The Breadth and Molecular Basis of Hcp-Driven Type VI Secretion System Effector Delivery

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ABSTRACT The type VI secretion system (T6SS) is a bacterial nanoscale weapon that delivers toxins into prey ranging from bacteria and fungi to animal hosts. The cytosolic contractile sheath of the system wraps around stacked hexameric rings of Hcp proteins, which form an inner tube. At the tip of this tube is a puncturing device comprising a trimeric VgrG topped by a monomeric PAAR protein. The number of toxins a single system delivers per firing event remains unknown, since effectors can be loaded on diverse sites of the T6SS apparatus, notably the inner tube and the puncturing device. Each VgrG or PAAR can bind one effector, and additional effector cargoes can be carried in the Hcp ring lumen. While many VgrG- and PAAR-bound toxins have been characterized, to date, very few Hcp-bound effectors are known. Here, we used 3 known Pseudomonas aeruginosa Hcp proteins (Hcp1 to -3), each of which associates with one of the three T6SSs in this organism (H1-T6SS, H2-T6SS, and H3-T6SS), to perform in vivo pulldown assays. We confirmed the known interactions of Hcp1 with Tse1 to -4, further copurified a Hcp1-Tse4 complex, and identified potential novel Hcp1-bound effectors. Moreover, we demonstrated that Hcp2 and Hcp3 can shuttle T6SS cargoes toxic to Escherichia coli. Finally, we used a Tse1-Bla chimera to probe the loading strategy for Hcp passengers and found that while large effectors can be loaded onto Hcp, the formed complex jams the system, abrogating T6SS function.

IMPORTANCE The type VI secretion system (T6SS) is an effective weapon used by bacteria to outgrow or kill competitors. It can be used by endogenous commensal microbiota to prevent invasion by pathogens or by pathogens to overcome resident flora and successfully colonize a host or a specific environmental niche. The T6SS is a key contributor to this continuous arms race between organisms as it delivers a multitude of toxins directed at essential processes, such as nucleic acid synthesis and replication, cell wall and membrane integrity, protein synthesis, or cofactor abundance. Many T6SS toxins with unknown function remain to be discovered, whose yet-uncharacterized targets could be exploited for antimicrobial drug design. The systematic search for these toxins is not facilitated by the presence of readily recognizable T6SS motifs, and unbiased screening approaches are thus required. Here, we successfully used a known shuttle for cargo T6SS effectors, Hcp, as bait to identify uncharacterized toxins.

KEYWORDS T6SS, Hcp, Pseudomonas, protein secretion, toxin

Bacterial secretion systems shuttle proteins to the extracellular milieu or into target cells (1, 2). The type VI secretion system (T6SS) functions as a contractile device that resembles a bacteriophage (3, 4). The contractile sheath assembles onto a
baseplate (TssKEFG) which is bound to the cytoplasmic membrane via an interaction with the membrane complex TssLMJ. The sheath made of TssBC protomers (5–8) extends in the cytosol while wrapping around a stack of hexameric Hcp rings (9) until it reaches the opposite side of the cell (10). The Hcp rings fit within the sheath lumen and pile on top of each other (11, 12). The formation of the Hcp tube is dependent on a spike structure, the VgrG trimer, whose large base provides the interface for assembly of the first Hcp ring (13). At the other end of VgrG sits a conical PAAR monomer (14). The PAAR-VgrG-Hcp could be compared to a spear, with the projectile point being the VgrG-PAAR complex and the shaft the Hcp tube. Sheath contraction results in spear ejection and, cryotomograms of the contracted sheath show hollow structures (10, 15).

The T6SS has been shown to be important in bacterium-host interaction but is also an antimicrobial weapon that injects toxins into bacterial or fungal prey cells (16, 17). The T6SS effectors have no recognizable signals, except on some occasions where they are found to contain MIX or FIX motifs (18, 19). These effectors are loaded on the PAAR, VgrG, or Hcp components of the T6SS apparatus. This can occur either through direct protein-protein interactions (20) or with the help of chaperones/adaptors such as EagR (21–23) or Tap proteins (24, 25). Effector proteins fused to PAAR, VgrG, or Hcp have also been identified and are termed specialized effectors (26–30) as opposed to cargo effectors associated with the T6SS via protein-protein interactions.

A large number of VgrG- and PAAR-associated effectors, either specialized or cargo, have been functionally characterized (28, 29, 31, 32). For Hcp, few specialized effectors have been reported for the Escherichia coli T6SS (33), while the Pseudomonas aeruginosa Tse2 cargo is loaded within the Hcp1 ring lumen (34). Hcp1 is not the only Hcp protein found in P. aeruginosa, and at least two additional Hcps, Hcp2 and Hcp3, could carry effector cargo. In this study, we used all three P. aeruginosa Hcp proteins to perform an in vivo pulldown approach. We confirmed that Hcp1 binds to Tse1 to -4, as reported previously (34, 35), and identified novel Hcp1-bound proteins, some in the size range of 10 to 30 kDa that might be suited to fit into the Hcp lumen. We were also able to identify Hcp2- and Hcp3-bound proteins, some of which exert toxicity when produced in the cytosol of Escherichia coli. Moreover, we found that conserved Hcp1 to -3 inner ring residues were required for successful interaction with both the known and the uncharacterized effectors. Moreover, we investigated the size limit for Hcp cargoes by using chimeric proteins. We observed that large chimeric effectors bound to Hcp can be recognized by the T6SS but jam the system, likely because of steric hindrance during the assembly of the T6SS sheath and tube. We conclude that Hcp-dependent transport is a widely used strategy for the delivery of T6SS cargo effectors but has limitations compared to VgrG-dependent transport, which allows for translocation of larger effectors.

RESULTS

Biochemical characterization of Tse1 to -4/Hcp interactions. Tse1 to -3 were the first T6SS antibacterial toxins identified (36) through a secretome analysis. It was later shown that the presence of Hcp1 can stabilize these effectors and proposed that Tse2 fills the lumen of the Hcp ring (34). Here, we produced a His-tagged version of Hcp1 (see Fig. S1 in the supplemental material) and hemagglutinin (HA)-tagged versions of the catalytically inactive Tse1 to -3 toxins, i.e., Tse1C30A, Tse2T79A/S80A, and Tse3E250Q (37, 38). We demonstrate that interaction could indeed be identified using our pulldown approach where the hcp1-encoding plasmid and one of the plasmid-encoding toxins were cotransformed in E. coli. Upon copurification using Hcp1 as a bait, elution from Ni-nitrilotriacetic acid (NTA) resin and size exclusion chromatography, Tse1, Tse2, and Tse3 were found in complex with Hcp1 (see Fig. S2).

The P. aeruginosa Tse4 toxin was discovered using quantitative cellular proteomics by comparing wild-type and hcp1 mutant strains and observing a low abundance of Tse4 in an hcp1 mutant background (35). Tse4 is a pore-forming toxin that allows small ion leakage (39). We assessed a direct interaction between Hcp1 and Tse4 by using the
pulldown approach. Overexpression of Tse4 in *E. coli* is toxic. We observed that adding a C-terminal HA tag, rather than an N-terminal tag, decreased the level of toxicity (see Fig. S3A). However, the solubility of Tse4-HA was limited, while copurification with Hcp1-His showed almost no interaction (Fig. S3B and C). We hypothesized that the C-terminal HA tag interferes with Tse4 activity but also impairs interaction with Hcp1. Instead, while an N-terminally tagged HA-Tse4 was not soluble, its solubility was improved upon coexpression with Hcp1-His, and copurification resulted in a Tse4-Hcp1 complex that was maintained after size exclusion chromatography (Fig. 1A and B).

Previous studies identified key residues in the Hcp1 inner ring surface that are important for secretion of Tse1 to -3, namely, S31, which is only important for secretion of Tse2, L28 and A29, which are important for secretion of Tse2 and Tse3, and T59, S115, and T122, which are important for secretion of Tse1, Tse2, and Tse3 (34). We confirmed that S31 is important for interaction with Tse2, while S115 is key for interaction with all 3 toxins, Tse1 to -3 (see Fig. S4). We tested whether these amino acids are also important for the Hcp1-Tse4 interaction and found that the Hcp1S31Q variant is still able to interact with Tse4, although only weakly since the complex is lost after gel filtration, whereas this interaction is totally abrogated when the Hcp1T59Q and Hcp1S115Q variants are used (Fig. 1C to E). We therefore conclude that S31 appears to

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**FIG 1** HA-Tse4 interacts specifically with Hcp1 inner ring residues. (A) Western blot of Hcp1-His and HA-Tse4 copurified via Ni resin and an SD200 SEC column; lanes labeled on the top are postexpression sample, insoluble and soluble samples after sonication and clarification, flowthrough (FT) from the Ni-NTA column, wash fraction before elution, Ni fractions corresponding to the elution peak, and SEC fractions corresponding to the gel filtration peak. The antibodies used are labeled on the right of the Western blot; top, anti-HA (BioLegend) at 1:1,000 concentration; bottom, anti-His (GenScript) at 1:1,000 concentration. Molecular weight standards are on the left. (B) SEC chromatograph of gel-filtrated Hcp1-His and HA-Tse4 copurification. Western blots of Hcp1-His inner ring mutants and HA-Tse4 copurifications with Hcp1S31Q-His (C), with Hcp1T59Q-His (D), and with Hcp1S115Q-His (E).
be specific for Tse2 interaction, while TS9 and S115 have a more general role, and their modification disrupts interaction with T6SS toxins.

