Membrane and Chaperone Recognition by the Major Translocator Protein PopB of the Type III Secretion System of Pseudomonas aeruginosa*

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Background: The type III secretion system allows bacteria to inject effectors directly into the host cytoplasm through a translocation pore.

Results: Translocator proteins were characterized in lipid-bound and chaperone-associated forms.

Conclusion: Pseudomonas translocators use different regions to recognize their common chaperone and the eukaryotic membrane.

Significance: Mapping of key regions in translocators provides a starting point for the potential development of novel antibacterials.

The type III secretion system is a widespread apparatus used by pathogenic bacteria to inject effectors directly into the cytoplasm of eukaryotic cells. A key component of this highly conserved system is the translocon, a pore formed in the host membrane that is essential for toxins to bypass this last physical barrier. In Pseudomonas aeruginosa the translocon is composed of PopB and PopD, both of which before secretion are stabilized within the bacterial cytoplasm by a common chaperone, PcrH.

In this work we characterize PopB, the major translocator, in both membrane-associated and PcrH-bound forms. By combining sucrose gradient centrifugation experiments, limited proteolysis, one-dimensional NMR, and β-lactamase reporter assays on eukaryotic cells, we show that PopB is stably inserted into bilayers with its flexible N-terminal domain and C-terminal tail exposed to the outside. In addition, we also report the crystal structure of the complex between PcrH and an N-terminal region of PopB (residues 51–59), which reveals that PopB lies within the concave face of PcrH, employing mostly backbone residues for contact. PcrH is thus the first chaperone whose structure has been solved in complex with both type III secretion systems translocators, revealing that both molecules employ the same surface for binding and excluding the possibility of formation of a ternary complex. The characterization of the major type III secretion system translocon complex in both membrane-bound and chaperone-bound forms is a key step for the eventual development of antibacterials that block translocon assembly.

Type III secretion systems (T3SSs) are complex macromolecular nanomachines that play key roles in the infectivity processes of a number of Gram-negative bacteria. These complex systems harbor >20 proteins that associate into a needle-like structure located directly on the bacterial surface. T3SSs encompass both bacterial membranes and serve as conduits to export specific effectors directly from the bacterial cytosol into target eukaryotic cells (1). Because the internal diameter of the needle is on the order of 25 Å, it is believed that effectors must travel in a semi-unfolded state (2, 3) due to energy provided by an ATPase located at the base of the system (4–6). Introduction of point mutations into key proteins of the T3SS have shown to be highly deleterious for the cytotoxicity potential of certain pathogens (7–11), and T3SS-deficient strains display attenuated infectivity in animal models (12–14), indicating that an understanding of T3SS formation and regulation mechanisms could lead to the development of strategies for infection control.

T3SS can be divided into three main sections: 1) a ring-like base structure that encompasses both bacterial membranes and bypasses the periplasm, 2) a hollow needle formed through the polymerization of a single protein through which effectors are thought to travel, and 3) the translocon, a pore formed on the target bilayer by two T3SS-translocated membrane proteins and one hydrophilic partner. The base and the needle are collectively known as the injectisome and have been elegantly

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3 The abbreviations used are: T3SS, type III secretion system; PS, α-phosphatidyl-d-serine; PC, 1,2-dipalmityl-sn-glycero-3-phosphocholine; β-Lac, β-lactamase reporter; RT, room temperature; DDM, n-Dodecyl-β-D-maltopyranoside; CMC, critical micelle concentration; MOI, multiplicity of infection.
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characterized through a combination of structural biology, biochemistry, and microbiology techniques (3, 10, 15–21). The translocon, however, remains less well studied despite the fact that it is essential for T3SS functionality (22–24).

