Review

The bacterial type VI secretion machine: yet another player for protein transport across membranes

Alain Filloux,1,2 Abderrahman Hachani1,2 and Sophie Bleves2

1Imperial College London, Division of Cell and Molecular Biology, Centre for Molecular Microbiology and Infection, South Kensington Campus, Flowers Building, London SW7 2AZ, UK
2Laboratoire d'ingénierie des systèmes macromoléculaires, UPR9027, CNRS-IBSM, 31 Chemin Joseph Aiguier, 13402 Marseille cedex 20, France

Several secretion systems have evolved that are widespread among Gram-negative bacteria. Recently, a new secretion system was recognized, which is named the type VI secretion system (T6SS). The T6SS components are encoded within clusters of genes initially identified as IAHP for IcmF-like homologous proteins, since they were all found to contain a gene encoding an IcmF-like component. IcmF was previously reported as a component of the type IV secretion system (T4SS). However, with the exception of DotU, other T4SS components are not encoded within T6SS loci. Thus, the T6SS is probably a novel kind of complex multi-component secretion machine, which is often involved in interaction with eukaryotic hosts, be it a pathogenic or a symbiotic relationship. The expression of T6SS genes has been reported to be mostly induced in vivo. Interestingly, expression and assembly of T6SSs are tightly controlled at both the transcriptional and the post-translational level. This may allow a timely control of T6SS assembly and function. Two types of proteins, generically named Hcp and VgrG, are secreted via these systems, but it is not entirely clear whether they are truly secreted effector proteins or are actually components of the T6SS. The precise role and mode of action of the T6SS is still unknown. This review describes current knowledge about the T6SS and summarizes its hallmarks and its differences from other secretion systems.

Introduction

Interaction between bacteria and hosts ranges from a commensal collaboration to a competition that may result in host death (Merrell & Falkow, 2004). Such interaction is guided by a communication/signalling game between the host and the pathogen, which, as in a game of chess, aims to influence the way you would like your opponent to play. Among the tools used by bacteria to influence the host response, secretion machines that deliver proteins and toxins into the environment and within a eukaryotic target cell are crucial for virulence and survival within hosts (Caron et al., 2006; Cossart & Sansonetti, 2004; Mota et al., 2005). These proteins are transported across the membranes of the bacteria, and eventually of the host, by means of specific devices called secretion systems (SSs). During the last 20 years, it has been found that Gram-negative bacteria have evolved several SSs, which have been identified by types (Fig. 1) (Filloux, 2004; Saier, 2006). Five types have been defined, i.e. type I to type V (T1SS to T5SS). The SSs vary in complexity but all use a single polypeptide or a supra-macromolecular complex to build a path through the bacterial cell envelope. SSs can be recognized by a set of core components used to build up the secretion device. Recent studies have led to the identification of a new type of SS, named T6SS. The T6SS components are encoded within gene clusters that vary in organization. These clusters were initially named IAHP, for IcmF-like homologous proteins, because they contain a gene encoding an IcmF-like component (Das & Chaudhuri, 2003). It thus all started with the finding that a known T4SS component, IcmF, was encoded within a conserved gene cluster among Gram-negative bacteria but whose other genes had no homology with T4SS components. These novel genes were likely to encode the components of a novel secretion machine or T6SS.

The type IVB secretion system

Legionella pneumophila is a facultative intracellular pathogen, and when it grows inside human cells or amoebae it is able to inhibit phagosome–lysosome fusion. L. pneumophila pathogenesis requires 26 dot (defect in organelle trafficking) and icm (intracellular multiplication) genes (Segal et al., 2005). These genes are essential for altering the endocytic pathway and for replication of L. pneumophila inside the host cells (Sexton & Vogel, 2002). The L. pneumophila dot/icm genes encode components of the type IVB SS, and share extensive similarities with the trb/tra genes found on the IncI plasmids and involved in bacterial
conjugation. The type IVA archetypes are the T-DNA transferring system called VirB in *Agrobacterium tumefaciens* and the Tra/Trb conjugative system from IncP plasmids (Christie *et al.*, 2005). The relationship between the B and the A subgroups of the T4SS is limited to a few components.

Whereas the Tra/Trb systems of the IncI plasmids are conjugation machines that deliver nucleoprotein complexes (Wilkins & Thomas, 2000), the *L. pneumophila* type IVB SS (Dot/Icm) is known to deliver proteins into target cells. In *L. pneumophila* Dot/Icm-dependent effectors have been characterized. Among them, RalF was shown to be required for the localization of ARF (ADP-ribosylation factor) on phagosomes containing *L. pneumophila* (Nagai *et al.*, 2002).

**The DotU/IcmF paradigm**

In contrast to most dot/icm genes, *icmF* is partially required for *L. pneumophila* replication in macrophages (Purcell & Shuman, 1998). The *icmF* gene is located at one end of the dot/icm cluster, downstream of a gene designated dotU or icmH. DotU and IcmF localize to the *L. pneumophila* inner membrane (Sexton *et al.*, 2004). IcmF contains several transmembrane domains and a putative Walker A nucleotide-binding motif whereas DotU contains a transmembrane segment in its C-terminal region. *dotU* and *icmF* mutants have similar intracellular growth phenotypes, i.e. partially defective in replication within macrophages (Sexton *et al.*, 2004; Zusman *et al.*, 2004), which suggested that the DotU and IcmF proteins worked together. Furthermore, the lack of IcmF resulted in a reduced level
of DotU protein, suggesting that the two proteins interact. It was also shown that the lack of DotU and/or IcmF affected the stability of three other Dot proteins. This suggests that DotU/IcmF assist in assembly and stability of a functional Dot–Icm complex and that in their absence the complex is not maintained in a fully active form (Sexton et al., 2004; VanRheenen et al., 2004).

