Multiple roles for a novel RND-type efflux system in Acinetobacter baumannii AB5075

Kyle A. Tipton, Emory University
Marjan Farokhyfar, Atlanta VA Medical Center
Philip Rather, Emory University

Journal Title: MicrobiologyOpen
Volume: Volume 6, Number 2
Publisher: Wiley Open Access | 2016-01-01, Pages e00418- e00418
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1002/mbo3.418
Permanent URL: https://pid.emory.edu/ark:/25593/s19nd

Final published version: http://dx.doi.org/10.1002/mbo3.418

Copyright information:
© 2016 Published by John Wiley & Sons Ltd.
This is an Open Access work distributed under the terms of the Creative Commons Attribution 4.0 International License (https://creativecommons.org/licenses/by/4.0/).

Accessed April 28, 2019 7:37 AM EDT
Multiple roles for a novel RND-type efflux system in Acinetobacter baumannii AB5075

Kyle A. Tipton | Marjan Farokhyfar | Philip N. Rather

1Department of Microbiology and Immunology, Emory University School of Medicine Atlanta, Atlanta, GA, USA
2Research Service, Atlanta VA Medical Center, Decatur, GA, USA
3Emory Antibiotic Resistance Center, Emory University School of Medicine, Atlanta, GA, USA

Correspondence
Philip N. Rather
Department of Microbiology and Immunology, Atlanta, GA, USA.
Email: prather@emory.edu

Funding Information
This work was funded by grant 1R21AI115183 from the National Institutes of Health. P. N. R. is also supported by grants from the Merit Review program and a Research Career Scientist award, both from the Department of Veterans Affairs.

Abstract
Colony opacity phase variation in Acinetobacter baumannii strain AB5075 is regulated by a reversible high-frequency switch. Transposon mutagenesis was used to generate mutations that decreased the opaque to translucent switch and a gene encoding a predicted periplasmic membrane fusion component of a resistance–nodulation–cell division (RND)-type efflux system was isolated. This gene was designated arpA and immediately downstream was a gene designated arpB that encodes a predicted membrane transporter of RND-type systems. A nonpolar, in-frame deletion in arpA resulted in a 70-fold decrease in the opaque to translucent switch. An arpB::Tc mutant exhibited a 769-fold decrease in the opaque to translucent switch. However, the translucent to opaque switch was largely unchanged in both the arpA and arpB mutants. The arpA and arpB mutants also exhibited increased surface motility in the opaque form and the arpB mutant exhibited increased susceptibility to aminoglycosides. The arpA and arpB mutants were both attenuated in a Galleria mellonella model of virulence. A divergently transcribed TetR-type regulator ArpR was capable of repressing the arpAB operon when this TetR regulator was overexpressed. The arpR gene was also involved in regulating the opaque to translucent switch as an in-frame arpR mutation decreased this switch by 1,916-fold.

KEYWORDS
Acinetobacter, phase variation, RND efflux system

1 | INTRODUCTION

Acinetobacter baumannii is a Gram-negative pathogen typically associated with infections in hospital settings (Bergogne-Berezin & Towner, 1996; Gootz & Marra, 2008; Joly-Guillou, 2005; Peleg, Seifert, & Paterson, 2008). Although most A. baumannii infections are seen in immunocompromised patients or those with severe injuries, community acquired infections and infections in otherwise healthy patients have increased in recent years (Antunes, Visca, & Towner, 2014; Charnot-Katsikas et al., 2009; Guerrero et al., 2010; Lowman, Kalk, Menezes, John, & Grobusch, 2008). This development, combined with the increasing frequency of multidrug resistance, has made A. baumannii an extremely problematic pathogen for clinicians to treat and mortality rates for these infections has approached 70% (Lee, Chen, Wu, Huang, & Chiu, 2014). It has become widely recognized that new therapies are needed to help combat these infections (Gootz & Marra, 2008; Hujer et al., 2006; Joly-Guillou, 2005; Scott et al., 2007).

Members of the resistance–nodulation–cell division (RND) class of efflux systems in Gram-negative bacteria are composed of three proteins: an inner membrane transporter, an outer membrane protein that serves as a pore, and a periplasmic adapter protein that interacts with both the inner and outer membrane proteins to form a conduit for the extrusion of small molecules. RND-type systems typically capture toxic compounds or metabolites and remove them from the cell, and because...
of this function they can be involved in resistance to antibiotics, disinfectants, and heavy metals (Alvarez-Ortega, Olives, & Martinez, 2013; Anes, McCusker, Fanning, & Martins, 2015; Delmar, Su, & Yu, 2014; Magnet, Courvalin, & Lambert, 2001; Routh et al., 2011; Venter, Mowla, Ohene-Agyei, & Ma, 2015). They have also been shown to have roles in virulence, resistance to host antimicrobial peptides, and in cellular homeostasis by removing excess metabolites (Helling et al., 2014; Warner, Folley, Shafer, & Jerse, 2007).