All three proteins that compose the translocon are encoded in a single operon that also carries genes that code for chaperones for the different molecules. In Pseudomonas aeruginosa, for example, the translocon is composed of membrane proteins PopB and PopD (chaperoned by PcrH) and the hydrophilic partner PcrV (chaperoned by PcrG), all of which are encoded within the genetic element pcrGVHpopBD (25). PopB is the “major translocator,” a 390-residue protein with two putative transmembrane regions, whereas PopD, the “minor translocator,” carries 295 amino acids and a single predicted transmembrane domain and makes a 1:1 complex with PcrH (26). The Yersinia proteins YopB and YopD display 64 and 60% sequence similarity with their counterparts from Pseudomonas, whereas Ipab and IpaC from Shigella spp. are considerably longer, harboring 580 and 363 amino acids, respectively. All major translocators studied to date display at least one predicted coiled coil region and two transmembrane domains, whereas minor translocators are considerably shorter molecules, mostly with one predicted coiled coil and transmembrane region (22). Understanding how translocon molecules interact with their respective chaperones within the bacterial cytoplasm and associate to the target membrane upon passage through the needle has been the objective of a number of laboratories, as both aspects have been shown to play key roles in bacterial pathogenesis.

P. aeruginosa is a major human pathogen that targets immuno-depressed and cystic fibrosis patients. Its type III secretion system plays an essential role in the initial phases of infection (12, 24, 27). In this work we characterize its major translocator PopB through lipid binding studies, nuclear magnetic resonance, x-ray crystallography, and cellular toxicity assays. The flexible N-terminal region does not become inserted into membranes upon lipid recognition and has characteristics of a molten globule, which could be important for its unfolding and passage through the needle. Notably, the high resolution structure of a complex between a peptide located within the N terminus of PopB and PcrH reveals that central hydrophobic residues of the peptide are important for chaperone recognition and excludes the formation of a 1:1:1 complex between PcrH and the two translocator molecules. These results shed light on common features of many of the characteristics of major translocon proteins in bacterial pathogens.

**EXPERIMENTAL PROCEDURES**

**Liposome Preparation and Sucrose Gradients**—Liposomes were prepared by initially preparing a lipidic biofilm using a 2:3 PS:PC (α-phosphatidyl-l-serine:1,2-dipalmitoyl-sn-glycero-3-phosphocholine) mixture in chloroform through evaporation under nitrogen for 30 min. The lipidic film was dissolved in 3 ml of PBS, pH 7.0, and 0.7% octyl-β-D-glucopyranoside by vortexing followed by repeated freeze/thaw cycles in liquid nitrogen. The mixture was stored at −80 °C. Subsequently, the lipidic mixture was thawed and extruded, producing liposomes with an average diameter of 100 μm. 5 mg of liposomes were incubated overnight at RT with 0.8 mg of PopB or 0.8 mg of PopD.

The two different proteoliposome mixtures were then incubated on a rotating wheel at RT with different agents: 1 m KCl, 6 mM urea, 0.425 mM D-Didodecyl-β-D-maltopyranoside (DDM) or PBS, pH 7.0, as a control. After 4 h of incubation the solutions were tested on a 0–45% sucrose gradient. Tubes were ultracentrifuged at 230,000 × g for 16 h in a SW41 Ti rotor. 1-ml fractions were recovered from the top to the bottom of the gradient and analyzed by Western blots, which were developed with either anti-PopB or anti-PopD rabbit antibodies.

**Trypsin Digestion**—PopB and PopD were purified by nickel affinity chromatography and gel filtration at pH 5.0 as described in Schoehn et al. (26). Liposomes were prepared by employing an extruder (Avanti Polar Lipids) using a PS:PC:sphingomyelin for PopD and PS:PC:sphingomyelin:cholesterol for PopB, with the same ratio for all lipids. Proteoliposomes were then prepared by incubation of liposomes at RT with PopD or PopB. Both proteoliposome samples were dialyzed into PBS, pH 7.0, before trypsin digestion. Samples were subsequently proteolyzed using trypsin:protein ratios of 1:500 and 1:100, respectively, at RT. Samples were taken at different times and loaded on SDS-PAGE and subsequently analyzed by Coomassie staining or Western blotting.