IcmF and DotU orthologues have been found in a wide range of Gram-negative bacterial species. In many cases, these genes are linked, but no other T4SS genes have been found in their vicinity. Instead, another set of conserved genes are systematically found, which were originally known as IAHP (Das & Chaudhuri, 2003). Each gene cluster encodes at least a dozen proteins with various degrees of conservation.

**The Rhizobium leguminosarum Imp system**

The first report describing genes belonging to IAHP clusters was published in 1997 (Roest et al., 1997). *Rhizobium leguminosarum* is a plant symbiont, which induces formation of nitrogen-fixing nodules. The host specificity is determined by the Sym plasmid (Van Brussel et al., 1986). The authors described a *R. leguminosarum* strain capable of nodulation on *Vicia sativa* but not on *Pisum sativum* and *Vicia hirsuta*. In these last species small nodules were formed on the plant roots, but they were not able to fix nitrogen. This strain was designated RBL5523. Tn5 mutagenesis on RBL5523 allowed the identification of a mutant, RBL5787, restored in its ability to nodulate *P. sativum* and *V. hirsuta*. Because the locus is important for nodulation or subsequent stages of symbiosis, it was named *imp* (impaired in nodulation). In 2003, another study reported the sequence of a 33 kb region around the Tn5 inserted in RBL5787 (Bladergroen et al., 2003), revealing a putative operon of 14 genes (Fig. 2) named *impA–impN*. These genes mostly encode proteins with unknown function, but ImpK and ImpL have similarities with DotU and IcmF, respectively. ImpK is similar to DotU but has a C-terminal extension with similarity to OmpA, DotU and IcmF, respectively. ImpK has similarities with DotU and IcmF and DotU homologues. It should be noted that the DotU homologue of *V. cholerae* (VCA0115/VasF) does not contain the OmpA/MotB C-terminal extension found in ImpK, which indicated that this is not a hallmark for DotU orthologues encoded within IAHP/T6SS gene clusters. All the hallmark features found within characterized T6SSs are described in Table 1. Pukatzki et al. (2006) investigated whether the Vas system may be involved in protein secretion. Analysis of *vas* mutants’ culture supernatant, and comparison with the parental strain, revealed the absence of a 28 kDa protein, which appeared to be Hcp, a ‘Haemolysin A co-regulated protein’ (Williams et al., 1996), and of proteins called VgrG. Analysis of an O1 serogroup strain sensitive to *Dictyostelium*, N16961, showed that it failed to secrete Hcp and VgrG proteins, similarly to the V52 *vas* mutants. *Dictyostelium* resistance thus appeared to be dependent on the secretion of Hcp and VgrG proteins. It should be noted that in the study by Pukatzki et al. (2006), secretion of VgrG1 and VgrG2 was shown to be Hcp-dependent, even though it was unclear how Hcp could serve this role. The VgrG1, VgrG2, VgrG3 and Hcp proteins do not contain an N-terminal signal peptide. Interestingly, secreted *V. cholerae* proteins such as chitinase, neuraminidase, PrtV and HlyA contain a signal peptide and are still secreted in the *vas* mutants. These observations suggested that the Vas/IAHP/T6SS is responsible for the secretion of proteins.
lacking signal peptides, like the T4SS and T3SS but in contrast to the T2SS (Filloux, 2004).

The T6SS in *Salmonella enterica*

Early in the discovery of the T6SS, a complete gene cluster, named *sci* (*Salmonella enterica* centisome 7 genomic island), was described in *S. enterica* (Folkesson *et al.*, 2002). The island is 47 kb long and harbours 37 genes. From *sciA* to *sciY*, many genes could be identified as core genes for the T6SS, including 9 of the 15 *vas* genes found in *V. cholerae*. The Sci island includes genes encoding an Hcp-like protein (*sciK*) and a VgrG-like protein (*vgrS*) (Table 1). In addition, genes involved in fimbrial assembly (*saf*) or invasin production (*pagN*) were found. A complete deletion of the Sci genomic island resulted in decreased ability of *S. enterica* to enter eukaryotic cells. This result is different from data obtained with a *sciS* (*icmF*-like) transposon mutant (Parsons & Heffron, 2005). In that case, it was observed that SciS limits intracellular growth in macrophages at late stages of infection and attenuates the lethality of *S. enterica* in a murine host. *sciS* was maximally expressed at a late stage of infection, and was shown to be negatively regulated by SsrB, part of the SsrA/SsrB two-
Table 1. Some features of T6SSs in various bacteria

