Bacterial Secretion Systems with an Emphasis on the Chlamydial Type III Secretion System

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Abstract

Numerous bacterial proteins exert their function outside the prokaryotic cell. To this end, both Gram-negative and Gram-positive bacteria have evolved specialized mechanisms to transport their proteins to the bacterial supernatant or host cell cytoplasm, so called secretion systems. These different strategies will be briefly discussed, followed by an in depth description of the Type III secretion system, an efficient molecular syringe assisting Gram-negative bacteria in entrance, growth and survival in eukaryotic host cells. Topics addressed include classification and role of multiple Type III secretion systems. the mechanism of protein translocation into the host cell as well as substrate recognition and chaperoning.

Chlamydiales have also been found to encode a Type III secretion system and associated effector proteins. In contrast to the genetic organization in other bacteria, the encoding genes are scattered throughout the genome. To date, no structural information is available on the chlamydial Type III secretion system. We therefore propose a model of the chlamydial Type III secretion system and summarize current knowledge on the role of Type III secretion in the different stages of the chlamydial developmental cycle.

Bacterial secretion systems

Introduction

Bacterial pathogens have evolved numerous strategies to efficiently infect and colonize their eukaryotic hosts. Most often, secreted or surface-exposed bacterial proteins play central roles in the interaction of these pathogens with their host cells, mediating diverse processes such as proteolysis, hemolysis, cytotoxicity and phosporylation or dephosphorylation of host cell proteins (Hueck, 1998). In order to fulfil their function they have to be transported from the bacterial cytoplasm to the supernatant or the bacterial surface, a process which is called secretion. In Gram-negative bacteria, the barrier to be crossed consists of an inner and an outer membrane, separated by the periplasmic space. Often, Sec or Tat systems transport unfolded and/or folded proteins first to the periplasmic space (export, see further), where they are processed (e.g. addition of disulphide bridges) to obtain their final three-dimensional structure, before being transported across the outer membrane. **Translocation** refers to the transport of proteins directly across the eukaryotic cell membrane into the host cell cytoplasm where they can interfere with cellular processes and suppress host defences (Buttner and Bonas, 2002). In addition, transport through the outer membrane requires ATP or other sources of energy, no indications of which exist that they are present at the outer membrane. Secretion systems must therefore be self-energized or harness energy at the cytoplasmic side of the inner membrane.

Until recently, six major pathways by which these proteins are transported from the bacterial cytoplasm to the extracellular space had been identified (Thanassi and Hultgren, 2000), four of which depend on the 'general secretory pathway' (GSP or Sec pathway): Type II and Type IV secretion, autotransporters (Type V secretion) and the chaperone/usher pathway. These Sec-dependent pathways export proteins with a short amino terminal signal sequence (~ 30 amino acids) through the inner membrane to the periplasmic space. where the largely hydrophobic signal sequence is cleaved off by a peptidase (Hueck, 1998; Thanassi and Hultgren, 2000). Alternatively, Type II secretion can also be Tat-dependent: proteins are targeted across the inner membrane in their fully folded conformation, by an amino terminal signal peptide containing two arginine residues (Panahandeh et al., 2009). Type I and Type III secretion do not depend on the Sec system to fulfil their function. Although first described in 1996, another protein secretion mechanism was recognized in 2006 as being the Type VI secretion mechanism (Mougous et al., 2006; Pukatzki et al., 2006). Figure 1 gives an overview of these seven major secretion mechanisms. which will be discussed briefly, except the Type III secretion system. Since Type III secretion is the main focus of this review, it will be described in more detail further on.

Protein export to the periplasm

Two parallel pathways for the export of proteins across the inner membrane are described in prokaryotes: the Sec-dependent pathway and the Tat (twin-arginine translocation) pathway. Both systems can deliver their substrates to the Type II secretion pathway.

