sterile spatula and were washed twice with 100 ml of saline (retaining the alginate-containing supernatant for subsequent precipitation). The cellular sediments were freeze-dried, and the final weight was determined. The alginate-containing supernatants were precipitated with 1 volume of ice-cold isopropanol, and the alginate was harvested and freeze-dried. For further purification, the precipitated alginate was redissolved in 0.05 M Tris-HCl–10 mM MgCl₂ (pH 7.4) to a final concentration of 0.5% (wt/vol), followed by incubation with 15 µg of DNase I/ml and 15 µg of RNase A/ml at 37°C for 6 h. Pronase E was added to a final concentration of 20 µg/ml, and this solution was incubated for a further 18 h at 37°C. The solutions were dialyzed against 5 liters of ultrapure H₂O for 48 h. Alginate was precipitated with 1 volume of ice-cold isopropanol and freeze-dried for quantification and uronic acid analysis.

The levels of free uronic acids (alginate degradation products) in the supernatants of 2 ml of overnight cultures were measured. The total uronic acid content of the supernatant was determined as described below; the supernatants were filtered with Amicon Ultra-0.5 (Millipore) centrifugal filter devices (nominal molecular mass cutoff, 10 kDa), and the flowthrough was collected. The uronic acid content of the flowthrough (which contained free uronic acids and short-length alginate degradation products) was determined as described below.

Uronic acid assay. The alginate concentrations were assayed by the uronic acid assay described previously (4) by using alginic acid from brown seaweed (Sigma-Aldrich) as the standard. Briefly, the alginate samples were dissolved in 200 µl ultrapure H₂O at concentrations of between 0.25 and 0.05 mg/ml. The sample was mixed with 1.2 ml tetraborate solution (0.0125 M disodium tetraborate in concentrated sulfuric acid), and the mixture was incubated on ice for 10 min. The mixtures were incubated at 100°C for 5 min and then cooled down on ice for a further 5 min. Twenty microliters of *m*-hydroxybiphenyl reagent (0.15% *m*-hydroxybiphenyl in 0.125 M NaOH) was added and the reaction mixtures were mixed for 1 min. For each sample or dilution, a negative control was assayed by using 0.0125 M NaOH instead of the hydroxybiphenyl reagent. The uronic acid concentrations were determined spectrophotometrically at a wavelength of 520 nm.

Purification of outer membranes. Strains of *P. aeruginosa* were grown overnight in LB medium containing the appropriate antibiotics. The cells were harvested by centrifugation (at 5,000 × *g* for 1 h) and washed twice with 1 volume of 10 mM HEPES (pH 7.4). The cells were placed in 15 ml 10 mM HEPES with Roche complete mini-EDTA-free protease inhibitor and sonicated on ice for 12 cycles of 15 s of sonication, followed by a 15-s cooldown. Cellular debris and the remaining intact cells were sedimented by centrifugation (at 5,000 × *g* for 1 h). The total membrane fraction was then isolated by centrifugation at 100,000 × *g* for 2 h. The supernatant (soluble fraction) was removed, the sediment was resuspended in 1 volume of 10 mM HEPES containing 0.7% (wt/vol) *N*-lauroylsarcosine, and the suspension was incubated at room temperature with shaking for 2 h to selectively solubilize the cytoplasmic membrane. This was then cen-

trifuged at 100,000 × *g* for 2 h; the resulting sediment represents the outer membrane fraction. The total protein concentration of the respective fractions was determined by using a Quant-iT protein assay kit (Invitrogen).