<table>
<thead>
<tr>
<th>Bacterium</th>
<th>T6SS genes</th>
<th>Secreted proteins</th>
<th>Putative secreted proteins</th>
<th>Regulatory aspects*</th>
<th>Notable features†</th>
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<tr>
<td><em>Burkholderia mallei</em></td>
<td><em>tss</em></td>
<td>Hcp1, VgrG1</td>
<td>TssB, TssM</td>
<td>VirAG two-component system</td>
<td>Attenuated virulence in hamsters</td>
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<td>BMAA1517 (AraC-type regulator)</td>
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<td><em>Edwardsiella ictaluri</em></td>
<td><em>eip</em></td>
<td>EvpC (Hcp)</td>
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<td>EsrB (response regulator, SsrB-like)</td>
<td>Anti-Eip produced during catfish infection</td>
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<td>evp</td>
<td>EvpP (Hcp-like)</td>
<td>Temperature-dependent</td>
<td>Attenuated virulence in blue gourami fish</td>
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<td><em>Escherichia coli</em> (EAEC)</td>
<td><em>sci-I</em></td>
<td>Hcp-like</td>
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<td><em>Francisella tularensis</em></td>
<td><em>sci-II</em></td>
<td>AaiC</td>
<td>AaiG (VgrG-like)</td>
<td>AggR (AraC-type regulator)</td>
<td>Required for intramacrophage growth; no ClpV-homologue</td>
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<td></td>
<td>PigG (VgrG-like)</td>
<td>MglA master regulator</td>
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<td><em>Pseudomonas aeruginosa</em></td>
<td><em>hsi-I</em></td>
<td>Hcp1</td>
<td>VgrG1</td>
<td>Two-component system sensor RetS</td>
<td>Attenuated virulence in rat lung infection; anti-Hcp1 produced in CF patients</td>
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<td>Two-component system sensor LadS</td>
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<td>PpkA/PppA S/T kinase-phosphatase</td>
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<td>Putative $\sigma^{54}$ activator (PA1663)</td>
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<td>Stk1/Stp1 S/T kinase-phosphatase</td>
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<td>Putative $\sigma^{54}$ activator (PA2359)</td>
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<td>No S/T kinase-phosphatase or Fha</td>
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<td>Induced by host-plant extract</td>
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<td>Temperature-dependent</td>
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<td>ImpM/ImpN S/T kinase-phosphatase</td>
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<td><em>Pectobacterium atrosepticum</em></td>
<td><em>hsi-II</em></td>
<td>Hcp1, -2, -3, -4, VgrG</td>
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<td><em>Rhizobium leguminosarum</em></td>
<td><em>imp</em></td>
<td>RbsB homologue</td>
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<td><em>Salmonella enterica</em> subspecies I</td>
<td><em>sci</em></td>
<td>SciK (Hcp), VgrS (VgrG)</td>
<td>Induced <em>in vivo</em> (macrophage and rabbit ileal loop)</td>
<td>Increased bacterial number in macrophages</td>
<td>Hypervirulent in mice</td>
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<td>Response regulator SsrB</td>
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<td><em>Vibrio cholerae</em></td>
<td><em>vas</em></td>
<td>Hcp1, -2, VgrG1, -2, -3</td>
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<td>Attenuated virulence on <em>Dictyostelium</em> and macrophages</td>
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<td>Induced <em>in vivo</em> (rabbit model for cholera)</td>
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<td>RpoN ($\sigma^{32}$)</td>
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<td>Putative $\sigma^{54}$ activator (VasH)</td>
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<td>Fha but no S/T kinase-phosphatase</td>
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*Regulatory components that control T6SS expression and function have been indicated as well as conditions in which T6SS genes are specifically controlled. The role of the putative $\sigma^{54}$ activator in expression of T6SS genes has not been demonstrated. However, in *V. cholerae*, a putative $\sigma^{54}$ activator is required for the T6SS-dependent phenotype as well as RpoN ($\sigma^{32}$).†The attenuation or hypervirulence is indicated when it has been observed for at least some of the described T6SS mutants.
component system. However, the two studies are not really comparable in terms of models (J774 macrophage vs Hep2 cells) and in the study by Folkesson et al. (2002) a strain with a complete deletion of the Sci genomic island was used. The Sci island also contains saf fimbral genes which may be required for invasion.

The T6SS in Pseudomonas aeruginosa

Three IAHP/T6SS clusters are found in Pseudomonas aeruginosa (Fig. 2, Table 1) (Das & Chaudhuri, 2003). It has been demonstrated that one of these clusters, named HSI-I (Hcp1 secretion island I) (Mougous et al., 2006) and spanning genes identified as PA0071 to PA0091, contributed to P. aeruginosa pathogenesis. Previous studies also found that P. aeruginosa PA01 mutants affected in PA0073, PA0077, PA0082, PA088 and PA0090 showed an attenuated virulence phenotype in a model of chronic infection in the rat lung (Potvin et al., 2003). This suggested that the HSI-I genes are induced in vivo, as is the case with the homologous V. cholerae vas genes. In laboratory growth conditions, the HSI-I genes could be induced by deleting the retS gene, encoding a two-component system sensor protein, which is part of a signalling pathway important for establishing chronic infections (Goodman et al., 2004). In a retS mutant, an Hcp-like protein could be detected in the culture supernatant. The gene encoding the Hcp protein is PA0085 and is part of the HSI-I gene cluster (Table 1). Mougous and collaborators further assessed whether mutation in HSI-I genes could interfere with Hcp secretion. The deletion of either PA0090 or PA0077 (icmF-like gene) genes resulted in a dramatic decrease of Hcp secretion. In V. cholerae, in addition to Hcp protein, VgrG proteins are secreted in a T6SS-dependent manner (Pukatzki et al., 2006). In the study by Mougous and collaborators, even though a VgrG protein is encoded within the HSI-I gene cluster (PA0091), no indication was obtained about its secretion (Table 1).

Some T6SS components

The T6SS machine, like other SSs, is defined by a set of original components. The T6SS gene clusters usually encode between 12 and 25 proteins, among which only DotU and IcmF-like proteins are found in another SS (T4SS). The cluster does not necessarily contain the genes encoding the Hcp and VgrG proteins, which may be distant from the T6SS locus. Most of the genes encode proteins with unknown function but some have known characteristics.

