The outer membrane protein OprQ and adherence of *Pseudomonas aeruginosa* to human fibronectin

Abraham Arhin and Cliff Boucher

The University of Texas at Tyler, 3900 University Blvd, Tyler, TX 75701, USA

Outer membrane proteins of the Gram-negative organism *Pseudomonas aeruginosa* play a significant role in membrane permeability, antibiotic resistance, nutrient uptake, and virulence in the infection site. In this study, we show that the *P. aeruginosa* outer membrane protein OprQ, a member of the OprD superfamily, is involved in the binding of human fibronectin (Fn). Some members of the OprD subfamily have been reported to be important in the uptake of nutrients from the environment. Comparison of wild-type and mutant strains of *P. aeruginosa* revealed that inactivation of the oprQ gene does not reduce the growth rate. Although it does not appear to be involved in nutrient uptake, an increased doubling time was reproducibly observed with the loss of OprQ in *P. aeruginosa*. Utilizing an oprQ–xylE transcriptional fusion, we determined that the PA2760 gene, encoding OprQ, was upregulated under conditions of decreased iron and magnesium. This upregulation appears to occur in early exponential phase. Insertional inactivation of PA2760 in the *P. aeruginosa* wild-type background did not produce a significant increase in resistance to any antibiotic tested, a phenotype that is typical of OprD family members. Interestingly, the in trans expression of OprQ in the ΔoprQ PAO1 mutant resulted in increased sensitivity to certain antibiotics. These findings suggest that OprQ is under dual regulation with other *P. aeruginosa* genes. Intact *P. aeruginosa* cells are capable of binding human Fn. We found that loss of OprQ resulted in a reduction of binding to plasmatic Fn in vitro. Finally, we present a discussion of the possible role of the *P. aeruginosa* outer membrane protein OprQ in adhesion to epithelial cells, thereby increasing colonization and subsequently enhancing lung destruction by *P. aeruginosa*.

INTRODUCTION

*Pseudomonas aeruginosa* is an opportunistic Gram-negative pathogen that causes infections in individuals with altered immune systems, such as burn, HIV, nosocomial and neutropenic patients (DiGiandomenico *et al.*, 2007; Sadikot *et al.*, 2005). This pathogen is also notorious for its role in infections of the respiratory tract of patients with cystic fibrosis (Wagner & Iglewski, 2008). The ability of *P. aeruginosa* to cause such a wide range of infections is, in part, due to the outer membrane proteins that it expresses (Wu *et al.*, 2005). One major class of outer membrane proteins (Oprs), containing members that function as virulence factors, is the OprD superfamily (Tamber *et al.*, 2006).

Outer membrane proteins are involved in many aspects of growth and development of the bacterial cell (Hancock & Brinkman, 2002). The outer membrane protein OprD superfamily consists of 19 members, all of which display at least 46% similarity at the amino acid level (Tamber *et al.*, 2006). The members of this family participate in amino acid and peptide transport, antibiotic uptake, and transport of carbon sources (Hancock & Brinkman, 2002). In addition, the permeability of the *P. aeruginosa* outer membrane has been attributed, at least in part, to the surface expression of the OprD class of proteins (Strateva & Yordanov, 2009). Although OprD homologues may perform multiple roles, it does not appear that the function of the channels of this family is non-specific. Each member of the OprD subfamily, at least those involved in nutrient uptake, seems to have a very narrow specificity (Tamber & Hancock, 2003). For example, *P. aeruginosa* OprD, a homologue of the *Escherichia coli* porin OmpF, exists in the outer membrane in a β-barrel configuration, and is known to play a significant role in the uptake of basic amino acids and resistance to imipenem and carbapenem (Hancock & Brinkman, 2002; Lister, 2002). The exact role for some members, such as OprQ and OprI, has yet to be determined. Recently, the group of Tamber has reported that a specific nutrient substrate for OprQ and OprI cannot be identified, at least in the *P. aeruginosa* PAK background (Hancock & Brinkman, 2002).