Analysis of outer membrane proteins. Fifteen micrograms of total protein was then separated by SDS-PAGE on 8% polyacrylamide gels. The resulting gels were then either stained with Coomassie brilliant blue or transferred to a nitrocellulose membrane by using an iBlot dry blotting system (Invitrogen). The nitrocellulose membrane was blocked with 5% (wt/vol) skim milk powder in Tris-buffered saline with 0.1% Tween 20 for 1 h and was subsequently probed with 1 µg/ml anti-FLAG (M2) tag antibody conjugated to horseradish peroxidase (HRP; Abcam, Cambridge, United Kingdom). The membrane was washed, and the bound antibodies were resolved with SuperSignal West Pico chemiluminescent substrate (Thermo Scientific, Rockford, IL) and developed on X-ray film. Bands suspected of being AlgE or the AlgE mutants were identified by tryptic peptide fingerprinting by matrix-assisted laser desorption/ionization–time of flight (MALDI-TOF) mass spectrometry (MS).

RESULTS

Construction of an isogenic knockout mutant of *algE*. To investigate the requirement for AlgE in alginate biosynthesis, a marker-free *algE* deletion mutant of alginate-overproducing strain *P. aeruginosa* PDO300 was generated. This mutant showed a nonmucoid phenotype when it was grown on solid medium. The outer membrane protein profiles showed that AlgE was absent from PDO300Δ*algE* (Fig. 1, left panel). It has been shown that the lack of mucoidity associated with some deletion mutants (*algK*, *algG*, and *algX*) is distinct from the biosynthesis/polymerization of alginate. These mutants secrete alginate degradation products (free uronic acids), which have been shown to be the products of alginate lyase, AlgL (19–21, 36). To address this for the Δ*algE* mutant, the culture supernatants of the respective mutants were filtered and the uronic acid contents of the filtrates (containing alginate degradation products) were determined. Alginate degradation products could be detected in PDO300Δ*algE* (pBBR1MCS-5) at levels 9.7 times that of PDO300 (pBBR1MCS-5) and 3.2 times that of the positive control, PDO300Δ*algX* (pBBR1MCS-5) (Table 2). Only small amounts of uronic acid were detected from the negative control, PDO300Δ*alg8* (pBBR1MCS-5) (Table 2). These results suggest that the Δ*algE* mutant is capable of the synthesis/polymerization of alginate but that it is subsequently degraded in the periplasm.

Complementation of the PDO300Δ*algE* isogenic knockout mutant. To verify that the observed loss of mucoidity was not due to polar effects on other genes in the alginate biosynthesis operon, a plasmid containing the *algE* ORF (pBBR1MCS-5: *algE*) was used to complement the mutant in *trans*. This plasmid was able to restore the mucoid phenotype as well as the production of alginate to levels beyond (five times greater) that of the parent strain containing the vector control PDO300 (pBBR1MCS-5) (Table 2). This indicates that there were no polar effects in the deletion mutant and demonstrates the requirement of AlgE for the mucoid phenotype and the formation of intact and extracellular alginate. Interestingly, complemented strain PDO300Δ*algE* (pBBR1MCS-5: *algE*) produced more alginate degradation products than PDO300 (pBBR1MCS-5) (53.6% and 8.4% total uronic acids, respectively) (Table 2). The presence of AlgE in the outer membrane was also restored (Fig. 1, left panel).

Predicted topology of AlgE. The PRED-TMBB transmembrane β-strands prediction program for Gram-negative bacte-

