**Pseudomonas aeruginosa** PilY1 Binds Integrin in an RGD- and Calcium-Dependent Manner

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**Abstract**

PILY1 is a type IV pilus (tfp)-associated protein from the opportunistic pathogen *Pseudomonas aeruginosa* that shares functional similarity with related proteins in infectious *Neisseria* and *Kingella* species. Previous data have shown that PILY1 acts as a calcium-dependent pili biogenesis factor necessary for twitching motility with a specific calcium binding site located at amino acids 850–859 in the 1,163 residue protein. In addition to motility, PILY1 is also thought to play an important role in the adhesion of *P. aeruginosa* tfp to host epithelial cells. Here, we show that PILY1 contains an integrin binding arginine-glycine-aspartic acid (RGD) motif located at residues 619–621 in the PILY1 from the PAK strain of *P. aeruginosa*; this motif is conserved in the PILY1s from the other *P. aeruginosa* strains of known sequence. We demonstrate that purified PILY1 binds integrin *in vitro* in an RGD-dependent manner. Furthermore, we identify a second calcium binding site (amino acids 600–608) located ten residues upstream of the RGD. Eliminating calcium binding from this site using a D608A mutation abolished integrin binding; in contrast, a calcium binding mimic (D608K) preserved integrin binding. Finally, we show that the previously established PILY1 calcium binding site at 851–859 also impacts the protein’s association with integrin. Taken together, these data indicate that PILY1 binds to integrin in an RGD- and calcium-dependent manner *in vitro*. As such, *P. aeruginosa* may employ these interactions to mediate host epithelial cell binding *in vivo*.

**Citation:** Johnson MDL, Garrett CK, Bond JE, Coggan KA, Wolfgang MC, et al. (2011) *Pseudomonas aeruginosa* PilY1 Binds Integrin in an RGD- and Calcium-Dependent Manner. *PLoS ONE* 6(12): e29629. doi:10.1371/journal.pone.0029629

**Editor:** Michael Hensel, University of Osnabrueck, Germany

**Received** August 30, 2011; **Accepted** December 2, 2011; **Published** December 29, 2011

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**Funding:** This work was supported by a Ruth Kirschstein Supplement (MDLJ) on the National Institutes of Health Grant AI078924 (MRR). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing Interests:** The authors have declared that no competing interests exist.

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**Introduction**

*Pseudomonas aeruginosa* is a Gram-negative, opportunistic pathogen prevalent in immunocompromised patients, burn victims, and people with cystic fibrosis. *P. aeruginosa* can infect any part of the body, but typically targets the respiratory tissues, skin abrasions, and the urinary tract [1]. This pathogen accounted for ~11% of the hospital-acquired infections reported in the U.S in 2002 [2,3], and has also been shown to infect other mammalian species, as well as insects and plants [4,5]. *P. aeruginosa* uses a range of methods to resist the effects of antibiotics, including efflux pumps, adaptive mutagenesis, and protective biofilms [6,7,8]. As such, *Pseudomonas aeruginosa* presents a significant challenge to human health.

*P. aeruginosa* employs type IV pili (tfp) for twitching motility and infection. The precise mechanism of host cell attachment has remained unclear, although evidence exists that loops of the major pilus structural subunit PilA exposed at the tip of the pilus fiber bind to gangliosides GM1 and GM2 [9]. In contrast, some data indicate that several *P. aeruginosa* clinical isolates, as well as laboratory strains, do not employ GM1 and GM2 during host cell attachment [10,11]. As such, it has been proposed that other factors on the *Pseudomonas* tfp are involved in binding to target cells [12,13].

Recent studies have suggested that host cell integrin proteins play a role in *Pseudomonas* infection. Anti-integrin antibodies were shown to reduce *P. aeruginosa* attachment to host cells [14]. Specifically, antibodies to the αVβ5 integrin and αVβ3 integrin were effective at disrupting *P. aeruginosa* binding to host cells, with antibodies to αVβ5 integrin having the most pronounced effect [14]. Integrins are present on the epithelial cell surface of tissues infected by *P. aeruginosa*; indeed, αVβ5 integrin is highly expressed in the lungs [15]. The presence of *P. aeruginosa* has also been shown to increase the expression of integrin subunits αV, α5, and β1 in epithelial cells [16,17]. Integrins and integrin-like proteins are found widely in animals, plants, and insects, all targets of *P. aeruginosa* infection [18]. Integrins and integrin-like proteins are responsible for a range of cellular processes, including cell-cell attachment, cellular signaling, and angiogenesis [19]. The most common ligands for integrins are proteins that contain an arginine-glycine-aspartic-acid (RGD) sequence, although other short peptides have also been found to mediate integrin-protein interactions (e.g., leucine-aspartic acid-valine or LDLV).