2.1 | Bacterial strains, plasmids, and growth conditions

Both A. baumannii and Escherichia coli were grown in modified Luria Broth containing 10 g tryptone, 5 g yeast extract, and 5 g NaCl per liter. For screening opaque and translucent colonies of AB5075, LB was prepared at 0.5× of the normal concentration with 8 g agar per liter. Escherichia coli transformants were selected with chloramphenicol (25 μg/ml), ampicillin (200 μg/ml), or kanamycin (20 μg/ml) when appropriate. Acinetobacter baumannii AB5075 transformants were selected with tetracycline (3 μg/ml). Plasmid pEX18Tc was used for allelic replacement in AB5075 (Hoang, Karkhoff-Schweizer, Kutcha, & Schweizer, 1998). For all experiments involving opaque and translucent variants, cells were obtained from freezer stocks that were grown to low density and contained ~0.5% of the opposite cell type. Strain AB00075 arpB::Tc was obtained from the A. baumannii transposon insertion library maintained by Dr. Colin Manoil’s laboratory at the University of Washington (Gallagher et al., 2015).

2.2 | Phase variation assays

Freezer stocks of each strain that were verified as 99.99% of a single colony type were grown in LB broth to an OD of 1.8–1.9. Colonies were resuspended in LB broth and serial dilutions were plated on 0.5× LB, 0.8% agar plates. After overnight growth, the total number of visible cells per ml was determined and the number of opaque or translucent colonies present was determined on plates with at least 200 colonies/plate by using oblique light to illuminate the colonies.

2.3 | Transposon mutagenesis

A culture of the A. baumannii AB5075 opaque variant was grown in 25 ml of LB broth at 37°C with vigorous shaking. Cells were harvested from cultures at an OD~0.5 by centrifugation at 4°C for 10 min. Pellets were washed twice with 10% glycerol to prepare electrocompetent cells. Transposon (0.1 pmole) and transposase (1 unit) components of an EZ-Tn5 <TET-1> kit (Epigen Biotechnologies) were combined with 10% glycerol and incubated at 37°C for approximately 45 min. Transposome complex mixture was then cooled on ice prior to electroporation. Aliquots of 1.25 μl transposome mixture and 1 μl of TypeOne Restriction Inhibitor (Epigen Biotechnologies) were electroporated into 60 μl of competent AB5075 O cells at 2.5 kV. Following electroporation, 1 ml of room temperature LB broth was added following electroporation. This cell suspension was transferred to tubes and incubated at 37°C with shaking for 1 hr. Mutagenized cells were plated in 100 μl aliquots on 0.5× LB, 0.8% agar plates supplemented with 3 μg/ml tetracycline to select for insertional mutants and plates were incubated at 37°C. Colonies were evaluated at 24 and 48 hr postplating for altered colony morphology or reduced colony sectoring when viewed under a dissecting scope with oblique illumination from below. Putative mutants were restreaked on 0.5× LB, 0.8% agar with Tet to ensure colony morphology was stable and that picked colonies were pure. Identification of insertion sites was accomplished by rescue cloning of the tetracycline resistance gene and DNA sequencing of plasmids generated from rescue cloning.

2.4 | Construction of in-frame deletions in arpA and arpR

An in-frame deletion in arpA was generated as described previously (Hoang et al., 1998) from AB5075 genomic DNA by PCR amplification of two approximately 1 kb fragments containing small portions of the arpA coding region using the primers arpA Up-1 (5′-AAAAGGATCCATACTACGGTACCATAC-3′), arpA...
Up-2 (5′-GTGAAAATTCAGGGAGCCA-3′), arpA Down-1 (5′-TTGTGTCGTAAGGTGTTG-3′), and arpA Down-2 (5′-AAAAAGGATCCCACTTGAATTACCT-3′). After digestion with BamH I restriction sites at the 5′ end and gel purification, digestion products were ligated with the shuttle vector pWH1266 (Hunger, Schmucker, Kishan, & Hillen, 1990) which had been digested with ScaI and treated with recombinant shrimp alkaline phosphatase (New England Biolabs). 