**One-dimensional NMR**—The one-dimensional proton spectrum was recorded at 25 °C with excitation-sculpting water suppression (28). The experiment was performed on an Agilent Direct Drive 800 MHz spectrometer equipped with a cryogenic triple resonance probe. The experimental time was set to 17 min using 1024 scans and a recovery time of 1 s.

**Plasmid Constructs**—The pET15b-PopB/PcrH vector was obtained by first cloning the two genes separately into two distinct pET15b plasmids. popB was cloned using NdeI/BamHI sites, whereas PcrH was cloned in the BamHI site of pET15b. Subsequently, pET15b-PcrH was digested with XbaI-BamHI to maintain the ribosomal binding site and cloned blunt-ended in the BamHI blunt-ended pET15b-PopB. The pETDuet-PopD/PcrH vector was obtained by cloning popD in the Ndel-Xhol sites present in the second multiple cloning site of pETDuet1-PcrH(1–160) vector, described previously (8).

**For the in vivo experiments**, the section of P. aeruginosa CHA operon containing pcrHpopBpopD was cloned into the Ndel-HindIII restriction sites of the plApG vector. Two different PopB mutants were generated with the QuikChange II site-directed mutagenesis kit (Stratagene), namely mutants L55D/P58D (named PopB-2D) and V53D/L55D/P57D/P58D (named PopB-4D). The presence of only these mutations was verified by DNA sequencing.

**Cytotoxicity toward Macrophages**—Macrophage cell line J774 was grown in DMEM/F-12 media with 20% of FCS and antibiotics. 2.5 × 10⁶ cells were separated into aliquots in 24-well plates and washed in DMEM without antibiotics before infection with a culture of P. aeruginosa at A₅₆₂₀ = 1.0. A multiplicity of infection (MOI) of 5 was used, and 30-μl aliquots of the supernatant were taken after 1, 2, 3, and 4 h of infection for quantification of LDH release with the LDH cytotoxicity kit (Roche Applied BioScience). The value of LDH release obtained after the addition of 0.5% Triton X-100 was considered as 100% cell death, and uninfected macrophages were used as negative controls.
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Only PopB Is Stably Inserted into Membranes—We previously determined that PopB-PcrH and PopD-PcrH complexes could be purified in soluble form upon overexpression in Escherichia coli. In addition, we showed that chaperone release could be accomplished by gel filtration of the complexes at acidic pH, leading to oligomerization of both PopB and PopD. However, Pop proteins could only form rings once in the presence of membranes, suggesting that a large conformational change could occur upon lipid binding (26).

To gain insight into the mode of association of PopB and PopD into target membranes, we initially sought to determine if they could be stably inserted into an artificial bilayer or if they remained superficially associated to it. We thus purified PopB-PcrH and PopD-PcrH by overexpressing the bicistronic forms in E. coli and isolated the Pop oligomers by gel filtration at pH 5.0, as previously described (26). These samples were subsequently incubated with artificial liposomes prepared with a 2:3 ratio of α-phosphatidyl-L-serine and 1,2-dipalmitoyl-sn-glycero-3-phosphocholine in the presence/absence of agents known to release peripheral membrane proteins (37, 38). Experiments were performed with (a) PBS, pH 7.0, as a negative control, (b) 1 M KCl, (c) 6 M urea, and (d) 0.425 mM DDM (which corresponds to 2.5 × critical micelle concentration (CMC)). We then separated soluble from membrane-bound proteins on sucrose gradients; fractions were analyzed by Western blotting by employing anti-PopB and anti-PopD antibodies (Fig. 1B).