The PilC class of proteins in *Neisseria gonorrhoeae*, *Neisseria meningitidis*, and *Kingella kingae* have been characterized as tfp biogenesis factors and as proteins involved in adhering to target
tissues [20,21,22]. Generally, PilC N-terminal regions are associated with adhesion domains, while the C-terminal domains regulate tfp biogenesis. *P. aeruginosa* PilY1 shares sequence homology with the C-terminal regions of the PilCs proteins, and this domain in PilY1 has been shown to be a calcium-mediated tfp biogenesis factor (Table S1) [23]. Indeed, PilY1 is required for both twitching and swarming motility and adhesion to host cells [23,24,25]. The N-terminal domains of PilY1 and the PilCs share low sequence similarity, which might be expected if these domains are involved in target cell-specific activities. The target tissues of the *Neisseria* strains and *Kingella* are likely to be distinct from those targeted by *P. aeruginosa*. Here, we focus on putative regions of PilY1 involved in host cell attachment and we show that purified PilY1 binds to integrins in an RGD- and calcium-dependent manner in vitro.

**Results**

PilY1 contains a conserved integrin binding motif

Given the importance of the tfp associated PilY1 protein in host tissue adherence and the ability of antibodies targeting the RGD binding domains of αVβ5 and αVβ3 integrins to reduce interactions between *P. aeruginosa* and host cells, we examined PilY1’s sequence to locate possible integrin binding motifs (RGD or LDV) [14,25]. PilY1 was found to contain a conserved RGD motif (Figure 1A, 1B). No other type IV pilus proteins or known surface proteins in *P. aeruginosa* contained an RGD or other putative integrin binding sequence [26,27]. One RGD motif was found in each of the PilY1s of *P. aeruginosa* strains of known sequence. For example, in the PilY1 from *P. aeruginosa* strain K (PAK), the RGD is located at residues 619–621; however, in the PAO1 *P. aeruginosa* strain PilY1, the RGD is at 657–659 (Figure 1B). Based on these observations, we hypothesized that PilY1 binds to integrin in an RGD-dependent manner.

PilY1 binds integrin in an RGD-dependent manner

To test the hypothesis that PilY1’s RGD motif mediates contacts with integrins, we created a purified form of PilY1 that starts at amino acid 532 and extends to C-terminus of the protein (residue 1163) and tested the ability of mid-nanomolar concentrations of this purified protein to specifically bind to αVβ5 integrin (Figure 2A). Next, we employed synthetic RGDS and GRADSP peptides to compete with PilY1 binding to αVβ5 integrin. We found that the RGDS peptide significantly reduced PilY1 binding to αVβ5 integrin (p<0.10), while the GRADSP control peptide had no effect (Figure 2B). Because previous reports also indicated that PAK binding to host cells can be reduced by using an anti-αVβ3 integrin antibody, we also examined PilY1 binding to αVβ3. We found that purified PilY1 binds to αVβ3 integrin in an RGD-dependent manner, although with a lower apparent affinity than observed with αVβ5 integrin (Figure 2C) [14]. We therefore used αVβ5 as the primary integrin of this study. Finally, we tested two RGD mutants of PilY1, D621A (RGA) and D619–621 (Drgd), for their ability to bind to αVβ5 integrin in vitro. Deleting the RGD residues or mutating the aspartic acid residue at position 621 to alanine significantly

![Figure 1](https://example.com/figure1.png)

**Figure 1.** *P. aeruginosa* PilY1 strains contain conserved integrin binding residues RGD and conserved putative calcium binding site.

(a) A bar representation of PilY1. The consensus C-terminal pilus biogenesis domain is in blue and the green represents the N-terminal addition to the previously examined construct. Calcium binding motifs are highlighted in yellow and the RGD is highlighted in orange. (b) Five varying strains of *P. aeruginosa* PilY1 were aligned using the biology workbench server [26,27]. Blue residues and an “*” corresponds to identical residues throughout the 5 strains, green residues and a “:” correspond to highly conserved residues, and navy residues and a “.” corresponds to mildly conserved residues (e.g. alanine and leucine). Reference numbers are for PAK_287 PilY1 (the strain used in this study).

doi:10.1371/journal.pone.0029629.g001
reduced the association of PilY1 with αvβ5 integrin (Figure 2D). Importantly, these mutations did not alter the global secondary structure or the melting temperature of purified 532–1163 PilY1 (Figure 2E, Table S2). Taken together, these data indicate that P. aeruginosa PilY1 binds to integrins in vitro in an RGD-dependent manner.