ATPases are frequently used by SSs to energize the transport process. In the case of the T6SS, a gene encoding a ClpB homologue is most frequently found. The Clp/ Hsp100 family belongs to the ring-forming AAA+ superfamily of ATPases (Neuwald et al., 1999). The ClpB homologues found in T6SS clusters could be distinguished as a subfamily, which was named ClpV. In prokaryotes, ClpB members are key players in protein quality control. They form hexameric rings and use ATP hydrolysis to insert substrates to be unfolded within their central channel, that is the proteolytic chamber, for further degradation (Weibezahn et al., 2004). Unlike ClpB, it was shown that ClpV from enterohaemorrhagic E. coli (EHEC) and Salmonella typhimurium failed to solubilize aggregated proteins (Schlieker et al., 2005). Moreover, substrates stimulating the activity of ClpB ATPases failed to do so with ClpV. Therefore, the function of ClpV might be different from that of other ClpB members, such as providing energy-dependent coupled transport of poly-peptides (Yeo & Waksman, 2004). Nevertheless, due to the similarity with the Clp/Hsp100 family, another possibility is that the ClpV component is required to unfold/fold components of the T6SS machine to be assembled or effectors to be secreted. A similar role was proposed for ATPases involved in the T3SS (Akeda & Galan, 2005). The energy specifically required for the movement of proteins through the T6SS could then be provided by other components and might alternatively involve the proton-motive force.

Apart from clpV, another gene is frequently found in T6SS clusters, which encodes a putative lipoprotein. This is for example the case for PA0080 (HSI-I) in P. aeruginosa or VCA0113 (VasD) in V. cholerae (Fig. 2). In most cases, the amino acid in position +2 after the putative cleavage site is a serine, indicating that it is a putative outer-membrane lipoprotein (Yamaguchi et al., 1988). Whether this protein contributes to pore formation in the membrane is unknown and proof for such a hypothesis may await the demonstration that it could be inserted into the outer membrane independently of its lipid anchor.

What is secreted, what is injected and what are the effectors needed for?

Most recent studies have suggested that proteins most likely to be secreted by the T6SS belong to the Hcp or VgrG families.

The Hcp family

The secretion of Hcp proteins in a T6SS-dependent manner was most significantly demonstrated in V. cholerae (Pukatzki et al., 2006). In this bacterium there are two hcp genes (hcpA, -B or hcp1, -2), encoding identical proteins (Table 1). The hcp gene products have a predicted size of about 28 kDa, but they show an aberrant mobility on gels since their apparent molecular mass is 18–19 kDa. The hcp2 gene (VC0017) is in the vicinity of the V. cholerae T6SS cluster (vas; VC0107–VC0123) whereas hcp1 is more distantly located (VC1415). The Hcp proteins do not contain a canonical signal peptide, indicating that they are not secreted in a Sec- or Tat-dependent manner, and probably cross the bacterial cell envelope in a single step (Pallen et al., 2003). Interestingly, an hcp1/hcp2 mutant is...
avirulent, whereas individual hcp1 or hcp2 single mutants retain virulence. This indicated that T6SS-dependent secretion of at least one Hcp protein is required and sufficient for virulence.

The secretion of Hcp protein was confirmed in *P. aeruginosa* (Mougous et al., 2006). Each of the *P. aeruginosa* T6SS clusters contains or is associated with hcp genes, hcp1 (PA0085/HSI-I), hcp2 (PA1512/HSI-II) or hcp3 (PA2367/HSI-III) (Fig. 2). The function of the HSI-I/T6SS system was studied in a retS background, which allowed overproduction of HSI-I genes and abundant secretion of Hcp1 (Mougous et al., 2007), suggesting that these proteins may also serve a T6SS function despite the fact that they appeared to be secreted substrates. Several vgrG genes located outside T6SS clusters are found in association with hcp genes. The Vgr proteins were initially considered as accessory components of the Rhs (recombination hot spot) family (Wang et al., 1998b). Rhs components are complex genetic composites, and in *E. coli*, eight components, RhsA–H, are known. Rhs components include a core ORF (G/C rich) and an adjacent core extension (A/T-rich) with linked downstream ORFs (A/T-rich). Two ORFs immediately upstream of the rhsG core were identified and named vgrG and hcp (Wang et al., 1998b). Although they are not homologous, vgrG and the RhsG core ORFs are similar in a number of respects. They are both large, their predicted products are hydrophilic, and both are characterized by a regularly repeated peptide motif. These motifs are YDxxGRL(I/T) for the RhsG core protein and a Val-Gly dipeptide motif (Valine glycine repeats) in the case of VgrG. In the case of the RhsE element, a vgr gene, vgrE, was also found upstream of the rhsE core ORF (Wang et al., 1998b). Interestingly, within the *S. enterica* T6SS gene cluster (sci), in addition to the vgrS gene, two genes were annotated rhs1 and rhs2, rhs2 encoding a protein with similarity to the *E. coli* RhsE protein (Folkesson et al., 2002).

Characterization of VgrG proteins revealed further remarkable features. In *V. cholerae*, the VgrG1 C terminus shares a domain (ACD) with the RtxA toxin, which mediates actin cross-linking (Sheahan et al., 2004). The functionality of the VgrG1 ACD domain was tested in COS-7 cells. As with RtxA ACD, ectopic expression of VgrG1 ACD produces cell rounding and actin cross-linking within the transfected cells. The activity of the VgrG1 ACD domain was confirmed *in vitro* using a purified VgrG1 protein, monomeric G-actin and cytoplasmic extracts of amoebae or macrophages (Pukatzki et al., 2007). The VgrG proteins

![Fig. 3.](Image)
containing a C-terminal extension were named ‘evolved VgrGs’. In *V. cholerae*, VgrG3 has a peptidoglycan-binding domain, whereas no extension was found in VgrG2.