The arpR in-frame deletion allele was generated from AB5075 genomic DNA by PCR amplification (Phusion Hot-Start Polymerase, Thermo Scientific) of two approximately 1-kbp fragments containing small portions of the arpR coding region. Oligonucleotide primers arpR Up-1 (5′-AAAAAGGATCCCGTGATAACCACAATACTTC-3′) and arpR Down-2 (5′-AAAAAGGATCCATGACATTAGTTTGAGTCGA-3′) were engineered to contain BamH I restriction sites at the 5′ end and were paired with arpR Up-2 (5′-CTCAAATATCGGCATTAAACC-3′) and arpR Down-1 (5′-ATTAACTGTTTGCACGAAAC-3′), respectively. The up- and downstream fragments were ligated (Fast-Link DNA Ligation Kit, Epicentre Biotechnologies) together to produce an approximately 2-kbp fragment containing the ΔarpR allele which was subsequently gel purified. The ΔarpR allele contains an in-frame deletion in the arpR coding sequence corresponding to amino acids 48–359 of the 366 (85% of coding region). Purified fragment was reamplified with outer primers (arpR Up-1 and arpR Down-2) and gel purified. The ΔarpR fragment and pEX18Tc were digested with BamH I and gel purified. Digested fragments were then ligated and transformed into competent E. coli DH5α cells. To transfer the mutant alleles to the chromosome of A. baumannii AB5075, the suicide vector containing the in-frame arpR deletion was electroporated into competent AB5075 cells which had been grown overnight in LB and washed with 300 mM sucrose as described previously (Choi & Schweizer, 2006). Integrants were selected on LB + tetracycline at 5 μg/ml. Counterselection to select for excision of the integrated plasmid was carried out at room temperature on LB without NaCl supplemented with 10% sucrose. Potential mutants were screened by PCR and confirmed by DNA sequencing.

To prepare conditioned media, cells were grown in 25 ml LB cultures at an optical density of 1.7. At this time, aliquots were restreaked to verify that the cultures remained at least 95% opaque or translucent. Cells were removed by centrifugation and the resulting media was adjusted to pH 7.0 with HCl and filter sterilized by passing through a 0.22-μm filter. Aliquots were frozen at −80°C and used within 2 weeks. To grow cells in conditioned media, a 2-ml aliquot was thawed and a 10× concentrate of tryptone and yeast extract (TY) was added back to a final concentration of 0.25×. Cells were grown by shaking at 270 rpm at 37°C to an optical density of 1.1 and dilutions were then plated to determine phase transition frequencies. Growth of cells in the LB control was done at 0.25× LB, a concentration that gave a similar growth rate as the TY-supplemented conditioned media.
produced the expression vector parpR. The vector was gel purified and ligated (Fast-Link Ligation Kit, Epicentre Biotechnologies) into pWH1266 that had been digested with Scal and treated with recombinant shrimp alkaline phosphatase (rSAP, New England Biolabs). The ligation was electroporated into competent E. coli EC100D Transformax cells (Epicentre Biotechnologies). Plasmids that conferred tetracycline resistance but not ampicillin resistance were purified and confirmed by DNA sequencing prior to introduction into A. baumannii. This produced the expression vector parpR.

### 2.9 Switching frequency determination

Cultures were grown to the OD\(_{600}\) ~0.7 or ~1.7 in LB medium, serially diluted in LB, and plated on 0.5\(\times\) LB, 0.8% agar. Plates were incubated at 37°C overnight. CFU per milliliter and number of variant colonies were determined for each strain and variant.

### 2.10 RNA isolation

Cultures of A. baumannii strain ABS075 and the ΔarpR mutant were grown in LB medium at 37°C with shaking to an OD\(_{600}\) ~0.7. Cultures of A. baumannii strain ABS075 harboring pWH1266 or parpR/WH1266 were grown in LB medium supplemented with tetracycline (5 μg/ml) at 37°C with shaking to an OD\(_{600}\) ~0.5. Cells were harvested from cultures by centrifugation and RNA was isolated with the MasterPure RNA Purification Kit according to the manufacturer’s protocol (Epicentre). Contaminating DNA was removed by treatment with TURBO DNA-free according to the manufacturer’s protocol (Ambion). DNA contamination was evaluated by PCR with purified RNA as template and RNA concentration was quantified with a NanoDrop ND-1000 spectrophotometer.