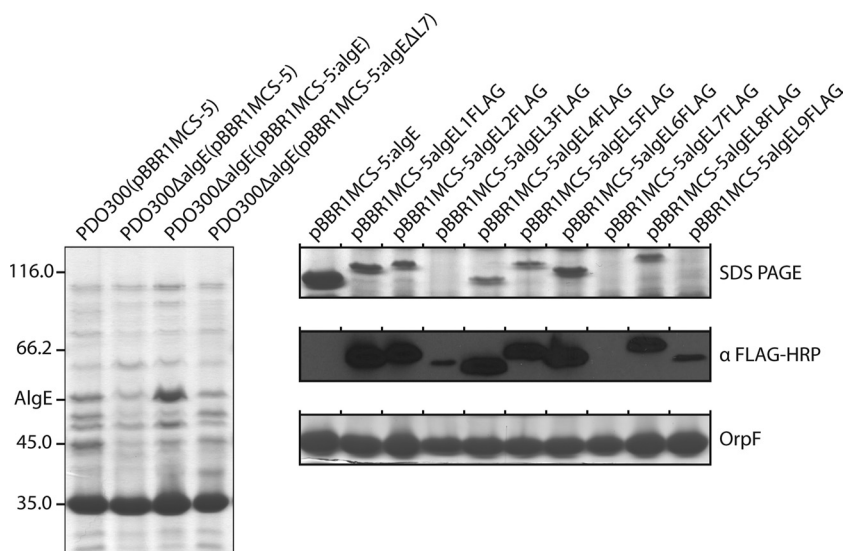


FIG. 1. (Left panel) Outer membrane profiles of the $\Delta algE$ mutant and the $\Delta algE$ mutant harboring various plasmids showing the absence or presence of AlgE. Faint bands with the same apparent molecular mass (indicated on the left, in thousands) as AlgE present in the PDO300 $\Delta algE$ (pBBR1MCS-5) and PDO300 $\Delta algE$ (pBBR1MCS-5:algE Δ L7) outer membrane profiles were identified as flagellin type B protein by MALDI-TOF MS. (Right panels) Presence of tagged AlgE proteins in the outer membrane, as indicated by Western immunoblotting (middle). Constitutively produced OrpF was used to standardize the samples (bottom).

ria outer membrane proteins (2) predicts that the mature AlgE protein has 18 transmembrane β strands with nine extended surface-exposed loops and eight short periplasmic turns (Fig. 2). This is similar to what was predicted previously (29, 30). A search through the HHPred program (40) did not reveal homology to any annotated protein families or domains with known structures with which to refine this model.

The extracellular loop 7 region of AlgE is proposed to be required for the translocation of the alginate polymer. The proposed surface-exposed loop 7 is significantly longer than the other loops and contains a high density of positively charged amino acid residues which are conserved among all *Pseudomonas* species in the *Pseudomonas* Genome Database sequenced (49) (data not shown), as well as AlgJ from *Azoto-*

bacter vinelandii (29). A conserved cluster of these positively charged residues is present at the apex of the loop region, suggesting that this region may be folded into the pore, facilitating the translocation of the anionic alginate chain through the pore. A vector encoding AlgE with an 18-amino-acid deletion in this conserved region at the apex of loop 7 (pBBR1MCS-5:algE Δ L7) was constructed in an attempt to analyze any role that this region may have in alginate translocation and/or protein folding/structure. This plasmid could not restore alginate production in the $\Delta algE$ mutant, nor could the AlgE variant be localized to the outer membrane (Table 2; Fig. 1). This strain produced free uronic acids, but at levels 1.6 times lower than those for the *algE* deletion mutant, PDO300 $\Delta algE$ (pBBR1MCS-5).

TABLE 2. Production of alginate and free uronic acids by $\Delta algE$ mutants

Strain	Mean alginate concn (g/g CDW ^a) \pm SD	Free uronic acids	
		Source ^b	Mean concn (g/g CDW) \pm SD
PDO300(pBBR1MCS-5)	0.216 \pm 0.027	Total	0.524 \pm 0.076
		Filtrate	0.044 \pm 0.009
PDO300 $\Delta algE$ (pBBR1MCS-5)	ND ^c	Total	0.486 \pm 0.050
		Filtrate	0.425 \pm 0.039
PDO300 $\Delta algE$ (pBBR1MCS-5:algE)	1.096 \pm 0.251	Total	0.500 \pm 0.014
		Filtrate	0.268 \pm 0.010
PDO300 $\Delta algE$ (pBBR1MCS-5:algE Δ L7)	ND	Total	0.249 \pm 0.007
		Filtrate	0.261 \pm 0.031
PDO300 $\Delta algX$ (pBBR1MCS-5)	ND	Total	0.138 \pm 0.002
		Filtrate	0.131 \pm 0.001
PDO300 $\Delta alg8$ (pBBR1MCS-5)	ND	Total	0.012 \pm 0.002
		Filtrate	0.013 \pm 0.001

^a CDW, cellular dry weight.

