



# Haemolysin Coregulated Protein Is an Exported Receptor and Chaperone of Type VI Secretion Substrates

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#### **SUMMARY**

Secretion systems require high-fidelity mechanisms to discriminate substrates among the vast cytoplasmic pool of proteins. Factors mediating substrate recognition by the type VI secretion system (T6SS) of Gram-negative bacteria, a widespread pathway that translocates effector proteins into target bacterial cells, have not been defined. We report that haemolysin coregulated protein (Hcp), a ring-shaped hexamer secreted by all characterized T6SSs, binds specifically to cognate effector molecules. Electron microscopy analysis of an Hcpeffector complex from Pseudomonas aeruginosa revealed the effector bound to the inner surface of Hcp. Further studies demonstrated that interaction with the Hcp pore is a general requirement for secretion of diverse effectors encompassing several enzymatic classes. Though previous models depict Hcp as a static conduit, our data indicate it is a chaperone and receptor of substrates. These unique functions of a secreted protein highlight fundamental differences between the export mechanism of T6 and other characterized secretory pathways.

# INTRODUCTION

Bacteria possess discrete pathways for delivering proteins to their surroundings. A diversity of secretion systems facilitates the necessary heterogeneity in size, structure, and function within the proteins exported by these cells (Economou et al., 2006). For example, the type V secretion system can export exceptionally large proteins, such as filamentous adhesins, to the cell surface or the extracellular milieu (Dautin and Bernstein, 2007; Wagner et al., 2011), whereas types III, IV, and VII systems exhibit the ability to directly translocate proteins into host cells (Abdallah et al., 2007; Fronzes et al., 2009; Galán, 2009; Izoré et al., 2011). Of the thousands of proteins present in a cell,

each secretion system acts with extraordinary fidelity to specifically recognize and release a distinct subset of substrates.

Substrate selection can be affected by several factors, including signal sequences, chaperones, and receptors. For example, type III secretion system (T3SS) substrates are recruited to the secretory apparatus in complex with specialized chaperones (Akeda and Galán, 2005; Stebbins and Galán, 2001). A T3S-associated ATPase then engages the substrate by binding its N-terminal export signal, releasing it from the chaperone and driving substrate unfolding and translocation through the apparatus. Similar mechanisms appear to drive substrate recognition and export by the T4S pathway (Cambronne and Roy, 2007; Sutherland et al., 2012). While the factors involved in substrate selection for T3SSs and T4SSs are in part defined, comparatively little is known about this process in a related but more recently described pathway, the type VI secretion system (T6SS).

The T6SS is a widely distributed protein translocation pathway that is able to directly target bacterial and eukaryotic cells (Silverman et al., 2012). The core T6 machinery consists of 13 essential subunits, encoded within large gene clusters on bacterial chromosomes. While the T6SS ultrastructure has not been determined, X-ray crystallographic and protein-protein interaction data suggest that an envelope-spanning complex and a bacteriophage tail-like structure are core components of the secretory machinery (Cascales and Cambillau, 2012). The phage-like complex consists of TssB (type six secretion B), TssC, Hcp (haemolysin coregulated protein), and VgrG (valine-glycine repeat protein G). Visualization of Vibrio cholerae cells by fluorescence microscopy showed that TssB and TssC form a dynamic filamentous complex within the cytoplasm, switching between extended and contracted states while remaining anchored at the inner membrane (Basler et al., 2012). Hcp and VgrG are secreted components of the T6SS with significant structural homology to T4 bacteriophage tail tube (gp19) and tail spike proteins (gp5-gp27), respectively. By analogy to contractile bacteriophage, TssB and TssC are proposed to form a sheath enclosing a tube composed of Hcp, with VgrG localized at the tip to allow membrane breaching (Kanamaru, 2009; Kapitein and Mogk, 2013). Contraction of the TssB-TssC complex is postulated to drive Hcp and VgrG out of the cell. A recent





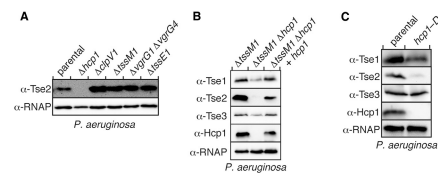


Figure 1. Tse2 Requires Hcp1 for Intracel**lular Accumulation** 

(A and B) Western blot analysis of intracellular levels of H1-T6S effectors (Tse1, Tse2, and Tse3) in the indicated P. aeruginosa backgrounds. RNA polymerase (RNAP) is included as a loading control. Unless otherwise indicated, the parental background in this and all subsequent figures is

(C) Western blot analysis of intracellular Hcp1 and H1-T6S effector levels in a ClpXP-dependent Hcp1 depletion assay. Samples were processed 90 min after induced sspB expression in P. aeruginosa strains lacking the native sspB gene and containing wild-type hcp1 or hcp1-D4.

report defining a complex composed of these proteins in Agrobacterium tumefaciens lends support to this model (Lin et al., 2013). Nevertheless, many aspects of the export mechanism for the T6SS remain undefined. In particular, it is not known how effectors are recognized for export and how they transit the T6S pathway.

Hcp is a central component of the bacteriophage-like model of T6S-dependent intercellular effector transport. Studies have demonstrated that this universally and abundantly T6S-exported protein is essential for both the assembly of the T6S apparatus and the export of its effectors (Hood et al., 2010; Mougous et al., 2006; Pukatzki et al., 2007). Nevertheless, the precise function of Hcp in the T6SS has not been addressed. Crystal structures of diverse Hcp homologs from several species show the protein adopts a homohexameric ring conformation with an ~40 Å internal diameter; furthermore, Hcp rings stack to form tubular structures within the crystal lattices (Jobichen et al., 2010; Mougous et al., 2006; Osipiuk et al., 2011). It has been speculated based on these data that the protein functions as a channel through which T6S effectors transit. However, Hcp tubes have only been observed under crystallographic conditions, or when stabilized in vitro with engineered disulfide bonds between ring subunits (Ballister et al., 2008). Also, since both stacking arrangements (head-to-head or head-to-tail) are found in Hcp X-ray crystal structures, the physiological significance of the Hcp tubule is uncertain.

The Hcp secretion island I (HSI-I)-encoded T6SS (H1-T6SS) of *P. aeruginosa* is a model system for studying the substrates, apparatus dynamics, and physiological significance of T6 (Basler and Mekalanos, 2012; Hood et al., 2010; LeRoux et al., 2012; Russell et al., 2011). The H1-T6SS delivers at least three toxic effectors, Tse1-3 (type VI secretion exported 1-3), into target bacterial cells. Tse1 and Tse3 are translocated into the periplasm of target cells, where they degrade peptidoglycan and cause cell lysis using amidase and muramidase activity, respectively (Russell et al., 2011). The precise molecular target of Tse2 is not yet known; however, this effector induces stasis and acts in the cytoplasm of recipient cells. Structure prediction algorithms indicate that Tse2 may function as a nuclease (Li et al., 2012). As the bacterial targeting activities of the H1-T6SS are also directed at kin, P. aeruginosa requires immunity proteins, Tsi1-3 (type VI secretion immunity 1-3), encoded alongside their cognate effectors in bicistrons. P. aeruginosa harbors two additional T6SSs, H2-T6SS and H3-T6SS. Interestingly, these systems appear to export nonoverlapping sets of effectors, though the factors mediating effector discrimination have not been elucidated (Russell et al., 2013).

