Crystallographic Analysis of the Pseudomonas aeruginosa Strain K122-4 Monomeric Pilin Reveals a Conserved Receptor-Binding Architecture†‡

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ABSTRACT: Adherence of pathogens to host cells is critical for the initiation of infection and is thus an attractive target for anti-infective therapeutics and vaccines. In the opportunistic human pathogen Pseudomonas aeruginosa, host-cell adherence is achieved predominantly by type IV pili. Analysis of several clinical strains of P. aeruginosa reveals poor sequence conservation between pilin genes, including the residues in the receptor-binding site. Interestingly, the receptor-binding sites appear to retain a conserved surface epitope because all Pseudomonas type IV pili recognize the same receptor on the host cell and cross-reactive antibodies specific for the receptor-binding site exist. Here, we present the crystallographic analysis of two crystal forms of truncated pilin from P. aeruginosa strain K122-4 (ΔK122-4) at 1.54 and 1.8 Å resolution, respectively. The ΔK122-4 structure is compared to other crystallographically determined type IV pilin structures and an NMR structure of ΔK122-4 pilin. A comparison with the structure of the highly divergent P. aeruginosa strain K (ΔPAK) pilin indicates that the receptor-binding loop in both pilins forms a shallow depression with a surface that is formed by main-chain atoms. Conservation of this putative binding site is independent of the sequence as long as the main-chain conformation is conserved and could therefore explain the shared receptor specificity and antibody cross reactivity of highly divergent Pseudomonas type IV pilins.

The opportunistic pathogen Pseudomonas aeruginosa is a significant cause of morbidity and mortality in clinical settings. Acute P. aeruginosa infections observed in ventilator-associated pneumonia (1–3) and burn wounds (4–6) have high mortality rates. Furthermore, in individuals with a compromised or suppressed immune system, such as patients with cancer (7), cystic fibrosis (8, 9) and HIV (10), P. aeruginosa can establish persistent infections that are frequently fatal through immune pathology, rather than the direct actions of the pathogen. Finally, the high innate and acquired antibiotic resistance of P. aeruginosa isolates complicate treatment (11, 12). Therefore, the development of anti-infective agents against targets that bypass known resistance mechanisms is of considerable interest.

P. aeruginosa infection begins with the adherence of the pathogen to the epithelium, which is primarily achieved through the type IV pilis. Consequently, type IV pilis are an important virulence factor of P. aeruginosa (13) and other Gram-negative pathogens (14, 15). Indeed, engineered strains of P. aeruginosa that lack functional pili exhibit reduced virulence (16, 17). Also, a humoral response toward P. aeruginosa pili is protective (18–20). Type IV pilis are fiber-like structures that are assembled from ~15 kDa pilin monomers (21, 22). Extensive studies on P. aeruginosa pilins have shown that each monomer contains a functional receptor-binding site within a disulfide-bound loop region (D-region; 16, 23–29). However, Lee and colleagues (27) demonstrated that receptor binding occurs only at the tip of the pilus, suggesting that the binding site is blocked during assembly except for the monomers exposed at the tip. In addition to cellular adhesion, the type IV pilus is involved in other processes such as twitching motility (30, 31), biofilm formation (32–34), induction of host-cell signals (35–37), DNA uptake during natural transformation (38), and bacteriophage infection (39).

While pilins from different P. aeruginosa strains exhibit high sequence variability (40, 41), it has been shown that they share specificity of the glycosphingolipids asialo-GM1 and asialo-GM2 that are present on the epithelial cells of the mucosal layer (27, 42). In particular, Sheth et al. (43) demonstrated that the minimal receptor recognized by the pilin is the disaccharide β-D-GalNAc(1–4)β-D-Gal, a substructure of both asialo-GM1 and asialo-GM2. Furthermore, immunization with a peptide corresponding to the D-region of the pilin yields cross-reactive antibodies (18). Antibody cross reactivity and shared receptor specificity suggest that the receptor-binding domains of all P. aeruginosa strains

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† Coordinates and structure factor amplitudes for ΔK122-4 pilin have been deposited in the RCSB Protein Data Bank with accession codes 1QVE and 1RG0 for the triclinic and monoclinic crystal forms, respectively.
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have retained a conserved surface epitope despite the poor sequence conservation.