Our results show that PopB remains bound to liposomes in all conditions, i.e. in control experiments as well as in the presence of high levels of urea, KCl, and DDM, indicating that it is stably inserted into membranes. PopD, however, although remaining associated to lipids in the control experiment, could be partly detached from liposomes in the presence of all three agents (Fig. 1B), suggesting that it is not stably/integrally associated to the membrane but could be superficially associated to it. This is in agreement with ex vivo observations made on the translocon-forming proteins of Shigella in which the major translocator IpaB is strongly associated to RBC membranes, whereas the minor translocator IpaC is associated to a lesser extent (39, 40). In addition, this observation also supports in vitro studies performed on other translocon molecules of Salmonella (40), Yersinia (41, 43), and Shigella (42). These results suggest that, in Pseudomonas, PopB could be the major translocon component that binds most stably to target membranes upon infection, whereas PopD remains superficially associated to the bilayer, a suggestion that is in agreement with model membrane studies which showed that PopB plays the central role in pore formation (44, 45).
The Exposed N Terminus of PopB Is a Molten Globule—To gain an understanding into the orientation of the molecules on target membranes, we isolated Pop-containing fractions after sucrose gradient centrifugation and submitted them to trypsinolysis. We reasoned that trypsin would only target the exposed regions of the molecules, allowing us to map the sequences that were inserted into the bilayer. Pop proteins purified by gel filtration but not submitted to interaction with membranes were used as a control. After cleavage, samples were loaded onto an SDS-PAGE gel and blotted onto PVDF membranes to allow for N-terminal sequencing of the resulting bands.

For PopD, trypsin cleaved at several sites within the protein sequence irrespective of whether PopD was bound to liposomes or not (Fig. 2, A and C). N-terminal sequencing of the bands produced identified that residues Gln-56, Met-170, Thr-156, Ile-183, Ala-205, and Asn-210 (Fig. 2, A and D) were immediately juxtaposed to the trypsin cleavage site, indicating that PopD was proteolyzed in its entirety both in the lipid-free and liposome-bound forms, potentially due to the fact that it is only superficially associated to the membrane (which would confirm the results presented in Fig. 1B). In the case of the lipid-free PopD, however, a cleavage site juxtaposed to Asp-96 could also be identified (diamond in Fig. 2, C and D), suggesting that there is a potential structural rearrangement of PopD upon lipid binding.

Although extensive proteolysis was also observed for a PopB sample trypsinized in the absence of liposomes (not shown), as expected for a molten globule protein (46), for the PopB sample in the lipid-bound form proteolysis was restricted to the N-terminal region (Fig. 2, B and D). After trypsin treatment, most of the protein still ran on SDS-PAGE with a molecular weight

**FIGURE 1.** PopB and PopD interact with artificial membranes. A, scheme of expression clones used in this work. Both Pop proteins are co-expressed with their common chaperone PcrH, which carries a thrombin-cleavable 6His tag, rbs, ribosomal binding site. B, liposomes containing PopB (left) or PopD (right) were treated with different agents that destabilize the association of proteins to membranes (1 M KCl, 6 M urea, 2.5 × CMC DDM) to characterize their association with membranes. After 4 h of incubation and sucrose gradient centrifugation, fractions were recovered from the top to the bottom of the gradient and were analyzed by Western blotting using anti-PopB and/or anti-PopD rabbit antibodies.

**FIGURE 2.** Trypsin digestion of PopB and PopD inserted into liposomes. A and C, purified PopD (in solution or inserted into liposomes) was digested with trypsin at RT at a molar ratio of (1:500), and samples were loaded on SDS-PAGE after different digestion times. n.d, non-digested sample. B, PopB proteoliposomes were digested with trypsin at the same ratio as described above; aliquots were loaded on SDS-PAGE, and the gel was developed by Western blotting using purified anti-PopB antibodies. D, representation of PopB and PopD with the positions of the residues recognized by trypsin and identified by N-terminal sequence are highlighted. The black rectangles represent the two transmembrane (TM) regions. Asterisks indicate bands analyzed by N-terminal sequencing for proteins in liposomes, and the diamond indicates a band seen for PopD only in solution. E, one-dimensional NMR spectrum of PopB (1–149) recorded at 25 °C on an 800 MHz spectrometer.
similar to that of the non-digested sample (Fig. 2B); N-terminal sequencing of the five minor bands in the sample treated with trypsin for 2 h revealed the same sequence for all of them (ESEEAAKE). The first Glu corresponds to residue 153, which is located in a position that is immediately N-terminal to the beginning of the first predicted transmembrane region (residues 170–192; Fig. 2D; Ref. 26). This result suggests that in the membrane-bound form the 152 N-terminal residues of PopB are trypsin-accessible and thus could be located toward the outside of the membrane, whereas sequences that immediately follow this region are stably imbedded in the bilayer. These results suggest that as opposed to PopD, which only associates to bilayers superficially, PopB is stably inserted through its predicted transmembrane region, and its N terminus is exposed.