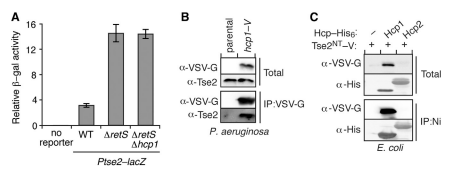
In this study, we report that Hcp is a chaperone and receptor of T6S effectors. Focusing on Tse2, we show that a direct and highly specific interaction with the pore of its cognate Hcp, Hcp1 of the H1-T6SS, is required for the stability and export of the protein. We further demonstrate that effectors with unrelated sequences, Tse1 and Tse3, also require direct interactions with the pore of Hcp1 for secretion. Certain amino acid substitutions within the Hcp1 pore differentially affect the secretion of the three Tse proteins, indicating the capacity of Hcp1 to bind diverse proteins via distinct epitopes. Finally, we probe the generality of our results and find that specific interactions between T6S effectors and cognate Hcp proteins occur in diverse bacteria. Overall, our findings reveal a new paradigm for substrate recognition by a complex bacterial protein translocation machine.

#### **RESULTS**

# **Hcp1 Stabilizes Tse2 Posttranslationally**

In the course of investigating factors that influence effector transport through the H1-T6SS of P. aeruginosa, we observed that strains lacking hcp1 displayed markedly reduced intracellular levels of Tse2 (Figure 1A). We conducted this and subsequent studies in the  $\Delta retS$  background, which is frequently employed for the study of the H1-T6SS (Basler and Mekalanos, 2012; Hachani et al., 2011; Mougous et al., 2006). The inactivation of retS overrides the requirement for cell-surface contact for T6S activation, and it results in elevated expression of both the H1-T6SS and its effectors, facilitating their detection via western blot (Hood et al., 2010; Silverman et al., 2011). Deletions of other essential T6SS components including clpV1, tssM1, tssE1, and the functionally redundant vgrG1 and vgrG4 genes did not lead to reduced Tse2 levels, indicating this phenotype is specific to the loss of hcp1 (Figure 1A). To measure more conclusively the effect of Hcp1 on overall Tse2 levels, we probed the consequences of hcp1 deletion in the ΔtssM1 background. The tssM1 gene encodes a core component of the secretory system; its deletion abrogates apparatus assembly





# Figure 2. Hcp1 Interacts Directly with Tse2

- (A) Relative levels of β-galactosidase activity from the indicated *P. aeruginosa* strains containing a chromosomally integrated *lacZ* reporter fused to the promoter region and first eight codons of *tse2* (*Ptse2–lacZ*). Error bars represent standard deviation based on three independent replicates.
- (B) Western blot analysis of Tse2 from total and bead-associated fractions of an anti-VSV-G immunoprecipitation from *P. aeruginosa* strains encoding Hcp1 or Hcp1-V.
- (C) Immunoblot detecting total and bead-

and effector export (Felisberto-Rodrigues et al., 2011; Mougous et al., 2006). Tse2 levels were also strongly reduced in the  $\Delta hcp1$   $\Delta tssM1$  background, suggesting that Hcp1 is required for maintenance of intracellular Tse2 (Figure 1B). We also examined whether the levels of Tse1 and Tse3 are sensitive to the presence of Hcp1. Though not as dramatic as the effect on Tse2, we detected a significant reduction of intracellular Tse1 and Tse3 upon deletion of hcp1 from the  $\Delta tssM1$  background (Figure 1B). The effects of  $\Delta hcp1$  on all effectors could be partially (Tse2) or fully (Tse1 and Tse3) genetically complemented with a plasmid expressing hcp1.

As an alternative approach for probing the influence of Hcp1 on effector levels, we used the ClpXP targeted degradation system to specifically deplete the protein from P. aeruginosa (Castang and Dove, 2012; McGinness et al., 2006). To accomplish this, we generated a P. aeruginosa strain wherein the native hcp1 open reading frame was modified to include a C-terminal fusion to the ssrA-like sequence, DAS+4 (Hcp1-D4). Consistent with our deletion studies, depletion of Hcp1-D4 resulted in a concomitant precipitous drop in cellular Tse2 levels (Figure 1C). Tse1 levels similarly decreased as observed in the Δhcp1 background, while Tse3 levels were not visibly affected by Hcp1 depletion. The cellular half-life of Tse3 may be too long to observe the relatively minor effect of Hcp1 during the 90 min depletion assay (Figures 1B and 1C). Together, these data demonstrate that the Hcp1 protein is essential for the maintenance of intracellular Tse2 levels. Though Tse1 and Tse3 levels are measurably influenced by hcp1 deletion, significant intracellular accumulation of these effectors occurs independent of the protein. To investigate the mechanism underlying the influence of Hcp1 on H1-T6S effectors, we initially focused on Tse2, where a highly distinct Hcp1-dependent phenotype was observed.

# **Hcp1 Interacts Directly with Tse2**

The dramatically lowered abundance of Tse2 in the absence of Hcp1 suggests that Hcp1—acting directly or indirectly—plays a critical role either in regulating *tse2* expression or in stabilizing Tse2 posttranslationally. To test the former, we generated *tse2* expression reporter strains containing a chromosomally integrated construct consisting of the predicted *tse2* promoter followed by an open reading frame consisting of the first eight codons of *tse2* fused to *lacZ*. Though the activity of the reporter

increased as expected in the  $\Delta retS$  background, a known posttranscriptional regulator of tse2, its activity was insensitive to the deletion of hcp1 (Figure 2A) (Brencic and Lory, 2009). Unable to implicate Hcp1 in a regulatory capacity, we probed for a physical interaction between the proteins using a coimmunoprecipitation (coIP) assay. Interestingly, Tse2 specifically precipitated with Hcp1, suggesting a physical association between this effector and Hcp1 might underlie the Hcp1 requirement for Tse2 stability (Figure 2B). To test whether the observed interaction requires other T6S-associated factors, we next performed coIP experiments from T6SS E. coli strains coexpressing C-terminally hexahistidine-tagged hcp1 (hcp1his<sub>6</sub>) and a vesicular stomatitis virus glycoprotein (VSV-G) epitope-tagged nontoxic allele of tse2 (tse2NT-V). In this heterologous host, the accumulation of Tse2 remained dependent upon the presence of Hcp1 (Figure 2C), Furthermore, coexpressed Tse2NT-V and Hcp1 associated tightly. To rule out nonspecific mechanisms that could explain our E. coli data, we tested the activity of another Hcp protein from P. aeruginosa, Hcp2, which does not participate in the H1-T6SS. Despite expression equivalent to or in excess of Hcp1, Hcp2 did not stabilize or interact with Tse2. From these data, we conclude that a direct interaction between Hcp1 and Tse2 likely promotes Tse2 stability.

# Tse2 Interacts with the Inner Surface of Hcp1 Rings

X-ray crystallographic and electron microscopic (EM) studies of Hcp proteins, including Hcp1, have consistently observed the protein in a hexameric ring configuration (Jobichen et al., 2010; Leiman et al., 2009; Mougous et al., 2006; Osipiuk et al., 2011). Using sedimentation velocity analytical ultracentrifugation at low Hcp1 concentrations (10.3–3.2  $\mu$ M), we further confirmed this quaternary state as the single detectable species in solution (see Figure S1 online). These observations, taken together with the large hydrophobic interface observed between Hcp protomers, argue that the hexameric assembly is the relevant physiological state of Hcp proteins.