Sec-dependent protein export

The Sec-pathway is evolutionarily conserved among bacteria, archaea and eukaryotes, mediating export of proteins across the bacterial inner membrane or endoplasmic reticulum (Stephenson, 2005). All Secexported proteins contain an amino terminal signal peptide (~30 mainly hydrophobic amino acids) targeting them to the membrane embedded SecYEG translocon, serving as a polypeptide conducting channel, through which no tightly folded preproteins can pass. Premature folding of a preprotein is avoided by interaction with specific targeting factors: 1) in the SRP pathway, SRP targets the preprotein to the translocon (adjacent to an SRP receptor) while still in its nascent, ribosome associated state, followed by integration in the lipid bilayer (co-translational targeting); 2) the other is the SecB pathway, with chaperone SecB binding to the full-length translated preprotein, targeting the protein to the SecA-SecYEG pore. Subsequently, the preprotein is exported for secretion in the periplasm (Ito and Mori, 2009). This post-translational mode of export is driven by the SecA ATPase, generating energy from ATP hydrolysis and the proton-motive force. The signal peptide is usually cleaved off on the periplasmic side by a signal peptidase (SPI), encoded by *lepB* in *E. coli*, releasing the mature protein into the periplasmic space (Rusch and Kendall, 2007). Certain archaeal Sec signal sequences contain a lipobox motif ([I/L/G/A]-[A/G/S]-C), ensuring recognition by the type II signal peptidase (SPII), and anchoring of the mature lipoprotein to the cytoplasmic or outer membrane (Pohlschröder and Dilks, 2007).

Twin-arginine translocation pathway

The twin-arginine translocation pathway is found in bacteria, archaea and eukaryotes to transport proteins

into or across the bacterial inner membrane (Berks et al., 2000; Wu et al., 2000) or across the chloroplast thylakoid membrane (Settles et al., 1997). Contrary to the Sec-pathway, proteins of variable dimensions are exported in their fully folded conformation across the membrane. without rendering cytoplasmic the membrane freely permeable to protons and other ions. Bacterial proteins are targeted to the Tat pathway by an amino terminal signal peptide containing an almost invariant twin-arginine sequence motif, SRRxFLK, where x is a polar amino acid, while Tat signals in chloroplasts are more variable (Panahandeh et al., 2009). The Tat pathway not only delivers proteins to the periplasm, but also allows integration of inner membrane proteins in e.g. E. coli (Hatzixanthis et al., 2003), Paracoccus denitrificans (Bachmann et al., 2006) and Legionella pneumophila (De Buck et al., 2007). The Tat translocase of E. coli consists of the membrane proteins TatA. TatB. and TatC (Sargent et al., 1998). The transmembrane TatBC complex binds Tat secretion substrates with TatC recognizing the precursor protein consensus signal peptide, followed by proton motive force dependent recruitment of TatA to form a fully assembled and functional Tat translocase. After export of the passenger domain across the membrane, without compromising the membrane potential, the TatABC complex dissociates again and the signal peptide is cleaved off by periplasmic signal peptidases (Panahandeh et al., 2009).



Figure 1: Major protein-secretion systems in Gram-negative bacteria. Four of these protein secretion pathways of depend on the Sec system for protein transport across the inner membrane. Type I, III and VI secretion are Sec-independent. Type I secretion systems predominantly secret toxins, proteases and lipases into the extracellular milieu, whereas Type III secretion systems also translocate virulence proteins into the host cell. Type VI secretion mediated by a novel kind of a complex multi-component secretion machine, is often involved in interaction with eukaryotic hosts. Type IV secretion systems mediate the transport of DNA and proteins across the double membrane to bacterial or eukaryotic cells, requiring direct cell-to-cell contact. Examples are the Bordetella pertussis toxin (Sec-dependent, a) or the Agrobacterium tumefaciens VirB/D4 sytem for transport of T-DNA-protein complexes (Sec-independent, b). Autotransporters form a pore through the outer membrane and are therefore classified among the Type V secretion systems. The chaperone/usher pathway (C/U) consists of an OM protein, termed an usher, and a periplasmic chaperone, guiding proper folding and preventing premature interactions. Complexes of 12-16 proteins mediate transport of extracellular enzymes and toxins in Type II secretion: a pilus-like structure of four IM proteins pushes the proteins, delivered to the periplasm by the Sec or Tat system, through an OM pore.