To gain insight into the fold of the exposed N terminus of PopB, we cloned residues 1–149 into a plasmid harboring an N-terminal His<sub>6</sub> coding sequence (pET15b) and expressed the protein in E. coli BL21 (DE3) cells. PopB (1–149) was purified by nickel chelation chromatography and eluted as a soluble monomer in gel filtration. Liposome flotation assays indicated that the protein does not have affinity for lipids, as it migrated at the bottom of the sucrose gradient (not shown). We subsequently performed a one-dimensional NMR spectrum of the sample (Fig. 2E), which shows poor dispersion of the amide resonances and broad line widths, suggesting that there may be exchange between the different conformations adopted by the protein. Notably, molten globule forms are folded proteins with extensive native-like secondary structures but that lack tertiary structures and are often characterized by spectra with extremely broad peaks with poor chemical shift dispersions. Thus, this is in agreement with studies that suggested that Pop proteins are molten globules (46), which could be a require-
ment for their passage through the interior of the T3SS needle and subsequent oligomerization and membrane interaction steps (26). These results collectively indicate that, upon activation of the T3SS and cellular recognition, PopB could be initially inserted into the membrane through its two transmembrane regions, whereas its N-terminal domain remains positioned toward the outside of the cell, where it could be recognized (and stabilized) by PopD, which is superficially positioned, and/or PcrV, which is potentially organized into a pentamer at the tip of the T3SS needle (47, 48). Notably, the presence of two predicted two transmembrane regions in PopB suggest that its C-terminal residues are also exposed to the outside of the cell (see below).

Both N and C Termini of PopB Are Exposed to the outside of the bilayer—To further explore the topology of PopB in an in vivo (infectious) setting, we generated a construct that expressed a fusion protein encompassing PopB followed by a β-lactamase reporter (β-Lac) at its C terminus (Fig. 3). The β-Lac reporter assay has been in the past successfully used to measure the level of injection and secretion of the T3SS toxins ExoS and ExoY by P. aeruginosa with two different molecules, CCF2 (30) and nitrocefin. In the first assay, if the C terminus of PopB were to be located within the target cell, the fluorescent substrate CCF2 would be recognized and cleaved by β-Lac, emitting a signal at 460 nm (blue, Fig. 3A). Conversely, location of the C terminus toward the outside of the cell would not generate CCF2 cleavage, causing a detection of a signal at 535 nm (green, Fig. 3B). Alternatively, β-Lac activity was also measured using the non-permeable, chromogenic β-lactamase substrate nitrocefin whose color was shifted from yellow to red (abs 486 nm) when it was cleaved by β-Lac (Fig. 3B).

We initially verified that the presence of the β-Lac located C-terminally to PopB did not interfere with its function in the P. aeruginosa clinical strain CHA. The PopB-β-Lac fusion proved to be functional, as it was secreted upon induction of T3SS in vitro (Fig. 4A) and was able to complement the P. aeruginosa PopB knock-out mutant (CHAΔBD (49)) in a cytotoxicity assay performed with J774 macrophages (Fig. 4B). Subsequently, to investigate whether the β-Lac reporter was found within the eukaryotic cell cytoplasm, A549 epithelial
Cells were infected with the CHA\textsubscript{ΔBD} strain supplemented with PopB-β-Lac and PopD. The cells were then incubated with CCF2, and the level of substrate cleavage was assessed by measuring the ratio of emission of light at 460 and 535 nm, as previously described (30). As a positive control we used a \textit{P. aeruginosa} CHA strain expressing an ExoS-β-Lac fusion that displayed high levels of CCF2 cleavage due to the fact that ExoS is translocated into the eukaryotic cytoplasm, and thus, CCF2 is recognized by the β-lactamase (Fig. 4, fourth bar). Notably, infection of A549 cells with the same strain expressing PopB-β-Lac instead of ExoS-β-Lac generated CCF2 cleavage levels that were comparable to background (Fig. 4, fifth bar), strongly suggesting that in the case of PopB the C terminus is not located within the cytoplasm of the eukaryotic cells.