### 2.11 Quantitative real-time PCR

Total RNA (1 μg) purified from strains ABS075 with or without plasmid and ΔarpR were converted into cDNA by using the iScript cDNA synthesis kit (Bio-Rad) with random primers and SuperScript III reverse transcriptase (Invitrogen). Cycling parameters for cDNA synthesis were as follows: 25°C for 5 min, 42°C for 45 min, and 85°C for 5 min. cDNA reactions were then diluted 1:10 with sterile H\(_2\)O supplemented with 10 μg/ml yeast t-RNA (Roche). Diluted cDNA was used as template for experimental reactions. Oligonucleotide primer pairs for quantitative real-time PCR (qRT-PCR) were generated by the Primer-BLAST program available at www.ncbi.nlm.nih.gov/tools/primer-blast/. Primers were designed to amplify a 145-bp fragment of clpX (cpx qRT For 5’-GGCTTGAAGTGGGCAAT-3’) and a 151-bp fragment of arpA (arpA qRT Rev 5’-CCATTCCAGGCCCACTCT-3’) and a 151-bp fragment of arpB (arpB qRT 5’-GCCGAAGCCTATCAACTG-3’). qRT-PCR was performed using IQ SYBR Green Supermix (Bio-Rad) with Bio-Rad CFX Connect cycler. The following cycle parameters were utilized to amplify and quantify fragments: 95°C for 3 min and then 95°C for 10 s, 55°C for 10 s, and 72°C for 20 s, repeated 40 times. Melt curve data were collected to ensure proper amplification of target genes. Data were generated from three separate RNA isolation and cDNA preparations and at least two technical replicates for each primer set. Relative expression of each gene was determined by comparing target gene expression with control gene (clpX) expression using the Pfaffl method (Pfaffl, 2001).

### 2.12 Antimicrobial resistance assays

Strains to be tested were grown to an OD\(_{600}\) of 0.3 and then diluted to an OD\(_{600}\) of 0.05 with sterile LB. The minimum inhibitory concentration for various antibiotics was determined using E-test strips on LB agar plates according to the manufacturer’s instructions (Biomerieux). The MICs for each antibiotic were determined after 12 hr of growth at 37°C. All susceptibility tests were done in duplicate.

### 3 RESULTS

#### 3.1 An RND-type efflux system is required for the opaque to translucent colony opacity phase variation

Previously, our laboratory reported on a phase variable mechanism in A. baumannii that results in the interconversion between opaque and translucent colonies (Tipton et al., 2015). This phase variation was stimulated at high density and when opaque colonies are grown for 36–48 hr, they become highly mottled in appearance due to translucent variants arising at high frequency within the colony (Figure 1a). This mottled appearance formed the basis for a genetic screen to identify mutants with a reduced frequency of phase variation in the opaque to translucent direction, as these mutants would have a reduction or absence in the mottled appearance of the colony. A library of EZ-Tn5 <Tet-1> insertions in ABS075 was screened for colonies that did not exhibit the mottled appearance after 36 hr of growth and mutants with this phenotype were isolated and confirmed by subsequent replating. One mutant 5075.8B (Figure 1a) was characterized further and the transposon insertion was mapped to nucleotide 64 of a 1,101 nucleotide open reading frame (ABUW_0034) encoding a putative periplasmic membrane fusion component of RND-type efflux systems (Figure 1b). This protein exhibited the highest degree of identity to annotated RND-type transporters from Pseudomonas mendocina (47% identity/69% similarity) and Pseudomonas pseudoalcaligenes (47% identity/68% similarity). A second open reading frame (ABUW_0035) was present beginning 3 bp downstream of ABUW_0034 that encoded a putative inner membrane transporter component of RND-type systems. This protein exhibited the highest identity to an annotated RND-type transporter from Bacillus mycoides (85% identity/93% similarity) and to Alkanindiges illinoisensis (75% identity/86% similarity). Given their close proximity, these two genes likely formed an operon and were previously identified in A. baumannii as a locus upregulated in the presence of farnesol and were
designated as acrAB (Kostoulas et al., 2016). However, the overall similarity of these proteins to the *E. coli* AcrA (23% identity/39 similarity) and AcrB (25% identity/43% similarity) was low and there are other RND-type systems in *E. coli* that exhibit greater identity to the *A. baumannii* proteins. In addition, as outlined below, this RND system confers aminoglycoside resistance, therefore, these genes were designated *arpA* and *arpB* (aminoglycoside resistance pump). The frequency of phase variation from opaque to translucent was quantitated in individual colonies of the 5075.8B mutant and wild-type cells after 48 hr of growth, which revealed an average 55-fold decrease in the opaque to translucent switch in the *arpA::EZ-Tn5 Tet-1* mutant 5075.8B.

While this analysis was being conducted, it was observed that there was significant colony to colony variation within the same strain with respect to phase variation frequencies, leading to large standard deviations. As a result of this, broth grown cells were used for subsequent experiments, which did reduce variability, although significant variability still existed. In broth cultures, the phase variation rate of the 5075.B mutant was 77-fold lower than wild type (Table 1).

To verify the role of the *arpA* gene in phase variation and to avoid polar effects on the downstream *arpB* gene that likely occurred from the *arpA::EZ-Tn5 Tet-1* insertion, an in-frame deletion in *arpA* (Δ*arpA2*) was constructed. This mutation had a similar effect on phase variation as the original *arpA::EZTN<Tc>* mutant, where the frequency of switching from opaque to translucent was reduced 71-fold in cultures at an OD<sub>600</sub> = 1.9 (Table 1). This Δ*arpA2* mutant strain was designated KT2.