It has been reported that as many as 100 outer membrane proteins are produced by *P. aeruginosa* (Hancock & Brinkman, 2002). Proteins localized to the outer membrane...
of *P. aeruginosa* have been shown not only to be involved in nutrient uptake and antibiotic resistance, but also to participate in this bacterium's pathogenicity. Some *P. aeruginosa* outer membrane virulence factors, such as XcpQ (Brok et al., 1999; Robert et al., 2005), QxhA (Martinez et al., 1998; Michel et al., 2007), EstA (Wilhelm et al., 1999), OprG (McPhee et al., 2009) and OprE (Yamano et al., 1998), have become areas of active research. However, there remain many Oprs that have not been investigated and their respective roles in *P. aeruginosa* have not yet been identified. One such potential virulence factor is the *P. aeruginosa* outer membrane protein OprQ. It has been reported that OprQ is a member of the OprD superfamily of proteins (Jaouen et al., 2006; Tamber et al., 2006). As stated above, initial studies of OprQ in *P. aeruginosa* did not reveal a specific role (Tamber et al., 2006). Studies of OprQ function in *Pseudomonas fluorescens* suggest that it plays a role significantly different from that of other members of the OprD subfamily. In this regard, OprQ in *P. fluorescens* has been shown to bind fibronectin (Fn) *in vitro* (Rebière-Huët et al., 2002). The function of *P. aeruginosa* OprQ in growth, antibiotic resistance and the colonization of the host is currently not known, but these studies suggest that OprQ is directly involved in the disease process.

In this study, we report the unexpected finding that loss of OprQ resulted in an increased rate of growth. We also found that in the inactivated oprQ background, the overexpression of OprQ may increase *P. aeruginosa* sensitivity to antibiotics. In addition, we determined that OprQ is upregulated under conditions of low iron and magnesium. Since these conditions are thought to mimic conditions at known sites of *P. aeruginosa* infection, it is possible that OprQ is needed for the establishment of disease. Finally, we report that OprQ in *P. aeruginosa* is potentially involved in adhering to Fn, thus resulting in the initial colonization of the host epithelial tissue.

### METHODS

**Bacterial plasmids, strains and culture conditions.** The strains, primers and plasmids used in this study are listed in Table 1. PAA2760 is a derivative of the *P. aeruginosa* strain PAO1. All strains were grown in the minimal medium M63, Pseudomonas Isolation Agar (PIA) or Luria–Bertani (LB; Difco) medium, with supplementation as indicated.

Using the PAO1 genomic database, oligonucleotides were designed to encompass the sequence between PA2760 and the upstream gene PA2759. Primers 2760F-3 and 2760R-1 for the DNA sequence of PA2760 were used in a PCR to amplify the gene for cloning. The PCR product was then cloned into the vector pCR2.1 to create the plasmid pAA2760Pr using the TOPO TA cloning kit (Invitrogen). This plasmid was then digested with the restriction enzyme EcoRI (Invitrogen), and the 232 bp promoter fragment was isolated and subsequently cloned into the EcoRI-digested vector pVDX18 (Konyecsni & Deretic, 1988), creating the fusion plasmid pAA2760-XylE.

To generate the suicide vector pAA2760Tc, the primers 2760F-1 and 2760R-2 were used in a PCR to amplify a 1.5 kb fragment of the

### Table 1. Bacterial strains, plasmids and primers

<table>
<thead>
<tr>
<th>Strain, primer or plasmid</th>
<th>Description</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>P. aeruginosa</strong> strains</td>
<td>Wild-type prototroph</td>
<td>B. Holloway (Monash University, Australia)</td>
</tr>
<tr>
<td>PAO1</td>
<td>PAO1 oprQ::Tc</td>
<td>This work</td>
</tr>
<tr>
<td>PAA2760</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>E. coli</strong> strain</td>
<td>lacZ+, recA+, Gal+</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>DH5a</td>
<td>5'-GTCAGGACAGTGTCCT-3'</td>
<td>This work</td>
</tr>
<tr>
<td><strong>Primers</strong></td>
<td>5'-GGACGGTGCGTGCG-3'</td>
<td>This work</td>
</tr>
<tr>
<td>2760F-1</td>
<td>5'-GGAAACGAGCAAGACGATG-3'</td>
<td>This work</td>
</tr>
<tr>
<td>2760F-2</td>
<td>5'-CGGGCACAACCGGCACC-3'</td>
<td>This work</td>
</tr>
<tr>
<td>2760F-3</td>
<td>5'-CAAGAGCGAGCGGCAAGGC-3'</td>
<td>This work</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
<td>tra+, mob+, Kan+</td>
<td>D. R. Helsinki, University of California, San Diego</td>
</tr>
<tr>
<td>pVDX18</td>
<td>IncQ, mob+, tac, lacPl</td>
<td>Deretic et al. (1987)</td>
</tr>
<tr>
<td>pUCtc</td>
<td>pCR2.1, partial oprQ coding sequence</td>
<td>This work</td>
</tr>
<tr>
<td>pAA2760</td>
<td>pCR2.1, oprQ::Tc</td>
<td>This work</td>
</tr>
<tr>
<td>pAA2760Tc</td>
<td>pCR2.1, complete oprQ coding sequence</td>
<td>This work</td>
</tr>
<tr>
<td>pAA00oprQ</td>
<td>pCR2.1, OprQ promoter region</td>
<td>This work</td>
</tr>
<tr>
<td>pAA2760WT</td>
<td>pVDCtac39 oprQ+</td>
<td>This work</td>
</tr>
<tr>
<td>pAA2760Pr</td>
<td>pCR2.1, OprQ promoter region</td>
<td>This work</td>
</tr>
<tr>
<td>pAA2760-XylE</td>
<td>pVDCtac39 oprQ::xylE</td>
<td>This work</td>
</tr>
</tbody>
</table>