^b Total, concentration of uronic acids in the unfiltered culture media (including alginate and free uronic acids); Filtrate, concentration of uronic acids in the filtered culture media (free uronic acids).

^c ND, none detected.

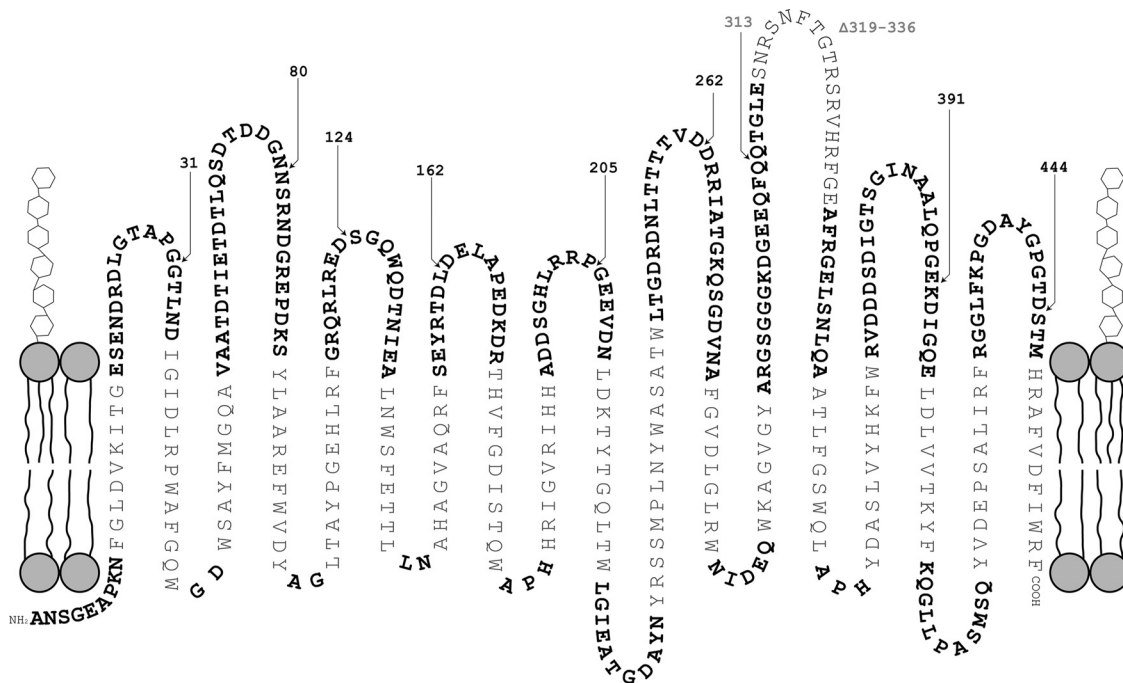


FIG. 2. Predicted topology of the outer membrane β -barrel AlgE. Predicted transmembrane β sheets are indicated in regular text (nonbold-face), extracellular loops at the top are indicated in boldface, and periplasmic turns at the bottom are indicated in boldface. Locations where the FLAG epitopes were inserted are indicated by triangles, and the positions (relative to the position of the cleavage of the signal peptide) are indicated; black numbers, the insertion was permissive, i.e., alginate production was restored; gray numbers, the insertion was nonpermissive. The deleted region of loop 7 is indicated by regular gray text (nonboldface).