Crystal structures of the full-length Neisseria gonorrhoeae strain MS11\(^{1}\) (44) and P. aeruginosa strain K (PAK; 45) pilins have been reported, but structure determination of full-length pilins is complicated by poor solubility. Full-length pilins contain a long N-terminal \(\alpha\)-helix (\(\alpha1\)). The first 28 residues (\(\alpha1-N\)) are exposed and highly hydrophobic, whereas residues 29–54 (\(\alpha1-C\)) pack onto the globular head domain of the pilin. Truncation of \(\alpha1-N\) results in a soluble pilin monomer that retains the receptor-binding characteristics of the intact pilin (41; Irvin, R. T., unpublished data). To date, crystal structures of the truncated PAK pilin (\(\Delta\)PAK; 40) and Vibrio cholerae toxin-coregulated pilin (45) have been reported, as well as an NMR structure of truncated pilin crystallized in the triclinic form but in the presence of the receptor analogue GalNAc(1-\(\beta\)-D-Gal-OMe) in 10 mM Tris at pH 7.4 (41).

However, the region of greatest interest, the C-terminal receptor-binding domain, could not be unambiguously assigned in the NMR study because of spectral overlap (41).

In this paper, we present the structures of two crystal forms of the \(\Delta\)K122-4 pilin refined to 1.54 and 1.8 Å resolution, respectively. We compare the \(\Delta\)K122-4 structure with other crystallographically determined type IV pilin structures and with the structure of \(\Delta\)K122-4 previously determined using NMR spectroscopy (\(\Delta\)K122-4\(^{\text{NMR}}\); 41). This presents the first opportunity to compare the receptor-binding sites of two highly divergent pilins that share receptor specificity (the V. cholerae pilin has no equivalent binding site, and to our knowledge, receptor specificity for MS11 pilin has not been reported). The comparison supports our earlier hypothesis that the type IV pilins of P. aeruginosa have a binding site consisting predominantly of main-chain atoms. Conservation of function is therefore independent of sequence as long as the main-chain conformation is retained.

### EXPERIMENTAL PROCEDURES

#### Crystallization and Data Collection

Purification, crystallization, and X-ray diffraction data collection for the triclinic form of \(\Delta\)K122-4 pilin has been reported previously (46). Briefly, \(\Delta\)K122-4 [\(\text{pilA}(\Delta1–28); 41\)] was expressed periplasmically in Escherichia coli as a maltose-binding-protein (MBP) fusion protein and purified using an amylose column (46). The purified MBP-K122-4 fusion protein was trypsinized to release \(\Delta\)K122-4 pilin from MBP. The monomeric \(\Delta\)K122-4 pilin contains four N-terminal residues (ISEF) from the expression construct followed by residues 29–150 of K122-4 pilin. Monomeric \(\Delta\)K122-4 was then purified by cation-exchange chromatography and crystallized as described (46). The monoclinic crystal form of \(\Delta\)K122-4 pilin crystallized under conditions similar to that of the triclinic form but in the presence of the receptor analogue \(\beta\)-d-GalNAc(1-\(\beta\)-d-Gal-OMe). Crystals were grown from 2-μL drops containing equal volumes of protein–carbohydrate solution (20 mg ml\(^{-1}\) \(\Delta\)K122-4 and 15.7 mM \(\beta\)-d-GalNAc(1-\(\beta\)-d-Gal-OMe) in 10 mM Tris at pH 7.4 and 100 mM sodium chloride) and reservoir solution (38% w/v PEG 4000, 100 mM sodium cacodylate at pH 5.8, and 100 mM monobasic potassium phosphate). Diffraction data were collected on beamline 8.3.1 at the Advanced Light Source and were processed using MOSFLM (47) and SCALA from the CCP4 suite (48). A summary of the diffraction and refinement statistics is shown in Table 1.

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\(^{a}\) Overall/highest resolution shell. \(^{b}\) \(R_{	ext{work}}\) is the unweighted R value between symmetry mates. \(R_{	ext{work}} = \sum_{hkl}[|F_{\text{calc}}(hkl)| - |F_{\text{act}}(hkl)|] / \sum_{hkl}|F_{\text{calc}}(hkl)|\). \(R_{\text{free}}\) is the cross validation R factor using 5% of reflections.