**Identification of novel T6SS effectors by in vivo Hcp cross-linking.** Having validated our pulldown approach using Hcp-His as a reliable bait, we used Hcp1 to -3 bearing a C-terminal FLAG tag for in vivo pulldown in *P. aeruginosa* PAO1 ΔrsmA. The rationale for using this strain is that an *rsmA* mutation increases expression of all three T6SSs in *P. aeruginosa* (40). Lysates of cells were treated with a dimethyl 3,3′-dithiobis-propionimidate (DTBP) cross-linker in order to capture transient or weak interactions between Hcp1 to -3 and new putative cargo proteins and were then applied to FLAG M2 magnetic beads; proteins that coeluted with each Hcp were identified by mass spectrometry. For each Hcp-FLAG pulldown, the identified proteins were cross-referenced against the results of two control pulldown experiments: one performed using untagged Hcp as the bait protein and the other using a FLAG-tagged version of each Hcp’s cognate VgrG protein (see Materials and Methods). Proteins found to be significantly enriched (≥3-fold; the level of enrichment of the known Hcp1 effector Tse1) over the untagged control, while at the same time not found to be enriched in the VgrG control pulldown, were considered putative Hcp-bound effectors. In each of the volcano plots depicting the results (Fig. 2), the tagged Hcp protein is, as expected, highly enriched. Initial inspection of the proteins identified in the Hcp pulldowns (Table 1) revealed 107 proteins enriched ≥3-fold by Hcp1, 32 of which are significant Hcp1-specific interacting partners. Of these proteins, 23 (71.88%) have an *M*<sub>ω</sub> equal to or below that of Tse3, which at 44.4 kDa is the largest of the known Hcp1-delivered effectors. Tse2, Tse3, and Tse4 appear as the top 3 hits, followed by the remaining Hcp-interacting proteins, some of which appear to be good candidates for novel Hcp1-bound effector proteins (such as PA3440, an 11.6-kDa protein of unknown function). Applying these same criteria to the Hcp2 and Hcp3 pulldowns revealed 30 proteins enriched ≥3-fold by Hcp2, 2 of which are significant Hcp2-specific interacting partners with molecular weights below that of Tse3 (PA2066 and PA2966) and 63 proteins enriched ≥3-fold by Hcp3, 9 of which are significant Hcp3-specific interacting partners with molecular weights below that of Tse3 (PA0256, PA3187, PA2372 [a 21-kDa protein of unknown function encoded within the H3-T6SS cluster], PA3076, PA0381, PA1373, PA3182, PA4920, and PA0771) (Tables 2 and 3). Finally, in order to ensure that our approach offers an unbiased method for identifying new Hcp-bound T6SS effectors, we compared the theoretical isoelectric point (pI) of the putative Hcp-bound effectors identified in our pulldown experiments to the theoretical average pI of the *P. aeruginosa* PAO1 proteome. The average pI of the Hcp-bound putative effectors identified here was 6.16 ± 1.29 (range, 3.79 to 9.92), while the average pI of the *P. aeruginosa* proteome is 7.12 ± 1.96 (range, 3.12 to 13.51). Thus, this approach has no obvious bias for proteins of a certain charge and is able to identify proteins from throughout the *P. aeruginosa* proteome.

**PA2066 and PA0256 are putative antibacterial T6SS toxins.** In contrast to Hcp1, no Hcp-bound effectors for Hcp2 and Hcp3 have been identified to date. For this reason, we investigated the two top hits from the Hcp2 and Hcp3 in vivo pulldown, i.e., PA2066 for Hcp2 (10-fold enrichment) and PA0256 for Hcp3 (8.4-fold enrichment). We validated the interaction by performing copurification experiments with C-terminally His-tagged Hcp (see Fig. S5A to D) and N- or C-terminally HA-tagged PA2066/PA0256. PA2066 was more soluble when tagged with HA at the C terminus (Fig. 3A), but both versions coeluted with Hcp2-His from the nickel column (Fig. 3B and C; data not shown for N terminus). The interaction is weak, since the complex was lost after size exclusion chromatography, but PA2066-HA was not retained on the nickel column if not coexpressed with Hcp2-His (see Fig. S6A), which suggests a specific interaction. The N-terminally HA-tagged version of PA0256 formed a robust complex with Hcp3-His (Fig. 3D to F), which was still present after size exclusion chromatography. We conclude that PA2066 and PA0256 interact with their cognate Hcp partners and that the strength of the interaction varies depending on the Hcp-effector pair analyzed. This is also seen for
Hcp1/2/3 pulldown hits. Volcano plots showing LFQ analysis of Hcp1-, Hcp2-, and Hcp3-FLAG pulldowns performed in *P. aeruginosa*. Results are from three independent experiments, and the full data sets are provided in Data Set S1 in the supplemental material. The x axes show the fold change in enrichment over the untagged Hcp control. The y axes depict the significance of the enrichment. Proteins plotted were not found to be enriched in the cognate VgrG-FLAG controls (Data Set S1). (A) Hcp1-FLAG pulldown results. The red dot is Tse1, a known Hcp1-delivered effector that was enriched 3-fold, but was below the significance cutoff. This was used as the enrichment baseline for possible hits. (B) Hcp2-FLAG pulldown results. (C) Hcp3-FLAG pulldown results. Dotted lines represent the enrichment cutoff of $\geq 3$-fold, as determined by Tse1 and the significance cutoff $-\log_{10} P$ value of $\geq 1.3$, which is equivalent to a $P$ value of $\leq 0.05$. Increasing significance on the y axis corresponds to a smaller $P$ value. Blue dots are proteins above the enrichment and significance cutoffs. Purple dots are the bait Hcp proteins.
Hcp1 interactions; Tse2 interacts strongly with Hcp1, while other Tse effectors display a weaker affinity.

The inner ring Hcp1 residues that were important for the interaction with Tse4 (Fig. 1C to E) were examined for their conservation in Hcp2 and Hcp3. T59 in Hcp1 is conserved in Hcp2 and Hcp3 (residues T71 and T60, respectively) and, in all cases, located on the inner ring (see Fig. S7). Hcp2 and Hcp3 substitutions of these residues impacted interaction with the putative effectors PA2066 and PA0256, respectively (compare Fig. 3D and G and Fig. 3D and H), suggesting a conserved role of this threonine residue in effector interaction.

Having shown that both PA2066 and PA0256 are putative Hcp cargoes, we assessed whether they can be putative T6SS antibacterial toxins by overexpressing the proteins in E. coli. A standard toxicity assay was performed by plating overnight-grown bacterial cultures on agar plates containing isopropyl-β-D-thiogalactopyranoside (IPTG) (toxin induction) or glucose (toxin repression), and the CFU was enumerated. These experiments revealed that PA2066 N-terminally HA-tagged is toxic in the E. coli cytoplasm (Fig. 4A). There is a 2-log₁₀ drop in survival, equivalent to 135 times less survival when comparing the mean survival of E. coli carrying pET22b to that carrying pET22b/HAPA2066. When harboring a C-terminal HA tag, PA2066 did not display any toxicity, which suggests that the tag in that position interferes with PA2066 activity. In the case