Legend: C, bacterial cytoplasm; IM, bacterial inner membrane; P, bacterial periplasm; OM, bacterial outer membrane; ECM, extracellular milieu. PM, host cell plasma membrane. When appropriate, coupling of ATP hydrolysis to transport is highlighted. Arrows indicate the route followed by transported proteins. Adapted from Buttner and Bonas (2002) and Filloux et al. (2008).

Type I secretion system

Type I secretion systems (T1SSs), also known as ATPbinding cassette (ABC) transporters, are used by a vast number of Gram-negative bacteria for the secretion of various molecules, ranging from ions and drugs to proteins. Substrates include toxins (leukotoxine of Pasteurella haemolvtica and hemolvsin A of E. coli). proteases (metalloproteases of Pseudomonas aeruginosa and Erwinia chrysanthemi), hemophores (hemoproteins of Serratia marcescens and Pseudomonas aeruginosa) and lipases (Serratia marcescens LipA). An overview of substrates for the T1SS can be found in Jenewein et al. (2009). The secreted molecules vary in size from the small E. coli peptide colicin V of 10 kDa (Gilson et al., 1990) to the Pseudomonas fluorescens cell adhesion protein LapA of 900 kDa (Hinsa et al., 2003). Type I secretion is also involved in export of non-proteinaceous substrates like cyclic β-glucans and polysaccharides (Jenewein et al., 2009). Both in eukaryotic and prokaryotic organisms, related ABC transporters are conserved, transporting different toxins and antibiotics (Andersen, 2003).

In Gram-negative bacteria, Type I secretion is carried out by a translocator made up of three proteins that span the cell envelope: an inner membrane ATPase (the actual ABC transporter), a specific outer membrane protein (OMP, exported through the Sec-pathway) and the so-called membrane fusion or adaptor protein (MFP), connected in the inner membrane and spanning the periplasmic space. The ABC protein ensures that only T1SS specific substrates are recognized via a carboxy terminal secretion signal, whereafter MFP binds the substrate on the cytoplasmic side. The MFP also links the inner membrane components to the OMP, forming a pore in the outer membrane. The secretion signal, mostly rich in the glycine-repeated motif GGxGxD, located within the carboxy terminal 50-60 amino acids, is specific for the different subfamilies of the T1SS. Type I secretion is Sec-independent and secretes the protein directly to the extracellular medium, without cleavage of the signal peptide (Delepelaire, 2004; Jenewein et al., 2009).

Type II secretion system

Type II secretion systems (T2SSs) are widely conserved and common, but by no means universal, among Gramnegative bacteria (α -, β -, γ -, δ - and ϵ -Proteobacteria). Predominantly extracellular toxins and hydrolytic enzymes (e.g. cellulases, elastases, amylases, proteases, phosphatases, lipases, nucleases,...) are secreted by the T2SS of bacterial pathogens of plants (Pseudomonas fluorescens, Erwinia spp., Xanthomonas spp.), animals (Aeromonas hydrophila) and humans (Klebsiella oxvtoca. Pseudomonas aeruginosa. Legionella pneumophila), as recently reviewed by Cianciotto (2005) and Michel and Voulhoux (2009). Before the discovery of the Tat pathway, the T2SS was regarded as the 'main terminal branch of the general secretory pathway', extending the Sec export pathway. This led to confusion when 'general secretory pathway' was also used to describe the whole process of Secdependent export followed by Type II secretion (T2S). With the Tat pathway appearing in the discussion, it was proposed to reserve the designation 'GSP' to describe the Sec-pathway, while retaining the name 'T2SS' to

describe a **two-step process** where the protein is first exported into the periplasmic space via a **translocase** (Sec or Tat, Table 1) and subsequently transported across the outer membrane by a dedicated secretion apparatus called the '**secreton**' (Desvaux et al., 2004b). The signal sequence in cleaved off by a periplasmic peptidase, thus releasing the mature protein.