The Hcp1 hexamer reveals several potential protein-protein interaction surfaces. Sequence analysis of Hcp homologs shows that residues on the "top" and "bottom" faces of Hcp rings are highly conserved and may be important for ringring interactions (Figure S2) (Mougous et al., 2006). Thus, we reasoned that Tse2 likely interacts with either the inside or



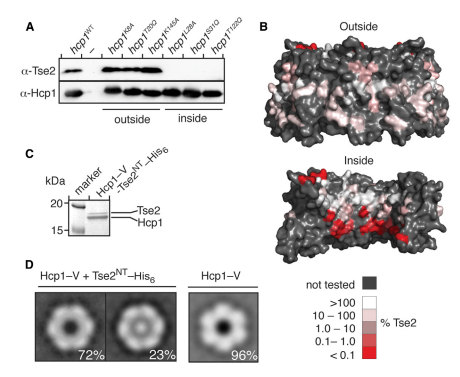


Figure 3. Tse2 Binds to the Pore of the Hcp1 Ring

(A) Representative western blots showing the effect of the indicated Hcp1 amino acid substitutions on intracellular levels of Tse2NT in E. coli. The localization of the substitutions to the inside or outside surface of the Hcp1 ring is noted.

(B) Surface representation of the Hcp1 ring colored to reflect Tse2 stabilization activity of each variant tested (white-red, 100%-0.01%; gray, not tested). The lower image depicts a cutaway view of the Hcp1 hexamer. Levels of Tse2 were calculated based on the ratio of band intensity of Tse2NT and Hcp1 point mutants, normalized to Tse2 and wild-type Hcp1 (Figure S3).

(C) Coomassie-stained SDS-PAGE gel of copurified Tse2<sup>NT</sup>-His<sub>6</sub> and Hcp1-V.

(D) Class averages with applied six-fold symmetry from analysis of transmission electron micrographs of Tse2NT-His6 -Hcp1-V and a Hcp1-V-only control. The percentage of particles represented by each class average is indicated in the corresponding frame. See also Figures S1-S4.

the outside face of the Hcp1 ring. To define the regions of Hcp1 involved in Tse2 interaction, we pursued a site-directed mutagenesis approach, targeting all residues with more than 60% solvent accessibility located on the inner or outer surface of the Hcp1 ring. To this end, we mutated 34 positions; nonpolar or small side chain amino acids were substituted with glutamine and those with a polar or large side chain with alanine. The relative effect of each Hcp1 point mutation on Tse2 stability was determined by coexpression of the hcp1 alleles with tse2-V in E. coli. We found that approximately half of the mutations localized to the inside surface of Hcp1 exhibit 1% or less Tse2 stabilization relative to wild-type Hcp1 (Figures 3A and 3B and Figure S3), whereas all Hcp1 variants with substitutions mapping to the outer surface of the ring displayed substantial Tse2-stabilization activity (1.0%-100%). Interestingly, the majority of substitutions that destabilize Tse2 over 100-fold map together within a discrete patch inside the Hcp1 ring (Figure 3B).

The ring shape of Hcp1 is readily discernable by negative stain transmission electron microscopy (TEM). Therefore, we reasoned that Hcp1 rings bound to Tse2 might be distinguished by TEM as "filled" particles. TEM analysis of approximately 3,000 randomly selected purified  $Hcp1-V-Tse2^{NT}-His_6$  single particles showed that filled class averages constituted a significant fraction of total Hcp1 particles (Figures 3C and 3D and Figure S4). In contrast, no class averages appeared filled in the control sample. The low fraction of filled rings observed in the Hcp1-Tse2 sample is likely explained by Tse2 degradation and precipitation during preparation (Li et al., 2012; Zou et al., 2012). Visualization of the Tse2-Hcp1 complex provides direct evidence that Tse2 interacts with the inner surface of Hcp1 rings.

# Tse2 Secretion Requires Interaction with Hcp1

Motivated by our in vitro findings, we next sought to determine the influence of the interaction between Tse2 and the inner surface of Hcp1 on Tse2 stability and secretion in vivo. Using allelic exchange, we generated a P. aeruginosa strain encoding Hcp1<sup>S31Q</sup> at the native *hcp1* locus. We selected this mutant from those identified by our in vitro studies, as this amino acid resides within the inner surface patch and strongly disrupts interaction with Tse2 via a relatively conservative substitution (small polar to large polar). Western blot analyses showed that although Hcp1<sup>S31Q</sup> is produced and secreted at levels comparable to the wild-type, strains bearing this mutation do not support Tse2 accumulation (Figure 4A). Furthermore, using interbacterial competition assays, we found that the hcp1 SaiQ background displays a marked defect in Tse2-dependent fitness (Figure 4B). The effects on H1-T6SS function appeared specific, as the levels of export and interbacterial delivery of Tse1 and Tse3 were not significantly impacted in the hcp1 S31Q background (Figures 4A and 4C). Overall these data demonstrate that Tse2 stability requires interaction with the inner surface of the Hcp1 ring.

Since Tse2 does not accumulate to detectable levels in  $\mathit{hcp1}^{\mathrm{S31Q}},$  we could not explicitly measure the requirement of interaction with Hcp1 for Tse2 secretion. However, a previous report from our laboratory demonstrated that overexpressed Tse2 can be detected in P. aeruginosa backgrounds lacking hcp1 (Hood et al., 2010). Therefore, to overcome the Tse2 detection limit, we monitored the export of Tse2-V expressed with high induction from the lacUV5 promoter in the  $hcp1^{S31Q}$  background. Western blot analysis confirmed that the Tse2-V protein is highly overproduced relative to endogenous Tse2 and that it is secreted in an Hcp1-dependent manner (Figure 4D). Interestingly, we found that despite intracellular accumulation, Tse2-V



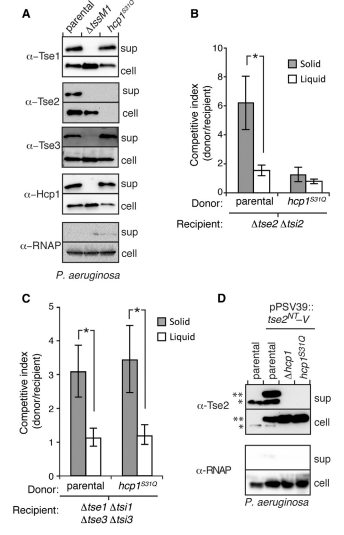


Figure 4. Tse2 Requires Interaction with Hcp1 for Secretion

(A) Western blot analysis of cell and supernatant fractions of Hcp1 and H1-T6S effectors in P. aeruginosa strains harboring wild-type hcp1 or hcp1 s31Q Equally exposed α-RNA polymerase (RNAP) blots of equivalent fractions of total cell and supernatant (sup) samples are included as loading and cytoplasmic leakage controls in this and subsequent secretion assays.

(B and C) Outcome of growth competition experiments between P. aeruginosa donor strains (parental or  $hcp1^{S31Q}$ ) and a Tse2-susceptible ( $\Delta tse2 \ \Delta tsi2$ ) (B) or a Tse1- and Tse3-susceptible recipient strain (Δtse1 Δtsi1 Δtse3 Δtsi3) (C) on solid (gray) or liquid (white) media. The competitive index is calculated as the change (final/initial) in ratio of donor to recipient cfu. Error bars represent standard deviation based on four replicates. Asterisks denote statistical significance using ANOVA and Tukey's post hoc test between the indicated conditions (p < 0.001).

(D) Western blot analysis of cell and supernatant-associated fractions of chromosomal, endogenous Tse2 (one asterisk) and ectopically expressed Tse2<sup>NT</sup>–V (two asterisks) in the indicated *P. aeruginosa* strains.

is not exported in the hcp1S31Q background. Thus, taken together with our observation that H1-T6SS function is generally preserved in the hcp1<sup>S31Q</sup> background, we conclude that interaction with Hcp1 is required both for the stabilization and the export of Tse2.

