# **BC** ARTICLE



# The structure of PilA from *Acinetobacter baumannii* AB5075 suggests a mechanism for functional specialization in *Acinetobacter* type IV pili

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Type IV pili (T4P) are bacterial appendages composed of protein subunits, called pilins, noncovalently assembled into helical fibers. T4P are essential, in many bacterial species, for processes as diverse as twitching motility, natural competence, biofilm or microcolony formation, and host cell adhesion. The genes encoding type IV pili are found universally in the Gram-negative, aerobic, nonflagellated, and pathogenic coccobacillus Acinetobacter baumannii, but there is considerable variation in PilA, the major protein subunit, both in amino acid sequence and in glycosylation patterns. Here we report the X-ray crystal structure of PilA from AB5075, a recently characterized, highly virulent isolate, at 1.9 Å resolution and compare it to homologues from A. baumannii strains ACICU and BIDMC57, which are C-terminally glycosylated. These structural comparisons revealed that  $m PilA^{AB5075}$  exhibits a distinctly electronegative surface chemistry. To understand the functional consequences of this change in surface electrostatics, we complemented a  $\Delta pilA$ knockout strain with divergent pilA genes from ACICU, BIDMC57, and AB5075. The resulting transgenic strains showed differential twitching motility and biofilm formation while maintaining the ability to adhere to epithelial cells. PilA<sup>AB5075</sup> and PilA<sup>ACICU</sup>, although structurally similar, promote different characteristics, favoring twitching motility and biofilm formation, respectively. These results support a model in which differences in pilus electrostatics affect the equilibrium of microcolony formation, which in turn alters the balance between motility and biofilm formation in Acinetobacter.

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This work is dedicated to our colleague and friend, Mark Shirtliff, who provided technical input on assaying biofilm formation early on in this project. We mourn his loss.

This article contains Table S1 and Figs. S1–S9.

The atomic coordinates and structure factors (code 5VAW) have been deposited in the Protein Data Bank (http://wwpdb.org/).

Type IV pili (T4P)<sup>2</sup> are bacterial appendages composed of protein subunits, called pilins, noncovalently assembled into helical fibers. These appendages are found in a wide range of eubacteria and are related structurally to type II secretion pseudopili, competence-induced pili, and archeal archaella (1-3). In many bacteria, these organelles are essential for processes as diverse as twitching motility, natural competence, biofilm or microcolony formation, and adhesion to biotic and abiotic surfaces. These systems are phylogenetically diverse and are commonly divided into type IVa (widely distributed), type IVb (primarily found in enteric bacteria), and, proposed recently, type IVc (tight adherence pili) (4). All type IV pilus systems contain genes encoding for a cytoplasmic AAA+ ATPase (PilB), an integral membrane protein (PilC), and at least two pilins (5-7). One pilin, the major pilin, makes up nearly the entirety ( $\sim$ 99%) of the pilus, with the other subunits, minor pilins, being incorporated either at the tip (8) or scattered along the length (9).

Type IV pilin gene products are easily identified by their combination of an N-terminal signal peptide (which is removed by a specific protease prior to their incorporation into the pilus fiber) followed by a hydrophobic  $\alpha$ -helix (serving as a transmembrane domain for pilins in the inner/plasma membrane) and finally a soluble region commonly referred to as the pilin headgroup. The vast majority of known pilin headgroup structures are similar; a single globular domain consisting of an N-terminal  $\alpha$ -helix with a C-terminal  $\beta$ -sheet packed against it (10). However, the amino acid sequences of these pilin headgroups are so diverse that even structurally similar proteins have insignificant ( $\sim$ 10–20%) sequence identities (11). Despite the variety of type IV pilin proteins, which include minor pilins with multidomain headgroups (8, 9), a given pilin gene within a given species is typically well-conserved with only the major pilin showing significant variation (12). Sequence diversity in the major pilin has typically been attributed to diversifying selection (13).

Acinetobacter baumannii, a Gram-negative, aerobic, nonflagellated coccobacillus, expresses type IV pili, which are essential for twitching motility and natural competence (14), and contribute to

<sup>&</sup>lt;sup>2</sup> The abbreviations used are: T4P, type IV pill; CLSM, confocal laser scanning microscopy.



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host cell adherence (15). T4P are ubiquitous within the Acintobacter genus, which contains environmental strains from soil and water, as well as commensal and pathogenic strains isolated from mammalian hosts, including humans (16-18). A. baumannii has recently gained notoriety as a source of hospital-acquired infections (19-21), particularly for military personal returning from the Middle East (hence the moniker "Iraqibacter") (22); however, several other species related to A. baumannii, often simply referred to as the "Ab group," are routinely isolated from nosocomial infections (23, 24). In particular the four related species of the Acinetobacter calcoaceticus-A. baumannii (Acb) complex (A. baumannii, A. calcoaceticus, Acinetobacter pitti, and Acinetobacter nosocomialis) are difficult to distinguish (25) and have all been found to be infectious in model systems (26).

The type IVa pili of *A. baumannii* and related species have diverse PilA proteins (both in amino acid sequence and O-linked glycosylation) that are unrelated to the overall taxonomy. In this paper, we compare the structure and function of pilin proteins from three strains of A. baumannii: (i) ACICU (also known as H34), an epidemic, multidrug-resistant strain belonging to the European clone II group that was isolated from cerebrospinal fluid in an outbreak in Rome in 2005 (27); (ii) BIDMC 57, a 2013 respiratory isolate from Beth Israel Deaconess Medical Center (Boston, MA) and sequenced at the Broad Institute (Cambridge, MA); and (iii) AB5075, which was isolated in 2008 from a osteomyelitis of the tibia by a group at the Walter Reed Army Institute of Research (Silver Spring, MD) (28) and found by those authors to be more virulent than other A. baumannii isolates. However, correlating pilus isotypes with phenotypic characteristics is complicated by the divergence between pilA genotype and Acinetobacter genetic diversity; by way of example, A. calcoaceticus PHEA-2 is an industrial wastewater isolate, but PilAPHEA2 is 94% identical to PilAACICU.