To confirm the outer membrane exposure of the C terminus of PopB, we again infected A549 epithelial cells with \textit{P. aeruginosa} mutant strains carrying PopB-β-Lac and measured the β-lactamase activity by adding nitrocefin to infected cells (Fig. 4D). Here, we employed two strains that insert PopB into the membrane: \textit{P. aeruginosa} ΔBD complemented with both PopB-β-Lac and PopD (ΔBD/β-Lac/D) and ΔPcrV complemented with the same constructs (ΔV/β-Lac/D). This latter strain, despite not being able to express PcrV, is able to form the translocon, although it is non-infectious (49). β-Lactamase cleavage of nitrocefin could be measured for both transfected constructs. Subsequently as the negative control, we complemented a strain that cannot form the secretion needle with the aforementioned constructs (ΔF/β-Lac/D). The observation that little activity was associated to cells infected with this strain is consistent with its inability to secrete or insert the translocators into the host cell plasma membrane. Collectively, these data suggest that the PopB translocator is inserted within the eukaryotic membrane with its flexible, N-terminal domain and its C-terminal tail exposed to the outside of the bilayer.

**The N Terminus of PopB Binds to the Concave Region of PcrH—**

After having mapped the architecture of PopB on bilayers, we thus set out to investigate its cytoplasmic form; that is, in a complex with its chaperone PcrH. We had previously shown through limited proteolysis that the 95 N-terminal residues of the minor translocator PopD could associate to PcrH with 1:1 stoichiometry (26, 46). By solving the high resolution crystal structure of PcrH, we had also shown that it folds into seven TPR motifs that generate a concave region that is capable of binding PopD residues 49–56 (8). Notably, IpgC, the type II chaperone of the type III secretion system of \textit{Shigella}, also recognizes a short peptide of the N terminus of the major translocator molecule (50, 51), and SycD from \textit{Yersinia enterocolitica} was also structurally characterized in complex with a peptide from minor translocator YopD (52).

By performing sequence alignments between PopB and PopD, we identified a short peptide in PopB (residues 51–59) within the N-terminal flexible region that displayed sequence
similarities with the minimal region in PopD that interacts with PcrH (residues 49–56, Fig. 5A; Ref. 8) and harbors the six-amino acid chaperone-binding consensus sequence initially identified by Kolbe and co-workers (50). This PopB peptide, \( \text{TGVALTPPS} \), was thus synthesized and incubated with PcrH before crystallization trials at 4 °C. Diffraction data were collected to a resolution of 2.25 Å at the European Synchrotron Radiation Facility in Grenoble, and crystals of the complex belonged to space group P21212 with two molecules per asymmetric unit. The structure was solved by molecular replacement using the structure of the apo form of PcrH as a search model (PDB 2XCC). Data collection and refinement statistics are shown in Table 1. Calculation of a difference Fourier map \( (F_o - F_c) \) revealed density corresponding to the peptide in both monomers, but only the peptide associated to monomer A could be traced unambiguously in its entire length (residues 51–59). We will thus describe the analysis of this monomer.