PilY1 has a second calcium binding site

A second potential calcium binding site was noted in the sequence of P. aeruginosa PilY1 between residues 600 and 608, which is in close proximity to the RGD motif (619–621) described above (Figure 1A). Recall that a calcium binding site at residues 851–859 in PilY1 was shown in previous work to be critical to tfp biogenesis and twitching motility [23]. The sequences of the two calcium binding sites are closely related (Figure 3A), and the putative 600–608 site is conserved in PilY1s of known sequence (Figure 1B). Indeed, in the P. aeruginosa PilY1 sequences examined, this second potential calcium binding site was always found 8–10 residues upstream of the conserved RGD motif (Figure 1B). To test the 600–608 site’s ability to bind calcium in purified 532–1163 PilY1, we eliminated the previously published calcium binding site using either a D859A or D859K mutation [23], and measured calcium binding. The D859A and D859K forms of 532–1163 PilY1 exhibited K_d’s for calcium binding of 412 and 266 nM, respectively (Figure 3B). Thus, the 600–608 site appeared to bind calcium. We next created corresponding mutations (D608A, D608K) in the 532–1163 PilY1 construct, and found that these variants bound calcium with 2.3–2.4 μM affinity, similar to that reported previously for the 851–859 PilY1 site (Figure 3C) [23]. Finally, we created a D608A/D859A double-mutant form of 532–1163 PilY1 and compared its calcium binding to wild-type 532–1163 PilY1. We found that the double-mutant exhibited only non-specific calcium binding, while wild-type 532–1163 PilY1 associated with calcium with a 700 nM affinity (Figure 3D). The variant forms of 532–1163 PilY1 examined in vitro in these studies maintained an overall wild-type fold and melting temperature, indicating that these changes did not alter the stability of the proteins (Figure S1A–E; Table S2). Additionally, mutations in the RGD sequence adjacent to the 600–608 site did not alter calcium binding by these purified forms of 532–1163 PilY1 (Figure S2A). The 532–1163 PilY1 construct demonstrated 1.3 μM and 600 nM affinity binding to Zn and Mn, respectively, and non-specific

Figure 2. Purified PilY1 binds to integrin in an RGD dependent manner. (a) PilY1 was titrated onto a solid phase binding assay keeping the concentration of αVβ5 integrin constant at 2 μg/mL. The curve was fit to the ligand binding, one site saturation equation f = Bmax*abs(x)/(Kd+abs(x)) with an r^2 = 0.98. The K_d was calculated at 164 nM±38. (b–d) Wild type PilY1 was established as a reference point and the value all measurements were compared to. (b) RGDS inhibitor peptide or GRADSP control peptide were added in 2, 100, and 250 fold molar excess to PilY1. ** corresponds to p<.1 compared to PilY1 binding to αVβ5 integrin. (c) PilY1 was added with or without 50 fold molar excess of the inhibitor RGDS peptide to 2 μg/mL of αVβ3 integrin. *** corresponds to p<.02 compared to PilY1 binding to αVβ3 integrin. (d) ΔRGD (A619–621) or D621A mutations of PilY1 were added to 2 μg/mL of αVβ5 integrin. **** corresponds to p<.01 compared to PilY1 binding to αVβ5 integrin. (e) Molar ellipticity values were calculated for the respective wavelength scans and compared. doi:10.1371/journal.pone.0029629.g002
However, due to PilY1’s structural similarity to established calcium binding sites, we consider calcium to be the primary metal bound to PilY1 [23,28,29,30]. Taken together, these observations support the conclusion that purified 532–1163 PilY1 contains two calcium binding sites, the previously characterized one at 851–859 and a newly identified one at 600–608 that is ten residues N-terminal to PilY1’s RGD motif.