Previously, we solved X-ray crystal structures of PilA, the major pilin protein, from two strains of A. baumannii: ACICU and BIDMC57 (15). PilAACICU and PilABIDMC57 were much more similar to pilin structures from Pseudomonas aeruginosa and Dichelobacter nodosus than to each other, seemingly products of convergent evolution. We have proposed that this divergence within A. baumannii pilA genes could potentially be explained by specialization for aerobic or anaerobic environments caused by differences in the distribution of disulfide bonds; the solved structures of FimA, the D. nodosus major pilin, and PilABIDMC57 lack disulfide bonds at the C terminus typically found in pilins from Gram-negative bacteria.

However, a phylogenetic analysis of amino acid sequences from the major pilins of A. baumannii, P. aeruginosa, and D. nodosus showed three clusters, each containing sequences from multiple species (15). We chose to examine the structure of PilA from A. baumannii AB5075, a recently characterized and unusually virulent clinical isolate (28), both as a representative of this third taxonic group and because unlike PilA<sup>ACICU</sup> and PilA<sup>BIDMC57</sup>, PilA<sup>AB5075</sup> is not natively C-terminally glycosylated. We have solved the X-ray crystal structure of PilAAB5075 and, after comparing it to the other Acinetobacter PilA structures, found differences in surface electrostatics,

which suggest a mechanism for functional differentiation in T4P based on the structure of PilA. To test our hypothesis, we have complemented a pilA knockout strain of a model strain, A. nosocomialis M2, with the pilin genes from A. baumannii AB5075, ACICU, and BIDMC57 to directly compare the resulting phenotypes. Our results, described below, suggest that a functional trade-off may exist between the ability of Acinetobacter T4P to promote biofilm formation and to function in twitching motility.

### Results

# Acinetobacter pilA is highly variable and shows evidence of convergent evolution with type IV pilins from other species.

Type IV pili are found in a wide variety of bacteria and have been widely studied in Gram-negative infectious strains, particularly Pseudomonas and Neisseria species, which produce copious amounts of pili under laboratory conditions (46-51). We previously noted the similarity in structure between A. baumannii PilA proteins and the equivalent major pilin proteins in P. aeruginosa and D. nodosus. All three of these gammaproteobacteria species have been isolated from mammalian hosts, as well as soil, and typically present with persistent opportunistic infections rather than acute bacteremia (26, 52, 53). These phenotypic similarities suggest that functional similarities may also exist between the three T4P systems.

Fig. 1A shows a phylogenic tree of 60 A. baumannii PilA, P. aeruginosa PilA, and D. nodosus FimA amino acid sequences excluding the N-terminal signal peptides (all sequences begin FTLIEL...). The three branches of the unrooted tree each contain sequences from multiple species. On the top left, the branch containing PilAACICU also contains P. aeruginosa PilAPAO1 and PilAPAK among others. Counterclockwise, the next branch, containing PilABIDMC57, also includes D. nodosus FimA sequences from the predominant serotypes (A-C and E–G). The final branch contains representatives from all three species, including PilAAB5075, D. nodosus FimA from serotypes D and H, and P. aeruginosa PilA<sup>1244</sup>. We noted the existence of this third branch in the dendogram previously (15). However, unlike the ACICU and BIDMC57 division, we could find no ready explanation for the division between the ACICU and AB5075 branches.

Notably, of the seven A. baumannii pilin genes, pilA alone shows this divergence, suggesting that the "machinery" of pilus assembly is conserved throughout. Fig. 1B shows a dendogram of the seven pilins from A. baumannii AB5075, ACICU, and BIDMC57. The pilA branch (circled in red) shows substantially more variation than the other six, which are well-conserved, particularly between ACICU and AB5075; in one case, PilV, the ACICU and AB5075 amino acid sequence are identical.

# High-resolution structure of PilA<sup>AB5075</sup>

We determined the structure of PilA from A. baumannii AB5075 as a C-terminal fusion to maltose-binding protein to a resolution of 1.9 Å (Table 1). PilA AB5075 possesses a typical type IVa pilin fold (Fig. 2A), beginning with an  $\alpha$ -helix ( $\alpha$ 1-C, the N-terminal portion, residues 1-22, is hydrophobic and was removed for expression and crystallization), which leads into an



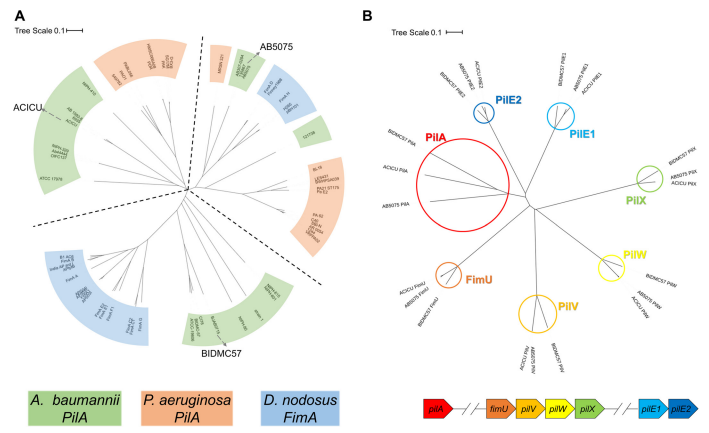


Figure 1. Phylogeny of A. baumannii type IV pilin genes. A, dendogram of major pilin amino acid sequences from 20 representative strains of A. baumannii (green), P. aeruginosa (orange), and D. nodosus (blue). The sequences of PilAABBOO75, PilAACICU, and PilABIDMC57 are marked on their respective branches (accession numbers in Table S1). B, dendogram of A. baumannii pilin amino acid sequences. Branches for each of the seven pilins are highlighted in red (PilA), orange (FimU), light orange (PilV), yellow (PilW), green (PilX), light blue (PilE1), and dark blue (PilE2); the genetic organization of the pilin genes is diagrammed below in the corresponding colors.