Downloaded from www.microbiologyresearch.org by IP: 54.70.40.11 On: Thu, 06 Dec 2018 01:20:21
PA2760 gene. This fragment was cloned into plasmid pCR2.1 (Invitrogen), creating plasmid pAA2760WT. This plasmid was double-digested with the restriction enzymes HindIII and XhoI (both from New England Biolabs), and the fragment containing the PA2760 gene was extracted from an agarose gel via a PureLink Quick Gel Extraction kit (Invitrogen). The plasmid vector pUC19 (New England Biolabs) was also double-digested with HindIII/XhoI and subsequently ligated with the PA2760 HindIII/XhoI fragment. E. coli strain DH5α, harbouring the newly constructed pUC2760 plasmid, was grown on agar (Difco plates containing 50 μg ampicillin ml⁻¹ (Invitrogen). The isolated pUC2760 plasmid DNA was further digested with the restriction enzyme BstEII (Invitrogen), purified and blunt-ended with large fragment Klenow (Invitrogen). The plasmid pUC19, containing a cassette that confers tetracycline resistance (TcR), was digested with PstI. The 1.5 kb tetracycline resistance gene cassette was isolated from a 1% agarose gel and blunt-ended. The vector (pUC2760) and insert (TcR cassette) were ligated in the presence of 5% PEG 1000 in order to concentrate the blunt-ended DNA fragments. Following ligation, DH5α cells were transformed with the ligated DNA and plated onto LB agar plates containing 10 μg tetracycline ml⁻¹ (Invitrogen), and the suicide vector pAA2760Tc containing the tetracycline insert was identified via digestion with HindIII and XhoI and visualization on a 1% agarose gel.

For complementation of the OprQ mutant, we created the plasmid pAA2760WT. Using the primers 2760F-1 and 2760R-3 in a PCR, we amplified the complete coding region of the oprQ gene. The amplified fragment was gel-purified and cloned into the EcoRI-digested plasmid pVDtac-39 (Deretic et al., 1987).

All bacterial triparental conjugations were carried out using the donor E. coli DH5α strain carrying the plasmids pAA2760-XylE and pAA2760WT with conjugates grown in LB supplemented with either ampicillin (50 μg ml⁻¹) or tetracycline (10 μg ml⁻¹), respectively. Recipient PAO1 was grown in LB and E. coli harbouring the helper plasmid pRK2013 (Figurski & Helinski, 1979) in LB with kanamycin (50 μg ml⁻¹). Each strain was grown overnight in 5 ml of medium, spun, washed in sterile saline and resuspended in 0.5 ml LB medium. The Pseudomonas recipient strain, donor DH5α, and the helper E. coli were combined (0.9 ml total) and passed through a 0.45 μm pore-size filter. The filter was placed on an LB plate and grown overnight at 37°C. The bacterial growth from the filter was plated on PIA plates with either 1000 μg carbenicillin ml⁻¹ or 150 μg tetracycline ml⁻¹. Growth of P. aeruginosa strains for doubling time determination was performed using both LB and M63 media supplemented with 1% glucose or glycerol (as carbon source), and either 0.2% potassium nitrate or ammonium sulfate (as nitrogen source). c.f.u. were determined by serial dilution in 0.8% sodium chloride and plating on LB agar plates containing 500 μg carbenicillin ml⁻¹. One millilitre of a P. aeruginosa overnight culture was used to inoculate a fresh solution of M63 in a 1:100 dilution, and the culture was grown at 37°C with shaking until the appropriate OD₆₀₀ was reached (see above). The concentrations of magnesium and iron used for each experimental condition were as follows: (i) regular magnesium, 1.0 mM MgSO₄·7H₂O; (ii) low magnesium, 0.2 mM MgSO₄·7H₂O; (iii) regular iron, 0.1 mM FeCl₃; and (iv) low iron, 0.01 mM FeCl₃. All cultures were centrifuged at 10000 r.p.m. for 10 min. The cells were then resuspended in 5 ml lysis buffer (50 mM potassium phosphate buffer, pH 7.5, 10% acetone; Sigma-Aldrich). Cells were sonicated with a Branson cell sonifier for 1 min on ice with no interruptions. The disrupted cells were centrifuged and protein concentration was determined by the Bradford method in a Beckman Coulter spectrophotometer at 375 nm. Samples were prepared by adding to a 5 ml cuvette (Bio-Rad) the following: 2.5 ml H₂O, 0.3 ml 0.5 M potassium phosphate buffer and 0.1 ml 10 mM catechol, all at room temperature. The correct amount of extract was added to equilibrate samples to 30 μg ml⁻¹, except the blank, which received no protein. Samples were added immediately and inverted two to three times to mix thoroughly. Readings were done at 375 nm, for 150 s with a 2.5 s interval. For each condition, regular and low iron or magnesium, a total of three independent sample protein extracts were analysed.