AlgE membrane topology analysis. To test the predicted AlgE model presented in Fig. 2, site-directed epitope insertion mutagenesis targeting the proposed extracellular loops was done. This method utilizes the observation that insertions in the hydrophilic loop regions distant from the β -barrel core are usually tolerated, permitting the correct formation of the β barrel required for insertion of the protein into the outer membrane (27, 32). The FLAG epitope was inserted into the predicted loop regions for loops 1 to 9 by SLIM (6) at the positions indicated in Fig. 2. Plasmids encoding these tagged proteins were introduced into the $\Delta algE$ mutant strain, the outer membrane protein profiles were obtained by SDS-PAGE, and the respective proteins were subjected to Western immunoblotting with anti-FLAG HRP-conjugated antibody. Insertion of the FLAG epitope into loops 1, 6, 8, and 9 were tolerated, and the protein could be detected in the respective outer membranes. Interestingly, although the tagged proteins have identical deduced molecular masses of 52.2 kDa, they migrated significantly differently by SDS-PAGE: insertions in loops 1, 6, and 9 at 53.5 kDa; loop 2 at 54.7 kDa; loop 3 at 52.3 kDa; loop 4 at 51.4 kDa; loop 5 at 55 kDa; and loop 8 at 56.5 kDa. Additionally, AlgEL3FLAG and AlgEL9FLAG appeared to be present at significantly smaller quantities than the other AlgE proteins (Fig. 1, right panels). AlgE variants harboring insertions in loops 1 to 6, 8, and 9 restored alginate production in the $\Delta algE$ mutant (Table 3), indicating that the core protein structure was not disrupted and that no functional regions essential for alginate formation were affected. Surprisingly, insertion of the epitope into the proposed loop 7 region was not tolerated.

This protein could not be detected in the outer membrane and did not restore alginate formation (Table 3).

DISCUSSION

The role of AlgE in alginate production and its membrane topology were investigated in the present study. The isogenic marker-free *algE* deletion mutant was nonmucoid and did not produce alginate (Table 2). Alginate secretion could be restored by introduction of only the *algE* gene in *trans*. This indicates that AlgE is essential for the mucoid phenotype. The *algE* deletion mutant secreted free uronic acids, indicating that alginate is still being polymerized but is presumably degraded

TABLE 3. Alginate production of $\Delta algE$ complemented with various epitope tagged AlgE variants

Strain	Mean alginate concn (g/g CDW ^a) \pm SD
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL1</i> FLAG)	0.728 \pm 0.080
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL2</i> FLAG)	1.261 \pm 0.236
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL3</i> FLAG)	0.874 \pm 0.081
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL4</i> FLAG)	0.835 \pm 0.067
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL5</i> FLAG)	1.091 \pm 0.053
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL6</i> FLAG)	0.796 \pm 0.146
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL7</i> FLAG)	ND ^b
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL8</i> FLAG)	0.654 \pm 0.105
PDO300 $\Delta algE$ (pBBR1MCS-5: <i>algEL9</i> FLAG)	0.892 \pm 0.158

^a CDW, cellular dry weight.

^b ND, none detected.

by AlgL in the periplasm (Table 2). Thus, like AlgK, AlgX, AlgG, and AlgL, AlgE is not required for the polymerization or translocation of alginate across the inner membrane but is required for the successful translocation of alginate through the periplasm and across the outer membrane (1, 14, 19–21, 36). This supports the hypothesis that alginate is guided through the periplasm by a multiprotein complex or scaffold containing periplasmic proteins AlgK, AlgX, AlgG, and AlgL and outer membrane protein AlgE. The loss of any of these proteins results in the lack of the integrity of this complex and degradation of the alginate chain. It is possible that the predicted protein scaffold is attached to the outer membrane via AlgE, perhaps via the tetratricopeptide repeat motif of AlgK often found in adaptor proteins involved in the assembly of multiprotein complexes and/or the membrane fusion domain of Alg44 (3, 33). Indeed, mutants lacking Alg44 show reduced levels of AlgE in the outer membrane (28).

The in *trans* complementation of the Δ algE mutant still resulted in the secretion of free uronic acids (albeit less than the level of secretion by the Δ algE mutant) (Table 2). Since AlgE is overproduced in the outer membrane (Fig. 1) of the complemented Δ algE mutant, it could interfere with the stoichiometry of proteins in the proposed scaffold complex. Hence, the presence of an increased AlgE copy number might, to a certain extent, increase the number of functional multiprotein complexes contributing to alginate overproduction, while the additional copy number might also lead to dysfunctional multiprotein complex formation, causing the degradation of some of the alginate as it traverses the periplasm.