| Protein | Fold change | Significance | Description of protein | Size (kDa) |
|---------|-------------|--------------|------------------------|------------|
| PA2702 Tse2 | 23.4 | 3.85 | Known Hcp1 effector | 17.7 |
| PA2774 Tse4 | 11.2 | 2.42 | Known Hcp1 effector | 19.2 |
| PA3484 Tse3 | 9.6 | 3.78 | Known Hcp1 effector | 44.4 |
| PA3296 PvdF | 9.2 | 2.88 | Pyoverdine synthetase F | 31 |
| PA2000 DhcB | 7.2 | 1.77 | Dehydroacornitine CoA transferase | 23.2 |
| PA0662 ArgC | 6.6 | 1.35 | N-Acetyl-gamma-glutamyl-phosphate reductase | 36.7 |
| PA4407 YdfG | 6.4 | 2.24 | Short-chain dehydrogenase | 27.4 |
| PA0075 PppA | 5.8 | 1.44 | Represses H1-T6SS | 25.9 |
| PA5220 | 5.3 | 1.32 | Hypothetical | 30.5 |
| PA4748 TpiA | 5.3 | 2.46 | Triosephosphate isomerase | 25.6 |
| PA0895 ArcC | 5 | 1.73 | Succinylmethylthine transaminase/acytrolmethylthine aminotransferase | 43.7 |
| PA3471 MaeA | 4.8 | 2.41 | NAD-dependent malic enzyme | 62.4 |
| PA1588 SucC | 4.8 | 2.73 | Succinate-CoA ligase (ADP-forming) subunit beta | 41.5 |
| PA3017 | 4.8 | 2.02 | 2-Hydroxyglutarate dehydrogenase | 51.3 |
| PA1589 SucD | 4.5 | 1.78 | Succinate-CoA ligase (ADP-forming) subunit alpha | 30.3 |
| PA5427 AdhA | 4.5 | 1.90 | Alcohol dehydrogenase | 35.9 |
| PA2553 | 4.5 | 2.42 | Probable acyl-CoA thiolase | 41.4 |
| PA4854 PurH | 4.4 | 2.02 | Bifunctional purine biosynthesis protein | 57.7 |
| PA6232 L Jad | 4.2 | 3.12 | Isocitrate dehydrogenase (NADP) | 45.6 |
| PA4464 PtsN | 4 | 1.30 | Nitrogen regulatory protein | 16.7 |
| PA2015 LiuA | 4 | 2.25 | Putative isovaleryl-CoA dehydrogenase | 42.2 |
| PA4483 GatA | 3.9 | 1.39 | Glutamyl-tRNA(Gln) amidotransferase subunit A | 51.9 |
| PA3183 Zwf | 3.9 | 2.06 | Glucose-6-phosphate 1-dehydrogenase | 55.6 |
| PA3635 Eno | 3.8 | 2.99 | Enolase | 45.2 |
| PA5460 IleS | 3.7 | 2.99 | Isoleucine-tRNA ligase | 105.5 |
| PA2624 Idh | 3.7 | 2.57 | Isocitrate dehydrogenase (NADP) | 81.6 |
| PA3529 | 3.6 | 1.98 | Alkyldihydroperoxide reductase C | 21.8 |
| PA0552 Pgk | 3.5 | 1.40 | Phosphoglycerate kinase | 40.4 |
| PA0130 BauC | 3.4 | 1.39 | Putative 3-oxopropanoate dehydrogenase | 53.3 |
| PA0382 TrmB | 3.4 | 1.39 | tRNA (guanine-N(7)-methyltransferase) | 27.6 |
| PA1838 CysL | 3.3 | 2.52 | Sulfite reductase | 62.1 |
| PA2513 AmtB | 3.3 | 1.30 | Anthranilate dioxygenase small subunit | 19.3 |
| PA3286 | 3.2 | 1.33 | Beta-ketodecanoyl-(acyl-carrier-protein) synthase | 38.2 |
| PA3440 | 3.2 | 1.40 | Hypothetical | 11.6 |
| PA4756 CarB | 3 | 1.56 | Carbamoyl-phosphate synthase large chain | 117.3 |
| PA1844 Tse1 | 3 | 0.85 | Known Hcp1 effector | 16.4 |

*Hits are ordered by fold change, which is the difference between the Hcp-FLAG pulldown and the Hcp untagged pulldown. All hits were not enriched in the respective VgrG-FLAG control.

*Significance is −log P value, where > 1.3 equals a P value of < 0.05.

*CoA, coenzyme A.
of PA0256, the tag position had no significant impact, since we observed a 2-log$_{10}$ drop in survival, equivalent to 135 and 109 times less survival when comparing the mean survival of *E. coli* harboring pET22b to that harboring pET22b/HA-PA0256 and pET22b/PA0256-HA, respectively (Fig. 4B). We also tested the toxicity of one of the low-molecular-weight hits of unknown function retrieved from the Hcp1 pulldown, i.e., PA3440, but found no difference in the CFU counts (Fig. 4C). Based on our data, we suggest that PA2066 and PA0256 are bona fide Hcp cargoes and might be H2- and H3-T6SS toxins with antibacterial activity, respectively.

While both proteins are uncharacterized, *in silico* analyses of PA2066 showed it to be a YdcF family protein with <70% identity to orthologues in other species. PA2066 contains a Rossmann-like alpha/beta/alpha sandwich fold which has been linked to cofactor binding. The neighboring gene of PA2066 is PA2067, which encodes a predicted HAD family phosphatase. While not studied experimentally, if PA2067 was the immunity protein of this putative effector, it would suggest a kinase-phosphatase toxin-immunity pair, with PA2066 being a kinase, as Rossmann-like alpha/beta/alpha sandwich folds have been found in a characterized family of kinases (41). Immunity proteins counteracting the enzymatic activity of their cognate toxin have been previously reported, for example, in the case of Tri1 that removes the FtsZ ADP ribose modification driven by the T6SS toxin Tre1 from *Serratia proteamaculans* (42).

PA0256 contains an uncharacterized DUF4347 domain and is primarily found in *P. aeruginosa* with the exception of a single strain of *Acinetobacter baumannii* (99.35% protein identity) and three strains of *Pseudomonas fluorescens* (60.97, 37.3, and 36.99% protein identity). The genomic organization does not suggest any candidate encoding an immunity protein; however, this effector is encoded in the vicinity of the vgrG2b orphan cluster (PA0259-PA0263), which still contains genes of unknown function (29).

### Tse4 features that modulate Hcp-dependent delivery.

We observed that the position at which a T6SS toxin is tagged, for example, addition of an HA tag at the C terminus of Tse4 (Fig. 5C), may interfere with the interaction between the effector and its cognate Hcp and subsequently secretion (Fig. 5A). We used Tse4 to investigate whether specific features of a T6SS-bound toxin might be required for T6SS-dependent delivery of Hcp cargoes and whether N- or C-terminal residues are important for recog-

### TABLE 2 Hcp2-FLAG top interaction hits$^a$

| Protein    | Fold change | Significance$^b$ | Description of protein                                   | Size (kDa) |
|------------|-------------|------------------|----------------------------------------------------------|------------|
| PA2066     | 10          | 2.71             | Hypothetical, YdcF family protein                        | 23.9       |
| PA4226 PchE | 4.4         | 1.52             | Dihydroaeruginic acid synthetase                         | 156.4      |
| PA2966 Arcp | 3.1         | 1.35             | Acyl carrier protein                                     | 8.7        |

$^a$Hits are ordered by fold change, which is the difference between the Hcp-FLAG pulldown and the Hcp untagged pulldown. All hits were not enriched in the respective VgrG-FLAG control.

$^b$Significance is $-\log P$ value, where $>1.3$ equals a $P$ value of $<0.05$.

### TABLE 3 Hcp3-FLAG top interaction hits$^a$

| Protein    | Fold change | Significance$^b$ | Description of protein                                   | Size (kDa) |
|------------|-------------|------------------|----------------------------------------------------------|------------|
| PA0256     | 8.4         | 1.76             | Hypothetical                                             | 33.5       |
| PA3187 GltK | 8           | 1.5              | ATP-binding component of ABC transporter                 | 42.2       |
| PA2372     | 5.8         | 1.60             | Hypothetical                                             | 21.1       |
| PA3076     | 4.3         | 1.52             | Hypothetical                                             | 39.4       |
| PA0381 ThiG | 3.9         | 1.77             | Thiazole synthase                                        | 28.2       |
| PA1373 FabF2 | 3.4       | 1.48             | 3-Oxoacyl-(acyl-carrier-protein) synthase 2              | 43.5       |
| PA3182 PgI  | 3.3         | 1.87             | 6-Phosphogluconolactonase                               | 25.6       |
| PA4920 NadE | 3           | 1.65             | NH$_2$-dependent NAD$^+$ synthetase                      | 29.7       |
| PA0771 Era  | 3           | 2.10             | GTPase                                                   | 34.5       |

$^a$Hits are ordered by fold change, which is the difference between the Hcp-FLAG pulldown and the Hcp untagged pulldown. All hits were not enriched in the respective VgrG-FLAG control.