PilY1 binds integrin in a calcium dependent manner

Because the 600–608 calcium binding site in PilY1 is close in sequence to the RGD motif, we next sought to examine whether calcium binding impacted the association between 532–1163 PilY1 and integrin. We found that D859A, D859K, and D608K mutant forms of 532–1163 PilY1 exhibit wild-type levels of integrin binding (Figure 4B). For example, the D608K/D859A variant of 532–1163 PilY1 shows nearly wild-type levels of integrin binding (Figure 4B); recall that the D608K form of PilY1 exhibited little integrin binding in vitro (Figure 4A). Thus, D859A appears to act as a dominant positive over the D608K mutation. Similarly, the D608K/D859A PilY1 shows increased integrin binding relative to wild-type (Figure 4B). In contrast, adding the D859A mutation to either D608A or D608K in PilY1 had no impact on integrin binding relative to the 608 mutants alone (Figure 4B). Taken together, these data suggest that both calcium binding sites in PilY1 impact integrin binding in vitro. Note that the single- and double-mutants considered here did not bind calcium (Figure S2B), and exhibited wild-type CD spectra and melting temperatures (Figure S1A–E, Table S2), indicating that the overall structure of these purified proteins was not altered.

The crystal structure of the C-terminal domain of PilY1 does not contain the 600–621 region; as such, it is not known whether the two calcium binding sites are in close structural proximity.

**Figure 3. PilY1 has two functional calcium binding sites.** (a) An alignment of the two calcium binding sites in PilY1 was performed as in figure 1. Calcium coordinating residues are underlined. (b, c, and d) A calcium competition binding assay using Oregon Green was performed with (b) D859A and D859K, (c) D608A and D608K, and (d) wild type and D608A/D859A. Binding curves were modeled to one-site competition (D859A, D859K, D608A, and D608K, and wild type), or linear line (D608A/D859A). Error represents standard error of the mean. doi:10.1371/journal.pone.0029629.g003
However, it is possible that they interact physically, and that the functional impact of one upon the other is via the swapping of an aspartic acid from one site (e.g., D851 or D855) into the other site (600–608). To test this hypothesis, we created two triple mutant forms of 532–1163 PilY1, D608A/D851A/D859A and D608A/D855A/D859A, and one quadruple mutant form, D608A/D851A/D855A/D859A. Both triple mutants and the quadruple mutant show wild type levels of binding (Figure S4). Thus, we conclude that an interaction between the two calcium binding sites in PilY1 is not primarily associated with aspartic acids in one site helping to coordinate calcium in the other site. However, taken together, these data indicated that a functional interaction between the two calcium binding sites in PilY1 impacts the ability of the RGD motif in this protein to bind to integrins in vitro.

Discussion

P. aeruginosa is an established and increasingly antibiotic resistant pathogen that predominantly infects patients with compromised defense mechanisms. Toward this end, we sought to understand how P. aeruginosa binds to host cells, a necessary first step in infection. We focused on the type IV pilus-associated protein PilY1, which is involved in bacterial motility and is related in sequence to factors in other bacteria required for host cell binding. We show that PilY1 contains an integrin binding RGD motif that mediates contact between purified PilY1 and integrin proteins in vitro. Additionally, we show that PilY1 contains two calcium binding sites and that each site impacts the protein’s ability to associate with integrin (Figure 4C). Both the calcium binding sites and the RGD motif are conserved in all PilY1 sequences from P. aeruginosa strains reported to date (Figure 1B).

It was not anticipated that the PilY1 calcium binding sites, particularly the one distal (851–859) to the RGD sequence, would impact integrin binding. In fact, D859A can be seen as a dominant positive with respect to integrin binding (Figure 4C). For example, while the D608A mutant form of PilY1 is significantly reduced in integrin binding, combining it with D859A to create a D608A/D859A double-mutant produces a protein that is capable of binding to integrin (Figure 4B). Further, a D608K/D859A double-mutant exhibits higher integrin binding than either single mutant or the wild-type protein (Figure 4B). In spite of these in vitro data, however, the molecular contacts involved in these effects are not clear, in part because the region between residues 600 and 643 in PilY1 has not been elucidated structurally. It is tempting to speculate that a physical association exists between the two calcium binding sites that impacts RGD presentation for integrin binding. Such speculation is supported by the data outlined here, but remains speculation until future studies to examine the structure of this region of PilY1 are complete.
Integrins and integrin-like proteins are present in plants, insects, and the animal kingdom. They are up-regulated during cellular stress and recovery [31,32,33]; indeed, the presence of \textit{P. aeruginosa} itself up-regulates the expression levels of the \( \alpha V \) integrin subunit [16]. In addition, integrins are established targets for bacterial pathogens. \textit{Bordetella pertussis} protein pertactin, containing two RGD motifs, has been shown to adhere to Chinese hamster ovary cells in an RGD-dependent manner [34,35]. Pertactin is a member of the auto-transporter family, many of which contain RGD motifs mediating adherence to integrins [36]. However, there are conflicting data on whether auto-transporter RGDs are relevant in mouse models of infection [37]. Still, other studies have shown that the following bacteria bind integrins in an RGD-dependent manner: \textit{Pyrenophora triticci-repentis} (a wheat pathogen), \textit{Mycoplasma conjunctivae} (sheep pathogen), and the mammalian pathogens \textit{Klebsiella pneumoniae} and \textit{Helicobacter pylori} [38,39,40]. Integrins are also known targets for viral pathogens. Proteins in hepatitis C virus, coxsackievirus A9, human herpes virus 8, Epstein-Barr virus, and adenovirus bind to host cells in an RGD dependent manner [41,42,44,45]. In summary, integrins have the potential to be important target proteins for PilY1-mediated attachment of \textit{P. aeruginosa}.