Deletion of the conserved extracellular loop 7 region likely interfered with the folding pathway of AlgE, as evidenced by its absence in the outer membrane and its inability to restore alginate production (Fig. 1; Table 3). This suggests that the deleted region is essential for the functional folding of AlgE. The deleted region contains a relatively high concentration of conserved (among all sequences of *Pseudomonas* spp. in the *Pseudomonas* Genome Database [49] as well as AlgJ of *Azotobacter vinelandii*) positively charged amino acid residues, and therefore, it was proposed that this region is involved in the active translocation of the anionic alginate chain through the outer membrane (Fig. 2). Alternatively, it cannot be ruled out that the topology prediction could be incorrect and that the deletion site may be in an unpredicted transmembrane region.

Site-directed epitope insertion mutagenesis of all predicted extracellular loop regions was completed to probe the predicted membrane topology of AlgE. Similar methods have been utilized to probe the topologies of various outer membrane proteins (27, 32, 46, 47). As expected, insertion of the FLAG epitope into eight of nine loop regions was tolerated and did not disrupt the β -barrel structure, allowing insertion into the outer membrane (Fig. 1). These proteins were also capable of complementing the Δ algE mutant by restoring alginate production (Table 3). Hence, supporting evidence that these tagged regions are loop regions not involved in the formation of the β barrel through which alginate is exported was provided. When the FLAG epitope was inserted into the predicted loop 7 region, it was not tolerated, even though this insertion site is the most distant from any predicted β sheets among the loop insertions. However, it was consistent with the finding that the deletion of an 18-amino-acid region 5 amino

acid residues upstream of the insertion site in this loop was not tolerated (Fig. 1; Table 3). Loop 7, which was proposed to be actively involved in the transport of alginate, might be folded and might bend into the hydrophobic β -barrel core, similar to the “eyelet” region described in other porins. Indeed, this loop 7 contains a concentration of charged residues similar to that found in other eyelet loops (5, 22, 42, 44, 45). Consequently, insertion of the highly hydrophilic FLAG tag into this loop could disrupt the hydrophobic core of the barrel and prevent folding.

Under the conditions used in this study, AlgE did not migrate aberrantly as a 54-kDa protein, as described elsewhere (8, 30), but migrated at its deduced molecular mass of 51.2 kDa. Additionally, even though they had identical deduced molecular masses, the epitope-tagged AlgE variants migrated differently, an observation similar to that reported previously for OprH (32) (Fig. 1). This can be explained by the observation that outer membrane proteins are known for their extraordinary stability and have been reported to exhibit heat-modifiable characteristics, resulting in differences in their migration patterns when they are subjected to SDS-PAGE (15).

In conclusion, experimental evidence was provided for the requirement of AlgE for the secretion of alginate, whereas it is not required for alginate polymerization and translocation into the periplasm. Furthermore, it was demonstrated that loop 7 is required for the functional folding of AlgE, while the other insertion mutants support the proposed membrane topology as a β barrel with 18 β strands and 9 loops. Overall, the data suggest that AlgE not only might be involved in the secretion of alginate but also contributes to the proposed protein scaffold, guiding and protecting the nascent alginate chain through the periplasm and outer membrane.

ACKNOWLEDGMENTS

This study was supported by research grants to B.H.A.R. from the Institute of Molecular BioSciences at Massey University and the Deutsche Forschungsgemeinschaft (Re 1097/6-1). I.D.H. is funded by a Massey University doctoral scholarship. Z.U.R. is supported by a doctoral scholarship and research grant from the Higher Education Commission of Pakistan.

We are grateful to Uwe Remminghorst for the construction of plasmid pEX100T: Δ algEGm.

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