Complementation analysis was performed to determine if the loss of *arpA* was responsible for the phase variation defect. When a plasmid containing the *arpA* gene was introduced into the *arpA* deletion mutant (KT2), colonies regained the ability to form the mottled phenotype due to translucent variants arising at high frequency (data not shown). In addition, the frequency of phase variation in 24 hr colonies was increased by 3,600-fold in the *arpA* deletion mutant containing the cloned *arpA* gene (parpA), relative to the *arpA* deletion mutant containing the pWH1266 vector alone (Figure S1, panel A).

**FIGURE 1** Decreased phase variation in an *arpA* mutant. (a) A typical wild-type opaque colony variant of AB5075 is shown compared to 5075.8B *arpA::EZ-Tn5 Tet-1* after 36 hr of growth on a 0.5× LB, 0.8% agar plate. The mottled appearance at the outside edge of the wild type is due to translucent variants arising within the opaque colony. (b) The organization of the *arpAB* region and the site of the *arpA::EZ-Tn5 Tet-1* insertion that blocks phase variation is depicted by an arrow. Proteins exhibiting the closest match to ArpA and ArpB are shown below each gene.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relative O to T phase variation frequency&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>AB5075 wild type</td>
<td>1&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>5075.8B <em>arpA::EZTN5Tc</em></td>
<td>0.013 ± 0.02 (77-fold decrease)</td>
</tr>
<tr>
<td>KT2 Δ<em>arpA2</em></td>
<td>0.014 ± 0.03 (71-fold decrease)</td>
</tr>
<tr>
<td>AB00075 <em>arpB::Tc</em></td>
<td>0.0013 ± 0.001 (769-fold decrease)</td>
</tr>
<tr>
<td>KT3 Δ<em>arpR</em></td>
<td>0.0009 ± 0.0002 (1,916-fold decrease)</td>
</tr>
</tbody>
</table>

<sup>a</sup>Determined in cultures at an OD<sub>600</sub> of 1.85–1.9.

<sup>b</sup>Phase variation frequency from O to T was typically 25%–35%.

<sup>c</sup>Phase variation frequency from T to O was typically 40%–50%.

**TABLE 1** Phase variation frequencies
3.2 | An arpB mutation also decreases the opaque to translucent phase variation

The gene located immediately downstream from arpA is predicted to encode a putative inner membrane transporter of RND efflux systems and likely functions with ArpA in this process. This gene was designated arpB. To determine if arpB also regulates phase variation, two separate transposon insertions in arpB (AB00075, AB00076) were obtained from the A. baumannii AB5075 transposon insertion library at the University of Washington (Gallagher et al., 2015). Like the arpA mutant, opaque colonies of the arpB mutants did not produce mottled colonies at 48 hr. The frequency of phase variation for AB00075 arpB::Tc in broth grown cells at an OD$_{600}$ of 1.9 was reduced an average of 769-fold compared to wild-type cells (Table 1).

3.3 | arpA and arpB mutations do not alter the translucent to opaque switch

The effect of the ΔarpA mutation on the reciprocal opacity switch, from translucent to opaque, was measured in KT2 grown to an OD of 1.9, where the rate of phase variation was 1.5-fold over wild type (Table 1). The arpB::Tc mutant AB00075 exhibited a phase variation rate for translucent to opaque that was 1.2-fold greater than wild type.

3.4 | An arpB mutant still produces an extracellular signal that stimulates phase variation

Previous work demonstrated that the increase in phase variation at high cell density is mediated, in part, by the accumulation of a secreted signal (Tipton et al., 2015). If this signal is dependent on ArpAB for secretion and sensed at the cell surface or the cytoplasmic membrane, then this could explain the reduction in phase variation in the arpA mutant. To test this possibility, conditioned media were prepared from opaque variants of wild-type and the ΔarpB mutant at an optical density of 1.7 and tested for the ability to stimulate phase variation in the opaque to translucent direction. Conditioned media from wild-type cells stimulated the opaque to translucent conversion 11-fold and the conditioned media from the arpB mutant stimulated phase variation ninefold (Table 2). There was not a statistically significant difference between these values ($p = .39$) indicating that the arpB mutant produced a similar level of extracellular signal activity as the wild-type parent.