Table 1 Crystallographic parameters for MBP-PiIA<sup>AB5075</sup> The values in parentheses are for the highest resolution shell.

F		
	Resolution range	29.21-1.9 (1.968-1.9)
	Space group	P 1 21 1
	Unit cell	39.557, 103.04, 56.195, 90, 98.947, 90
	Total reflections	156,212 (15709)
	Unique reflections	34,906 (3467)
	Multiplicity	4.5 (4.5)
	Completeness (%)	99.64 (99.77)
	Mean $I/\sigma(I)$	14.63 (2.44)
	Wilson B-factor	28.44
	$R_{\text{merge}}$	0.05718 (0.5332)
	$R_{\text{pim}}$	0.02973 (0.28)
	R <sub>pim</sub> CC <sub>1/2</sub>	0.999 (0.849)
	CC*	1 (0.958)
	$R_{\text{work}}$	0.2002 (0.3110)
	$R_{\text{free}}$	0.2432 (0.3382)
	Root mean square	
	Bonds	0.005
	Angles	0.73
	Ramachandran (%)	
	Favored	97.70
	Allowed	2.30
	Outliers	0.00
	Clashscore	4.04
	Average B-factor	41.53
	Macromolecules	41.45
	Ligands	53.97
	Solvent	42.23

extended loop (the  $\alpha\beta$ -loop, see below) and then a central β-sheet packed against the helix. Like PilAACICU (and the majority of solved type IV pilin structures), one of its disulfide bonds is at the C terminus between the final two  $\beta$ -strands of

the central  $\beta$ -sheet, between residues 135 and 148. This similarity in disulfide bonding may explain why, despite the poor sequence conservation between the PilA<sup>ACICU</sup> and PilA<sup>AB5075</sup> C termini, their D-regions (the loops bound by the C-terminal disulfide bonds) are superimposable (Fig. 2C), unlike the equivalent C-terminal region of PilA BIDMC57 (Fig. 2D). This supports our hypothesis that the C-terminal structure of PilABIDMC57 and the structurally similar D. nodosus FimA (serotype A, PDB ID: 3SOK) (54) contain C-terminal helices and hydrophobic regions to stabilize them in the absence of the disulfide bond. Previously we showed that alanine mutations of those C-terminal hydrophobic residues destabilized PilA BIDMC57 (15). Fig. 2B shows a sequence alignment of PilAAB5075, PilAACICU, and PilA<sup>BIDMC57</sup>; despite the structural conservation in the central β-sheet, the sequence similarity is low:  $\sim$ 35% for any of the three to either of the other two, excluding the N-terminal transmembrane helix (residues 1-22).

The most striking feature of the PilAAB5075 structure is the  $\alpha\beta$ -loop (from the end of the central  $\alpha$ -helix to the start of the first  $\beta$ -strand in the central  $\beta$ -sheet); bounded by a *dashed gray line* in Fig. 2*A*. Approximately 15 residues longer than the  $\alpha\beta$ -loop of PilA ACICU, it contains a disulfide bond to the central  $\alpha$ -helix (residues 50 and 65) and twists back over itself twice, extending out from the center of the headgroup. It contains none of the  $\alpha$ -character found in the ACICU  $\alpha\beta$ -loop but has some positional overlap with the BIDMC57  $\alpha\beta$ -loop (Fig. 2)

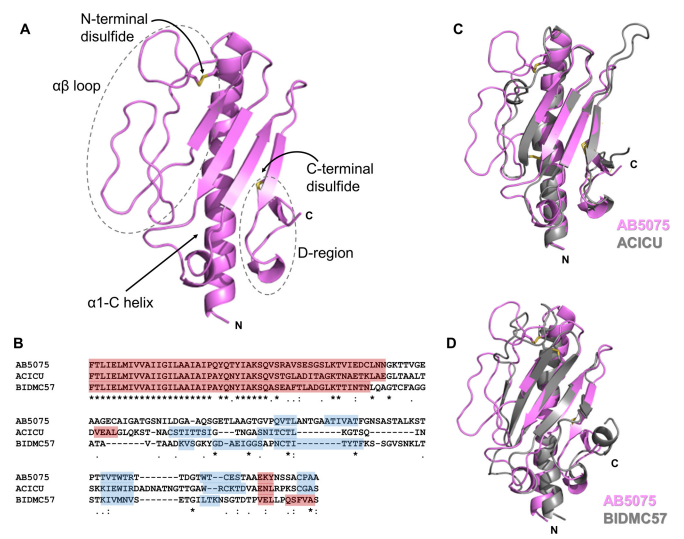


Figure 2. Structure of PilA<sup>AB5075</sup>. *A*, cartoon representation of PilA<sup>AB5075</sup> headgroup; disulfide bonds are displayed in *yellow*. *B*, sequence alignment of PilA<sup>AB5075</sup>, PilA<sup>ACICU</sup> and PilA<sup>BIDMC57</sup>. α-Helices are highlighted in *red*, and β-strands are in *blue*. Sequence identity (\*), close similarity (:), and similarity (.) are indicated below. *C*, superimposition of PilA<sup>AB5075</sup> (*pink*) and PilA<sup>ACICU</sup> (*gray*). *D*, superimposition of PilA<sup>AB5075</sup> (*pink*) and PilA<sup>BIDMC57</sup> (*gray*).