Structure Solution and Refinement. The structures of both crystal forms of the \(\Delta\)K122-4 pilin were solved using molecular replacement (MR) and employed the AMoRe software package (49) of the CCP4 suite (48). For the triclinic crystal form, the molecular replacement search model was the globular domain of the MS11 pilin (PDB ID 1AY2; 44), from which nonconserved regions were removed. Analysis of the self-rotation function revealed a single peak 13σ above the background (\(κ = 180^\circ\)), indicating that there is a \(\Delta\)K122-4 dimer in the unit cell (46). To determine the initial positions of the two \(\Delta\)K122-4 molecules, the top rotation peak was interpreted as the first \(\Delta\)K122-4 molecule and then fixed. With the top peak fixed, the subsequent 10 rotation peaks were employed in the translation search to determine the position of the second \(\Delta\)K122-4 pilin molecule relative to the first. After the initial coordinates for both molecules were obtained, rigid-body refinement in AMoRe resulted in a correlation coefficient and R value of 0.490 and 0.468, respectively.

The MR solution was used as input for ARP/wARP (50, 51) to fill in the nonconserved regions that were removed from the MS11 search model. After removal of the dummy atoms generated by ARP/wARP, examination of an initial \(σ_A\)-weighted (52) \(2F_o - F_c\) electron density map in Xfit (53) allowed for the building of all residues with the exception of Ile 25 and Ser 26 of chain A and Thr 149 and Pro 150 of chain B. These residues, with the exception of Pro 150B, were located during subsequent rounds of refinement. Residue 36 has been reported as Arg in the sequence database (gi|77636; 54). However, our sequencing demonstrated that residue 36 was in fact an Ala. Alanine is also more consistent with the preference for a small residue at this position in other pilin sequences (40), and an Arg would be incompatible because of steric constraints. Accordingly, we feel that an Ala is the correct residue at this position. The model was

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\(^{1}\) Abbreviations: \(\Delta\)K122-4, the truncated Pseudomonas aeruginosa strain K122-4 pilin; \(\Delta\)K122-4\(^{\text{NMR}}\), the truncated K122-4 pilin solved using NMR; MS11, the Neisseria gonorrhoeae strain MS11 pilin; PAK, the P. aeruginosa strain K pilin; \(\Delta\)PAK, the truncated PAK pilin; rmsd, root-mean-square deviation; NCS, noncrystallographic symmetry; NMR, nuclear magnetic resonance spectroscopy.
refined using REFMAC (55), and TLS parameters (56) were refined prior to refining atomic positions and isotropic B factors. All data were used during the refinement, with 1336 reflections being set aside for $R_{	ext{free}}$ estimation. Engh and Huber (58) stereochemical restraints and loose noncrystallographic symmetry (NCS) restraints were employed throughout the refinement. Ronds of refinement were followed by model building in Xfit using $\sigma_A$-weighted density maps. Solvent atoms were located using ARP/wARP and confirmed with visual inspection. Several residues were modeled with alternative side-chain conformations based on the shape of the electron density; the occupancies of these alternative conformations were refined using SHELXL (59). Restrained individual anisotropic $B$ values were refined in the final cycle. The validity of the anisotropic B value refinement at this resolution was supported by the $R_{	ext{free}}$ statistic, which dropped from 18.1 to 17.8%. The refinement statistics are summarized in Table 1.

The structure of the monoclinic form of the ΔK122-4 pilin was solved using the refined structure of triclinic ΔK122-4 pilin (protein only) as the MR search model. The correlation coefficient and $R$ value following molecular replacement were 0.721 and 0.312, respectively. The model was refined using a similar strategy as that for the triclinic crystal, with 661 reflections being set aside for $R_{	ext{free}}$ calculation and $\sigma_A$ estimation. Occupancies for residues with alternative conformations were refined using SHELXL. A summary of refinement statistics is shown in Table 1.

RESULTS AND DISCUSSION

Quality of the Final Models. The structure of the triclinic crystal form of ΔK122-4 pilin was refined to 1.54 Å resolution, with a NCS-related pilin dimer (chains A and B) in the P1 unit cell. $R_{	ext{work}}$ and $R_{	ext{free}}$ were refined to 13.8 and 17.6%, respectively. In the final model, all residues except the C-terminal residue (Pro 150) of chain B were observed and 289 water molecules could be modeled in the electron density. Five residues were modeled with dual side-chain conformations. These residues are Ser 34, Thr 45, Ser 48, and 289 water molecules could be modeled in the electron density and may exhibit some disorder that could not be modeled at the present resolution. Diffraction and refinement statistics for both crystal forms are summarized in Table 1.