3.5 | Role of ArpAB in antimicrobial resistance

It is well established that RND-type systems can export antimicrobials and confer higher levels of resistance (Alvarez-Ortega et al., 2013; Anes et al., 2015; Coyne et al., 2011; Magnet et al., 2001; Rosenfeld et al., 2012; Routh et al., 2011; Venter et al., 2015). Therefore, both the arpA and arpB mutants were tested for levels of resistance to various antibiotics. For this analysis, both the opaque and translucent variants of each mutant and wild-type cells were tested (Table 3). Due to the multiple resistances that already exist in AB5075, the number of antibiotics that could be tested was limited. There were no significant differences between wild-type and the arpA mutant for any of the antibiotics tested (Table 3), although there were subtle differences between opaque and translucent variants of wild type as reported previously (Tipton et al., 2015). However, the arpB mutant AB00075 was more sensitive to the two aminoglycosides that were tested, amikacin and tobramycin (Table 3). This sensitivity was seen in both the opaque and translucent variants of AB00075 arpB::Tc.

3.6 | arpA and arpB mutations selectively increase surface motility in the opaque variants

Both the arpA and arpB mutants formed irregular, slightly spreading colonies on 0.8% agar plates that were distinct from wild-type colonies that contained smooth rounded edges. This observation suggested that motility may be enhanced by the arpA and arpB mutations. To test this hypothesis, the motility of both opaque and translucent arpA and arpB mutants were compared to opaque and translucent wild-type AB5075 on 0.3% Eiken agar plates. In opaque colonies, both the arpA and arpB mutants exhibited increased motility, 53 mm and 75 mm, respectively, over wild-type value of 33 mm (Figure 2). However, in the translucent form, there was no significant difference in motility between wild-type and either the arpA or arpB mutants (Figure 2). The increased motility observed in the arpA mutant was restored back to wild-type levels by parpA containing the cloned arpA gene relative to cells containing the pWH1266 vector alone (Figure S1, panel B).

**Table 2** Effect of conditioned media on relative phase variation frequency

<table>
<thead>
<tr>
<th>Growth condition</th>
<th>Opaque to translucent switching frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>LB broth</td>
<td>0.35% ± 0.29</td>
</tr>
<tr>
<td>Wild-type opaque conditioned media</td>
<td>3.9% ± 2.8 (11-fold increase)</td>
</tr>
<tr>
<td>ΔarpB opaque conditioned media</td>
<td>3.2% ± 2.7 (9-fold increase)</td>
</tr>
</tbody>
</table>

*a Determined in cells grown to an optical density $A_{600}$ of 1.0.

**Table 3** Minimum inhibitory concentrations for antibiotics

<table>
<thead>
<tr>
<th>Strain</th>
<th>COL</th>
<th>TET</th>
<th>AK</th>
<th>RIF</th>
<th>TIG</th>
<th>TOB</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT opaque</td>
<td>0.38</td>
<td>3</td>
<td>128</td>
<td>3</td>
<td>3</td>
<td>96</td>
</tr>
<tr>
<td>ΔarpA opaque</td>
<td>0.38</td>
<td>3</td>
<td>128</td>
<td>3</td>
<td>3</td>
<td>96</td>
</tr>
<tr>
<td>arpB::Tc opaque</td>
<td>0.38</td>
<td>ND</td>
<td>64</td>
<td>3</td>
<td>3</td>
<td>24</td>
</tr>
<tr>
<td>WT translucent</td>
<td>0.38</td>
<td>3</td>
<td>64</td>
<td>4</td>
<td>3</td>
<td>24</td>
</tr>
<tr>
<td>ΔarpA translucent</td>
<td>0.38</td>
<td>3</td>
<td>64</td>
<td>4</td>
<td>3</td>
<td>24</td>
</tr>
<tr>
<td>arpB::Tc translucent</td>
<td>0.38</td>
<td>ND</td>
<td>16</td>
<td>4</td>
<td>3</td>
<td>8</td>
</tr>
</tbody>
</table>

COL, colistin; TET, tetracycline; AK, amikacin; RIF, rifampicin; TIG, tigecycline; TOB, tobramycin.
3.7 | ArpAB is required for virulence in *Galleria mellonella*

RND-type efflux systems have been shown to have roles in virulence in a number of bacterial pathogens (Alvarez-Ortega et al., 2013; Delmar et al., 2014; Routh et al., 2011; Taylor, Bina, & Bina, 2012; Venter et al., 2015; Warner et al., 2007; Yoon et al., 2015). Therefore, the impact of *arpA* and *arpB* mutations on virulence were examined in a *Galleria mellonella* waxworm model that has previously been shown to be a useful model of virulence in *A. baumannii* (Esterly et al., 2014; Gaddy et al., 2012; Heindorf, Kadari, Heider, Skiebe, & Wilharm, 2014; Iwashkiw et al., 2012; Jacobs et al., 2014; Nwugo et al., 2012; Peleg et al., 2009; Repizo et al., 2015; Stahl, Bergmann, Gottig, Ebersberger, & Averhoff, 2015; Tipton et al., 2015). Since previous work indicated that the opaque form was more virulent, these studies were done with the opaque forms of wild-type AB5075, Δ*arpA* and the *arpB::Tc* mutant (Tipton et al., 2015). Relative to wild-type AB5075, both the Δ*arpA* and *arpB* mutations substantially reduced the ability of *A. baumannii* to kill *G. mellonella* waxworms (Figure 3a). The decreased virulence exhibited by the *arpA* mutant was restored by parpA containing the cloned *arpA* gene relative to cells containing the pWH1266 vector alone (Figure S1, panel C).