and the pilin of P. aeruginosa PAK (36). Despite the long stretches with no  $\alpha$ - or  $\beta$ -structure, the loop conformation can be unambiguously determined because of the well-defined electron density in this region (Fig. S1). An analysis of relative B-factors for PilA $^{\rm AB5075}$ , PilA $^{\rm ACICU}$ , and PilA $^{\rm BIDMC57}$ suggests that the internal dynamics of PilA<sup>AB5075</sup> and PilA<sup>ACICU</sup> may be similar as well (Fig. S2). Both have relatively low b-factors for the  $\alpha$ 1-C helix and central  $\beta$ -sheet with intermediate relative b-factors for the  $\alpha\beta$ -loop (though higher for two of the loops in  $PilA^{AB5075}$ ) and the highest relative b-factors at the C terminus. Conversely, PilABIDMC57 has a gradient of relative b-factors running along the α1-C helix from the N-terminal portion (low) to the tip and  $\alpha\beta$ -loop (high), whereas the C terminus is relatively well-ordered.

### Surface electrostatics of A. baumannii PilA variants

Variation in the  $\alpha\beta$ -loop of pilin proteins is well-documented (10), but the unusual structure of the  $\alpha\beta$ -loop in PilAAB5075 suggested some functional role to us. An examination of the surface electrostatics of PilAAB5075 showed an unusual concentration of acidic groups at the surface, particularly in the  $\alpha\beta$ -loop itself (Fig. 3A). The contrast is particularly striking with PilA<sup>ACICU</sup>. Calculations of theoretical isoelectric points for the two headgroups (i.e. excluding residues 1–22) give values of 4.73 for PilAAB5075 and 8.43 for PilAACICU. To evaluate the implications of the extended  $\alpha\beta$ -loop and its electronegative surface for a native pilus, we created models of fulllength PilAAB5075 (that is modeling the transmembrane helix spanning residues 1–22) (Fig. S3A) and an assembled AB5075 type IV pilus based on the 2006 model Neisseria gonorrhoeae pilus (37) (Fig. S3B) and the higher-resolution 2017 P. aeruginosa model from Wang et al. (55) (Fig. 3B and Fig. S3C). Electronegativity clearly predominates, with the few electropositive regions on the surface confined to a recessed groove, which follows the helical symmetry axis around the pilus fiber (Fig. 3C) and Fig. S3D).

If we compare the charged residues that are exposed on the surface of the AB5075 and ACICU pilus models (Fig. S4), each has six negatively charged (aspartate or glutamate) residues, but the ACICU surface also contains seven positively charged (lysine or arginine) residues, whereas none are found on the



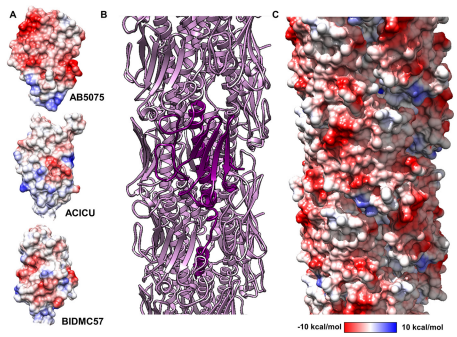


Figure 3. Model of the A. baumannii AB5075 type IV pilus. A, columbic surfaces of the PilAAB5075, PilAACICU, and PilABIDMC57 headgroups. B, cartoon representation of pilus model (pink). A single modeled full-length pilin is depicted in violet. C, columbic electrostatic surface depiction of the AB5075 pilus. Electrostatic potential key for A and C is shown below.

equivalent AB5075 surface (lysine 117 can be found in the previously mentioned groove). Two of these basic surface residues form unambiguous salt bridges with acidic residues (lysine 69/glutamate 63 and arginine 122/aspartate 109), whereas three other pairs are within 3 Å but ambiguous from the sidechain density (lysine 102/aspartate 126, lysine 103/glutamate 105, and arginine 132/glutamate 129). One logical consequence of an increased electronegativity of the pilus surface is that electrostatic repulsion would then disfavor pilus-pilus contacts, including the formation of pilus bundles, which have been demonstrated to promote microcolony formation in type IVb pili (56, 57).

### Twitching motility by $\Delta pilA$ complements

Phenotypic comparisons of A. baumannii have been undertaken by several other groups previously (58-60). Eijkelkamp et al. (60), in particular, examined the correlation between pilA sequence and motility and biofilm phenotypes, finding a link between pilA sequence and twitching motility. In this study, because we wished to isolate the effects of variation in PilA from other factors, we introduced pilA genes from A. baumannii AB5075, ACICU, and BIDMC57 into a  $\Delta pilA$  strain of A. nosocomialis M2. This strain, originally described as A. baumannii M2 (14), has been used as a model system for studies of multiple aspects of Acinetobacter pathogenesis (15, 21, 61).

We measured the ability of A. nosocomialis M2  $\Delta pilA$  complemented with plasmids containing pilA from A. baumannii AB5075, ACICU, and BIDMC57 (Fig. S5), as well as positive and negative controls to move at the interface between nutrient agar and polystyrene using standard methods (14). To quantitate the extent of twitching motility, we used crystal violet to stain the bacteria and image analysis software to distinguish bacteria from background and using WT,  $\Delta pilA$ ,  $\Delta pilT$ , and

complements for validation (Fig. S6). The results from 1% MacConkey agar plates (Fig. 4) show that the AB5075 and BIDMC57 complements were able to complement the  $\Delta pilA$ phenotype with much greater effectiveness than the ACICU mutant. The ACICU mutant did show significantly more movement than  $\Delta pilA$  (p = 0.035) but was at least an order of magnitude worse than the WT, native complement, AB5075 complement, and BIDMC57 complement.