Independently of C-terminal extensions, VgrGs share a conserved region, which contains two domains that showed similarities with the gp5 and gp27 proteins that constitute the bacteriophage T4 tail spike. Dimer of trimers of gp5 and gp27 constitutes the tail spike, which is used for puncturing the bacterial envelope and allows DNA injection into the bacterial cytoplasm (Kanamaru et al., 2002; Rossmann et al., 2004). Pull-down experiments using *V. cholerae* VgrG-directed antibodies revealed that these proteins form homotrimeric or heterotrimeric complexes (Pukatzki et al., 2007). The assembly of the VgrG proteins into a putative tail-spike-like structure suggests that the resulting structure may be used as a puncturing device, to allow perforation of the bacterial cell envelope or of the host cell membrane, or both. In any case, VgrGs appear to be not simply secreted proteins but structural components of the T6SS machine. This is reminiscent of the other family of secreted proteins, Hcp, which form hexameric rings with a central channel of 40 Å (4 nm) (Mougous et al., 2006). The tube formed by the bacteriophage T4 gp5/gp27 complex is about 30 Å (3 nm) in diameter (Kanamaru et al., 2002). Pukatzki and colleagues speculated that the VgrG complex may adopt a similar structure and could be surrounded by the Hcp rings, both forming the tube for T6SS substrates. If that were the case, Hcps and VgrGs may be appended at the bacterial surface and possibly released into the culture supernatant by shearing.

The presence of catalytic domains in some VgrG proteins does not support the idea of VgrGs being solely T6SS components. Considering the example of the VgrG1 ACD domain, one has to think that it should be translocated into host cells. Specifically, the ACD domain is conserved in VgrGs' V. cholerae homologues containing (pertussis toxin) and signal-peptide-lacking substrates (*Helicobacter pylori* CagA) are transported in a T4SS-dependent manner (Backert & Meyer, 2006). Whether RbsB secretion in *R. leguminosarum* is significant remains unclear, but strikingly, a gene encoding an RbsB homologue is found downstream of the vgrG3 gene in the *V. cholerae* T6SS cluster (Bladergroen et al., 2003).

### The key role of threonine phosphorylation in type VI secretion

Another remarkable feature of T6SS gene clusters is that they generally encode Ser/Thr kinases and phosphatases (Fig. 2, Table 1). Mukhopadhyay et al. (1999) reported that bacteria carrying a gene encoding an IcmF-like protein, which is now recognized as part of the HSI-II/T6SS system of *P. aeruginosa*. In *Yersinia*, the Ser/Thr kinase YpkA is involved in T3SS virulence (Cornelis, 1998), and it is considered that some bacterial Ser/Thr kinases have host targets due to their similarity with mammalian proteins. Mukhopadhyay et al. (1999) suggested that Stk1 and Stp1 could be translocated in host cells by a mechanism involving IcmF and thus the T6SS. It is now clear that it is not the case.

In *P. aeruginosa*, in addition to stk1 and stp1, genes encoding Ser/Thr kinases and phosphatases, ppkA and pppA, respectively, are found in the T6SS/HSI-I cluster. The ppkA gene was previously found specifically induced within the host and required for virulence in neutropenic mice (Wang et al., 1998a; Motley & Lory, 1999). Since the T6SS/HSI-I gene cluster is upregulated in the retS background, Mougous and collaborators could monitor T6SS function when introducing mutations in either ppkA or pppA (Mougous et al., 2007). They showed that PpkA and PppA have antagonistic activities since a ppkA mutation resulted in defective Hcp1 secretion, whereas mutation in pppA resulted in increased Hcp1 secretion. The antagonistic effect of PpkA and PppA on T6SS function was confirmed by investigating T6SS localization. Using a ClpV1-GFP chimera, it was shown that the *P. aeruginosa* ClpV1 protein is localized in foci in the bacterial cell (Mougous et al., 2006). However, ClpV1 was found evenly distributed throughout the cell in *hcp1* or *icmF1* mutants, indicating that the T6SS machine assembles as a macromolecular complex. Interestingly, the localization of ClpV1 into foci was observed in a pppA mutant, but not in a ppkA mutant. This suggested that both T6SS assembly and Hcp1 secretion require phosphorylation by PpkA and are prevented by dephosphorylation through PppA. Mougous and colleagues recognized a gene from the T6SS/HSI-I cluster encoding a protein with an FHA (forkhead-associated) domain. Proteins with FHA domains have affinity for phosphothreonine and have been...
implicated in phosphorylation-dependent signalling (Pallen et al., 2002). The gene was named fha1 and a mutation in this gene resulted in defective Hcp1 secretion. The phosphorylation status of Fha1 was shown both in vitro and in vivo to be PpkA/PppA-dependent and occurred on Thr-362. Overall, Mougous and collaborators suggest that, in standard conditions, the level of Fha1 phosphorylation is kept low by PppA. However, upon sensing of unknown environmental cues by PpkA, the kinase may overrule PppA activity and Fha phosphorylation initiates a signal transduction cascade that results in T6SS assembly and function. In conclusion, PpkA/PppA/Fha1 play a crucial role in controlling activity of the P. aeruginosa T6SS/HSI-I at the post-translational level. Whether this is applicable to all known T6SSs is questionable. Some clusters do not contain Ser/Thr kinase-phosphatase and Fha homologues (Fig. 2, Table 1). That is true for the P. aeruginosa T6SS/HSI-III but one may suggest that it uses heterologous components from the HSI-I or HSI-II systems. In the case of V. cholerae, the T6SS gene cluster encodes an Fha homologue (VCA0112) but no Ser/Thr kinase-phosphatase (Table 1). This could suggest that Fha could be phosphorylated by a kinase encoded by a gene outside the T6SS gene cluster.