3.8 | A divergently transcribed gene encoding a TetR-type regulator represses *arpAB* and is required for the opaque to translucent switch

Adjacent to the *arpAB* genes in *A. baumannii* was a divergently transcribed gene encoding a predicted TetR-type repressor of the AcrR family (Figure 1). In *E. coli*, the AcrR protein acts to repress the *acrAB* operon, which is organized in a similar manner as *arpAB*. To determine if a similar function was present in *A. baumannii*, an in-frame deletion was constructed in this gene, designated *arpR*. The Δ*arpR* deletion did not have a significant impact on *arpAB* expression, with the levels of expression 1.1-fold higher in the mutant (Figure 4). However, the *arpR* mutation reduced the frequency of the opaque to translucent conversion by 1,916-fold (Table 1).

As a second method to determine if ArpR could function as a repressor, the *arpR* gene was overexpressed by cloning the gene into pWH1266, where expression was driven from the promoter for the β-lactamase gene. When this plasmid was introduced into wild-type AB5075, the colony morphology was altered with irregular slightly spreading colonies that were similar to *arpA* and *arpB* mutants,
suggested that expression of both arpA and arpB were reduced when ArpR was overexpressed. To investigate this possibility, expression of the first gene arpA was examined by qRT-PCR and was found to be 7.3-fold lower in cells with parpR versus the pWH1266 vector alone (Figure 4). Next, we investigated if the reduced ArpAB expression resulting from ArpR overexpression impacted the opaque to translucent switch. In cells overexpressing the arpR gene, the frequency of phase variation was reduced threefold, relative to cells containing the vector only (data not shown).

4 | DISCUSSION

In this study, a previously undescribed RND-type efflux system in A. baumannii was found to regulate phase variation in the opaque to translucent direction. The original mutation defining this phenotype was a transposon insertion in arpA, a gene encoding a membrane fusion component of a putative RND-type efflux system. This arpA mutation resulted in a 77-fold decrease in the frequency of phase variation from the opaque to translucent colony opacity phenotype (Table 1). The role of arpA in this process was confirmed by the construction of a nonpolar, in-frame deletion within arpA, which decreased the conversion to translucent colonies by 71-fold, and by complementation analysis with the wild-type arpA gene. In addition, a transposon insertion in the downstream arpB gene obtained from the A. baumannii AB5075 transposon library at the University of Washington (Gallagher et al., 2015) exhibited a stronger effect with a 769-fold reduction in phase variation (Table 1). Interestingly, neither the arpA or arpB mutations had a significant effect on phase variation in the translucent to opaque direction. This provides strong evidence that a distinct mechanism regulates phase variation in each direction.

Two additional phenotypes were altered by the loss of arpA and arpB. First, surface motility of the opaque variants on 0.3% agar plates was increased, with the arpB mutant exhibiting a more substantial increase than the arpA mutant (Figure 2). Interestingly, in translucent variants, the motility of both the arpA and arpB mutants was similar to wild-type. The basis for the selective motility increase in opaque variants is unknown, but previously it was noted that in wild-type cells, the opaque variants were more motile (Tipton et al., 2015). Although the actual mechanism responsible for motility in AB5075 has not been identified, if this mechanism was only expressed in opaque cells and also repressed by ArpAB, it could explain the selective increase in motility seen in arpA or arpB mutants in the opaque background. Second, the arpA mutation decreased the ability of A. baumannii AB5075 to kill G. mellonella waxworms. Additional examples exist where the loss of an RND-type system decreases virulence (Alvarez-Ortega et al., 2013; Delmar et al., 2014; Routh et al., 2011; Taylor et al., 2012; Venter et al., 2015; Warner et al., 2007; Yoon et al., 2015). The decreased virulence may be due to a role for ArpAB in efflux of host antimicrobial peptides or other antibacterial compounds present inside G. mellonella.

The arpB mutant consistently exhibited stronger effects than the arpA mutant for phase variation (Table 1), surface motility (Figure 2), and antibiotic sensitivity (Table 3). It is unlikely that this results from the arpA mutation being leaky, as the in-frame deletion removed most of the coding region. The weaker phenotype of the arpA mutation may indicate that a periplasmic fusion protein from another RND-type system can partially substitute for ArpA. The stronger effect of a arpB mutation suggests that ArpB is unable to be substituted for by another RND component and has a more critical role than ArpA. However, this hypothesis is speculative at the current time. In addition, the outer membrane channel that works with ArpAB is unknown, but AbuO, a TolC-like protein is a possible candidate (Srinivasan, Vaidyanathan, & Rajamohan, 2015).