### Adhesion to A549 and Detroit 562 cells

Previously we reported that the  $\Delta pilA$  mutant of A. nosocomialis M2 showed a defect in adhesion to A549 cells, an immortalized cell line derived from lung epithelial cells, which was restored in the complemented strain (15). Additionally, adhesion was significantly increased in the  $\Delta pilT$ mutant, which is incapable of retracting type IV pili. Based on these results, we reasoned that if the pilA<sup>ACICU</sup> complement was poorly motile because it produced few T4P, it would correspondingly be a poor complement for the native pilin in these host cell adhesion assays. If, however, the pilA<sup>ACICU</sup> complement was capable of normal pilus biogenesis but incapable of retraction, similar to what was observed previously by Rogers et al. (79), we would expect it to adhere to A549 cells more strongly than the pilAAB5075 and  $pilA^{BIDMC57}$  complements.

These assays were performed as described previously (15) with the exception that the bacteria were grown in MacConkey medium rather than Luria broth (see "Experimental procedures"). This change was prompted by our observation that type IV pilus expression by A. nosocomialis M2, as evidenced by twitching motility, is significantly greater in MacConkey medium than in Luria broth (Fig. S7). The results (Fig. 5) show that under these conditions, robust binding to both A549

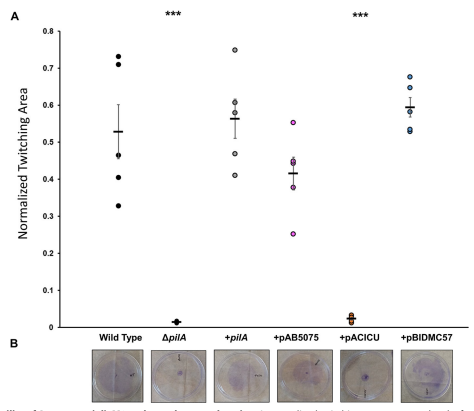


Figure 4. Twitching motility of A. nosocomialis M2 and complemented strains. A, normalized twitching area, expressed as the fraction of the plate covered. Error bars represent standard error. \*\*\*, p < 0.001. B, representative images of twitching results after staining with crystal violet.

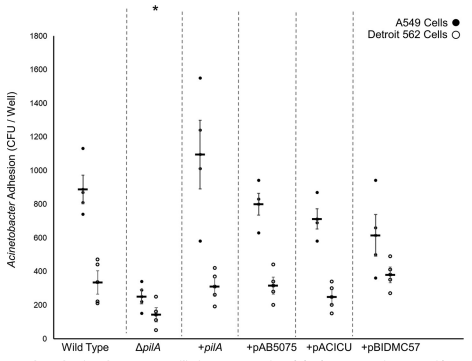


Figure 5. Host cell adherence through Acinetobacter type IV pili. The average number of cfu of A. nosocomialis recovered from a binding experiment with either A549 cells (black circles) or Detroit 562 cells (white circles) is shown. Error bars represent standard error. Significant (p < 0.05) reduction for both cell lines is marked with an asterisk (\*).

and Detroit 562 (nasopharyngeal) cells is observed for the WT strain and significantly decreased in the  $\Delta pilA$  mutant and that all three of *A. baumannii pilA* complements restore adhesion with no significant differences between them.

These data indicate that all three pilA complements are capable of both extension and retraction of T4P, and we find no relationship between PilA sequence and the relative binding to these two cell types.



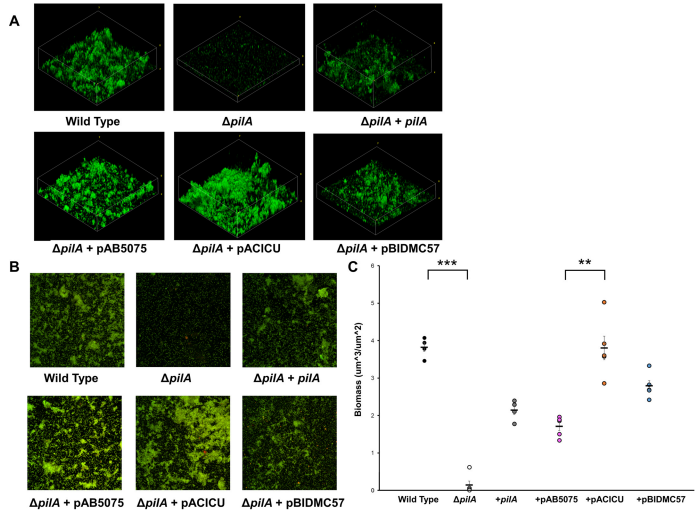


Figure 6. Acinetobacter biofilm formation on stainless steel. A, 3D reconstructions of biofilms imaged by CLSM. B, 2D, top-down CLSM images. C, biofilm biomass calculated for A. nosocomialis M2,  $\Delta pilA$ , and complements. Error bars represent standard error. \*\*, p < 0.01; \*\*\*, p < 0.01.