**Electroporation and generation of the insertional mutant PAA2760.** The newly created pAA2760Tc plasmid was used via electroporation to transform the wild-type P. aeruginosa strain PAO1 using the method of Smith & Iglewski (1989). Briefly, to create electrocompotent P. aeruginosa cells, 500 μl of 300 mM sucrose was added to a microcentrifuge tube and approximately 3 mg of PAO1 cells, grown overnight in LB at 37°C, was suspended in the sucrose solution. The P. aeruginosa cells were washed twice, and the final pellet was resuspended in 40 μl 300 mM sucrose. The electroporation-competent P. aeruginosa was kept on ice for the remainder of the procedure. Five microlitres of pAA2760Tc DNA was added to the PAO1 cell pulsed at 3.0 kV mm⁻¹ using a Bio-Rad Electropulsor, 1 ml of LB broth was added, and the cells were incubated at 37°C with shaking for 1 h. Cells were then plated (100 μl) onto LB agar plates containing 50 μg tetracycline ml⁻¹ and incubated overnight at 37°C. Possible PA2760 knockout mutants were screened by replica plating on LB agar plates containing 500 μg carbenicillin ml⁻¹. Since the pUC19 plasmid is non-replicative in PAO1, sensitivity to carbenicillin (CarbR) and resistance to tetracycline (TetR) were used to ensure that the oprQ gene was knocked out during electroporation. Carbenicillin sensitivity indicates the loss of the pUC19 plasmid. The PAO1 2760::TetR carb conjugant was verified for proper insertion of the genetic cassette by PCR using the primers 2760F-2 and 2760R-2, which flank the tetracycline insertion site.

**Xyle assays.** The catechol-2,3-dioxigenase (XylE) activity of the OprQ transcriptional fusion was assayed as described by Schweizer & Hoang (1995). Three separate protein extracts were prepared by growing conjugated cells overnight in M63 medium supplemented with 300 μg carbenicillin ml⁻¹. One millilitre of a P. aeruginosa overnight culture was used to inoculate a fresh solution of M63 in a 1:100 dilution, and the culture was grown at 37°C with shaking until the appropriate OD₆₀₀ was reached (see above). The concentrations of magnesium and iron used for each experimental condition were as follows: (i) regular magnesium, 1.0 mM MgSO₄·7H₂O; (ii) low magnesium, 0.2 mM MgSO₄·7H₂O; (iii) regular iron, 0.1 mM FeCl₃; and (iv) low iron, 0.01 mM FeCl₃. All cultures were centrifuged at 10000 r.p.m. for 10 min. The cells were then resuspended in 5 ml lysis buffer (50 mM potassium phosphate buffer, pH 7.5, 10% acetone; Sigma-Aldrich). Cells were sonicated with a Branson cell sonifier for 1 min on ice with no interruptions. The disrupted cells were centrifuged and protein concentration was determined by the Bradford method in a Beckman Coulter spectrophotometer at 375 nm. Samples were prepared by adding to a 5 ml cuvette (Bio-Rad) the following: 2.5 ml H₂O, 0.3 ml 0.5 M potassium phosphate buffer and 0.1 ml 10 mM catechol, all at room temperature. The correct amount of extract was added to equilibrate samples to 30 μg ml⁻¹, except the blank, which received no protein. Samples were added immediately and inverted two to three times to mix thoroughly. Readings were done at 375 nm, for 150 s with a 2.5 s interval. For each condition, regular and low iron or magnesium, a total of three independent sample protein extracts were analysed.