A TetR-type transcriptional regulator was encoded adjacent to the arpAB genes and divergently transcribed, a genetic organization similar to that of acrR-acrAB in E. coli and mexR-mexAB in P. aeruginosa. A ΔarpR mutation did not alter arpAB expression, however, overexpressing the arpR gene decreased arpAB expression by 7.3-fold. In addition, cells overexpressing arpR exhibited phenotypes that at least partially mimicked the arpAB mutations, as the colonies formed the irregular spreading appearance and the frequency of the opaque to translucent conversion was reduced threefold. It is unclear why the effects of ArpR on arpAB expression are seen only when it is overexpressed, but under the laboratory growth conditions we tested, it does not have a role in arpAB regulation, possibly because it is not expressed at high enough levels to mediate repression. If certain environmental conditions increased arpR expression, this condition would be predicted to repress arpAB and reduce the opaque to translucent conversion.

To our knowledge, this represents the first report of an RND-type efflux system regulating bacterial phase variation. A direct role for ArpAB in this process is unlikely and loss of ArpAB may indirectly modulate a regulatory pathway controlling phase variation. At the present time, the regulatory mechanism regulating the interconversion between opaque and translucent colonies is unknown. However, whole genome sequencing suggests that nucleotide changes are not responsible. This information, together with the observation that phase variation between colony opacity phenotypes is essentially undetectable in cells at low density, but is sharply activated at high density and reaches frequencies above 10% (Table 1) suggests a nonmutational mechanism is involved.

**FIGURE 4** Effect of ArpR on arpA expression. The levels of arpA expression were determined by quantitative RT-PCR and the values shown are relative to the control gene clpX. The values for the left side represent the levels of arpA expression in the arpR mutant relative to wild type. The values on the right side represent the levels of arpA expression in cells overexpressing arpR from a plasmid relative to cells containing the vector alone.
A possible mechanism to control phase variation is via a bistable switch involving one or more regulatory proteins (Casadesus & Low, 2013; Chang et al., 2010; Dubnau & Losick, 2006; Ferrell, 2002; Maamar & Dubnau, 2005; Mitropahanov & Groisman, 2008; Piggot, 2010; Turner, Vallet-Gely, & Dove, 2009; Veening, Smits, & Kuipers, 2008). Bistability has previously been shown to regulate colony opacity in Pseudomonas fluorescens (Gallie et al., 2015). Based on this information, there are several possibilities for how ArpAB might regulate phase variation. First, the extracellular signal that stimulates the opaque to translucent phase variation might require ArpAB for secretion. This would imply that the signal works via a sensing mechanism that operates at the cell surface or at the cytoplasmic membrane and then regulates the bistable switch controlling the phase variation. However, spent culture supernatants from an arpB mutant and wild-type cells appear to activate the opaque to translucent switch at equal frequencies (Table 2). A second possibility is that a metabolite normally secreted by the ArpAB system accumulates in arpB mutants and this alters a regulatory pathway that controls phase variation. The isolation of both extragenic and high-copy suppressors that restore phase variation to an arpA or arpB mutant should help uncover the role of this efflux system in regulating the process of colony opacity phase variation in A. baumannii and these studies are in progress.

ACKNOWLEDGMENTS

This work was funded by grant 1R21AI115183 from the National Institutes of Health. P. N. R. is also supported by grants from the Merit Review program and a Research Career Scientist award, both from the Department of Veterans Affairs. We are grateful to Dr. Colin Manoil for providing mutants from the University of Washington A. baumannii transposon library. In addition, we thank Chui-Yoke Chin for help with the Kaplan–Meier plots.

CONFLICT OF INTEREST

None declared.

REFERENCES

calcoaceticus and its use for Escherichia coli shuttle plasmids. Gene, 87, 45–51.


Jacobs, A. C., Thompson, M.G., Black, C. C., Kessler, J. L., Clark, L. P., McQueary, C. N., … Zurawski, D. V. (2014). AB5075, a highly virulent isolate of Acinetobacter baumannii, as a model strain for the evaluation of pathogenesis and antimicrobial treatments. MBio, 5, e01076–01014.


Yoon, E. J., Chabane, Y. N., Goussard, S., Snesrud, E., Courvalin, P., De, E., & Veening, J. W., Smits, W. K., & Kuipers, O. P. (2008). Additional Supporting Information may be found online in the supporting information tab for this article.

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.