# Recognition of Tse2-like Effectors by Hcp Is General

To determine the generality of our findings concerning P. aeruginosa Tse2 and Hcp1, we sought to identify and characterize functionally analogous proteins in other bacterial species. Protein BLAST analyses revealed Tse2 homologs within several T6SS+ Gram-negative bacteria, including Methylomonas methanica, Shewanella frigidimarina, Burkholderia ambifaria, and Pseudoalteromonas sp. (Figure 5A). Sequence alignments of the homologs highlighted the conservation of several motifs, including a putative aspartic acid-containing nuclease-related catalytic site (Asp63 in Tse2) identified using protein structure prediction algorithms (Figure S5) (Li et al., 2012). As additional evidence of the functional relationship between Tse2 and its homologs, we found each homolog encoded in an apparent bicistron with a smaller gene encoding a protein with attributes similar to Tsi2, including length (Tsi2, 77; homologs, 75-79 amino acids) and an acidic pl (Tsi2, 3.9; homologs, 4.0-4.5) (Figure 5A).

Confident we had identified bona fide Tse2 and Tsi2-related proteins, we selected M. methanica Tse2 (Tse2MM), which shares 71% identity with P. aeruginosa Tse2, for more detailed analyses. The genome of M. methanica contains a single T6S gene cluster and encodes a single clear homolog of  $P.\ aeruginosa\ Hcp1,\ Hcp1^{MM},\ making\ Tse2^{MM}\ well\ suited\ for$ our study. To first establish that the M. methanica tse2 tsi2 pair is functionally orthologous to P. aeruginosa tse2 tsi2, we conducted toxicity studies in E. coli. Similar to tse2, expression of tse2MM in E. coli caused a dramatic drop in recovered cfu (Figure 5B). Moreover, this toxicity was observed for Tse2<sup>MM(D63N)</sup>, and it was inhibited by coexpression with  $tsi2^{MM}$ .

We next tested whether Tse2<sup>MM</sup>, like Tse2, interacts with the Hcp1 homolog, Hcp1<sup>MM</sup>, encoded within the putative T6SS of *M. methanica*. Indeed, we found Tse2<sup>MM</sup> interacts directly with Hcp1<sup>MM</sup> and, additionally, requires Hcp1<sup>MM</sup> for intracellular accumulation (Figure 5C). Interestingly, we found that Tse2<sup>MM</sup> also interacts with and is stabilized by P. aeruginosa Hcp1, and likewise for the P. aeruginosa Tse2-Hcp1<sup>MM</sup> pair (Figures 5C and 5D). The specificity of these interactions is underscored by the observation that neither Tse2 nor Tse2<sup>MM</sup> interacts with Hcp1 from Pseudomonas protegens (Hcp1PP), despite the fact that this Hcp1 homolog is nearly equally divergent from P. aeruginosa Hcp1 as Hcp1<sup>MM</sup> (Hcp1<sup>PP</sup>, 76%; Hcp1<sup>MM</sup>, 77%). Notably, Hcp1<sup>PP</sup> is encoded within an HSI-I-like gene cluster, yet P. protegens does not have a Tse2 homolog encoded within its genome. Finally, consistent with the hypothesis that interaction with Hcp is a key determinant for export via the T6S pathway, we observed efficient release of Tse2<sup>MM</sup> via the H1-T6SS (Figure 5E).

# **Amidase and Muramidase T6S Effector Classes Recognize Cognate Hcp Proteins**

In the course of studying the behavior of Hcp1 point mutants in P. aeruginosa, we observed that while the effects of Hcp1<sup>S31Q</sup> are limited to Tse2, other Tse2-destabilizing Hcp1 pore substitutions also reduce the export of Tse1 and Tse3 (Figure 6A). These Hcp1 variants are themselves secreted, suggesting their effect on Tse1 and Tse3 export is not a consequence of misfolding or a failure to be recognized by the secretory apparatus. Moreover, certain substitutions exhibited differential effects on the



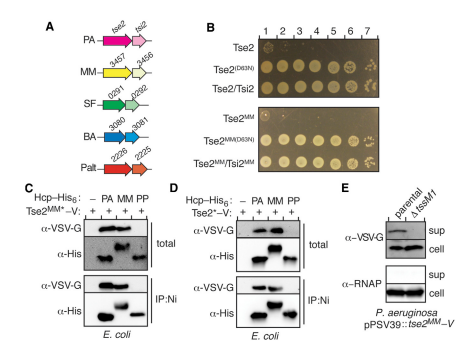


Figure 5. Stabilization by Cognate Hcp Is a General Feature of Tse2-like Effectors

(A) Genomic organization of tse2 and tsi2 homologs from *P. aeruginosa* (PA) *M. methanica* (MM), *S. frigidimarina* (SF), *B. ambifaria* (BA), and *Pseudoalteromonas* sp. (Palt). Abbreviated locus tag numbers are indicated for tse2 (darker shade) and tsi2 (lighter shade) homologs from each organism. See also Figure S5.

(B) Growth of *E. coli* containing plasmids with inducible expression of *tse2* or *tsi2* homologs from *P. aeruginosa* (upper panel) or *M. methanica* (lower panel). Serial 10-fold dilutions are indicated by numbers.

(C and D) Western blot results of coIP assays from  $E.\ coli$  coexpressing  $hcp-his_6$  homologs from  $P.\ aeruginosa$  (PA),  $M.\ methanica$  (MM), or  $P.\ protegens$  (PP) with  $tse2^{(D63N)}$ –V (mutation denoted with asterisk) from  $M.\ methanica$  (C) or  $P.\ aeruginosa$  (D).

(E) Analysis of cell and supernatant-associated fractions of the indicated *P. aeruginosa* strains expressing Tse2<sup>MM</sup>–V ectopically.

secretion of Tse1 and Tse3. For example, Hcp1<sup>L28A</sup> inhibits Tse3 secretion, but not that of Tse1. On the contrary, Hcp1<sup>S115Q</sup> strongly affects both Tse1 and Tse3 secretion. Together with our observation that Tse1 and Tse3 display decreased abundance in strains lacking Hcp1 (Figure 1B), our findings led us to hypothesize that non-Tse2-type effectors also engage in specific interactions with the inner surface of the Hcp1 pore, and that these are required for H1-T6SS-dependent secretion.

To determine whether Hcp1 directly interacts with Tse1 and Tse3, we performed coIP assays using proteins coexpressed in E. coli. As a means of linking potential interactions to the secretion phenotypes observed, we utilized the secretion-defective hcp1<sup>S115Q</sup> pore substitution mutant in addition to wild-type hcp1. Interbacterial competition assays confirmed this mutation functionally disrupts Tse1- and Tse3-dependent intoxication (Figure S6). While Tse1 and Tse3 did not efficiently coIP with Hcp1<sup>S115Q</sup>, both proteins were detected in association with the wild-type protein (Figures 6B and 6C). From these data, we conclude that interaction with the inner surface of Hcp1 is a common requirement for the secretion of the three known H1-T6SS effectors. The observation that Hcp1 substitutions can have variable effects on the Tse proteins suggests that, consistent with their divergent sequences, the proteins each form a unique network of interactions with Hcp1.