**Antibiotic disk diffusion assay.** Disks saturated separately with cefotaxime (30 μg), gentamicin (10 μg), amoxicillin/clavulanic acid (30 μg), erythromycin (15 μg), vancomycin (30 μg), levofloxacin (5 μg), ampicillin (10 μg), piperacillin (100 μg), chloramphenicol (2 μg), neomycin (30 μg), meropenem (10 μg) and aztreonam (30 μg) were applied to Mueller–Hinton agar plates containing P. aeruginosa strain PAO1, PAA2760 or PAA2760 carrying wild-type oprQ. The plates were incubated for 18 h at 37°C. At the end of the incubation period, the diameters (mm) of the zones of growth inhibition were measured. Replicates for antibiotic susceptibility determination were performed for each strain using separate inoculations from individual cultures.

**Determination of MIC.** The MICs of colistin, ciprofloxacin, imipenem monohydrate, tetracycline and tobramycin sulfate at starting concentrations of 16, 10, 50 and 4 μg ml⁻¹, respectively, were determined by twofold serial dilution using a sterile 96-well flat-bottomed microtitre plate, as described by Andrews (2001). The MICs of each individual strain were determined from at least three independent assays.

http://mic.sgmjournals.org
Fn-binding assay. Fn-binding assays were performed in 96-well sterile microtitre plates. Wells were coated overnight at 8 °C with 1.25 μg plasmonic Fn (Sigma-Aldrich) in sterile PBS (125 μl per well of 10 μg Fn ml⁻¹). The wells were washed four times with 125 μl of 1% (w/v) BSA in PBS. Then wells were blocked with 125 μl of 1% (w/v) BSA in PBS for 1 h. Just before adding bacteria, wells were washed six times with sterile prewarmed PBS.

P. aeruginosa strains PAO1, PAA2760 and PAA2760 pAA2760WT were grown overnight at 37 °C on LB plates. Bacteria were resuspended in PBS and collected by centrifugation. The bacteria were washed twice with prewarmed PBS and resuspended in the same sterile buffer at OD₆₀₀ 0.4. A total of 50 μl of bacterial suspension was added to the Fn-coated wells and incubated at room temperature for 3 h. Non-adherent bacteria were removed by washing six times, using vigorous pipetting, with sterile PBS. Adherent bacteria were then collected with 100 μl sterile PBS containing 0.5% Triton X-100 under manual desorption. Bacteria were plated by 10-fold serial dilutions onto LB plates, incubated overnight at 37 °C, and quantified the following day. A minimum of three replicates for Fn binding were performed for each strain using separate inoculations from individual cultures.

Quantification of pyocyanin. For quantification of pyocyanin levels in PAO1, PAA2760 and PAA2760 carrying the wild-type oprQ in trans, strains were grown in LB medium for 48 h at 30 °C. Cultures were centrifuged at 7 000 g for 15 min. The pyocyanin pigment was extracted using chloroform/0.2 M HCl (1:1, v/v). Quantification was performed by the measurement of A₅₃₀ on a Beckman DU 800 spectrophotometer (Beckman Coulter).

Statistical analysis. Statistical analysis of the results of the growth analysis, xylE assay, antibiotic susceptibility and Fn binding were conducted by analysis of variance (Tukey’s post test) using Systat (Systat Software).

RESULTS

Growth rate increase upon loss of OprQ

The OprD family of porins is diverse, with a few members participating in nutrient acquisition from the environment (Hancock & Brinkman, 2002; Tamber & Hancock, 2003). As a member of this OprD subfamily, it is possible that the P. aeruginosa OprQ participates in nutrient uptake. One earlier study using a transposon oprQ mutant in the P. aeruginosa PAK background was unable to determine a specific amino acid transported by PA2760 (Tamber et al., 2006). It remains possible that OprQ is involved in the overall growth of P. aeruginosa. To further investigate the growth phenotype for OprQ in the P. aeruginosa PAO1 wild-type background, we insertionally inactivated the oprQ gene using a tetracycline resistance gene cassette. The parent and mutant strains were grown on minimal M63 medium to determine whether any phenotypic changes could be observed. Interestingly, P. aeruginosa strain PAA2760, in which OprQ expression was disrupted, had a significant increase in growth rate as compared with the PAO1 parent when cultured in LB medium (Fig. 1). As seen in Fig. 1, the faster doubling time for PAA2760, 29.9 min, was retarded back to the wild-type rate of 44 min when pAA2760WT was introduced into the OprQ mutant.