Interestingly, calcium has also been shown to play a vital role in host RGD substrate:integrin binding events (Figure 5). Calcium binding helps in the folding of the blood clotting factor fibrinogen, and it protects against cleavage and degradation of the hemostasis protein von Willebrand factor and the elastin scaffold protein fibrillin (Figure 5) [46,47,48,49]. Some integrin substrates, such as thrombospondin, use cooperative calcium binding at multiple sites to facilitate integrin binding, while calcium increases adhesion in tenasin, vitronectin, and osteopontin to target integrins (Figure 5) [50,51,52,53,54]. Thus, there is precedent for calcium regulation of RGD-mediated contacts with integrins.

PilY1 is an established pilus biogenesis factor [23] and has been implicated in \textit{P. aeruginosa} adhesion to host cells by functional studies [25] and due to its homology to the PilCs adhesin proteins of \textit{Neisseria} [55]. Here, we provide evidence that PilY1 serves as an adhesin that binds to integrins in an RGD-dependent manner. Furthermore, we show that integrin binding is regulated by two calcium binding sites in PilY1. These data provide an advance in our understanding of PilY1 structure and function. It will be of interest to determine in future studies whether PilY1 and integrins are involved in \textit{P. aeruginosa} attachment to host cells in \textit{vivo}.

### Methods

#### \textit{P. aeruginosa} PilY1 Binds Integrin

**PilY1 constructs and protein purification**

Site-directed mutagenesis was performed to produce D608A, D608K, D621A, D619-621, D859A, D859K, D608A/D859A, D608K/D859A, D608K/D859K, D608A/D859A/D859K, D608A/D859A/D859K/D859A mutants in a pDONR vector containing PilY1. Amino acids 532–1163 were cloned out of the pDONR vector for entry into pMCSG7 for protein expression. Vectors were transformed into BL21 Gold (Stratagene) on ampicillin plates overnight and a single colony was used to inoculate a 100 mL LB flask overnight containing 50 \( \mu \)M/mL ampicillin. Cell cultures were centrifuged at 3000 \( \times \)g and the supernatant was discarded. The resultant pellet was used to inoculate a 1.5 L shaker flask of Terrific Broth with 50 \( \mu \)L of anti-cob (Sigma-Aldrich) and 50 \( \mu \)M/mL ampicillin. Cells were grown at 37°C until OD\(_{600}\) reached 0.6–0.8. Temperature was reduced to 18°C and protein expression was induced with 0.5 mM IPTG. Cells were grown overnight and harvested by centrifugation 6000 \( \times \)g at for 15 minutes at 4°C, and pellets were stored at \(-80°C\).

Cells pellets were thawed using buffer consisting of 25 mM HEPES pH 7.5, 150 mM NaCl, 10 mM imidazole, 5% glycerol, DNase and protease inhibitor tablets (Roche). Cells were sonicated and cell lysate was separated into soluble and insoluble fractions using high-speed centrifugation. The soluble fraction was filtered then nicked purified, buffer exchanged, and separated using an S200 gel filtration column on an AKTAexpress™ (GE Health-Care). If necessary, protein and storage buffers were chelated by Chelex-100 to remove bound calcium (Bio-Rad Laboratories). Purified proteins were concentrated to \( \sim \)100 \( \mu \)M, frozen, and stored at \(-80°C\).