Controlling the expression of T6SS genes

Independently of post-translational control of T6SS function, expression of T6SS genes is highly regulated. In R. leguminosarum, T6SS secretion is temperature-dependent, since putative T6SS effectors were optimally secreted at 24 °C and only weakly at 20 °C (Bladergroen et al., 2003). Other observations have revealed that the T6SS could be induced under in vitro conditions. That was the case for V. cholerae (Das et al., 2000), for which induction of the icmF gene (vasK) was shown by comparing the global transcription response of V. cholerae cells grown overnight in rabbit ileal loops (in vivo) and in rich media (in vitro).

In S. enterica, induction of T6SS genes, and more precisely sciS, encoding the IcmF-like protein, was shown to occur inside macrophages (Parsons & Heffron, 2005). Expression of a sciS–lacZ transcriptional fusion was not observed when Salmonella was grown in vitro. Salmonella invasion and multiplication in macrophages relies on two T3SSs. T3SS-1 is involved in earlier killing, whereas T3SS-2 mediates host cell death at later stage (18–24 h post-infection). However, bacteria can persist in macrophages beyond 24 h, and this seems to be dependent on silencing the T6SS. Indeed, a sciS mutant is able to survive and multiply in macrophages at a late stage post-infection without lysis of the host cell. The response regulator SsrB, part of the two-component system SsrA/SsrB, negatively controls expression of sciS. In contrast, it was shown that SsrA/SsrB positively controls genes encoded on Salmonella pathogenicity island 2 (SPI-2), including the T3SS-2 genes (Garmendia et al., 2003). Finally, in contrast to T3SS, the Salmonella T6SS favours bacterial persistence, and limits intracellular replication and bacterial load in host tissues, which is consistent with the hypervirulent phenotype of a sciC mutant in mice (Parsons & Heffron, 2005).

In P. aeruginosa, a balance between expression of genes involved in bacterial persistence and chronic infection versus genes involved in cytotoxicity and acute infection is nicely documented. As previously mentioned, expression of the T6SS/HSI-I gene cluster from P. aeruginosa is induced in vitro in a retS mutant. RetS is a hybrid sensor, which was identified as positively controlling expression of genes required for cytotoxicity (T3SS and exoenzymes) and negatively controlling genes required for biofilm formation (exopolysaccharide biogenesis gene clusters pel and psl) (Goodman et al., 2004). The negative control of the P. aeruginosa T6SS by RetS is reminiscent of the negative control of the Salmonella T6SS by SsrB. It is also important to recall that antibodies against T6SS components were found in sera of CF patients chronically infected with P. aeruginosa (Mougous et al., 2006), which supports the idea that T6SS favours persistence. Interestingly, in P. aeruginosa another hybrid sensor, LadS, is antagonist to RetS and acts positively on T6SS gene expression (Ventre et al., 2006). Since T6SS gene expression could not be seen in vitro except in a retS mutant, one may speculate that the T6SS genes may be controlled in vivo upon recognition of specific host signals by the LadS/RetS pathway.

In P. aeruginosa, LadS and RetS do not influence expression of T6SS genes from HSI-II and HSI-III. Within each cluster, one gene encodes a σ54 activator: PA1663 for HSI-II and PA2359 for HSI-III (Fig. 2, Table 1). Activators of bacterial σ54-RNA polymerase holoenzyme use ATP hydrolysis to activate transcription by promoting the transition from a closed RNA polymerase (RNAP) promoter complex to a transcriptionally competent open complex (Rappas et al., 2005). In the V. cholerae T6SS gene cluster, a σ54 activator is encoded by vasH. A mutant in vasH was attenuated in Dictyostelium, as was a mutant affected in rpoN (Pukatzki et al., 2006), the gene encoding the σ54 alternative subunit of RNAP, which suggests that VasH plays a role in the likely σ54-dependent expression of V. cholerae T6SS genes.

Overall, regulatory processes appear to tightly control T6SS gene expression during the host–pathogen interaction and prevent premature or inappropriate expression of these genes, which may negatively affect in vivo survival.

The T6SSs in other Gram-negative bacteria: hallmarks and differences?

Besides the early discovered, or archetypal, T6SSs of R. leguminosarum, V. cholerae, S. enterica and P. aeruginosa, our knowledge about T6SSs in other bacteria is fragmentary. The data collected do not really contribute to further understanding of the T6SSs but point to a certain degree of variability.
Burkholderia mallei, Burkholderia pseudomallei and Burkholderia thailandensis possess multiple T6SS clusters, i.e. four, six and five, respectively. In B. mallei, one T6SS gene cluster (tssA–tssN) is upregulated upon overproduction of the VirAG two-component system (Schell et al., 2007), which resulted in an increased T6SS-dependent secretion of the Hcp1 protein. A B. mallei hcp mutant is attenuated in the hamster infection model (Schell et al., 2007). As with P. aeruginosa Hcp1, antibodies against B. mallei Hcp1 were found in sera of infected animals. Although VgrG proteins are encoded within Burkholderia T6SS clusters, their secretion has not been demonstrated. Interestingly, a non-Hcp and non-VgrG putative T6SS substrate was proposed to be the B. mallei TssB protein. The tssB gene is part of the T6SS cluster, and whereas a hcp mutant was not impaired in Hcp1 secretion it was attenuated for virulence in hamsters, similarly to a hcp mutant (Table 1). This observation suggested that if TssB is not part of the T6SS machine, it is involved in a T6SS-dependent phenotype and might be a T6SS effector (Schell et al., 2007). Finally, tssM encodes a protein containing an ubiquitin-specific protease domain (Schell et al., 2007). Since ubiquitination of proteins does not occur in bacteria, one can speculate that TssM is injected into the host cell.