$^b$Significance is $-\log P$ value, where $>1.3$ equals a $P$ value of $<0.05$. 
FIG 3 PA2066 and PA0256 are putative Hcp cargoes. (A) Solubility test for PA2066; E. coli BL21 cells expressed either HA-tagged PA2066 alone or with Hcp2-His. After induction, cells were grown at 18°C overnight, harvested, and resuspended in purification resuspension buffer, sonicated, and centrifuged. Western blot of postexpression, insoluble, and soluble samples. (B) Western blot of Hcp2-His and PA2066-HA copurification; lanes labeled at the top are postexpression sample, insoluble and soluble samples after sonication and clarification, flowthrough (FT) from the Ni-NTA column, wash fraction before elution, Ni fractions corresponding to the elution peak, and SEC fractions corresponding to the gel filtration peak. The Hcp2-His with PA2066-HA copurification Western blot membrane was cut so that the wash-SEC fractions could be exposed longer to check that the wash fraction was clear and if there were bands detectable in the SEC peak fractions with anti-HA antibody. (C) SEC chromatograph of purified Hcp2-His and PA2066-HA. Hcp3-His and HA-PA0256 copurification Western blot (D), SEC chromatograph (E), and Coomassie blue stain (F). (G) Copurification of Hcp2T71Q-His and PA2066-HA. (H) Copurification of Hcp3T60Q-His and HA-PA0256. The antibodies used are labeled on the right: top, anti-HA antibody (BioLegend) at 1:1,000 concentration; bottom, either anti-Hcp2 antibody (Eurogentec) at 1:500 concentration or anti-His antibody (GenScript) at 1:1,000 concentration. Molecular weight standards are on the left.
nition by Hcp. Tse4 is a pore-forming protein (39), and putative transmembrane domains are present near both the N- and C-terminal regions of the protein (Fig. 5B). We engineered truncations of Tse4 still harboring an N-terminal HA tag and assessed secretion of these variants. The intact and various truncated Tse4 constructs were expressed in the H1-T6SS active \textit{P. aeruginosa} strain PAO1\textsubscript{D}rsmA; while all constructs were expressed, HA-NT1 seemed less stable (Fig. 5A). All other variants were secreted, including those carrying deletions at the C terminus. This indicates that the presence of the tag at the C terminus does not necessarily obscure key residues for Hcp recognition, but instead, the tag may block Hcp interaction by altering the overall fold of the protein. This is supported by our observation that Tse4 toxicity in \textit{E. coli} was lost for Tse4-HA but not for the HA-Tse4 version (Fig. S3A). We conclude that specific regions or residues of Hcp cargoes, notably within the N- and C-terminal regions, are not a major determinant of Hcp-dependent recognition. Instead, the effector size and the steric fit with key residues within the Hcp lumen are likely two of the main drivers for the loading of effectors into the T6SS tube.

**Large T6SS chimeric effectors block T6SS function when bound to Hcp.** We examined whether the addition of a large extension to an Hcp cargo might interfere with its secretion. When using the Hcp1 Tse1 toxin fused with beta-lactamase (Bla), we observed that they copuriﬁed and formed a stable complex (Fig. 6A and B). We tested whether the Hcp1-bound chimera was delivered into target cells by performing competition assays using green ﬂuorescent protein (GFP)-tagged \textit{E. coli} as prey for the \textit{P. aeruginosa} attacker. The tssB1 or hcp1 mutant was less effective at killing \textit{E. coli} (Fig. 6C), since an 11-fold higher ﬂuorescence level than for the wild type was detected. Deletion of tse1 did not have a drastic effect on prey killing, since other H1-T6SS-dependent toxins were still delivered. When the tse1 gene was replaced on the chromosome with the gene encoding the Tse1-Bla chimera, killing was drastically reduced (9-fold ﬂuorescence increase), nearly as much as with the inactive T6SS mutants. The lack of killing is not linked to the presence of a tag at the C terminus, since the addition of a small HA tag (Tse1-HA construct) did not interfere with killing. Instead, the addition of another large protein, mScarlet-I, also abrogated killing. To conﬁrm that a large
Tse1 chimera blocks T6SS firing because it binds to Hcp1, we used a chromosomally encoded inner ring mutant of Hcp1, Hcp1^S115Q, which we previously showed lacks interaction with Tse1 (Fig. S4A). An attacker strain P. aeruginosa PAO1 ΔrsmA hcp1^S115Q was still able to kill E. coli (Fig. 6C), showing that the hcp1 mutation itself does not stop T6SS activity. Finally, when this strain expressed Tse1-HA-Bla, E. coli was killed to the same level as with the parental attacker, showing that in the absence of interaction between Tse1-HA-Bla and Hcp1^S115Q, there is no longer blocking of the T6SS.

From the above-described experiments, we conclude that unusual Hcp-effector complexes, notably, if the effector is large, may block Hcp ring stacking and therefore extension of the T6SS sheath. This was further validated by monitoring T6SS sheath formation using a P. aeruginosa strain carrying a double rsmA pppA mutation and expressing a GFP-tagged version of TssB. The pppA mutation results in a larger number of T6SS firing events and allows robust quantification of the number of T6SS sheaths formed in each strain (43). We used fluorescence microscopy to assess the number of T6SS firing events per cell for 1 min for each strain (Fig. 7). Firing events were determined by elongation of fluorescent foci into a sheath and then contraction back into a focus and sometimes disassembly (disappearance of foci). In the parental P. aeruginosa PAO1 ΔrsmA ΔpppA strain, ~5.5% of the cells showed a firing event within 1 min, while in the strain lacking hcp1, no firing events were observed after examining 31,336 cells. When the Tse1-HA variant was expressed instead of Tse1, there was still ~5.5% of cells that showed a firing event within 1 min; however, when Tse1-HA-mScarlet-I was expressed, this number decreased to ~0.7% (~8-fold fewer firing events). Together, these data show that loading large chimeric effectors onto Hcp1 results in a lack of delivery of the chimera and impairs delivery of any other T6SS effectors by preventing normal assembly and contraction of the T6SS sheath.

**Phylogenetic analysis of Hcp domains does not discriminate specialized versus nonspecialized proteins.** The amino acid sequence of Hcp proteins and their predicted structures are highly conserved. Therefore, it may be hard to identify key residues in the Hcp protein which may account for specific interactions with a given effector. This seems to be true for the S31 residue in Hcp1, which is involved in the interaction with Tse2 but does not play a role in the interaction with Tse1, Tse3, or Tse4 (34, 35). Other inner ring residues, such as S115 and T59, have a broader impact, and substitution of those amino acids results in loss of interaction with all known Hcp cargoes.

In specialized Hcps which contain a C-terminal domain accounting for the toxic activity, specific interaction might not be required. Therefore, if key residues required for cargo recognition existed, they might be unimportant in this context. We constructed a phylogenetic tree using ~61,461 Hcp proteins retrieved from the Integrated Microbial
Genomes (IMG) database (44) (Fig. 8A) and including core Hcps, such as Hcp1 to -3 from *P. aeruginosa* (33). The core Hcp domain is 160 amino acids long, and so we categorized Hcp proteins with an extra C-terminal extension of 60 amino acids long as specialized. These specialized Hcps are colored in black in the first layer of the tree. The second layer denotes the experimentally validated toxic domains for specialized Hcps (33). As seen from the distribution, there is no clustering of specialized Hcp proteins and core Hcps, which suggests that there are no major differences in sequence or structure between unspecialized and specialized Hcp core domains. Conserved residues in both categories of Hcp core domains do not indicate any obvious differences (Fig. 8B), which mean that those residues are most likely involved in the folding of the proteins and

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**FIG 6** Large Tse1 chimera can bind Hcp1 but block H1-T6SS-dependent killing. (A) Western blot of Hcp1-His and Tse1<sup>C30A</sup>-HA-Bl<sub>TEM-1</sub> copurification; lanes labeled on the top are postexpression sample, insoluble and soluble samples after sonication and clarification, flowthrough (FT) from the Ni-NTA column, wash fraction before elution, Ni fractions corresponding to the elution peak, and SEC fractions corresponding to the gel filtration peak. The antibodies used are labeled on the right: top, anti-HA antibody (BioLegend) at 1:1,000 concentration; bottom, anti-His antibody (GenScript) at 1:1000 concentration. Molecular weight standards are on the left. Asterisk denotes the correct size of Tse1<sup>C30A</sup>-HA-Bl<sub>TEM-1</sub>, at 46.4 kDa. (B) SEC chromatograph of purified Hcp1-His and Tse1<sup>C30A</sup>-HA-Bl<sub>TEM-1</sub>. (C) Competition assay of *P. aeruginosa* PAO1 ΔrsmA attackers against *E. coli*-GFP prey after a 13-h competition. GFP output represents the number of *E. coli* cells in the sample. Statistical testing was conducted by one-way ANOVA with Dunnett multiple-comparison test and family-wise significance and confidence level set to *P* = 0.05; each strain compared to parental *P. aeruginosa* PAO1 ΔrsmA strain. ****, *P* < 0.0001; ***, *P* < 0.001; ns, not significant. Mean and SEMs of *n* biological replicates: *n* = 6, 6, 4, 6, 6, 6, 1, 5, and 3, in order of strains.
formation of Hcp rings. Overall, this suggests that interaction between cargo effectors and core Hcp domains relies on an overall structural fold and fit within the inner ring rather than on specific motif recognition.