The four ΔK122-4 monomers in our crystal forms show minimal differences, with root-mean-square deviations (rmsd’s) between all Cα positions ranging from 0.22 to 0.67 Å. Unless otherwise specified, all analyses presented below are based on chain A of the high-resolution triclinic crystal form, which is most clearly defined by its electron density. Both crystal forms of ΔK122-4 contain equivalent NCS-related dimers in the asymmetric unit. Pilin dimers have also been observed in the crystal structures of MS11 (44) and PAK (45); however, these dimers are distinct from each other and from our ΔK122-4 dimers. Also, there is no evidence for a functional role for type IV pilin dimers, and dimerization of ΔK122-4 was not observed in NMR or equilibrium centrifugation studies (41). Therefore, the observed ΔK122-4 dimers appear to have no biological relevance and likely arise because of crystallization conditions.

Common Architecture of the Type IVa Pilins. The structures of four type IV pilins have been described in the literature. Three have been examined crystallographically: the N. gonorrhoeae strain MS11 pilin (44), the truncated toxin-coregulated pilin from V. cholerae (45) and the P. aeruginosa strain K pilin, both as truncated (ΔPAK; 40) and full-length (PAK; 45) pilins. The fourth type IV pilin, the ΔK122-4 pilin, has been examined via NMR methods (ΔK122-4NMR; 41). The toxin-coregulated pilin, classified as a type IVb pilin, differs significantly in size, leader sequence, monomer sequence, and structure from the Pseudomonas and Neisseria type IVa pilins (45, 62). Accordingly, it is not included in the present analysis.

The ΔK122-4 crystal structure exhibits the characteristic type IVa pilin fold, with the N-terminal α-helix (α1-1C) packed onto a four-stranded antiparallel β-sheet (Figure 1a). The relative positions of the core secondary structure elements are well-conserved among the crystal structures (Figure 1b), and structural superimposition gives Cα rmsd values of 1.0–1.4 Å (Table 1). Interestingly, ΔK122-4 most closely resembles MS11 rather than ΔPAK even though MS11 pilin is of Neisserial rather than Pseudomonal origin. A closer phylogenetic relationship between K122-4 and MS11 pilin is also evident at the amino acid sequence level with 46.1% sequence identity between K122-4 and MS11.
versus 38.8% sequence identity between K122-4 and PAK (these values are 36.3 and 26.1%, respectively, for the globular domain sequence starting at residue 29; Table 2). These observations indicate that the large sequence variations seen between pilin genes in different P. aeruginosa isolates originate at least in part from horizontal gene transfer.

Surprisingly, a global rmsd could not be calculated between the ΔK122-4 structures determined by crystallography and NMR because of differential packing of the helix and sheet in the two structures (Figure 2). If we first superimpose the average ΔK122-4NMR structure onto ΔK122-4 based on their β-sheets, 37 residues superimpose with an rmsd of 1.2 Å (Table 2). An additional 44.5° rotation and 3.9 Å translation are required to subsequently superimpose the α helices. Analysis of the 10 individual ΔK122-4NMR models in the NMR ensemble gave basically the same results. It has been speculated that the disulfide bond between Cys 57 and Cys 93 may have caused the different helix–sheet packing (41), but this disulfide bond in both X-ray and NMR structures is actually well-conserved (Figure 2). Because neither cysteine is part of the α-helix, it is not obvious from the structure how this disulfide would change the helix packing.

In ΔK122-4NMR, the α-helix is shifted down by one turn and is deflected away from the β-sheet, especially at the N-terminal end of the α-helix (Figure 2). As a consequence, the packing of the hydrophobic core is less tight in ΔK122-4NMR and a conserved hydrogen bond between the Oε of Gln 32 and the amide nitrogen of Ala 105 is missing. Because all crystal structures, with several determined at high resolution, reveal a consistent helix–sheet packing, we believe that they represent an accurate and relevant conformation of the pilin. The deviation observed in ΔK122-4NMR could be due to one or more incorrect NOE assignments. However, a comparison of the NMR restraint data with both the NMR and X-ray structures suggests that this is not the case. There are also data that indicate structural flexibility of the pilin monomer. Molecular models of type IV pili generated using the crystal structures of PAK and MS11 only explain X-ray fiber diffraction patterns when the globular domain is omitted from calculations of simulated diffraction patterns (63). This suggests a disorder of the globular domain relative to the helical backbone. Furthermore, Neisseria pilins can be proteolytically cleaved in vivo after residue 39 to release a soluble pilin monomer (64). Residue 39 is in the middle of the α-helix (α1), and this peptide bond would not be accessible for proteolytic cleavage when α1 is tightly packed against the β-sheet as seen in the crystal structures. It is therefore possible that under physiological conditions the β-sheet is able to move relative to the α-helix, whereas the high concentrations of precipitants used for crystallization may have stabilized the tighter packing. Further research will be required to directly address this question.