### A. baumannii biofilm formation on stainless steel

If the divergence in PilA sequence is the result of functional specialization, we would expect that a defect in one function would be compensated for by a gain in another. Correspondingly, we also measured the ability of our complements to form biofilm in a vertical biofilm formation assay similar to a CDC biofilm reactor (62). Briefly, sterilized stainless steel coupons were placed upright in culture tubes and immersed in MacConkey medium, and the medium was inoculated with bacterial cultures from saturation growths. Our experimental design was influenced by several factors, including our prior results growing biofilms horizontally on glass surfaces (15). Previously we found no significant difference between A. noso*comialis* M2 WT,  $\Delta pilA$ , and the native complement in biofilm formation on horizontal glass surfaces grown in Luria broth. However, as noted above, we now expected stronger T4P-dependent phenotypes in MacConkey medium. Similarly, Acinetobacter adhesion to stainless steel is well-characterized and robust compared with untreated glass (63). We attribute the growth medium dependence of T4P expression to differences in the production of quorum-sensing molecules, consistent with the observation that virstatin (an inhibitor of AnoR/AnoI)

reduces both surface motility and biofilm formation in A. baumannii (64, 65).

After fixation and fluorescent staining with FM 1-43, biofilm formation was assessed using confocal laser scanning microscopy (CLSM) (Fig. 6). Calculated biomass shows a significant phenotype for the  $\Delta pilA$  mutant, which can be complemented by the native M2 pilA gene as well as the pilA genes of AB5075, ACICU, and BIDMC57. Additionally, the ACICU complement formed significantly greater biomass than the AB5075 complement (Fig. 6C). We attribute this difference in biomass to differences in bacterial aggregation because all strains, including  $\Delta pilA$ , were able to adhere in a monolayer to the stainless steel surface (Fig. 6B). Additionally, we found that biofilm formation was accompanied by the formation of crosslinking pilus-like fibers for all strains (Fig. S8), which we attribute to chaperone-usher (Csu) pili, consistent with their role in Acinetobacter biofilm formation (66, 67).

### Discussion

Variation in the genetics of *A. baumannii* virulence factors, including type IV pili, is well-established, and with the improvements in metagenomics sequencing technology, the acquisition

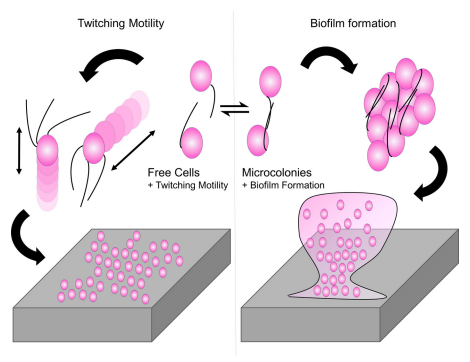


Figure 7. Schematic model for specialization in Acinetobacter type IV pili. A potential tradeoff between biofilm formation and twitching motility based solely on PilA structure is depicted based on the equilibrium between singled and bundled T4P.

of genomic data continues to accelerate. In this work we examine the relationships between genetic variation in the major subunit of type IV pili, PilA, the molecular structure of type IV pili, and the resulting bacterial phenotypes. We observed differences in surface chemistry between PilAAB5075 and PilAACICU even as the secondary structure was largely conserved and found that the two proteins promoted different bacterial behavior as well, with the AB5075 pilin favoring motility and the ACICU pilin favoring biofilm formation. We propose that these results can be explained by the equilibrium between single and bundled type IV pili.

Type IV pili mediate diverse functions in a variety of bacteria (10), but all of the known functions rely on some interplay between two general characteristics: adhesion (whether to DNA, biotic or abiotic surfaces, or each other) (56) and retraction (which is essential for twitching motility and natural competence) (14). The stark contrast in surface electrostatics between PilA<sup>AB5075</sup> and PilA<sup>ACICU</sup>, given the overall similarity in fold, suggested to us that fibers formed from PilAAB5075 would be more prone to electrostatic repulsion and hence adhere less to each other.

Pilus bundling has also been shown to be a function of pilin surface chemistry; Neisseriae type IV pili are less bundled when glycosylated at serine 63 (68, 69). Because, unlike A. baumannii AB5075, the ACICU and BIDMC57 strains C-terminally glycosylate PilA, they may reduce pilus bundling through glycosylation without the need for electrostatic repulsion. However, it is important to distinguish between bundling between the pili of a single cell (cis-bundling) and bundling between the pili of adjacent cells (trans-bundling). cis-Bundling is no detriment to motility and in fact has been shown to dramatically increase force of pilus retraction and to increase the persistence of twitching motility (70, 71).

Trans-bundling promotes microcolony formation in enteropathogenic Escherichia coli and Vibrio cholerae (56, 57), a precursor to biofilm formation. The balance between cis- and trans-bundling is also dependent upon the number of pili per cell, which is dramatically lower in Acinetobacter than Neisseriae and Pseudomonas (14, 37, 55).

An increase in microcolony formation could explain both the greater biofilm formation of the ACICU complement and its poor motility. A scheme describing this model is shown in Fig. 7. Single cells are free to move across the surface, resulting in an overall increase in twitching motility, whereas cells joined together into microcolonies are less motile but serve as nucleants for the formation of biofilm.

Inverse relationships between twitching motility and biofilm formation have been observed previously in P. aeruginosa  $\Delta pilT$ mutants (38) and correlatively in clinical isolates (72). In cases where  $\Delta pilT$  mutants are hyperpilated, increases in retraction-independent adhesive functions can be explained simply through the increase in adhesive molecules on the surface. By way of example, we previously reported increased host cell adhesion for the A. nosocomialis M2  $\Delta pilT$  mutant (15). However, we observe no hyperpilation of the pilA<sup>ACICU</sup> complement, either directly by TEM (Fig. S9) or in increased host cell adhesion (Fig. 5), implying that the defect in motility does not stem directly from decreased pilus retraction.