**DISCUSSION**

The idea of Hcp being a dual component with structural and effector function is akin to what has been shown for VgrG, which is a puncturing device but also interacts directly with effectors or carries effector extensions. However, there was very little functional demonstration of evolved Hcps, except for a few in *E. coli* (33). In addition, there were very few effectors identified that directly interact with Hcp, except for Tse1 to -3 from *P. aeruginosa* (34). Here, by using an *in vivo* pulldown approach, we showed that Hcp-bound proteins are probably as abundant as VgrG-bound effectors. Using 3 distinct Hcps from *P. aeruginosa*, we pulled down several effector candidates, two of which, PA2066 and PA0256, were validated for their direct interaction with Hcp and displayed antibacterial activity. The prominent role of Hcp as a shuttle for effectors also has important implications for the number of effectors which could be injected by one T6SS device upon a single contraction. Indeed, the trimeric VgrG spike likely carries at most 3 effectors, one per monomer, while one additional effector could be attached to the monomeric PAAR protein; this would result in the delivery of a total of 4 effectors per T6SS expulsion. Considering the sheath wraps around the Hcp stack and expands across the cytoplasm, one would expect more than 100 Hcp rings to be released from each firing event (45). If every ring was loaded with an effector, it becomes obvious that the main T6SS toxic payload is linked with Hcp and not VgrG or PAAR.

One observation we made when performing the pulldown experiments with *P. aeruginosa* Hcp1, Hcp2, and Hcp3 was that there were no common hits between the lists of identified putative effectors, which suggests a specificity in the Hcp-effector interaction. Also of note is the fact that there seem to be few Hcp2 and Hcp3 effectors...
compared to the large number of hits pulled down with Hcp1. Sequence and structural comparisons between the 3 Hcp proteins showed they all have the same highly conserved residues, which likely account for their conserved structure. Yet some residues might be described as a key feature for Hcp-effector interaction. For example, a conserved threonine residue within the inner Hcp ring is shown, when substituted for a glutamine, to abrogate interaction between all 3 Hcps and their cognate effectors (34). This of course may not account for specificity but rather for a good steric match.

FIG 8 Phyllogenetic tree of Hcp and HMM of specialized (evolved) and unspecialized Hcp. (A) The 61,461 Hcp domains were clustered and used to construct the phylogenetic tree. Each leaf represents an Hcp cluster representative. In the first layer, specialized Hcps are colored black. Second layer shows known C-terminal toxic Pfam domains in specialized Hcps. If there is no recognizable effector domain, they are shown without color, and they are colored green, red, or blue if they have a pyocin S-type, DUF2235 phospholipase, or colicin DNase domain, respectively. (B) HMM logo of cluster representatives of specialized and unspecialized Hcps.
between the Hcp and its substrate. In terms of specificity, one key Hcp residue, S31, has been proven to impact Hcp1 interaction with Tse2 only but not with other Hcp1 partners (34). It is obvious that one would need to have a better grasp of how Tse2, and possibly Tse1 and Tse3, fits within the Hcp1 ring to assess how S31 would only impact the interaction with Tse2. The existence of specific residues within these effectors is thus yet to be clearly proven.

Understanding the factors that drive the recognition of a cargo by Hcp is one pending question, while another key question regarding Hcp-dependent delivery is how the effector interacts with Hcp without disturbing its packing, which is crucial for sheath polymerization (46, 47). From previous studies, it is proposed that the effector may just fit into the ring, as for Hcp1-Tse2 (34). The presence of the effector within a Hcp tube has been shown using cryo-electron tomography not only in isolated Hcp rings but within an intact T6SS-like system, called a metamorphosis-associated contractile structure (MACS), in which the presence of the effector Mif1 was detected as a filled density in the structure as opposed to the condition where Mif1 was lacking (15, 48). To match the inner diameter of the ring and not protrude far out, the size of the effector might be a limiting factor. Here, we showed that addition of a small tag on Tse1 does not interfere with secretion, but addition of a large protein of 30 kDa, despite not hindering the interaction of the chimera with Hcp, arrests sheath assembly and secretion, suggesting that Hcp1-Tse1 complexes no longer stack on top of each other. One possibility which cannot be ruled out is that regions of larger effectors protruding from the Hcp lumen may be lodged at the interface between the Hcp ring and the surrounding sheath (49) but may need these domains to be unfolded.

In our study, we have shown that the breadth of Hcp-effector pairs is likely far beyond our current knowledge. Understanding how this partnership occurs specifically and how these assemblies fit into the T6SS sheath would likely call for detailed structural analyses of the Hcp-effector complexes. This is an area that deserves thorough investigation, since Hcp proteins not only are central to the T6SS mechanism and effector delivery but are antigenic (9) and good targets for potential anti- \textit{P. aeruginosa} vaccine development (50).

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and growth conditions.** The bacterial strains used in this study are listed in Table S1 in the supplemental material. Plasmids used in this study are listed in Table S2. Bacteria were cultured in lysogeny broth (LB) (Miller) or tryptic soy broth (TSB) (Sigma) or on LB agar (Miller) plates. Unless mentioned otherwise, bacteria were grown at 37°C with agitation. Media were supplemented with ampicillin (50 \( \mu \)g/ml) and chloramphenicol (34 \( \mu \)g/ml) for \textit{E. coli} and carbenicillin (100 \( \mu \)g/ml) streptomycin (2,000 \( \mu \)g/ml), and tetracycline (100 \( \mu \)g/ml) for \textit{P. aeruginosa}. Isopropyl-\( \beta \)-D-thiogalactopyranoside (IPTG) was used for \textit{E. coli} at 50, 100, or 500 \( \mu \)M, where stated, and for \textit{P. aeruginosa} at 250 \( \mu \)M.

The \textit{E. coli} DH5\( _a \) strain was used as the cloning host, and \textit{E. coli} BL21 (DE3) was for protein expression. \textit{P. aeruginosa} chromosomal mutants were constructed as previously described (51). pKNG101 was transferred via three-partner conjugation with \textit{E. coli} CC118 \( \Delta pir \) donor and \textit{E. coli} 1047/prK2013 helper strains with counterselection by growth on 20\% (wt/vol) sucrose.

**Western blot analysis.** SDS-PAGE and Western blot were conducted as previously described (29). Antibodies used in this study are provided in Table 4.

**Protein purification.** To purify or copurify Hcp with potential effectors, Hcp proteins were expressed from pACYCDuet-1 and effectors from pET22b. Cells were grown in 1 liter TSB supplemented with appropriate antibiotics to an optical density at 600 nm (OD\(_{600}\) of 0.6, expression from the plasmids was induced with 500 \( \mu \)M IPTG, and cells were grown at 18°C. Cells were harvested and resuspended in 40 ml buffer (50 mM Tris, 500 mM NaCl, 20 mM imidazole, pH 8) supplemented with 1 tablet of cOmplete protease inhibitor cocktail (Roche). The resuspended cell pellet was sonicated, and insoluble and soluble samples were separated by centrifugation (18,000 \( \times \) g, 4°C, 45 min). The supernatant was filtered (0.45 \( \mu \)m) and applied to Ni-NTA resin columns (HisTrap Fast Flow, 5 ml; GE Healthcare) pre- and copurified with resuspension buffer, using an ÄKTA prime plus system (GE Healthcare). The column was washed with 150 ml resuspension buffer before elution with 40 ml elution buffer (50 mM Tris, 500 mM NaCl, 400 mM imidazole, pH 8). Size exclusion chromatography (SEC) of the eluted fractions corresponding to the peak was performed using a Superdex 200 10/300 column (GE Healthcare) pre-equilibrated in 50 mM Tris, 150 or 250 mM NaCl, pH 8 (SEC of Hcp proteins alone and of Hcp1-His with HA-Tse2 was performed using 250 mM NaCl, whereas SEC of other Hcp-effector combinations was performed using 150 mM NaCl). Presence of both Hcp and effector was assessed by Coomassie stain and Western blot analysis.

**Electron microscopy and image processing.** Ten microliters of the Hcp1 sample diluted to 0.01 mg/ml was deposited onto glow-discharged copper 400 mesh grids with a continuous carbon film
TABLE 4 Antibodies used in this study

| Antibody* | Host | Serum      | Dilution | Source          |
|-----------|------|------------|----------|-----------------|
| Anti-HA   | Mouse| Monoclonal | 1:1,000  | BioLegend       |
| Anti-HisA | Mouse| Monoclonal | 1:1,000  | GenScript       |
| Anti-Hcp1 | Rabbit| Polyclonal | 1:500    | Laboratory collection |
| Anti-Hcp2 | Rabbit| Polyclonal | 1:500    | Laboratory collection |
| Anti-RpoB | Mouse| Monoclonal | 1:5,000  | NeoClone       |
| HRP-conjugated anti-mouse | Rabbit| Polyclonal | 1:5,000  | Sigma-Aldrich |
| HRP-conjugated anti-rabbit | Goat | Polyclonal | 1:5,000  | Sigma-Aldrich |

*HRP, horseradish peroxidase.