The loop connecting α1-C to the β-sheet (the αβ-loop) is highly variable in sequence and structure. In PAK, a small β-sheet follows α1-C, whereas in MS11, the equivalent space is occupied by a single helical turn that is glycosylated at Ser 63. In models for the type IV pilus fiber (40, 44, 62), these structural features block the receptor-binding site of a pilin monomer in the preceding turn of the pilus fiber. It has been proposed that this may explain why receptor binding only occurs at the tip of the pilus (40). ΔK122-4 pilin does not have the small β-sheet of PAK pilin, and while Thr 64 of ΔK122-4 is within 5 Å of the glycosylated Ser 63 of MS11, P. aeruginosa strain K122-4 pilin is not glycosylated (65; Irvin, R. T., unpublished results). Therefore, the loop of ΔK122-4 does not appear to provide a bulky group that can block receptor-binding sites along the flanks of the pilus. Two alternative, though speculative, mechanisms for occluding the receptor-binding domain in the pilus fiber can be envisioned. The first involves the C-terminal extension. While other P. aeruginosa pilins have only two to three residues following the C-terminal cysteine, K122-4 has eight residues.
Interestingly, the a somewhat different relative orientation of the a turn that exposes only main-chain atoms, and the relative orientation of the two a turns is such that a shallow solvent-exposed pocket is formed that is dominated by main-chain atoms (40; Figures 1 and 3). Several carbonyl oxygen and amide nitrogen atoms of the main chain point up into the pocket and could potentially act as hydrogen-bonding partners for hydroxyl groups of the carbohydrate ligand. This led to the hypothesis that the pocket formed by the two a turns forms a unique main-chain-atom-dominated receptor-binding site (40). Shared receptor specificity and antibody cross reactivity by highly divergent pilins could then result from conservation of the main-chain structure because of evolutionary constraints.

To better understand the conformational determinants of the D-region, we looked at conserved properties of the P. aeruginosa pilin sequences. There are seven strictly conserved residues in the globular domain of Pseudomonas pilins (40); in K122-4, these residues are Arg 30, Lys 44, Thr 98, Trp 127, Cys 129, Pro 139, and Cys 142. Within the D-region, only the two cysteines that form the disulfide and proline 139 that starts the second a turn are strictly conserved. Disulfide bonds and proline residues restrict the conformational flexibility of peptides and therefore likely play a structural role in defining the conformation of the D-region. Studies have indeed shown the importance of the disulfide bond for structure and function of synthetic receptor-binding loop peptides (28, 29, 66–69). In the pilin structures, the side chains of the cysteine and proline residues are buried between the loop and the core of the protein. This also points to a structural rather than a functional role because the residues cannot directly interact with the receptor but instead define the loop–protein packing interface. Tyr 137 also contributes to this packing interface, and an aromatic amino acid is conserved at this position in the loop (40). Furthermore, in the AK122-4 structure, the strictly conserved Arg 30 stabilizes the D-region through hydrogen bonds between its N° and N°° atoms and the carbonyl oxygen of Lys 136 (Figure 3a). The conserved Trp 127 and Lys 44 interact with each other, forming a hydrophobic stacking interaction between the a-helix and a-sheet. This interaction is important to stabilize the globular domain structure but should not directly influence the D-region structure. Finally, Thr 98 is exposed to the solvent and not close to the D-region. Its

The region was first studied by NMR using synthetic peptides with sequences derived from several highly divergent P. aeruginosa strains (28, 65, 66). These studies showed that all peptides contained two a turns and a type I turn (residues 134–137) followed by a type II turn (residues 139–142). NMR studies of a D-region peptide in complex with a receptor analogue or a cross-reactive antibody indicated little structural rearrangement upon complex formation (29). Apparently, these peptides have an “intrinsic propensity” to adopt a native-like conformation despite the high sequence divergence.