Bioinformatic work conducted by our laboratory has found evidence of a nonrandom association between *hcp* genes and predicted T6S amidase effectors (Tae) (Figure S7) (Russell et al., 2012). Based on this observed genetic link, our finding that direct interactions between Tse1-3 and Hcp1 are required for secretion, and the results of a prior study demonstrating the interaction of a putative *Edwardsiella tarda* T6S effector with an Hcp homolog (Zheng and Leung, 2007), we posited that the recognition of effectors by cognate Hcp proteins might be a general determinant of substrate selectivity by the T6SS. As a first

step toward validating this model, we selected two predicted cognate Tae-Hcp pairs based on genetic linkage (Salmonella enterica serovar Typhi Tae2) or homology to Tse1 (Burkholderia phytofirmans Tae1). As predicted, both effectors bound cognate, but not noncognate, Hcp proteins (Figures 6D and 6E). In total, our data suggest that the interaction between effectors and cognate Hcp proteins is a general phenomenon that is likely to—at least in part—define the particular effectors transported through a given T6SS.

## **DISCUSSION**

We have identified Hcp as a critical gatekeeper protein of the T6SS. As a chaperone, a substrate receptor, and a secreted protein itself, the multipurpose nature of Hcp grants it a unique position among secretory system-associated proteins (Figure 7). This implies that the mechanism of substrate export by the T6SS is fundamentally different from other characterized secretion systems. Arguably, T3S and T4S, which both translocate macromolecules in a concerted fashion into target cells, are the most analogous secretion pathways to the T6SS (Cornelis, 2006; Fronzes et al., 2009). Indeed, T6S and T4S share two homologous proteins, TssL (DotU) and TssM (IcmF) (Cascales and Cambillau, 2012). In light of this relatedness, it is interesting that effector recognition by the T6SS occurs via a principally different process than these related pathways, which both utilize ATPases to directly engage substrates and provide the energy necessary for export (Akeda and Galán, 2005; Christie et al., 2005). The fact that the T6S pathway does not appear to directly link effector recognition to energizing secretion may be explained by its putative bacteriophage origins (Kanamaru, 2009). The energy driving unidirectional protein transport in the T6S pathway is thought to be stored in a filamentous structure composed of TssB and



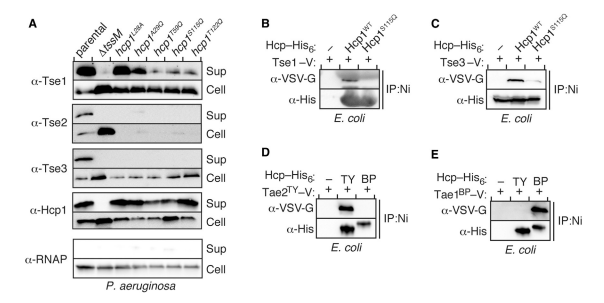


Figure 6. Binding to the Pore of Cognate Hcp Proteins Is Required for Export of Effectors with Amidase and Muramidase Activities
(A) Western blot analysis of cell and supernatant fractions of Hcp1 and H1-T6S effectors in *P. aeruginosa* strains wherein the native *hcp1* allele is substituted with the indicated mutant.

(B and C) Western blot analysis of bead-associated fractions of a coIP from *E. coli* coexpressing the indicated  $hcp-his_6$  alleles with Tse1–V (B) or Tse3–V (C). (D and E) Immunoblot detecting bead-associated fractions of a coIP from *E. coli* coexpressing  $hcp-his_6$  homologs (TY, S. Typhi; BP, *B. phytofirmans*) and  $tae2^{TY}-V$  (D) or  $tae1^{BP}-V$  (E). See also Figures S6 and S7.

TssC, which contracts and is then recycled by a ClpB-like AAA+ family ATPase, ClpV (Basler et al., 2012; Bönemann et al., 2009).

Prior models of the T6SS depict Hcp as a passive conduit for effectors (Basler et al., 2012; Bönemann et al., 2010; Cascales and Cambillau, 2012; Silverman et al., 2012). However, our data suggest that Hcp associates with effectors, protects them from proteolysis, and likely traffics with effectors during transport (Figure 7). Stoichiometric, or near-stoichiometric, release of Hcp hexamers with effectors reconciles the massive accumulation of Hcp in the culture supernatants of T6S-activated strains (Mougous et al., 2006; Pieper et al., 2009; Pukatzki et al., 2006; Wu et al., 2008). Whether effectors in complex with Hcp are folded cannot be definitively established from our data. The structures of several amidase effectors indicate the proteins are of a size that could be accommodated within the Hcp pore (Chou et al., 2012; Dong et al., 2013a; Zhang et al., 2013). However, based on the instability of effectors in the absence of cognate Hcp proteins, we speculate they do not adopt their final folded configuration until, or accompanying, release from Hcp. It is of note that intramolecular disulfide bonds have been observed in the structures of amidase effectors. The failure of these bonds to form in the cytoplasm could explain the instability of the proteins in this compartment, while their formation in the periplasm could drive release from Hcp. Interestingly, Tse2 lacks cysteine residues and ultimately accesses the cytoplasm of the recipient. In this case, interaction with the outer face of the inner membrane, or a receptor associated with the membrane, might drive effector release. It is also possible that the T6S apparatus remodels or modifies proteins during export to disrupt Hcp-substrate interactions.

Our data suggest that interactions with the pore of Hcp are, broadly, a critical determinant for the secretion of T6 effectors. However, we posit that other mechanisms may facilitate the recognition and export of disparate effector classes. Rather than a genetic linkage with hcp genes, Tle (type VI secretion lipase effector) and Rhs (recombination hot spot) effectors are encoded in association with vgrG genes (Koskiniemi et al., 2013; Russell et al., 2013). VgrG proteins often facilitate intercel-Jular T6S-dependent delivery of effectors present as C-terminal fusion to their structural domain (Brooks et al., 2013; Pukatzki et al., 2007; Suarez et al., 2010). It is conceivable that Tle and Rhs effectors, which are considerably larger than Tse-type effectors, might gain access to recipient cells by noncovalently associating with VgrG proteins. A Tle protein from V. cholerae was found in the immunoprecipitate of a VgrG protein; however, the genetically linked VgrG was not tested (Dong et al., 2013b).

While we have delineated a critical step in the process of effector secretion by the T6SS, many aspects of the overall structure-function model for the system remain untested. So far, the localization of Hcp inside the TssB-TssC sheath-like complex has not been demonstrated. Furthermore, there is some evidence suggesting that Hcp may not associate within the TssB-TssC tubule. For instance, Hcp proteins bearing large C-terminal fusions have been identified, and it is difficult to envision how these much larger proteins would be accommodated inside the TssB-TssC structure (Blondel et al., 2009; Parret and De Mot, 2002). Irrespective of the association of Hcp-effector complexes with TssB-TssC, it also remains unresolved how Hcp-substrate complexes cross the inner and outer bacterial membranes. An envelope-spanning complex consisting of integral inner membrane proteins and an outer membrane



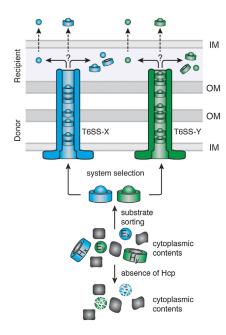


Figure 7. Model Depicting the Role of Hcp in T6S Effector Recognition and Export

The schematic depicts the junction between two Gram-negative bacterial cells (OM, outer membrane; IM, inner membrane), a donor cell harboring two T6SSs T6SS-X (blue) and T6SS-Y (green), and a recipient cell that is targeted by these systems. T6S effectors  $(E_x,\,E_y)$  are sorted from the cytoplasmic pool of proteins via interactions with cognate Hcp proteins  $(H_x,\,H_y)$ . Interaction with Hcp prevents effector degradation. Hcp-effector complexes are recognized and transported through the appropriate T6SS via an unknown mechanism. The Hcp-effector complex may be translocated into the periplasm intact, or the effector may dissociate from Hcp prior to periplasmic delivery. Likewise, effectors destined for the cytoplasm may be transported in complex or in isolation.

lipoprotein has been identified; however, the ultrastructure of this novel complex awaits visualization (Felisberto-Rodrigues et al., 2011).