(Agar Scientific). After incubation for 2 min, the sample was washed twice with water before being stained with 2% (wt/vol) uranyl acetate for 1 min. Two hundred micrographs were collected with a 4 k by 4 k TVIPS camera on an FEI CM200 microscope operating at 200 kV at a magnification of 30,000× with a defocus range of −1.5 to −3.0 μm. All subsequent data processing steps were performed with cryoSPARC v3.0.1 (52). The contrast transfer function parameters for each micrograph were calculated with CTFFIND4 (53). Approximately 1,500 particles were manually picked to generate representative two-dimensional (2D) class averages for automated particle picking of the entire data set. A total of 85,598 particles were autopicked and extracted using a 256-by 256-pixel-size box. After several rounds of 2D classification, 56,350 particles were kept in good classes showing top and side views.

**Bacterial toxicity assays.** E. coli BL21(DE3) cells expressing effectors from pET22b and Hcp proteins from pACYCDuet-1 were grown in LB supplemented with appropriate antibiotics and 2% glucose. Cells were normalized to an OD_{600} of 1 before serial dilution in phosphate-buffered saline (PBS). Twenty microliters of each dilution was spotted on LB agar containing the appropriate antibiotics and inducer (50 μg/ml IPTG for Tse4 and 100 μg/ml IPTG for pulldown hits) or repressor (1% glucose for Tse4 and 2% glucose for pulldown hits). Plates were grown overnight, and CFU were quantified. The number of biological replicates is given in each figure legend.

**Bacterial competition assays.** For *P. aeruginosa* competition assays against E. coli Top10/pRL662.5: gfp, overnight cultures in LB were washed with PBS and normalized to an OD_{600} of 3. One hundred microliters of each *P. aeruginosa* attacker and E. coli prey were mixed at a 1:1 ratio. Five microliters of the attacker-prey mix was spotted on LB agar, dried, and incubated for 13 h. CFU of the input bacteria were checked to confirm the same numbers of prey and attacker were in each competition. After incubation, competition spots were scraped into 1 ml PBS, resuspended, and serially diluted. Triplicate serial dilutions were spotted on LB agar plates and grown overnight. The 10^{-2} dilution spots were scraped into 1 ml PBS and resuspended. The GFP fluorescence intensity in each triplicate competition spot, which represents the number of E. coli/pRL662.5:gfp present, was measured using a microplate reader (BMG Labtech). The number of biological replicates is given in each figure legend.

**FLAG-bead pulldown.** *P. aeruginosa* PA01 ΔrsmA strains expressing Hcp1/2/3/FLAG, Hcp1/2/3 (no tag), and VgrG1b/2b/3-FLAG from pME6032 were grown overnight in TSB supplemented with 100 μg/ml tetracycline and then subcultured to OD_{600} of 0.1 in TSB supplemented with 30 μg/ml tetracycline. After growth to OD_{600} of 0.3, expression was induced with 250 μg/ml IPTG. For the H1-T6SS proteins (Hcp1-FLAG, Hcp1, and VgrG1b-FLAG), cultures were grown at 37°C for 5.5 h. For the H2-T6SS proteins (Hcp2-FLAG, Hcp2, and VgrG2b-FLAG), cultures were grown at 25°C for 13 h. For the H3-T6SS proteins (Hcp3-FLAG, Hcp3, and VgrG3-FLAG), cultures were grown at 25°C for 24 h. Cultures at an OD_{600} of 200 were harvested by centrifugation (4,000 × g, 4°C, 20 min), and the pellet was resuspended in 50 ml PBS. The cells were cross-linked using dimethyl 3,3′-dithiobispropionimidate (DTBP) (Thermo Fisher Scientific) to a final concentration of 5 mM, and the mix was incubated at room temperature for 45 min. Tris-HCl (pH 7.6) was added to a final concentration of 20 mM for 15 min to quench the remaining DTBP. The cross-linked cells were centrifuged (4,000 × g, 4°C, 20 min), washed with PBS, and centrifuged. The pellet was resuspended in 35 ml PBS and sonicated. This sample was centrifuged (18,000 × g, 4°C, 45 min) to separate the soluble and insoluble proteins, and the supernatant was collected. Forty microliters anti-FLAG M2 magnetic beads (Millipore) (prewashed in PBS) was added to each supernatant and mixed with gentle agitation at 4°C for 4 h to overnight. Beads and bound protein were separated from unbound protein using a magnetic rack by carrying out 8 washes in 1 ml PBS. The beads and bound protein were resuspended in 20 μl PBS and frozen using liquid nitrogen. Three independent experiments for each pulldown were performed.

**Mass spectrometry analysis.** Mass spectrometry analysis was conducted by the Plateforme Protéomique Structurale et Fonctionnelle at the Institut Jacques Monod, Paris. Briefly, samples underwent on-bead digestion with 12.5 mg/ml sequencing-grade trypsin (Promega) before peptide analysis on an Orbitrap Fusion Tribrid mass spectrometer coupled to an Easy-spray nanoelectrospray ion source and an Easy nano-LC Prowess Xevo 1000 liquid chromatography system (Thermo Scientific). Chromatographic separation of the peptides was achieved by an Acclaim PepMap 100 C_{18} precolumn and a PepMap-RSLC Prowess C_{18} column at a flow rate of 300 μl/min. The solvent gradient consisted of 95% solvent A (water, 0.1% [vol/vol] formic acid) to 35% solvent B (100% acetonitrile, 0.1%[vol/vol] formic acid) over 98 min. An Orbitrap mass spectrometer analyzed the peptides in full ion scan mode, with the resolution set at 120,000 at a m/z mass range of 350 to 1550. High energy collision-induced dissociation activation with a collisional energy of 28% permitted fragment acquisition with the quadrupole isolation width of 15
cargo proteins were retrieved from https://www.pseudomonas.com. VgrG1b; Hcp2, VgrG2b; Hcp3, VgrG3) using the same cutoff. Isoelectric point data for the putative Hcp1-3 cargo proteins were retrieved from http://isoelectricpointdb.org/.

Control immunoprecipitation performed using a FLAG-tagged version of the Hcp

with 90% acetone, air dried, and resuspended to an OD600 of 10 in Laemmli buffer. Whole-cell lysate and supernatant samples were assessed by SDS-PAGE and Western blotting.

sequence conservation was assessed and displayed in Jalview version 2.10.1. (61) using percentage identity. Hcp1 to -3 sequence alignment was performed using Clustal Omega multiple sequence alignment, and each HMM was made and visualized using Skylign (67). Clustal Omega (64), and the phylogenetic tree was constructed using FastTree (65). The phylogenetic tree was visualized using iTOL (66).

Secretion assays. P. aeruginosa PAO1 expressing Tse4 variants from pMMB67HE were grown overnight in TSB supplemented with 100 μg/ml carbenicillin and then subcultured to an OD600 of 0.1 in 25 ml TSB supplemented with 50 μg/ml carbenicillin. Strains were grown to an OD600 of 0.3 for induction with 250 μM IPTG and then grown a further 4 h. Cells at an OD600 of 1 were harvested for the whole-cell lysate sample and resuspended in 100 μl Laemmli buffer. The supernatant was clarified by centrifugation (4,000 × g, 4°C, 10 min), and proteins were precipitated with 10% trichloroacetic acid (TCA), washed with 90% acetone, air dried, and resuspended to an OD600 of 10 in Laemmli buffer. Whole-cell lysate and supernatant samples were assessed by SDS-PAGE and Western blotting.

Microscopy. P. aeruginosa strains were grown in TSB and then subcultured to an OD600 of 0.1 in TSB and grown for 5 h. One microliter of each culture was spotted onto a glass-bottomed 35-mm petri dish (μ-Dish 35 mm, high glass bottom; Ibidi) and covered with a 1% (wt/vol) UltraPure agarose (Invitrogen) dissolved in PBS) pad. Imaging was performed using an Axio Observer Z1 (Zeiss) inverted widefield microscope with a Plan-Apochromat 100×, 1.4 oil Ph3 M27 lens objective (Zeiss), a SpectraX light-emitting diode (LED) light engine (Lumencore), an ORCA-Flash 4.0 digital complementary metal oxide semiconductor (CMOS) camera (Hamamatsu), and an environmental control system and microscope chamber set to 37°C. Phase contrast and fluorescence (exposure time of 200 ms for superfolder GFP [sfGFP]) images were taken to calculate the total cells and number of foci, followed by a fluorescence time-series with images acquired every 2 s for a total of 180 s. Microscopy was performed at Facility for Imaging by Light Microscopy (FILM) Imperial College London. Microscopy image analysis was performed in FIJI Imagej version 2.1.0/1.53c (60). “Threshold” was used to highlight cells and then “Analyze Particles” was used to count the cells in each image. The “Multi-point” tool was used to manually count the number of foci in an image. Bleach correction was applied to time-series experiments; each extension-contraction event was counted manually using the Multi-point tool, and the final number of events per minute was calculated.