#### Integrin binding assay

Costar ELA/RIA stripwell high binding plates were coated with either 2 \( \mu \)g of purified \( \alpha V \beta 5 \) integrin or 2 \( \mu \)g of purified \( \alpha V \beta 3 \) integrin (Millipore) for the experimental wells, or the molar equivalent of BSA for the control wells. Wells were blocked with 1% BSA in PBS for one hour at room temperature. Plates were washed 1 \( \times \) with 300 \( \mu \)L of PBS. Wells were then blocked with 1% BSA in PBS for one hour at room temperature. Plates were washed 1 \( \times \) with PBS and 100 \( \mu \)g/L of PilY1 (Wild Type, D621A, D619-621, D608A, D608K, D859A, D859K, D608A/D859A, D608A/D859K, D608K/D859A, D608K/D859K, D608A/D859A/D859K, D608A/D859K/D859A/D859K mutants) in Terrific Broth with 50 \( \mu \)L/mL of antifoam (Sigma-Aldrich) and 50 \( \mu \)M/mL ampicillin. The resultant pellet was used to inoculate a 1.5 L shaker flask of Terrific Broth with 50 \( \mu \)L of anti-cob (Sigma-Aldrich) and 50 \( \mu \)M/mL ampicillin. Cells were grown at 37°C until OD\(_{600}\) reached 0.6–0.8. Temperature was reduced to 18°C and protein expression was induced with 0.5 mM IPTG. Cells were grown overnight and harvested by centrifugation 6000 \( \times \)g at for 15 minutes at 4°C, and pellets were stored at \(-80°C\).

Cells pellets were thawed using buffer consisting of 25 mM HEPES pH 7.5, 150 mM NaCl, 10 mM imidazole, 5% glycerol, DNase and protease inhibitor tablets (Roche). Cells were sonicated and cell lysate was separated into soluble and insoluble fractions using high-speed centrifugation. The soluble fraction was filtered then nicked purified, buffer exchanged, and separated using an S200 gel filtration column on an AKTAexpress™ (GE Health-Care). If necessary, protein and storage buffers were chelated by Chelex-100 to remove bound calcium (Bio-Rad Laboratories). Purified proteins were concentrated to \( \sim \)100 \( \mu \)M, frozen, and stored at \(-80°C\).

**Figure 5. Calcium effects on RGD binding proteins.**

doi:10.1371/journal.pone.0029629.g005

| Protein | Binds calcium | Known calcium binding effects | RGD binding integrin target(s) [17] |
|---------|---------------|-------------------------------|----------------------------------|
| vWF [49] | unknown | impedes unfolding/protect from cleavage | \( \alpha V \beta 3 \) and \( \alpha I \beta 3 \) |
| Thrombospondin [50] | yes | needed for integrin binding | \( \alpha V \beta 3 \), \( \alpha I \beta 3 \), \( \alpha V \beta 6 \), \( \alpha V \beta 1 \), \( \alpha V \beta 1 \), and \( \alpha I \beta 1 \) |
| Fibrinectin [46] | yes | unknown | \( \alpha V \beta 3 \), \( \alpha V \beta 5 \), \( \alpha V \beta 6 \), \( \alpha V \beta 1 \), \( \alpha V \beta 1 \), and \( \alpha I \beta 1 \) |
| Osteopontin [52, 53] | yes | increased cell adhesion | \( \alpha V \beta 3 \), \( \alpha V \beta 5 \), \( \alpha V \beta 6 \), \( \alpha V \beta 1 \), \( \alpha V \beta 1 \), and \( \alpha I \beta 1 \) |
| Fibrinogen [47, 49] | yes | affects folding | \( \alpha V \beta 3 \) and \( \alpha I \beta 3 \) |
| Fibrillin [48] | yes | helps avoid degradation | \( \alpha V \beta 3 \) |
| Tenascin [54] | unknown | activates cell-adhesion | \( \alpha V \beta 3 \) and \( \alpha I \beta 1 \) |
| Vitronectin [51] | unknown | increased cell adhesion | \( \alpha V \beta 3 \), \( \alpha V \beta 5 \), \( \alpha V \beta 1 \), and \( \alpha I \beta 1 \) |
D851A/D859A, D608A/D855A/D859A, or D608A/D851A/D855A/D859A in binding buffer (50 mM Tris pH 7.8, 100 mM NaCl 2 mM CaCl₂, 1 mM MgCl₂, 1 mM MnCl₂, and 0.1% Triton) was added to each well including the BSA control wells for 6 hours at 16 °C with or without RGD peptide (inhibitor), RADSP peptide (control), or no peptide. Metals were added in the binding buffer for integrin structural integrity. Plates were then washed 3× with PBST and 3× with PBS before adding rabbit anti-rabbit (Invitrogen) 1:5000 in 1% BSA in PBS protected from light. Melting values and the errors were compounded using the equation

\[ \text{experimental SEM}^2 = \text{(control SEM)}^2 + \text{(experimental SEM)}^2 \]