Immunity to Tse2 is provided by the Tsi2 protein, a small acidic cytoplasmic protein that directly binds and inactivates the toxin. The current study adds previously unrecognized complexity to the subject of effector immunity, particularly cytoplasmic effectors like Tse2 that interact with cognate immunity proteins prior to export. Based on the studies reported herein and the observation that Tsi2 is an essential gene, it is apparent that significant Tse2 immunity is not provided by Hcp1 (Hood et al., 2010; Jacobs et al., 2003). It is conceivable that Tsi2 interacts with Tse2 in the context of Tse2–Hcp1 complexes; however, we did not detect this complex and favor the hypothesis that Tsi2 functions by scavenging free Tse2, including endogenous Tse2 and Tse2 delivered by other *P. aeruginosa* cells in *trans*.

Our discovery that effectors interact with the pore of Hcp may have implications for the identification of novel T6S substrates. The discovery of effectors by virtue of interaction with cognate Hcp proteins stands to overcome several difficulties encountered in such efforts. For instance, this method would not depend upon effector export, which is often repressed under in vitro culturing conditions (Silverman et al., 2012). Our findings

may also guide efforts to engineer nonnative substrate export into the T6S pathway. This approach could provide new ways to modulate the outcome of interbacterial interactions.

#### **EXPERIMENTAL PROCEDURES**

## **Bacterial Strains, Plasmids, and Growth Conditions**

All P. aeruginosa strains were derived from the sequenced strain PAO1. Genomic DNA from M. methanica MC09, S. Typhi Ty2, P. protegens Pf-5, or P. aeruginosa PAO1 was used to amplify effector, immunity, and hcp genes (Boden et al., 2011; Deng et al., 2003; Paulsen et al., 2005; Stover et al., 2000). E. coli strains DH5 $\alpha$ , SM10, and BL21 were used for plasmid maintenance, conjugative transfer, and gene expression, respectively. Plasmids used in this study are described and referenced in Table S1. See the Supplemental Experimental Procedures for specific growth conditions and cloning procedures.

#### **Hcp1 Depletion Assays**

We used a controllable protein degradation system originally developed for use in *E. coli* but which has also been optimized for use in *P. aeruginosa* (Castang et al., 2008; McGinness et al., 2006). Full details are in the Supplemental Experimental Procedures.

#### **β-Galactosidase Assay**

β-galactosidase assays were performed as described previously with minor modifications. A chromosomal integration vector (mini-CTX-*lacZ*) carrying a fusion of *lacZ* to the putative promoter region and first eight codon of *tse2* was used to generate strains for analysis. The plasmid was introduced into *P. aeruginosa* by conjugation (Hoang et al., 2000). Full details are in the Supplemental Experimental Procedures.

#### **Coimmunoprecipitation Assays**

Full details for coIP assays from *P. aeruginosa* or *E. coli* can be found in the Supplemental Experimental Procedures. Briefly, cell lysates were applied to anti-VSV-G agarose beads or nickel-NTA Sepharose beads and incubated for 1 hr at 4°C. Beads were washed three times with buffer before resuspending in SDS-PAGE buffer.

#### Tse2 Stability Assays with Hcp1 Point Mutants

*E. coli* BL21 plysS with plasmids containing each *hcp1* allele (pET29b) and  $tse2^{NT}$  (pSCRhaB2-CV) were cotransformed into *E. coli*. Overnight cultures were subinoculated (1:1,000) into 3 ml of 2xYT and grown to an OD<sub>600</sub> of 0.6. Cultures were induced with 0.1% rhamnose and 10 μM IPTG and harvested after 4 hr. From each sample, 300 μl were centrifuged at 16,000 × g, and pellets were resuspended in 30 μl buffer 1 and 30 μl of SDS-PAGE loading dye, boiled and analyzed by western blot. Densitometry was performed using AlphaViewQ software (ProteinSimple). The percentage of total accumulated Tse2 was determined by the following equation: (([Tse2\*]/[Tse2])/([Hcp1\*]/[Hcp1])\*100, where Tse2\* is Tse2 accumulated in the presence of each Hcp1 point mutation (Hcp1\*).

#### **Secretion Assays**

Overnight cultures of *P. aeruginosa* strains were used to inoculate 2 ml of LB (1:1,000) supplemented with 0.5 mM IPTG. Cultures were grown at 37°C with shaking to mid-log phase, and cell and supernatant fractions were processed as previously described (Hood et al., 2010).

## **Interbacterial Competition Assays**

Competition assays between *P. aeruginosa* strains were performed as previously described with minor variations that are described in the Supplemental Experimental Procedures (Hood et al., 2010; Russell et al., 2011).

# Purification of Tse2-Hcp1 Complex for Transmission Electron Microscopy

The Tse2-Hcp1 complex was purified from *E. coli* containing plysS, pET29b::*tse2*<sup>NT</sup>-His<sub>6</sub>, and pPSV35::*hcp1*-VSV-G using a two-step affinity



chromatography method. Full details are in the Supplemental Experimental Procedures.

# Single-Particle Electron Microscopy of Hcp Rings and Hcp1-Tse2

Hcp1 rings and the Hcp1-Tse2 complex purified from E. coli were negatively stained by 0.75% uranyl formate as described previously (Ohi et al., 2004). Images were collected on a transmission electron microscope T12 (FEI) at room temperature under 120 kV and recorded at a magnification of 67,000× on a 4k × 4k Gatan CCD rendering a final pixel size of 1.65 Å on the specimen level. Particles were selected and windowed into 120 x 120 pixel images using WEB (Frank et al., 1996). Full details are in the Supplemental Experimental Procedures.

#### E. coli Toxicity Measurements

E. coli toxicity assays were performed as previously described with some variations described in the Supplemental Experimental Procedures (Russell et al., 2011).

#### SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures, one table, Supplemental Experimental Procedures, and Supplemental References and can be found with this article at http://dx.doi.org/10.1016/j.molcel.2013.07.025.

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#### **REFERENCES**

Abdallah, A.M., Gey van Pittius, N.C., Champion, P.A., Cox, J., Luirink, J., Vandenbroucke-Grauls, C.M., Appelmelk, B.J., and Bitter, W. (2007). Type VII secretion—mycobacteria show the way. Nat. Rev. Microbiol. 5, 883–891.

Akeda, Y., and Galán, J.E. (2005). Chaperone release and unfolding of substrates in type III secretion. Nature 437, 911-915.

Ballister, E.R., Lai, A.H., Zuckermann, R.N., Cheng, Y., and Mougous, J.D. (2008). In vitro self-assembly of tailorable nanotubes from a simple protein building block. Proc. Natl. Acad. Sci. USA 105, 3733-3738.

Basler, M., and Mekalanos, J.J. (2012). Type 6 secretion dynamics within and between bacterial cells. Science 337, 815.

Basler, M., Pilhofer, M., Henderson, G.P., Jensen, G.J., and Mekalanos, J.J. (2012). Type VI secretion requires a dynamic contractile phage tail-like structure. Nature 483, 182-186.

Blondel, C.J., Jiménez, J.C., Contreras, I., and Santiviago, C.A. (2009). Comparative genomic analysis uncovers 3 novel loci encoding type six secretion systems differentially distributed in Salmonella serotypes. BMC Genomics 10, 354, http://dx.doi.org/10.1186/1471-2164-10-354.

Boden, R., Cunliffe, M., Scanlan, J., Moussard, H., Kits, K.D., Klotz, M.G., Jetten, M.S., Vuilleumier, S., Han, J., Peters, L., et al. (2011). Complete genome sequence of the aerobic marine methanotroph Methylomonas methanica MC09. J. Bacteriol. 193, 7001-7002.