The APAK crystal structure confirmed the presence of the two a turns in the D-region with the main difference being a somewhat different relative orientation of the a turns. Interestingly, the a turn geometry creates one face of the...
FIGURE 3: Receptor-binding domain of the type IVa pilins. (a) Stereoview of the ΔK122-4 receptor-binding domain (boxed in Figure 1a). All residues of the receptor-binding loop and the conserved Arg 30 are shown in a ball-and-stick representation. (b) Stereorepresentation of the superimposed receptor-binding loop of ΔK122-4, ΔPAK, and MS11 pilins (boxed in Figure 1b). Backbone atom colors are yellow, magenta, and orange, respectively. The two β turns are labeled I and II, and the peptide flip at residue 141 in ΔPAK is labeled with an asterisk. (c) CPK representation of the PAK and K122-4 D-regions showing the surface-exposed atoms. Main-chain atoms of the residues that form the two β turns are in darker colors, highlighting the conserved surface of the proposed receptor-binding site. The peptide flip has again been highlighted with an asterisk.
role in the structure or function of pilin, if any, is at this moment unclear.

The D-region of ΔK122-4 exhibits, as expected, a main-chain structure that is largely conserved when compared with that of ΔPAK (Figure 3). The main difference between ΔPAK and ΔK122-4 is a flip of the peptide plane between residues 140 and 141 in ΔK122-4. This changes the type II β turn into a type III β turn. In ΔPAK, residue 141 adopts a distorted left-handed helical conformation that is only allowed for glycine (present in PAK, PAO, and KB7 pilins) and, to a lesser extent, asparagine (present in P1 pilin; 40). In contrast, the threonine present in K122-4 and the Neisseria pilins is not compatible with a left-handed helical conformation, resulting in the peptide flip. This is a very local structural change that may not affect receptor binding if the peptide plane carbonyl oxygen does not contact the receptor. It is also of interest to note that a water molecule in optimal hydrogen-bonding geometry with the main-chain nitrogen of Gly 141 in ΔPAK would have a position that coincides with the carbonyl oxygen of Lys 140 in ΔK122-4. Accordingly, an interaction by the carbonyl oxygen in ΔK122-4 could be replaced by a water-mediated interaction in ΔPAK. Finally, although Thr 141 prevents ΔK122-4 from adopting a ΔPAK-like β turn, it is possible that ΔPAK can adopt a type III β-turn conformation upon receptor binding. The structure of a pilin in complex with a carbohydrate analogue is required to delineate the receptor-binding determinants of the binding pocket and to fully evaluate the impact of the peptide flip on receptor binding.

The ΔPAK and ΔK122-4 receptor-binding pockets also show one difference with respect to amino acid side chains. Whereas the binding pocket is devoid of any side chains in the ΔPAK structure, the pocket is occupied by the side chain of Gln 143 in the ΔK122-4 structure (Figure 3b). However, Gln 143 is part of the C-terminal extension that was found to be highly flexible in the NMR structure (41). Indeed, in the NMR ensemble of structures, Gln 143 typically does not block the receptor-binding pocket. It is tempting to hypothesize that the NMR structure resembles a tip-exposed binding site, whereas the crystal structure reflects the situation for nontip-exposed monomers where protein–protein interactions cause the C terminus to block the binding pocket. However, because the crystal-packing interface will differ from the protein–protein interactions in the pilus fiber, this hypothesis remains to be confirmed.

We have solved the crystal structure of ΔK122-4 pilin to determine the features in the receptor-binding site that have been conserved between the ΔK122-4 and ΔPAK pilins. We found that the main-chain structure of the D-region was conserved between both pilins. This supports our earlier hypothesis that highly divergent P. aeruginosa type IV pilins share a receptor-binding site that is formed by main-chain atoms and that receptor specificity and antigenic characteristics are conserved by conservation of the main-chain structure. Interestingly, the D-region in Neisseria MS11 pilin is very similar to especially the ΔK122-4 pilin (Figure 3b) even though it has to accommodate a 17-residue insertion. This suggests that the D-region in Neisseria and Pseudomonas may share a common function. Such a function does not have to be limited to receptor binding because type IVa pilin is associated with many other functions such as twitching motility, biofilm formation, and DNA binding (31, 33, 62). A potential role for the D-region in these additional pilus-associated functions is currently under investigation.

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