The secreton, spanning the periplasm and connecting the inner and outer bacterial membranes is encoded by a cluster of 12 to 16 *gsp* genes, depending on the species. Peabody et al. (2003) noticed that some bacteria seem to have a significant, but incomplete, number of T2S core proteins (e.g. *Aquifex aeolicus* encodes *gspDEFGO*, *Thermotoga maritima gspDEFG*, and *Chlamydiaceae* harbor *gspCDEFG*), which was confirmed by Cianciotto (2005). Further research is needed to determine whether these T2S homologs encode a functional secreton.

Exoproteins that use the T2SS are characterized by the presence of a leader peptide (or signal peptide) at their N terminus. The T2SS is unique in its ability to secrete large multimeric proteins that are folded in the periplasm and species specificity is mainly related to the gatekeeper proteins GspC and GspD (Michel and Voulhoux, 2009). GspG, H, I and J encode pseudopilins, pushing the substrate through GspD in the outer membrane. GspE contains a conserved ATP binding motif and exhibits autokinase activity, thus providing energy and regulating the whole secretion process (Thanassi and Hultgren, 2000).

At least four constituents of the T2SS (GspE, F, D and O) share homology with components of Type IV pili (T4P) of *Pseudomonas aeruginosa* and other bacteria (Hueck, 1998; Peabody et al., 2003) and of DNA transfer systems of e.g. *Haemophilus influenzae* and *Bacillus subtilis* [reviewed in Averhoff and Friedrich (2003)]. The T2S homologs identified in some bacteria might therefore also function in piliation or DNA uptake.

Recently, a novel protein targeting system was identified in Dickeya dadantii, formerly classified as Erwinia chrysanthemi. A pectin lyase, PnIH, contains a wellconserved Tat signal sequence, but lacks any obvious signal peptidase cleavage site, and thus is not processed in the periplasmic space. Subsequently, the protein is targeted to the outer membrane by a novel T2SS, where it remains localized on the surface of the cell, probably because the uncleaved amino terminal Tat signal acts as an outer membrane anchor. The authors hypothize that this is likely to be a general mechanism as E. coli was able to successfully integrate proteins bearing the D. dadantii signal peptide into its own outer membrane. This second T2SS is encoded on a cluster of 11 stt (second type two) genes, all of which share homology to the standard T2SS, and its substrate pnIH is encoded immediately downstream of this stt cluster (Ferrandez and Condemine, 2008).

Type IV secretion system

Type IV secretion systems (T4SSs) are macromolecular nanomachines which transport macromolecules (proteins, DNA or DNA-protein complexes) across the bacterial cell envelope of Gram-negative and Grampositive bacteria into the extracellular milieu or directly into recipient cells. They can be classified according to sequence homology into 1) T4SSs related to the

Agrobacterium VirB/D4 archetypal tumefaciens mechanism (Type IVA), 2) Legionella (L.) pneumophila Dot/Icm related T4SSs (Type IVB, some of which are classified as Type VI secretion. described further on) and 3) other T4SS not yet fully characterized. Alternatively, a classification according to function can be made: 1) conjugative transfer of plasmids and other mobile DNA elements to recipient bacteria, 2) delivery of protein or substrates to eukaryotic target cells ('effector translocator' systems), 3) exchange of DNA substrates with the extracellular milieu ('DNA release or uptake systems') or 4) exclusive transport of proteins to the exterior medium or directly into recipient cells [reviewed in Christie et al. (2005) and Baron (2009)].

The best studied model of a T4SS is the T-DNA transfer system of Agrobacterium tumefaciens. It comprises 12 proteins named VirB1 through 11 and VirD4. VirB4. VirB11 and VirD4 contain conserved Walker A nucleotide-binding/hydrolysis motifs, energizing the assembly of the T4SS and/or substrate translocation. However, VirD4 appears to function as a general receptor for DNA and protein, operating independently from ATP energy (Christie and Cascales, 2005). Core components VirB6, VirB7, VirB8, VirB9 and VirB10 likely form the translocation channel, while the T-pilus is built up by VirB2 and VirB5, the major and minor surface-exposed pilus subunits, respectively. These pili mediate cell-to-cell contact and substrate translocation. but it is unclear whether substrates actually move through the pili (Baron, 2005; 2009).