\( \text{TGVALTPPS} \) binds in an extended conformation within a cleft in the concave region of PcrH, and it presents an antiparallel orientation in respect to the first helix of the chaperone (Fig. 5, B and D). Notably, residues Val-53, Leu-55, and Pro-58 (the prime indicates PopB peptide residues), which form the conserved translocator-chaperone binding motif \( P'/V'XL'XXP' \) (50–52), fit into PcrH surface cleft and interact mostly with conserved residues (Tyr-40, Ala-41, Phe-44, Phe-59, Leu-74, Gly-75, Ala-78, Tyr-93, His-109). Additionally, the main chain atoms of Thr-51, Gly-52, Val-53, and Leu-55 establish hydrogen bonds with the side chain atoms of Glu-112, Arg-146, His-109, Gln-48, and Tyr-40 from PcrH intermediated by water molecules, whereas the main chain atoms of Val-53, Ala-54, Thr-56, and Ser-59 establish direct hydrogen bonds with the Tyr-47, Arg-105, Tyr-40, and Glu-37 side chains of PcrH (Fig. 5, B–D). Only one peptide side chain, the hydroxyl group of Thr-56, makes a direct hydrogen bond with PcrH (with the side chain of Arg-105). Last, the side chains of residues Thr-51, Ala-54, and Pro-57 point out to the solvent. These observations indicate that the concave region of PcrH acts as the palm of a hand, “grasping” a common region of the Pop molecules (Fig. 5C), with the rest of the protein potentially wrapping around the chaperone and also interacting with the convex region of PcrH.

Other structures of T3SS type II chaperones have already been solved in complex with peptides corresponding to the

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**TABLE 1**

Data collection, molecular replacement and structure refinement statistics

| Data collection | Values | r.m.s.d., root mean square deviation. |
|-----------------|--------|-------------------------------------|
| Wavelength (Å)  | 0.977  |                                     |
| Space group     | P2,22  |                                     |
| Unit cell parameters (Å) | \( a = 45.10, b = 73.29, c = 98.94 \) | |
| Resolution (Å)  | 2.25 (2.25–2.39) | |
| Number of observed/unique reflections | 113,610/16,247 | |
| Completeness (%) | 99.4 (96.8) | |
| \( R_{	ext{measured}} \)  | 5.1 (44.7) | |
| \( R_{	ext{merge}}-F_o \)  | 6.6 (39.8) | |
| I/\( I_o \) | 29.7 (4.8) | |

| Refinement | Values | r.m.s.d., root mean square deviation. |
|------------|--------|-------------------------------------|
| Rwork      | 0.214  |                                     |
| \( R_m \)  | 0.264  |                                     |
| r.m.s.d.\( \text{bond} \) | 0.008 | |
| r.m.s.d.\( \text{angle} \) | 1.069 | |
| Mean B factor (Å²) | 43.242 | |
| No. of protein atoms | 2,028 | |
| No. of solvent atoms | 117 | |

| Ramachandran analysis (%) | Values | r.m.s.d., root mean square deviation. |
|---------------------------|--------|-------------------------------------|
| Residues in favored regions | 98.1   |                                     |
| Residues in allowed regions | 1.9    |                                     |
| Residues in outlier regions | 0     |                                     |

* \( R_{	ext{merge}}-F_o \) is defined in Ref. 55.
chaperone binding region of their respective translocators, as mentioned above. The comparison of these structures with our present model reveals that all peptides bind in a very similar fashion to the concave region of their respective chaperones, with their N termini toward helix 1 and their C termini toward helix 5 (Fig. 6). Notably, within the region that encompasses the common motif P’N’LXLXXP’, the conformation of the main chain peptide atoms is very similar, and the side chains of the conserved residues Pro/Val, Leu, and Pro fit deeply in the cleft of the chaperones. The most dissimilar conformations are observed for the residues located at the extremities of the peptides, not included within the conserved motif (Fig. 6).

PcrH is now the only type II chaperone that has been structurally characterized in complex with peptides from both major and minor translocator partners, unequivocally showing that the concave region of the chaperone serves as the binding site for both proteins. It is of note that PopD and PopB share and minor translocator partners, unequivocally showing that the chaperones. The most dissimilar conformations are observed for the residues located at the extremities of the peptides, not included within the conserved motif (Fig. 6).