**Calcium binding assay**

A binding curve for Oregon Green® 488 BAPTA-5N, hexapotassium salt (Invitrogen) in 25 mM HEPES pH 7.5, 250 mM NaCl, 10 mM imidazole, 5% glycerol was measured on a PHERAstar (BMG LabTech). The experimental and control well averages and standard error of the mean (SEM) were calculated, and then the control average was subtracted from the experimental values and the errors were compounded using the equation

\[ \text{experimental SEM}^2 = \text{(control SEM)}^2 + \text{(experimental SEM)}^2 \]

**Circular dichroism and thermal denaturation**

Protein samples were buffer exchanged into chelated 10 mM K2HPO4 50 mM NaF pH 7.7 buffer and brought to 50 mM with or without the addition of CaCl₂. A wavelength scan from 200–260 nm was performed on a Circular Dichroism Spectrometer 620 DS (Aviv) at 16°C with a 10 second averaging time. Melting temperatures were measured at λ 214 from 3°C to 95°C at one degree increments with a 10 second averaging time.

**Supporting Information**

**Figure S1** Circular dichroism for calcium binding mutants. A-E Molar ellipticity values were calculated for the respective wavelength scans and compared. (TIFF)

**References**

1. Richards MJ, Edwards JR, Culver DH, Gaynes RP (1999) Nosocomial infections in pediatric intensive care units in the United States. National Nosocomial Infections Surveillance System. Pediatrics 103: 339–345.
2. Klevens RM, Edwards JR, Richards CL, Jr., Horan TC, Gaynes RP, et al. (2007) Estimating health care-associated infections and deaths in U.S. hospitals, 2002. Public Health Rep 122: 160–166.
3. Schaberg DR, Culver DH, Gaynes RP (1991) Major trends in the microbial etiology of nosocomial infection. Am J Med 91: 725–738.
4. Apidianakis Y, Rahme LG (2009) Drosophila melanogaster as a model host for studying Pseudomonas aeruginosa infection. Nat Protoc 4: 1285–1294.
5. Starkey M, Rahme LG (2009) Modeling Pseudomonas aeruginosa pathogenesis in plant hosts. Nat Protoc 4: 117–124.
6. Holy F, Ciofu O, Bjarnsholt T (2010) Pseudomonas aeruginosa biofilm formation and cell culture. Microbiol Pathog 33: 109–114.
7. Lallebou E, Kim ET, Kim KC (2002) Identification of Pseudomonas aeruginosa flagellin as an adhesin for Muc1 mucin. Am J Physiol Lung Cell Mol Physiol 282: L751–756.
8. Mohn TF, O’Toole GA (2001) Mechanisms of biofilm resistance to antimicrobial agents. Trends Microbiol 9: 34–39.
9. Poole K (2004) Efflux-mediated multidrug resistance in Gram-negative bacteria. Clin Microbiol Infect 10: 12–26.
15. Goldman MJ, Wilson JM (1995) Expression of alpha v beta 5 integrin is necessary for efficient adenovirus-mediated gene transfer in the human airway. J Virol 69: 5951-5958.

16. Gravelle S, Barnes R, Havidon N, Shewchuk L, Ejl J, et al. (2010) Up-regulation of integrin expression in lung adenocarcinoma cells caused by bacterial infection: in vitro study. Innate Immun 16: 14-26.

17. Humphries JD, Byron A, Humphries MJ (2006) Integrin ligands at a glance. Journal of cell science 119: 3901-3903.

18. Swatrzell LJ, Edelmann RE, Makaroff CA, Kis JZ (1999) Integrin-like proteins are localized to plasma membrane fractions, not plastids, in Arabidopsis. Plant Cell Physiol 40: 173-183.

19. Sheppard D (2004) Roles of alphav integrins in vascular biology and pulmonary pathology. Curr Opin Cell Biol 16: 552-557.

20. Nasiil X, Beretti LJ, Loew, J, Stenberg P, O’Gara A, et al. (1994) Roles of pili and PIC in adhesion of Neisseria meningitidis to human epithelial and endothelial cells. Proc Natl Acad Sci U S A 91: 3769-3773.

21. Kalittrum H, Liszczewki MK, Atkinson JP, Jonsson AB (1997) Membrane cofactor protein (MCP or CD46) is a cellular pilus receptor for pathogenic Neisseria. Mol Microbiol 25: 639-647.

22. Kirschner M, Heuer D, Meyer TF (2005) CD46-independent binding of neisserial type IV pilus and the major pilus adhesin, PIC, to human epithelial cells. Infect Immun 73: 3072-3082.