In the case of Edwardsiella tarda, the T6SS cluster was named evp and contains 16 genes (Zheng et al., 2005; Zheng & Leung, 2007). Mutations in evp genes resulted in attenuated virulence in blue gourami fish (Table 1). A previous report on Edwardsiella ictaluri (Moore et al., 2002) identified eip genes similar to the imp genes from Rhizobium, and eip gene products were recognized by the catfish immune system during infection, indicating that they are produced in vivo. In Ed. tarda, it was shown that 13 eip genes were required for secretion of EvpC, an Hcp homologue (Rao et al., 2004), and EvpP, a secreted protein with no homology with either VgrG or Hcp proteins (Zheng & Leung, 2007) (Table 1). Sequence variation between Hcps is high, and EvpP shares several features with this family. It has no canonical signal peptide and runs aberrantly on polyacrylamide gels. Such aberrant migration was observed with V. cholerae Hcps. The formation of a T6SS complex involving the IcmF-like component EvpO was tested using a two-hybrid approach. Relevant interactions with EvpA, EvpL (lipoprotein) and EvpN (DotU) were found. No interaction between EvpH (ClpV) and any other Evp components was found, contradictory to results in P. aeruginosa, which suggests an interaction between ClpV1, Hcp1 and IcmF1 (Mougous et al., 2006).

In the case of enteraggregative E. coli (EAEC), expression of chromosomal genes that are part of a T6SS cluster (aai–aaiY; AggR-activated island) is under the control of the virulence-plasmid-encoded AggR (aggregative adherence) transcriptional regulator (Dudley et al., 2006), a member of the AraC family (Sheikh et al., 2002). This plasmid encodes additional virulence factors such as the Pet enterotoxins, the aggregative adherence fimbriae (aafDA) (Nataro et al., 1994) and the dispersin (aap) (Sheikh et al., 2002). The aai cluster is located in an EAEC pathogenicity island (117 kb) and is not found on the E. coli K-12 genome. It was shown that the aai cluster is required for secretion of AaiC. AaiC has no similarity with Hcp proteins, but like EvpP from Ed. tarda, has a size of about 18–19 kDa and does not contain a signal peptide. The aai cluster contains a gene encoding a VgrG-like protein but its secretion was not observed (Table 1). A second AggR-independent T6SS was found on the EAEC genome. The cluster consists of 21 genes and was named EAECSci-II, whereas the aai cluster was named EAECSci-I. In the EAECSci/I/T6SS cluster an hcp-like gene encodes a product that is secreted in a T6SS-dependent manner (Table 1).

In Pectobacterium atrosepticum, which causes diseases in potatoes and other plants, the production of four Hcps was observed (Mattinen et al., 2007). Mutation in one hcp gene did not impair virulence but overproduction of Hcp increased virulence. In vivo induction of T6SS gene expression is likely to occur in P. atrosepticum, since secretion of VgrG and Hcp proteins was induced by addition of potato tuber and stem extracts to the bacterial cultures (Mattinen et al., 2007).

In Francisella tularensis, a gene cluster encompassing 16–19 genes and located on a pathogenicity island (FPI) is needed for intramacrophage growth and virulence in chicken embryos (de Bruin et al., 2007; Nano et al., 2004; Nano & Schmerk, 2007). The organization of these genes is conserved in Francisella species, but is distinct from the well-characterized T6SS organization in other species. Two genes, iglA and iglB, have similarities to impB and impC, respectively. Expression of the iglAB genes is induced during growth in macrophages and is under the positive control of MglA (de Bruin et al., 2007) (Table 1), previously described as a master regulator for Francisella virulence (Lauriano et al., 2004). MglA is similar to the E. coli stringent starvation protein A (SspA), an RNAp-associated protein (Hansen et al., 2005). The mglA gene is linked with mglB, which is also required for intracellular growth of Francisella, and which encodes a protein similar to SspB from E. coli (Baron & Nano, 1998). The pdpB (pathogenicity determinant protein) gene encodes an IcmF-like protein, whereas the pigF (pathogenicity island gene) gene has similarities to dotU. The pigB gene encodes a protein with similarity to the VgrG-family of proteins. The iglD gene encodes a protein with some similarities to ImpI, particularly in the N-terminal domain. The iglD gene is essential for intracellular replication in primary human monocyte-derived macrophages (Santic et al., 2007). Other genes, iglC, pigC and pigG, seem to be unique to Francisella. Finally, no gene encoding a homologue to ClpV protein is found, which overall makes this system at the limits of variation to classify it as a T6SS.

**Evolutionary aspects of T6SSs**

The screening of databases shows that there are almost 100 different bacterial species with a T6SS but it has been...
experimentally studied in only a few. Whereas it is mainly found in pathogenic or symbiotic bacteria of the phylum Proteobacteria, the T6SS is also found in non-pathogenic soil bacteria such as *Pseudomonas putida* and *Myxococcus xanthus* (Bingle et al., 2008). Recently, a phylogenetic analysis distinguished four branches (A–D) within the T6SS tree (Bingle et al., 2008). Interestingly, multiple T6SS clusters from a single bacterium are found in different groups. For example, *P. aeruginosa* HSI-I is in group A (together with *B. pseudomallei* and *S. typhimurium*), HSI-II is in group B (together with *R. leguminosarum*) and HSI-III is in group D (together with *V. cholerae*). The three T6SS clusters are found in all *P. aeruginosa* genome sequences that are available, i.e. PAO1, PA14 and PA7 (http://www.pseudomonas.com). *Yersinia pestis* has five T6SS loci, which are spread between the A, C and D groups. This observation indicates that multiple T6SS loci may not have arisen through duplication but rather been acquired by horizontal gene transfer. The T6SS from *F. tularensis* is phylogenetically distant from the other T6SSs, and constitutes an outward fifth branch. This confirms the weak sequence similarities with other T6SS components that we have described above.