Bönemann, G., Pietrosiuk, A., Diemand, A., Zentgraf, H., and Mogk, A. (2009). Remodelling of VipA/VipB tubules by ClpV-mediated threading is crucial for type VI protein secretion. EMBO J. 28, 315-325.

Bönemann, G., Pietrosiuk, A., and Mogk, A. (2010). Tubules and donuts: a type VI secretion story. Mol. Microbiol. 76, 815-821.

Brencic, A., and Lory, S. (2009). Determination of the regulon and identification of novel mRNA targets of Pseudomonas aeruginosa RsmA. Mol. Microbiol. 72, 612-632.

Brooks, T.M., Unterweger, D., Bachmann, V., Kostiuk, B., and Pukatzki, S. (2013). Lytic activity of the Vibrio cholerae type VI secretion toxin VgrG-3 is inhibited by the antitoxin TsaB. J. Biol. Chem. 288, 7618-7625.

Cambronne, E.D., and Roy, C.R. (2007). The Legionella pneumophila IcmSW complex interacts with multiple Dot/Icm effectors to facilitate type IV translocation. PLoS Pathog. 3, e188, http://dx.doi.org/10.1371/journal.ppat.

Cascales, E., and Cambillau, C. (2012). Structural biology of type VI secretion systems. Philos. Trans. R. Soc. Lond. B Biol. Sci. 367, 1102-1111.

Castang, S., and Dove, S.L. (2012). Basis for the essentiality of H-NS family members in Pseudomonas aeruginosa. J. Bacteriol. 194, 5101-5109.

Castang, S., McManus, H.R., Turner, K.H., and Dove, S.L. (2008). H-NS family members function coordinately in an opportunistic pathogen. Proc. Natl. Acad. Sci. USA 105, 18947-18952.

Chou, S., Bui, N.K., Russell, A.B., Lexa, K.W., Gardiner, T.E., LeRoux, M., Vollmer, W., and Mougous, J.D. (2012). Structure of a peptidoglycan amidase effector targeted to Gram-negative bacteria by the type VI secretion system. Cell Rep. 1, 656-664.

Christie, P.J., Atmakuri, K., Krishnamoorthy, V., Jakubowski, S., and Cascales, E. (2005). Biogenesis, architecture, and function of bacterial type IV secretion systems. Annu. Rev. Microbiol. 59, 451-485.

Cornelis, G.R. (2006). The type III secretion injectisome. Nat. Rev. Microbiol. 4, 811-825.

Dautin, N., and Bernstein, H.D. (2007). Protein secretion in gram-negative bacteria via the autotransporter pathway. Annu. Rev. Microbiol. 61, 89-112.

Deng, W., Liou, S.R., Plunkett, G., 3rd, Mayhew, G.F., Rose, D.J., Burland, V., Kodoyianni, V., Schwartz, D.C., and Blattner, F.R. (2003). Comparative genomics of Salmonella enterica serovar Typhi strains Ty2 and CT18. J. Bacteriol.

Dong, C., Zhang, H., Gao, Z.Q., Wang, W.J., She, Z., Liu, G.F., Shen, Y.Q., Su, X.D., and Dong, Y.H. (2013a). Structural insights into the inhibition of type VI effector Tae3 by its immunity protein Tai3. Biochem. J. 454, 59-68.

Dong, T.G., Ho, B.T., Yoder-Himes, D.R., and Mekalanos, J.J. (2013b). Identification of T6SS-dependent effector and immunity proteins by Tn-seq in Vibrio cholerae. Proc. Natl. Acad. Sci. USA 110, 2623-2628.

Economou, A., Christie, P.J., Fernandez, R.C., Palmer, T., Plano, G.V., and Pugsley, A.P. (2006). Secretion by numbers: protein traffic in prokaryotes. Mol. Microbiol. 62, 308-319.

Felisberto-Rodrigues, C., Durand, E., Aschtgen, M.S., Blangy, S., Ortiz-Lombardia, M., Douzi, B., Cambillau, C., and Cascales, E. (2011). Towards a structural comprehension of bacterial type VI secretion systems: characterization of the TssJ-TssM complex of an Escherichia coli pathovar. PLoS Pathog. 7, e1002386, http://dx.doi.org/10.1371/journal.ppat.1002386.

Frank, J., Radermacher, M., Penczek, P., Zhu, J., Li, Y., Ladjadj, M., and Leith, A. (1996). SPIDER and WEB: processing and visualization of images in 3D electron microscopy and related fields. J. Struct. Biol. 116, 190-199.

Fronzes, R., Christie, P.J., and Waksman, G. (2009). The structural biology of type IV secretion systems. Nat. Rev. Microbiol. 7, 703-714.

Galán, J.E. (2009). Common themes in the design and function of bacterial effectors. Cell Host Microbe 5, 571-579.



Hachani, A., Lossi, N.S., Hamilton, A., Jones, C., Bleves, S., Albesa-Jové, D., and Filloux, A. (2011). Type VI secretion system in Pseudomonas aeruginosa: secretion and multimerization of VgrG proteins. J. Biol. Chem. 286, 12317-12327.

Hoang, T.T., Kutchma, A.J., Becher, A., and Schweizer, H.P. (2000). Integration-proficient plasmids for Pseudomonas aeruginosa: site-specific integration and use for engineering of reporter and expression strains. Plasmid 43, 59-72.

Hood, R.D., Singh, P., Hsu, F., Güvener, T., Carl, M.A., Trinidad, R.R., Silverman, J.M., Ohlson, B.B., Hicks, K.G., Plemel, R.L., et al. (2010). A type VI secretion system of Pseudomonas aeruginosa targets a toxin to bacteria. Cell Host Microbe 7, 25-37.

Izoré, T., Job, V., and Dessen, A. (2011). Biogenesis, regulation, and targeting of the type III secretion system. Structure 19, 603-612.

Jacobs, M.A., Alwood, A., Thaipisuttikul, I., Spencer, D., Haugen, E., Ernst, S., Will, O., Kaul, R., Raymond, C., Levy, R., et al. (2003). Comprehensive transposon mutant library of Pseudomonas aeruginosa. Proc. Natl. Acad. Sci. USA 100, 14339-14344.

Jobichen, C., Chakraborty, S., Li, M., Zheng, J., Joseph, L., Mok, Y.K., Leung, K.Y., and Sivaraman, J. (2010). Structural basis for the secretion of EvpC: a key type VI secretion system protein from Edwardsiella tarda. PLoS ONE 5, e12910, http://dx.doi.org/10.1371/journal.pone.0012910.

Kanamaru, S. (2009). Structural similarity of tailed phages and pathogenic bacterial secretion systems. Proc. Natl. Acad. Sci. USA 106, 4067-4068.

Kapitein, N., and Mogk, A. (2013). Deadly syringes: type VI secretion system activities in pathogenicity and interbacterial competition. Curr. Opin. Microbiol. 16. 52-58.

Koskiniemi, S., Lamoureux, J.G., Nikolakakis, K.C., t'Kint de Roodenbeke, C., Kaplan, M.D., Low, D.A., and Hayes, C.S. (2013). Rhs proteins from diverse bacteria mediate intercellular competition. Proc. Nat. Acad. Sci. USA 110, 7032-7037.

Leiman, P.G., Basler, M., Ramagopal, U.A., Bonanno, J.B., Sauder, J.M., Pukatzki, S., Burley, S.K., Almo, S.C., and Mekalanos, J.J. (2009). Type VI secretion apparatus and phage tail-associated protein complexes share a common evolutionary origin. Proc. Natl. Acad. Sci. USA 106, 4154-4159.