Returning to prior studies comparing the phenotypes of A. baumannii strains, we examined the degree to which the results reported here were consistent with the behavior of the native bacteria. Eijkelkamp et al. (60) compared the motility and adhesion characteristics of a wide variety of clinical isolates and reported that only 3 of 32 international clone II isolates (which includes ACICU) showed twitching motility, in contrast to international clone I (which have pilA sequences similar to



AB5075), in which 8 of 8 were motile; this correlation between PilA sequence and motility was also noted by the authors. However, although these results are consistent with what we observe with the pilA AB5075 and pilA ACICU complements, we note that American Type Culture Collection 19606, which has a pilA sequence very similar to BIDMC57, was found to be nonmotile in that study. More recently and also consistent with our findings, Sahl et al. (73) found greater biofilm formation by ACICU than AYE (which has a PilA sequence identical to AB5075).

If a tradeoff between twitching motility and biofilm formation can explain the differentiation of Acinetobacter type IV pili, what evolutionary pressures favor one over the other? Because the three pilin proteins in this study are from A. baumannii clinical isolates, one obvious possibility is that differences in infection sites led to specialist pathogens. This view is supported by the observation by Vijayakumar et al. (72) that A. baumannii isolates from the sputum formed more biofilm in *vitro* than isolates from the blood, whereas the reverse was true for twitching motility. However, Acinetobacter strains with similar type IV pilins have also been isolated environmentally; A. calcoaceticus PHEA-2 has a PilA sequence nearly identical to that of A. baumannii ACICU (94% amino acid identity) but was isolated from industrial wastewater (74). It is possible that a distinction exists between the T4P of environmentally adapted strains, which retain some infectivity and the T4P of specialized pathogenic strains; Wang et al. (75) observed a trend associating biofilm-forming strains with better clinical outcome. However, more work comparing clinical, commensal, and environmental isolates in controlled studies, particularly models of infection, remains to be done before we can draw such a conclusion.

The ability of all three A. baumannii pilA complements to adhere to host cells in a similar manner despite their differences in structure and surface chemistry is consistent with our prior observation that the removal of a C-terminal pentasaccharide from the A. nosocomialis PilA protein also has no effect on binding to A549 or Detroit 562 cells. We hypothesize that the relevant adhesin in this case is not PilA itself but a minor pilin subunit (consistent with the conservation we see of the minor A. baumannii minor pilins in Fig. 1) or a protein that interacts with the pilus, either constitutively at the tip (76) or as a secreted factor, as was recently shown in the type IVb pili of ETEC (77).

Recently, Harvey et al. (78) reported that pilus glycosylation can inhibit the binding of phage to the type IV pili of *P. aerugi*nosa, providing the most compelling explanation to date for the wide prevalence of pilin glycosylation. With that in mind, we considered the possibility that the electronegative surface chemistry of PilAAB5075 could also be explained in terms of defense against phage (PilAAB5075 is not C-terminally glycosylated, unlike PilA<sup>ACICU</sup> and PilA<sup>BIDMC57</sup>). However, based on the available structures of *Pseudomonas* PilA proteins that are not natively glycosylated, pronounced surface electronegativity does not appear to be a general feature of pilins lacking C-terminal glycans.

In conclusion, the results here demonstrate that subsets of A. baumannii produce type IV pili with markedly different molecular structure, and this variation, particularly in terms of surface chemistry, can result in phenotypic differences in motility and biofilm formation. The prevalence of type IV pili, in general, and the homology between A. baumannii type IV pili and those of phenotypically similar species such as P. aeruginosa, in particular, imply that these findings may be generalizable to other biofilm-forming bacteria.

### **Experimental procedures**

### Protein expression and purification

PilAAB5075 was expressed and purified as described previously (15). Briefly, the codon-optimized sequence, starting with alanine 23, was cloned into a maltose-binding fusion vector under a T7 promoter, making use of previously described surface entropy reduction mutations (pMal E) (29). A C-terminal His, tag was included to ease purification. This plasmid was transformed into BL21 (DE3) pLysS cells and grown to saturation overnight with shaking at 37 °C in LB medium with 50 μg/ml ampicillin. These saturation cultures were then diluted into fresh LB-ampicillin and grown with shaking to an optical density of 0.5 at 37 °C. The flasks were cooled to 18 °C before induction with 30 mm isopropyl  $\beta$ -D-1-thiogalactopyranoside and allowed to grow overnight with shaking before being harvested by centrifugation at 7,500  $\times$  g for 10 min. The cells were lysed using lysozyme (0.25 mg/ml final concentration), DNase (0.02 mg/ml) and Triton X-100 (0.5%) for 10 min, and the resulting lysate was centrifuged again, this time at 20,000  $\times$  g for 30 min. The supernatant was purified using a nickelnitrilotriacetic acid column, and the elution was further purified through size-exclusion chromatography over a GE S200 Superdex column using an Äkta Purifier FPLC.

### Structure determination and refinement

Maltose-binding protein-PilAAB5075 crystallization conditions were screened by sitting-drop vapor diffusion at a concentration of 20 mg/ml in 20 mm Bis-Tris (pH 6.0), with and without 50 mm maltose. A crystallization condition was found, without the addition of maltose, in the Morpheus screen (Molecular Dimensions), (H4), 12.5% (w/v) PEG 1000, 12.5% (w/v) PEG 3350, 12.5% (v/v) 2-methyl-2,4-pentanediol, 0.02 м of amino acid mix (0.2 M DL-glutamatic acid monohydrate, 0.2 M DL-alanine, 0.2 M glycine, 0.2 M DL-lysine monohydrochloride, and 0.2 M DL-serine), 0.1 M MES/imidazole, pH 6.5. Crystals were grown in hanging drops at room temperature and took ~48 h to grow at a protein concentration of 10 mg/ml. They were then harvested and flash-cooled in the mother liquor supplemented with 20% ethylene glycol. The data were collected at the Advanced Photon Source, GM/CA, Beamline 23ID-D. The General Medical Sciences and Cancer Institutes of Structural Biology Facility at the Advanced Photon Source (GM/CA @ APS) is a part of the X-ray Science Division at APS, Argonne National Laboratory (ANL).