23. Orani J, Johnson MD, Coggan KA, Sperlaaza JR, Heiniger RW, et al. (2010) Crystal structure analysis reveals Pseudomonas PilY1 as an essential calcium-dependent regulator of bacterial surface motility. Proc Natl Acad Sci U S A 107: 1065-1070.

24. Kuchma SL, Ballok AE, Merritt JH, Hammond JH, Lu W, et al. (2010) Cyclic-di-GMP-mediated repression of swarming motility by Pseudomonas aeruginosa: the pilY1 gene and its impact on surface-associated behaviors. Journal of bacteriology 192: 2950-2964.

25. Hearing RW, Winther-Larsen HC, Pickles RJ, Koomey M, Wolfgang MC (2010) Infection of human mucosal tissue by Pseudomonas aeruginosa requires sequential and mutually dependent virulence factors and a novel pilus-associated adhesin. Cell Microbiol 12: 1158-1173.

26. Higgins DG, Bidstr A, Fuchs R (1992) CLUSTAL V: improved software for multiple sequence alignment. Comput Appl Biosci 8: 189–191.

27. Thompson JD, Higgins DG, Gibson TJ (1994) CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res 22: 4673-4680.

28. Rigden DJ, Galperin MY (2004) The DxDxDG motif for calcium binding: multiple structural contexts and implications for evolution. J Mol Biol 343: 971-984.

29. Chattopadhyaya R, Meador WE, Means AR, Quiocio FA (1992) Calmodulin structure refined at 1.7 A resolution. J Mol Biol 228: 1177-1192.

30. Weber C, Lee VD, Chazin WJ, Huang B (1994) High level expression in Escherichia coli and characterization of the EF-hand calcium-binding protein structure refined at 1.7 A resolution. J Mol Biol 228: 1177-1192.

31. Manning SA, Hamilton SM, Karplus PA, Caffetti LM (2008) The Arg-Gly-Asp-containing, solvent-exposed loop of PyTonA is required for internalization. Mol Plant Microbe Interact 21: 315–325.

32. Wang FZ, Akula SM, Sharma-Walia N, Zeng L, Chandran B (2003) Human herpesvirus 8 envelope glycoprotein B mediates cell adhesion via its RGD sequence. J Virol 77: 3111–3117.

33. Xiao J, Palefsky JM, Herrera B, Berline TJ, Tugizov SM (2008) The Epstein-Barr virus BRLF2 protein facilitates virus attachment to oral epithelial cells. Virology 370: 430–442.

34. Chittukuntawar AV, Zhou X, Rajayya J, Choobesh J (2010) Viral capsid is a pathogenicity-associated molecular pattern in adenovirus keratitis. PLoS Pathog 6(10): e1000841.

35. Harvala H, Kalimo H, Stanway G, Hyytiä T (2003) Pathogenesis of coxsackievirus A9 in mice: role of the viral arginine-glycine-aspartic acid acid motif. J Gen Virol 84: 2375-2379.

36. Amphlett GW, Hrnida ME (1983) The binding of calcium to human fibronectin. Biochim Biophys Acta 747: 131-140.

37. Reinhardt DP, Oso RN, Sakai LY (1997) Calcium stabilizes fibrillin-1 against proteolytic degradation. J Biol Chem 272: 1231–1236.

38. Harfenist EJ, Packham MA, Kinloch-Rathbone KL, Cattano M, Mustard JF (1995) Effect of calcium ions on concentration of the ability of fibronectin and von Willebrand factor to support the ADP-induced aggregation of human platelets. Blood 80: 827-831.

39. Lawler J, Weinstein R, Hynes RO (1988) Cell attachment to thrombospondin: the role of ARG-GLY-ASP, calcium, and integrin receptors. J Cell Biol 107: 2351-2361.

40. Fusi FM, Bernocchi N, Ferrari A, Bronson RA (1996) Is vitronectin the velcro that binds the gametes together? Mol Hum Reprod 2: 859-866.

41. He B, Mirza M, Weber GF (2008) An osteopontin splice variant induces anchorage independence in human breast cancer cells. Oncogene 25: 2192-2202.

42. Singh K, Deonarine D, Shanmugam V, Senger DR, Mukherjee AB, et al. (1993) Calcium-binding properties of osteopontin derived from non-osteogenic sources. J Biochem 114: 702-707.

43. Slater M, Murphy CR, Barden JA (2002) Taraxacum, E-cartharin and P2X calcium channel receptor expression is increased during rat blastocyst implantation. Histochem J 34: 15-19.

44. Matsick JS (2002) Type IV pilus and twitching motility. Annu Rev Microbiol 56: 289-314.