LeRoux, M., De Leon, J.A., Kuwada, N.J., Russell, A.B., Pinto-Santini, D., Hood, R.D., Agnello, D.M., Robertson, S.M., Wiggins, P.A., and Mougous, J.D. (2012). Quantitative single-cell characterization of bacterial interactions reveals type VI secretion is a double-edged sword. Proc. Natl. Acad. Sci. USA 109. 19804-19809.

Li, M., Le Trong, I., Carl, M.A., Larson, E.T., Chou, S., De Leon, J.A., Dove, S.L., Stenkamp, R.E., and Mougous, J.D. (2012). Structural basis for type VI secretion effector recognition by a cognate immunity protein. PLoS Pathog. 8, e1002613, http://dx.doi.org/10.1371/journal.ppat.1002613.

Lin, J.S., Ma, L.S., and Lai, E.M. (2013). Systematic dissection of the Agrobacterium type VI secretion system reveals machinery and secreted components for subcomplex formation. PLoS ONE 8, e67647, http://dx.doi.org/10. 1371/journal.pone.0067647.

McGinness, K.E., Baker, T.A., and Sauer, R.T. (2006). Engineering controllable protein degradation. Mol. Cell 22, 701-707.

Mougous, J.D., Cuff, M.E., Raunser, S., Shen, A., Zhou, M., Gifford, C.A., Goodman, A.L., Joachimiak, G., Ordoñez, C.L., Lory, S., et al. (2006). A virulence locus of Pseudomonas aeruginosa encodes a protein secretion apparatus. Science 312, 1526-1530.

Ohi, M., Li, Y., Cheng, Y., and Walz, T. (2004). Negative staining and image classification-powerful tools in modern electron microscopy. Biol. Proced. Online 6, 23-34.

Osipiuk, J., Xu, X., Cui, H., Savchenko, A., Edwards, A., and Joachimiak, A. (2011). Crystal structure of secretory protein Hcp3 from Pseudomonas aeruginosa. J. Struct. Funct. Genomics 12, 21-26.

Parret, A.H., and De Mot, R. (2002). Escherichia coli's uropathogenicspecific protein: a bacteriocin promoting infectivity? Microbiology 148, 1604-1606.

Paulsen, I.T., Press, C.M., Ravel, J., Kobayashi, D.Y., Myers, G.S., Mavrodi, D.V., DeBoy, R.T., Seshadri, R., Ren, Q., Madupu, R., et al. (2005). Complete genome sequence of the plant commensal Pseudomonas fluorescens Pf-5. Nat. Biotechnol. 23, 873-878.

Pieper, R., Huang, S.T., Robinson, J.M., Clark, D.J., Alami, H., Parmar, P.P., Perry, R.D., Fleischmann, R.D., and Peterson, S.N. (2009). Temperature and growth phase influence the outer-membrane proteome and the expression of a type VI secretion system in Yersinia pestis. Microbiology 155, 498–512.

Pukatzki, S., Ma, A.T., Sturtevant, D., Krastins, B., Sarracino, D., Nelson, W.C., Heidelberg, J.F., and Mekalanos, J.J. (2006). Identification of a conserved bacterial protein secretion system in Vibrio cholerae using the Dictyostelium host model system. Proc. Natl. Acad. Sci. USA 103, 1528-1533.

Pukatzki, S., Ma, A.T., Revel, A.T., Sturtevant, D., and Mekalanos, J.J. (2007). Type VI secretion system translocates a phage tail spike-like protein into target cells where it cross-links actin. Proc. Natl. Acad. Sci. USA 104, 15508-15513.

Russell, A.B., Hood, R.D., Bui, N.K., LeRoux, M., Vollmer, W., and Mougous, J.D. (2011). Type VI secretion delivers bacteriolytic effectors to target cells. Nature 475, 343-347.

Russell, A.B., Singh, P., Brittnacher, M., Bui, N.K., Hood, R.D., Carl, M.A., Agnello, D.M., Schwarz, S., Goodlett, D.R., Vollmer, W., and Mougous, J.D. (2012). A widespread bacterial type VI secretion effector superfamily identified using a heuristic approach. Cell Host Microbe 11, 538-549.

Russell, A.B., LeRoux, M., Hathazi, K., Agnello, D.M., Ishikawa, T., Wiggins, P.A., Wai, S.N., and Mougous, J.D. (2013). Diverse type VI secretion phospholipases are functionally plastic antibacterial effectors. Nature 496, 508-512.

Silverman, J.M., Austin, L.S., Hsu, F., Hicks, K.G., Hood, R.D., and Mougous, J.D. (2011). Separate inputs modulate phosphorylation-dependent and -independent type VI secretion activation. Mol. Microbiol. 82, 1277-1290.

Silverman, J.M., Brunet, Y.R., Cascales, E., and Mougous, J.D. (2012). Structure and regulation of the type VI secretion system. Annu. Rev. Microbiol. 66, 453-472.

Stebbins, C.E., and Galán, J.E. (2001). Maintenance of an unfolded polypeptide by a cognate chaperone in bacterial type III secretion. Nature 414, 77-81.

Stover, C.K., Pham, X.Q., Erwin, A.L., Mizoguchi, S.D., Warrener, P., Hickey, M.J., Brinkman, F.S., Hufnagle, W.O., Kowalik, D.J., Lagrou, M., et al. (2000). Complete genome sequence of Pseudomonas aeruginosa PAO1, an opportunistic pathogen. Nature 406, 959-964.

Suarez, G., Sierra, J.C., Erova, T.E., Sha, J., Horneman, A.J., and Chopra, A.K. (2010). A type VI secretion system effector protein, VgrG1, from Aeromonas hydrophila that induces host cell toxicity by ADP ribosylation of actin. J. Bacteriol. 192, 155-168.

Sutherland, M.C., Nguyen, T.L., Tseng, V., and Vogel, J.P. (2012). The Legionella IcmSW complex directly interacts with DotL to mediate translocation of adaptor-dependent substrates. PLoS Pathog. 8, e1002910, http://dx. doi.org/10.1371/journal.ppat.1002910.

Wagner, C., Polke, M., Gerlach, R.G., Linke, D., Stierhof, Y.D., Schwarz, H., and Hensel, M. (2011). Functional dissection of SiiE, a giant non-fimbrial adhesin of Salmonella enterica. Cell. Microbiol. 13, 1286-1301.

Wu, H.Y., Chung, P.C., Shih, H.W., Wen, S.R., and Lai, E.M. (2008). Secretome analysis uncovers an Hcp-family protein secreted via a type VI secretion system in Agrobacterium tumefaciens, J. Bacteriol, 190, 2841-2850,

Zhang, H., Zhang, H., Gao, Z.Q., Wang, W.J., Liu, G.F., Xu, J.H., Su, X.D., and Dong, Y.H. (2013). Structure of the type VI effector-immunity complex (Tae4-Tai4) provides novel insights into the inhibition mechanism of the effector by its immunity protein. J. Biol. Chem. 288, 5928-5939.

Zheng, J., and Leung, K.Y. (2007). Dissection of a type VI secretion system in Edwardsiella tarda. Mol. Microbiol. 66, 1192-1206.

Zou, T., Yao, X., Qin, B., Zhang, M., Cai, L., Shang, W., Svergun, D.I., Wang, M., Cui, S., and Jin, Q. (2012). Crystal structure of Pseudomonas aeruginosa Tsi2 reveals a stably folded superhelical antitoxin. J. Mol. Biol. 417, 351-361.