The resulting data set was processed with XDS. Molecular replacement was carried out by Phaser (30) using a sequential search of (i) maltose-binding protein and (ii) PilA from A. baumannii ACICU (15). Phenix and Coot were used for phasing, building, and refinement (31-34). The crystallographic parameters of the refined data are summarized in Table 1.

### **Electrostatic calculations**

Columbic surfaces were calculated using UCSF Chimera (35) using a distance-dependent dielectric and a dielectric constant of 4.0, 1.4 Å from the surface. Theoretical polypeptide isoelectric points were calculated using The Swiss Institute of Bioinformatics (ExPASy) ProtParam server (https://web.expasy.org/protparam/).

### Pilus modeling

Full-length PilA<sup>AB5075</sup> was modeled based on the structure of the full-length P. aeruginosa PAK pilin (36). The initial model of the pilus was created by superimposition onto a model of the N. gonorrhoeae type IV pilus filament (Protein Data Bank code 2HIL) (37). The resulting model was then minimized using UCSF Chimera (35).

### Complementation of pilA mutant

pilA genes from A. baumannii AB5075, ACICU, and BIDMC57 were synthesized (Genscript) and ligated into pUCP20GM (38) using BamH1 and HindIII restriction sites (accession numbers in Table S1). The resulting vectors were electroporated into A. nosocomialis M2 ΔpilA (14) using standard protocols (39). The presence of the plasmids was confirmed by both resistance to gentamycin and PCR of the pilin genes.

### Twitching motility

A. nosocomialis M2 (including mutants and complement strains) was grown on 1.5% MacConkey agar plates overnight. Colonies were selected and stabbed through the centers of 1% agar plates in polystyrene Petri dishes. The plates were incubated in sealed bags at 37 °C for 3 days. The agar was then removed, and the bacteria which adhered to the polystyrene Petri dish were stained with 0.1% crystal violet for 5 min. Excess crystal violet was removed by gentle washing with deionized water. The subsurface twitching area on each plate was accessed using GIMP imaging software. Statistics were calculated for five replicates and significance determined by Student's t test.

### **Biofilm formation**

All strains were grown on 1.5% MacConkey agar plates, supplemented when necessary with gentamycin for plasmid maintenance. Overnight cultures were grown from these plates in MacConkey medium and diluted 1:10 into fresh MacConkey medium in 10-cm<sup>2</sup> flat tissue culture tubes (TPP Techno Plastic Products AG) containing upright  $1/8 \times 1$ -inch untreated stainless steel fender washers (Everbilt). After 72 h of shaking (50 rpm) at room temperature, the washers were removed to 6-well cell-culture plates, gently washed with PBS, stained, and covered in aluminum foil, with FM 1-43 dye (1:1000 in PBS) for 15 min at room temperature. The samples were then washed with PBS and fixed overnight with 4% paraformaldehyde in PBS at 4C. The fixed samples were stored at 4 °C in PBS until being imaged as described below.

### Confocal laser scanning microscopy

Biofilms were grown on stainless steel surfaces and prepared as described above. Each stainless steel washer was covered

with a 18 × 18-mm glass coverslip and read using a Nikon A1 confocal laser scanning microscope and accompanying software (Nikon, Tokyo, Japan). Z-stacks were acquired for each strain. The structural organization of the biofilms was analyzed using the Comstat2 software package (http://www.comstat. dk)<sup>3</sup> (40). The 3D representations of the biofilms were generated using the 3D viewer plugin for the FIJI distribution of ImageJ (http://3dviewer.neurofly.de)3 (41).

### Cell adhesion

A549 human airway adenocarincoma cells (52) (American Type Culture Collection, CCL 185) or Detroit 562 pharyngeal carcinoma cells (53) (American Type Culture Collection, CCL 138) were seeded in 24-well culture plates and cultured at 37 °C, 5% CO<sub>2</sub> to  $2.0 \times 10^5$  cells/well in Dulbecco's modified Eagle's medium containing 10% fetal bovine serum, 2.0 mm glutamine, 100 units/ml penicillin, and 100 μg/ml streptomycin. The cells were washed twice with PBS, pH 7.2, fixed for 10 min at room temperature with 2.5% (v/v) glutaraldehyde in PBS, pH 7.2, and washed three times with PBS, pH 7.2, as described (42, 43). A. nosocomialis M2 (including mutants and complements) was cultured overnight in MacConkey broth, washed twice with PBS, pH 7.2, resuspended in PBS, pH 7.2 containing 2.0 mg/ml glucose, and quantified spectrophotometrically at  $A_{600}$ . Fixed A549 or Detroit 562 cells ( $2.0 \times 10^5$ /well) were incubated with  $2.0 \times 10^7$  cfu/well of *A. nosocomialis* M2 in 0.5 ml for 40 min at 37 °C and washed three times with PBS, pH 7.2. Bound bacteria were released with 0.05% trypsin-EDTA, and bound colonyforming units were quantified on Luria Bertani agar plates, as described (42, 43). Significance was determined by Student's t

All depictions of protein structures were created using PyMOL (Schrödinger) or UCSF Chimera (35). Sequence alignments were made using Clustal Omega (44), and phylogenic trees were diagrammed using Interactive tree of life (iTOL) from EMBL (45).

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<sup>&</sup>lt;sup>3</sup> Please note that the JBC is not responsible for the long-term archiving and maintenance of this site or any other third party hosted site.

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