Substitution of glucose and potassium nitrate with glycerol and ammonium sulfate resulted in the same growth phenotype, suggesting that the increased doubling time is independent of the carbon or nitrogen source. These results indicate that OprQ is not required for the uptake of essential nutrients for growth. Furthermore, the observation of an increased growth rate suggests that OprQ is involved, directly or indirectly, in the regulation of other proteins/porins that are important in P. aeruginosa growth.

Transcription and expression of OprQ is increased under stress conditions

Under conditions of poor oxygenation, Jaouen et al. (2006) revealed, using RT-PCR, that the P. fluorescens oprQ gene is overexpressed. Oxygen stress and the induction of an OprD family outer membrane protein are not without precedents. It has been reported that OprE in P. aeruginosa is similarly induced in oxygen-depleted environments (Yamano et al., 1993). In order to identify other potential stressors that may induce the expression of the oprQ gene, the PA2760 promoter region, an intergenic sequence containing a potential promoter-binding site of the stress response sigma factor RpoH, was cloned into a xylE vector to analyse the expression of OprQ under varying conditions, such as that which might be encountered in the infection site. Comparisons were made between PAO1, harbouring the pAA2760-XylE plasmid, grown in M63 medium supplemented with regular magnesium (1.0 mM MgSO₄·7H₂O) and iron (0.1 mM FeCl₃) and M63 medium supplemented with low magnesium (0.2 mM MgSO₄·7H₂O) and iron (0.01 mM FeCl₃). Exposure to conditions of low magnesium resulted in a fivefold increase in transcription of the oprQ gene (Fig. 2). An even greater effect on promoter activity was observed when the strains were placed under low-iron stress. Under this condition, a 13-fold increase in oprQ promoter activity was observed (Fig. 2). These results suggest that OprQ is required for survival under conditions where P. aeruginosa is exposed to adverse growth conditions.

OprQ and antibiotic resistance

The ability of P. aeruginosa to be resistant to such a wide range of antibiotics is due to the outer membrane porins that it produces (Strateva & Yordanov, 2009). In particular, members of the OprD outer membrane protein superfamily have been shown to function in antibiotic resistance (Tamber et al., 2006). Since OprQ does not appear to play a significant role in growth, and given the potential role in antibiotic resistance, we hypothesized that OprQ affects the sensitivity of P. aeruginosa to these antimicrobials. It has been reported that OprQ is not important for carbapenem resistance (Okamoto et al., 1999). Therefore, we expanded this analysis of P. aeruginosa sensitivity upon inactivation of the oprQ gene to include 18 different antibiotics (gentamicin, neomycin, amoxicillin/clavulanic acid, ampicillin, piperacillin, levofloxacin, ciprofloxacin, meropenem, imipenem, cefsulodin, cefotaxime, colistin, vancomycin, tetracycline,
tobramycin, chloramphenicol, erythromycin and aztreonam) from 12 different classes (aminoglycosides, penicillins, quinolones, carbapenems, cephalosporins, polymyxins, glycopeptides, tetracyclines, aminoglycosides, amphenicols, macrolides and monobactams). Analysis of susceptibilities of the wild-type, PAA2760 and PAA2760 harbouring the pAA2760WT plasmid strains to these antibiotics was performed using the Kirby–Bauer and MIC methods (Andrews, 2001; Traub & Leonhard, 1994). Comparison of the zone of inhibition on disk diffusion for PAO1 and PAA2760 showed no significant statistical difference (Fig. 3).

Interestingly, it was observed that when the PAA2760 strain expressed wild-type OprQ, from the pAA2760WT plasmid, there was a reproducible increase in sensitivity to gentami-

\[\text{Fig. 1. Growth curve of PAO1 wild-type (}),\text{ PAA2760 (oprQ::Tcr) (})\text{ and PAA2760 harbouring the plasmid pAA2760WT (oprQ')} (\text{}).\text{ (a) Strains grown at 37 °C in LB medium. (b) Strains grown at 37 °C in M63 minimal medium supplemented with 1% glucose and 0.2% potassium nitrate. Asterisks indicate statistical difference between PAO1/PAA2760 harbouring pAA2760WT and PAA2760 as determined by Tukey’s post test (ANOVA).}\]