Bioinformatic analysis. Statistical analyses were undertaken in GraphPad Prism 8 version 8.4.3. Hcp1 to -3 sequence alignment was performed using Clustal Omega multiple sequence alignment, and sequence conservation was assessed and displayed in Jalview version 2.10.1. (61) using percentage identity with the conservation threshold set to 50%. Phyre2 (62) was used to predict the structure of Hcp3. Hcp1 to -3 protein structures were analyzed and annotated in PyMOL molecular graphics system.

Phylogenetic tree construction. All genes with Hcp Pfam domain (PF05638) were collected from the IMG database (44) from Gram-negative bacteria and metagenomes (~267,000 genes were collected). Hcps found in small metagenomic scaffolds of < 5 kb were removed as well as genes containing the PEP-CTERM sorting signal (PF07589). Additionally, Hcp domains shorter than 70 amino acids were also removed. Based on the Hcp hidden Markov model (HMM) (PF05638), Hcp envelope domain boundary was determined using HMMER and then clustered at 60% identity and 80% length using CD-HIT (63) into 2,340 unique clusters in order to prevent redundancy. Cluster representatives were aligned using Clustal Omega (64), and the phylogenetic tree was constructed using FastTree (65). The phylogenetic tree was visualized using iTOL (66).

HMM’s of specialized and unspecialized Hcps. Cluster representatives used for the construction of the phylogenetic tree were divided into representatives that came from a specialized Hcp (2,226 cluster representatives) and unspecialized Hcp (114 cluster representatives). These were aligned using Clustal Omega, and each HMM was made and visualized using Skyline (67).
SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

DATA SET S1, XLSX file, 1 MB.
FIG S1, TIFF file, 2 MB.
FIG S2, TIFF file, 1.3 MB.
FIG S3, TIFF file, 0.9 MB.
FIG S4, TIFF file, 1.1 MB.
FIG S5, TIFF file, 1.1 MB.
FIG S6, TIFF file, 0.5 MB.
FIG S7, TIFF file, 1 MB.
TABLE S1, DOCX file, 0.1 MB.
TABLE S2, DOCX file, 0.1 MB.

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Sophie A. Howard performed the main experiments, Dora Bonini performed some of the bacterial genetic experiments, Himani Amin performed the electron microscopy, Christopher D. Furniss performed the mass spectrometry data analyses, Patricia Paracuellos performed the protein structure modeling, and David Zlotkin performed the phylogenetic analyses. Alain Filloux and Sophie A. Howard designed the whole study, Despoina A. I. Mavridou designed the biochemical experiments, Tiago R. D. Costa designed the electron microscopy studies, and Asaf Levy designed the phylogeny studies.

REFERENCES

1. Costa TR, Felisberto-Rodrigues C, Meir A, Prevost MS, Redzej A, Trokter M, Waksman G. 2015. Secretion systems in Gram-negative bacteria: structural and mechanistic insights. Nat Rev Microbiol 13:343–359. https://doi.org/10.1038/nrmicro3456.
2. Green ER, Mecas J. 2016. Bacterial secretion systems: an overview. Microbiol Spectr 4:VMBF-0012-2015. https://doi.org/10.1128/microbiolspec.VMBF-0012-2015.
3. Leiman PG, Arisaka F, van Raaij MJ, Kostyuchenko VA, Aksyuk AA, Kanamaru S, Rossmann MG. 2010. Morphogenesis of the T4 tail and tail fibers. Virol J 7:355. https://doi.org/10.1186/1743-422X-7-355.
4. Veesler D, Cambillau C. 2011. A common evolutionary origin for tailed-bacteriophage functional modules and bacterial machineries. Microbiol Mol Biol Rev 75:423–433. https://doi.org/10.1128/MMBR.00014-11.
5. Clemens DL, Ge P, Lee BY, Horwitz MA, Zhou ZH. 2015. Atomic structure of T6SS reveals interlaced array essential to function. Cell 160:940–951. https://doi.org/10.1016/j.cell.2015.02.005.
6. Kudryashev M, Wang RY, Brackmann M, Scherer S, Maier T, Baker D, DiMaio F, Stahlberg H, Egelman EH, Basler M. 2015. Structure of the type VI secretion system contractile sheath. Cell 160:952–962. https://doi.org/10.1016/j.cell.2015.01.037.
7. Salih O, He S, Planamente S, Stach L, MacDonald JT, Manoli E, Scheres SHW, Filloux A, Freemont PS. 2018. Atomic structure of type VI contractile sheath from Pseudomonas aeruginosa. Structure 26:329.e3–336.e3. https://doi.org/10.1016/j.str.2017.12.005.
8. Wang J, Brackmann M, Castano-Diez D, Kudryashev M, Goldie KN, Maier T, Stahlberg H, Basler M. 2017. Cryo-EM structure of the extended type VI secretion system sheath-tube complex. Nat Microbiol 2:1507–1512. https://doi.org/10.1038/s41564-017-0020-7.
9. Mouguous JD, Cuff ME, Raunser S, Shen A, Zhou M, Gifford CA, Goodman AL, Joachimiak G, Ordenez CL, Lory S, Walz T, Joachimiak A, Mekalanos JJ. 2006. A virulence locus of Pseudomonas aeruginosa encodes a protein secretion apparatus. Science 312:1526–1530. https://doi.org/10.1126/science.1128393.
10. Basler M, Pilhofer M, Henderson GP, Jensen GJ, Mekalanos JJ. 2012. Type VI secretion requires a dynamic contractile phage tail-like structure. Nature 483:182–186. https://doi.org/10.1038/nature10846.
11. Ballister ER, Lai AH, Zuckermann RN, Cheng Y, Mougous JD. 2008. In vitro self-assembly of tallarable nanotubes from a simple protein building block. Proc Natl Acad Sci U S A 105:3733–3738. https://doi.org/10.1073/pnas.0712247105.
12. Brunet YR, Henin J, Celis H, Cascales E. 2014. Type VI secretion and bacteriophage tail tubes share a common assembly pathway. EMBO Rep 15:315–321. https://doi.org/10.1002/embr.201337936.
13. Renault MG, Zamarreno Beas J, Douzi B, Chabalier M, Zoued A, Brunet YR, Cambillau C, Journet L, Cascales E. 2018. The gp27-like hub of VgrG serves as adaptor to promote Hcp tube assembly. J Mol Biol 430:3143–3156. https://doi.org/10.1016/j.jmb.2018.07.018.
14. Shneider MM, Buth SA, Ho BT, Basler M, Mekalanos JJ, Leiman PG. 2013. PAAR-repeat proteins sharpen and diversify the type VI secretion system spike. Nature 500:350–353. https://doi.org/10.1038/nature12453.
15. Howard SA, Filloux A. 2019. Looking inside an injection system. Elife 8: e50815. https://doi.org/10.7554/eLife.50815.
16. Russell AB, Peterson SB, Mogus JD. 2014. Type VI secretion system effectors: poisons with a purpose. Nat Rev Microbiol 12:137–148. https://doi.org/10.1038/nrmicro3185.

17. Trunk K, Pellet J, Liu YC, Dill BD, Walker L, Gou NAR, Stark MJR, Quinn J, Stahl H, Trost M, Couthurst SJ. 2018. The type VI secretion system deploys antifungal effectors against microbial competitors. Nat Microbiol 3:920–931. https://doi.org/10.1038/s41564-018-0191-x.

18. Jana B, Fridman CM, Bosi E, Salomon D. 2019. A modular effector with a DnaE-like domain and a marker for T6SS substrates. Nat Commun 10:3395. https://doi.org/10.1038/s41467-019-11546-6.

19. Salomon D, Kinch LN, Trudgian DC, Guo X, Klimko JA, Grishin NV, Mirzaei H, Orth K. 2014. Marker for type VI secretion system effectors. Proc Natl Acad Sci U S A 111:9271–9276. https://doi.org/10.1073/pnas.140611011.

20. Hachani A, Allopp LP, Oduyo Y, Filouix A. 2014. The VgrG proteins are “a la carte” delivery systems for bacterial type VI effectors. J Biol Chem 289:17872–17880. https://doi.org/10.1074/jbc.M114.563429.