To test if the interactions between PcrH and the N-terminal peptide of PopB involving residues 51–59 are important for functionality of the T3SS, we modified the highly cytotoxic P. aeruginosa CHA strain (originally isolated from a cystic fibrosis patient) to carry mutations within this motif in PopB (Fig. 5A). We tested these mutants for (i) expression and secretion of PopB mutants and (ii) cytotoxicity toward macrophages. A mutant strain carrying a PopB variant with point mutations in the 53TGVALTTPPS59 region, in which two (L55D/P58D; PopB-2D) or four (V53D/L55D/P57D/P58D; PopB-4D) of the central hydrophobic residues were replaced by Asp allowed us to verify the importance of hydrophobic interactions within the PopB-PcrH complex. Plasmids expressing the mutant pcrH-popBpopD section of the operon were introduced individually into the P. aeruginosa CHA strain carrying deletions of popB and popD (ΔpopBD) before testing.

### FIGURE 7. Characterization of PopB mutants in P. aeruginosa. A, P. aeruginosa strains carrying the two different sets of mutations in PopB, L55D/P58D (PopB-2D) and V53D/L55D/P57D/P58D (PopB-4D), were grown in T3SS-inducing conditions. When the cultures reached an A600nm value of 1.0 absorbance units, 100 μl of cultures were centrifuged, and the pellet (Expression) and the supernatant (Secretion) were analyzed by Western blotting with anti-PopB, anti-PopD, and anti-PcrH antibodies. B, cytotoxicity of PopB mutants. Macrophages were infected with cultures of P. aeruginosa CHA depleted for popD and popD genes (ΔBD) or complemented with PopDwt and PopBwt, PopB-2D, or PopB-4D using a multiplicity of infection of 5. A non-infected culture was used as negative control. After 1–4 h of infection, LDH release into the supernatant was measured to determine the % of cell death. C, stability of PopB mutants was followed for up to 3 h after the addition of chloramphenicol, a protein synthesis inhibitor. The cell contents were analyzed by Western blotting with anti-PopB antibodies and anti-PcrV as a control.

P. aeruginosa strains expressing the two PopB mutants were able to express and secrete proteins at comparable levels (Fig. 7A); in addition, the PopB mutants were stable for up to 3 h in T3SS-inducing conditions, suggesting that PcrH binding is maintained (Fig. 7C). Finally the two mutant strains displayed levels of cytotoxicity toward macrophages that were similar to that displayed by the wild type strain (Fig. 7B), suggesting that the mutations within the PopB region that interacts with the PcrH “palm” are not sufficient to disrupt protein-protein interactions. This is distinct from cytotoxicity profile of mutants of the comparable region in PopD, which displayed no cytotoxicity toward macrophages (8), and indicates that Pop proteins display distinct affinities for their common chaperone. This could be explained by the fact that PopB (being a bigger mole-
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cule) may remain partly bound to PcrH despite the mutations, suggesting that other regions of the translocator protein may also contribute to chaperone binding.

The translocon is a key element of the pathogenicity arsenal of T3SS-carrying bacteria, and understanding the details of the individual steps required for its formation, including how proteins associate both within the cytoplasm and on the target bilayer, is essential for its characterization. The collective results presented here indicate that (i) within the cytoplasm both major and minor translocators use the concave region of their common chaperone, PcrH, for binding. This excludes the possibility of the formation of a 1:1:1 complex and suggests that the functional units are most likely 1:1 associations. (ii) Upon translocation, presumably through the T3SS needle, PopB becomes stably associated into the target bilayer with both N- and C-terminal regions directed toward the outside of the cell. These regions could potentially be the recognition sites for PopD, which is only superficially associated to the membrane, and could subsequently play the role of “bridge” with the tip of the T3SS needle (as suggested for the Yersinia spp YopD homolog; Ref. 54). Detailed information regarding the distinct steps involved in translocon formation will be essential for the exploitation of such molecules as potential targets for the development of novel antibacterials.

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