\[\text{Fig. 2. Transcriptional activation of the P. aeruginosa oprQ gene under stress conditions. The units of catechol-2,3-dioxygenase (CDO) are specific activity per milligram of protein. Bacterial growth in M63 was supplemented with regular iron (0.1 mM FeCl}_3; \text{Reg Fe}), low iron (0.01 mM FeCl}_3; \text{Low Fe}), regular magnesium (1.0 mM MgSO}_4, 7H}_2O; \text{Reg Mg}) or low magnesium (0.2 mM MgSO}_4, 7H}_2O; \text{Low Mg}) as indicated. The statistical significance for regular versus low iron was } P=0.001. \text{ An asterisk indicates statistical significance for regular versus low magnesium as determined by ANOVA (Tukey’s post test, SYSTAT).}\]
cin, neomycin and meropenem (Fig. 3). When we assayed for antibiotic resistance using the MIC method, the same sensitivity phenotype was observed for the antibiotics colistin, imipenem, tetracycline and tobramycin when wild-type OprQ was placed in trans (Fig. 4). This suggests that increased expression and outer membrane localization of OprQ lead to the repression of proteins required for P. aeruginosa resistance to these antibiotics.

**Determination of Fn-binding activity**

The binding of human Fn and its relationship to infection for outer membrane proteins of P. fluorescens have been established. A total of six P. fluorescens Fn-binding outer membrane proteins (FnBPs) have been identified (Rebière-Huët et al., 2002). One of these proteins, identified as a 44 kDa outer membrane protein, is homologous to the P. aeruginosa outer membrane protein OprQ (Jaouen et al., 2006). Studies presented here have excluded OprQ from a role in nutrient uptake and resistance to many classes of antibiotics. However, it remains possible that OprQ is indirectly involved in P. aeruginosa sensitivity to certain antibiotics. Therefore, based on the reports of P. fluorescens and the similarity of OprQ to these FnBPs, we hypothesized that OprQ could play a role in P. aeruginosa binding to the extracellular matrix via Fn. In order to test the ability of P. aeruginosa to bind plasmatic Fn, we compared wild-type PAO1 with the PAO1 ΔoprQ strain PAA2760. Addition of whole-cell P. aeruginosa PAO1 to plates coated with human Fn resulted in adsorption (Fig. 4). Upon addition of PAA2760 to Fn in vitro, there was a significant drop, 88-fold, in the ability of P. aeruginosa to adhere to the extracellular matrix protein (Fig. 4). The introduction of the plasmid pAA2760WT into PAA2760 led to a 41-fold
Therefore, it is expected that this outer membrane protein specifically involved in nutrient uptake and resistance regulation with other proteins, possibly Oprs that are proteins. The regulation of function in roles common to this class of outer membrane proteins may extend beyond those involved in metabolism, growth and antibiotic sensitivity. In addition to causing the effects previously discussed, OprQ appears to have an effect on the production of P. aeruginosa pyocyanin. Previous studies on pyocyanin have revealed that this secreted phenazine is important in P. aeruginosa virulence. It has been speculated that this type of gene regulation could be widely utilized by prokaryotes (Schmitz & McDonald, 1984). Likewise, the increased sensitivity to certain antibiotics may be controlled at the transcriptional level. Seven clinical isolates of P. aeruginosa from a hospital in the USA have been found to overexpress the mexXY genes (Wolter et al., 2004). In all seven cases, the transcription of the oprD gene was repressed, resulting in a decrease in outer membrane OprD. Upon entrance into the infection site, stress conditions encountered by P. aeruginosa may lead to an increased expression of OprQ and enhanced binding to host tissue via Fn. This would be in conjunction with a decrease in growth rate, a phenotype that could aid in this organism’s ability to survive in the host.

Furthermore, it was observed that the overexpression of OprQ could lead to increased sensitivity to certain classes of antibiotics. This suggests that expression of OprQ could be tied to the regulation of optimal growth characteristics for P. aeruginosa that favour the prevailing environmental condition. A model has been proposed for the outer membrane OmpC in E. coli whereby its transcriptional activation is reciprocal to that of the ompF gene (Ozawa & Mizushima, 1983). Early studies described this regulatory phenotype as being closed. Under conditions that favour upregulation of the E. coli ompF gene, OmpF expression is decreased (Schmitz & McDonald, 1984). Likewise, when ompF transcription is activated there is a reciprocal repression of the ompC gene. This phenotype is also apparent when a mutation in OmpF is created in E. coli. The expression of OmpC in this background is derepressed (Ozawa & Mizushima, 1983). It is now understood that this reciprocal regulation is controlled by the transcriptional factor OmpR (Yoshida et al., 2006). It has been speculated that this type of gene regulation could be widely utilized by prokaryotes (Schmitz & McDonald, 1984). Likewise, the increased sensitivity to certain antibiotics may be controlled at the transcriptional level. Seven clinical isolates of P. aeruginosa from a hospital in the USA have been found to overexpress the mexXY genes (Wolter et al., 2004). In all seven cases, the transcription of the oprD gene was repressed, resulting in a decrease in outer membrane OprD. Upon entrance into the infection site, stress conditions encountered by P. aeruginosa may lead to an increased expression of OprQ and enhanced binding to host tissue via Fn. This would be in conjunction with a decrease in growth rate, a phenotype that could aid in this organism’s ability to survive in the host.