21. Ahmad S, Tsang KK, Sachar K, Quentin D, Tashin TM, Bullen NP, Raunser S, Mougous JD. 2015. VgrG and PAAR proteins define distinct versions of a functional type VI secretion system. PLoS Pathog 12:e1005735. https://doi.org/10.1371/journal.ppat.1005735.

22. Alcoforado Diniz J, Coulthurst SJ. 2015. Intraspecies competition in Serratia marcescens is mediated by type VI-secreted Rhs effectors and a conserved effector-associated accessory protein. J Bacteriol 197:2350–2360. https://doi.org/10.1128/JB.00199-15.

23. Cianfanelli FR, Alcoforado Diniz J, Guo M, De Cesare V, Trost M, Coulthurst SJ. 2016. Francisella tularensis IglG belongs to a novel family of PAAR-like T6SS proteins and harbors a unique N-terminal extension required for virulence. PLoS Pathog 12:e1005821. https://doi.org/10.1371/journal.ppat.1005821.

24. Bondage DD, Lin JS, Ma LS, Kuo CH, Lai EM. 2016. VgrG C terminus confers the type VI effector transporter specificity and is required for binding with PAAR and adaptor-effector complex. Proc Natl Acad Sci U S A 113:3931–3936. https://doi.org/10.1073/pnas.1600428113.

25. Unterweger D, Kostiuk B, Otjengerdes R, Wilton A, Diaz-Satizabal L, Pukatzki S. 2015. Chimeric adaptor proteins translocate diverse type VI secretion system effectors in Vibrio cholerae. EMBO J 34:2198–2210. https://doi.org/10.15252/embj.201591163.

26. Rigard M, Boms JE, Mosnier A, Hologne M, Martin A, Lindgren L, Punginelli C, Labs C, Walker O, Charbit A, Pelouk P, Conlan W, Terradot L, Sjostedt A, Henry T. 2016. Towards a complete structural deciphering of type VI secretion system components measured by targeted proteomics. Nat Commun 10:3595. https://doi.org/10.1038/s41467-019-11548-6.

27. Whitney JC, Quentin D, Sawai S, LeRoux M, Harding BN, Ledvina HE, Tran BQ, Robinson H, Goo YA, Goodlett DR, Meraz AJ, Goodlett DR, Mogus JD. 2010. A type VI secretion system of Pseudomonas aeruginosa targets a toxin to bacteria. Cell Host Microbe 7:25–37. https://doi.org/10.1016/j.chom.2009.12.007.

28. Ahmad S, Tsang KK, Sachar K, Quentin D, Tashin TM, Bullen NP, Raunser S. 2015. The Hcp proteins fused to diverse extended-domains represent a novel pattern of anti-bacterial effectors in type VI secretion systems. Mol Cell 51:584–593. https://doi.org/10.1016/j.molcel.2013.07.025.

29. Whitney JC, Beck CM, Goo YA, Russell AB, Harding BN, De Leon JA, Cunningham DA, Tran BQ, Low DA, Goodlett DR, Hayes CS, Mogus JD. 2014. Genetically distinct pathways guide effector export through the type VI secretion system. Mol Microbiol 92:529–542. https://doi.org/10.1111/mmi.12571.

30. Ting SY, Bosch DE, Mangiameli SM, Radey MC, Huang S, Park YJ, Kelly KA, Filippo SK, Goo YA, Eng JK, Allaire M, Veesler D, Wiggins PA, Peterson SB, Mogus JD. 2018. Bifunctional immunity proteins protect bacteria against FtsZ-targeting ADP-ribosylating toxins. Cell 175:1380.e14–1392.e14. https://doi.org/10.1016/j.cell.2018.09.037.

31. Basler M, Ho BT, Mekalanos JJ. 2013. Tip-for-tat: type VI secretion system counterattack during bacterial cell-cell interactions. Cell 152:884–894. https://doi.org/10.1016/j.cell.2013.01.042.

32. Markowitz VW, Chen IM, Chu K, Szeto E, Palaniappan K, Grechkin Y, Ratner A, Babi B, But A, Huntemann M, Liolios K, Pagani I, Anderson I, Kanevski M, Ivanova NN, Kyrpides NC. 2012. IMG/M, the integrated metagenome data management and comparative analysis system. Nucleic Acids Res 40:D123–D129. https://doi.org/10.1093/nar/gkr975.

33. Lin L, Lezan E, Schmidt A, Basler M. 2019. Abundance of bacterial type VI secretion system components measured by targeted proteomics. Nat Commun 10:2584. https://doi.org/10.1038/s41467-019-10466-9.

34. Silverman JM, Agnellu DM, Zheng H, Andrews BT, Li M, Catalano CE, Genten T, Mogus JD. 2013. Human and corneal epithelial protein is an exported receptor and chaperone of type VI secretion substrates. Mol Cell 51:584–593. https://doi.org/10.1016/j.molcel.2013.07.025.

35. Bailey M, Steiner L, DiPaolo SF, Brubaker MA. 2017. cyoSPARC: algorithms for rapid unsupervised cyo-EM structure determination. Nat Methods 14:290–296. https://doi.org/10.1038/nmeth.4169.
53. Rohou A, Grigorieff N. 2015. CTFIND4: fast and accurate defocus estimation from electron micrographs. J Struct Biol 192:216–221. https://doi.org/10.1016/j.jsb.2015.08.008.

54. Tyanova S, Temu T, Cox J. 2016. The MaxQuant computational platform for mass spectrometry-based shotgun proteomics. Nat Protoc 11:2301–2319. https://doi.org/10.1038/nprot.2016.136.

55. Cox J, Neuhauer N, Michalski A, Scheltema RA, Olsen JV, Mann M. 2011. Andromeda: a peptide search engine integrated into the MaxQuant environment. J Proteome Res 10:1794–1805. https://doi.org/10.1021/pr101065j.

56. Cox J, Hein MY, Luber CA, Paron I, Nagaraj N, Mann M. 2014. Accurate proteome-wide label-free quantification by delayed normalization and maximal peptide ratio extraction, termed MaxLFQ. Mol Cell Proteomics 13:2513–2526. https://doi.org/10.1074/mcp.M113.031591.

57. Winsor GL, Griffiths EJ, Lo R, Dhillon BK, Shay JA, Brinkman FS. 2016. Enhanced annotations and features for comparing thousands of Pseudomonas genomes in the Pseudomonas genome database. Nucleic Acids Res 44:D646–S3. https://doi.org/10.1093/nar/gkw1227.

58. Kozlowski LP. 2017. Proteome-pI: proteome isoelectric point database. Nucleic Acids Res 45:D1112–D1116. https://doi.org/10.1093/nar/gkw978.

59. Rice P, Longden I, Bleasby A. 2000. EMBOSS: the European Molecular Biology Open Software Suite. Trends Genet 16:276–277. https://doi.org/10.1016/S0168-9525(00)02024-2.

60. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch S, Rueden C, Saalfeld S, Schmid B, Tinevez JY, White DJ, Hartenstein V, Eliceiri K, Tomancak P, Cardona A. 2012. Fiji: an open-source platform for biological-image analysis. Nat Methods 9:676–682. https://doi.org/10.1038/nmeth.1909.

61. Waterhouse AM, Procter JB, Martin DM, Clamp M, Barton GJ. 2009. Jalview Version 2--a multiple sequence alignment editor and analysis workbench. Bioinformatics 25:1189–1191. https://doi.org/10.1093/bioinformatics/btp033.

62. Kelley LA, Mezulis S, Yates CM, Wass MN, Sternberg MJ. 2015. The Phyre2 web portal for protein modeling, prediction and analysis. Nat Protoc 10:845–858. https://doi.org/10.1038/nprot.2015.053.

63. Li W, Godzik A. 2006. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. Bioinformatics 22:1658–1659. https://doi.org/10.1093/bioinformatics/btl158.

64. Madeira F, Park YM, Lee J, Buso N, Gur T, Madhusoodanan N, Basutkar P, Tiwey ARN, Potter SC, Finn RD, Lopez R. 2019. The EMBL-EBI search and sequence analysis tools APIs in 2019. Nucleic Acids Res 47:W636–W641. https://doi.org/10.1093/nar/gkz268.

65. Price MN, Dehal PS, Arkin AP. 2010. FastTree 2--approximately maximum-likelihood trees for large alignments. PLoS One 5:e9490. https://doi.org/10.1371/journal.pone.0009490.

66. Letunic I, Bork P. 2007. Interactive Tree Of Life (ITOL): an online tool for phylogenetic tree display and annotation. Bioinformatics 23:127–128. https://doi.org/10.1093/bioinformatics/btm529.

67. Wheeler TJ, Clements J, Finn RD. 2014. Skylign: a tool for creating informative, interactive logos representing sequence alignments and profile hidden Markov models. BMC Bioinformatics 15:7. https://doi.org/10.1186/1471-2105-15-7.