**Conclusion**

The hallmark of the T6SS is a set of approximately 15 genes found within clusters, which may vary in terms of organization. It is thought that T6SS genes encode components of a secretion machine and specific effectors. In several species, multiple T6SS loci can be found, which in some cases are incomplete. It is not known whether those loci that are most incomplete cooperate with other T6SSs. Consequently, it is difficult to unambiguously define the core components for a functional system. T6SS components that have been discussed are presented in a speculative model in Fig. 4.

Most T6SS clusters contain a gene encoding an AAA\(^+\) ATPase, ClpV, which may be the T6SS motor. ClpV should be considered as a core component; however, in *R. leguminosarum*, which is a T6SS archetype, no clpV-like gene is found in the *imp* cluster. One possibility is that some T6SSs use a protein encoded by a *clpV* gene located elsewhere on the chromosome. In order to cross the bacterial cell envelope, a membrane channel is necessary. Apart from DotU and IcmF none of the T6SS components possess transmembrane domains. There is a putative outer-membrane lipoprotein but this is unlikely to form a pore in the membrane if it is only anchored via the lipid modification.

The VgrG proteins from *V. cholerae* may form a puncturing device similar to the bacteriophage tail spike. This device may be used to perforate the bacterial envelope, and further the host cell membrane, to transport effectors all the way. Another class of proteins, the Hcps, have been shown to form hexameric rings, which may form an extracellular conduit that extends or surrounds the tube formed by the VgrG proteins. These observations are difficult to reconcile with the idea that Hcps and VgrGs are the T6SS substrates. One may compare this with the T3SS, which transports the translocator proteins to be inserted into the host cell membrane and subsequently the effectors to be translocated into the host cell cytosol (Edqvist et al., 2007). T6SSs could contribute to assembling VgrGs and Hcps outside the bacterial cell, into a putative conduit that...
may be further used by as yet unknown T6SS effectors to be injected into host cells.

Some VgrGs may have a dual role, somewhat similar to what has been described for autotransporters, in which both the transporter and the catalytic domain are carried on a single polypeptide (Henderson et al., 2004). Indeed, some VgrGs possess a C-terminal extension, such as an actin cross-linking domain in the V. cholerae VgrG1. This domain could be transferred across the bacterial and host membranes through the VgrG puncturing device, and eventually released by proteolytic cleavage into the host cell (Fig. 4). The presence of a tropomysosin-like domain, which putatively manipulates actin filaments, at the C-terminus of one of the Yersinia VgrGs also supports the idea that the T6SS is an injection machine (Pukatzki et al., 2007). The presence of domains at the C-terminus of VgrG proteins, which have homology with bacterial adhesins, such as YadA, pertactin, mannose-binding domain or fibronectin-like domain, suggests that other transported domains may not be injected but remain attached at the bacterial cell surface (Pukatzki et al., 2007). The identification of Hcps and VgrGs in the supernatant may be an artefact due to mechanical release of a surface-exposed structure into the medium, as with bacterial flagellin (Fernandez & Berenguer, 2000).

Whether substrates other than Hcps or VgrGs could travel through T6SSs is under debate and few candidates have been proposed. Whether all T6SS substrates are translocated into host cells or whether some are released into the extracellular medium is under debate too. One such controversy is illustrated by the finding that, in R. leguminosarum, extracellular addition of T6SS-dependent secreted protein prevents the formation of nodules on pea plants (Bladergroen et al., 2003). In other words T6SS substrates may act even though added exogenously. In the case of V. cholerae this was shown not to be the case and instead, as with the T3SS or T4SS, bacterial–host contact is needed to transport T6SS effectors into the cell. Too little is known at the moment to overspeculate and it will be wise to await the elucidation of the biochemical function and target of the putative T6SS secreted effectors before we reconsider what could be their likely final localization.

T6SS genes are mostly not induced in laboratory growth conditions but are induced in vivo during infection. Several two-component systems have been identified that may play a crucial role in inducing/repressing expression of the T6SS genes while the bacteria are colonizing the host and encountering different environmental conditions. The T6SS seems to be involved in the intracellular behaviour of bacteria and allows chronic and persistent infection (Yahr, 2006). Furthermore, in some cases, the activity of the T6SS has been shown to be dependent on the phosphorylation status of Fha, which is subject to the antagonistic activity of Ser/Thr kinase-phosphatase. The Ser/Thr kinase-phosphatase/Fha system is lacking in the T6SS of some bacterial species or is incomplete (Table 1). The presence of such a phosphorylation system that times T6SS assembly and function (Kulasekara & Miller, 2007) may not be mandatory, but may provide a useful switch to control delivery of effectors only upon detection of specific environmental signals. In bacterial species lacking such post-translational control, the T6SS may function constitutively once assembled.

The function and characteristics of the T6SS are far from being understood but the system appears as a novel key player in bacterial pathogenesis and bacteria–host interaction. It may become in the future another target in the hunt for novel antimicrobials, which will help fight human morbidity and mortality due to persistent bacterial infections.

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References


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