**DISCUSSION**

In this study, we provide evidence that the P. aeruginosa OprQ protein, a member of the OprD superfamily, does not function in roles common to this class of outer membrane proteins. The regulation of oprQ expression may be in dual regulation with other proteins, possibly Ops that are specifically involved in nutrient uptake and resistance to antibiotics, since the PAO1 Δ oprQ strain displays an increased growth rate under stress conditions.

**Fig. 4.** Effect of OprQ on P. aeruginosa binding to human Fn. Binding of P. aeruginosa to Fn is reported as c.f.u. per well. The statistical significance between PAO1 and PAA2760 was P=0.0153. Overexpression of the oprQ gene was achieved by addition of 1 mM IPTG to the growth medium. This concentration of IPTG was maintained during incubation of PAA2760 harbouring the pAA2760WT plasmid with Fn. Asterisks indicate statistical significance between PAO1 and PAA2760 as determined by ANOVA (Tukey’s post test, SYSTAT).

Increase in adsorption as compared with PAA2760 alone (Fig. 4). The restored binding to human Fn in vitro was still significantly lower than that observed for the P. aeruginosa parent strain PAO1. The complementation plasmid pAA2760WT contains an IPTG-inducible promoter. Addition of 1 mM IPTG to PAA2760 carrying the pAA2760WT plasmid resulted in complete restoration of the binding by P. aeruginosa to levels indistinguishable from those of PAO1 (Fig. 4). These results suggest that OprQ is important in binding of the extracellular matrix protein Fn. Therefore, it is expected that this outer membrane protein will prove to be important for P. aeruginosa in causing disease. Furthermore, the level of adsorption to human Fn is determined by the amount of OprQ expressed by P. aeruginosa.

In this study, we provide evidence that the P. aeruginosa OprQ protein, a member of the OprD superfamily, does not function in roles common to this class of outer membrane proteins. The regulation of oprQ expression may be in dual regulation with other proteins, possibly Ops that are specifically involved in nutrient uptake and resistance to antibiotics, since the PAO1 Δ oprQ strain displays an increased growth rate under stress conditions.
pyocyanin from PAA2760 by A$_{530}$ revealed at least a threefold increase over P. aeruginosa PAO1 (0.05 vs 0.15). Data presented here seem to suggest that the early expression of OprQ on the outer membrane of P. aeruginosa may be important for colonization. However, upon establishing disease and the formation of an environment that favours diminished expression of OprQ, pyocyanin production may be increased. This would allow for the expression of the full complement of virulence factors and the establishment of more substantial disease. Further investigation into the relationship between OprQ and pyocyanin production will be required to determine the exact interaction.

Most pathogens have been reported to bind host tissue via the protein Fn, which aids in colonization and infection (Joh et al., 1999; Rebière-Huët et al., 1999). The expression of Fn in epithelial cells has been positively correlated with the degree of injury in affected tissues (Roman et al., 2006). It is possible that the damaged cells of patients that express high levels of Fn provide a means for P. aeruginosa to adhere to and colonize the patient. In addition, it has been demonstrated that in the host respiratory epithelium, the innate mechanisms that protect against invading pathogens can be compromised when levels of iron are altered (Ganz, 2009). It is thought that the decreased level of iron in the respiratory tract is important as an innate mechanism against infection (Ganz, 2009). In this environment, for a true opportunistic pathogen such as P. aeruginosa, the ability to overcome the iron-reduced condition is crucial. The upregulation of virulence factors, especially in the initial stage of infection, would aid in P. aeruginosa colonization in this stressful host environment. This hypothesis is in agreement with the induction of the oprQ gene as presented in this study. We have determined that under conditions of reduced iron and magnesium, such as those described above, OprQ expression increases. Upon activation of oprQ, P. aeruginosa adherence to the host extracellular matrix would be enhanced. This phase of colonization to host tissue would lead to further tissue damage and the subsequent increased expression of Fn and thus more P. aeruginosa binding. This suggests that OprQ would be an important virulence factor involved in P. aeruginosa infections.

**ACKNOWLEDGEMENTS**

This research was supported by grant BOUCHERUTT3 from The University Of Texas at Tyler faculty research program.


Edited by: P. van der Ley