Only limited information is available regarding protein secretion by the T4SS. In Helicobacter pylori, CagA is translocated through the T4SS to the host cells where it induces disruption of cell-to-cell junctions, actin cytoskeletal rearrangement as well as proliferative, proinflammatory and anti-apoptotic nuclear responses (Brandt et al., 2005; Backert and Selbach, 2008). Bordetella pertussis toxin (Ptx), involved in whooping cough, is also secreted through a T4SS, but instead of directly entering a target cell like most other effector proteins, it is secreted into the extracellular medium. Ptx is composed of five different protein subunits (S1-S5), which are first exported into the bacterial periplasm by the Sec-pathway (Table 1). After cleavage of the signal peptides, components assemble to form the holotoxin, followed by extracellular secretion. Once secreted, Ptx alone mediates host cell binding and delivery of its catalytic S1 subunit into the host cytosol. Thus, protein secretion and host cell translocation of the effector are not linked in this particular T4SS (Rambow-Larsen and Weiss, 2004). The L. pneumophila Type IVB SS (Dot/Icm) is known to deliver proteins into target cells, e.g. RalF required for the localization of ARF (ADPribosylation factor) on phagosomes containing L. pneumophila (Nagai et al., 2002). T4SSs in other bacteria can display additional

components or have homologs for only a subset of those proteins described in this section. A recent review on the topic is published by Baron (2009).

Table 1: Protein export, secretion and translocation systems in bacteria. G⁺, Gram-negative; G⁺, Gram-positive. Adapted from Economou et al. (2006).

(2000).					
Name	Process	Sec/Tat dependence ^a	G	G⁺	Remarks
Sec (SRP-SecB)	Export	NA	Yes	Yes	
Tat	Export	NA	Yes	Yes	
T1SS	Secretion	No	Yes	Yes	Generally small substrates in G ⁺
T2SS	Secretion	Yes	Yes	No	
T3SS	Translocation	No	Yes	No	
Flagellum	Secretion	No	Yes	No	Equivalent to T3SS
T4SS	Translocation	Variable	Yes	Yes	Conjugation system
T4P	Secretion	Yes	Yes	Yes	Equivalent to T2SS
T5SS	Secretion	Yes	Yes	No	
T6SS	Translocation	No	Yes	No	
C/U	Secretion	Yes	Yes	No	Assembly of pili and fimbriae
^a For translagated prot	ain: accombly of moch	inony dooo roquiro Coo			

^a For translocated protein; assembly of machinery does require Sec

NA: not applicable

Type V secretion systems: autotransporters and twopartner secretion

Type V secretion systems (T5SSs) are present in a wide range of Gram-negative bacteria for secretion of large virulence proteins across the outer membrane. Until now, three families have been recognized: the **autotransporter** (AT) system (Type V_a or AT-1), the **two-partner secretion pathway** (Type V_b), and the more recently described **Type** V_c system (also termed AT-2) (Desvaux et al., 2004a). Proteins secreted via these pathways have similarities in their primary structures as well as striking similarities in their modes of biogenesis. Shared characteristics include: 1) an amino terminal **signal sequence** for inner membrane export via the Sec machinery (Table 1), 2) a functional **passenger domain** that can be surface-exposed or released into the extracellular milieu, 3) a **linker region** necessary for translocation of the passenger domain through the outer membrane, 4) a carboxy terminal region involved in the formation of a **transmembrane pore**. Once through the inner membrane, the signal sequence is cleaved and a β -domain inserts into the outer membrane to form a β -barrel pore. The passenger domain is then translocated to the bacterial cell surface via the pore, where it may or may not undergo further processing. This process is assisted by an internal autochaperone sequence in the passenger domain (Henderson et al., 2004; Scott-Tucker and Henderson, 2009). However, recent studies reveal the probable

involvement of accessory proteins such as periplasmic chaperones and BamA OM protein as well [overview in Scott-Tucker and Henderson (2009)].

The <u>autotransporters</u> are most prevalent in the phylum Proteobacteria, including the α -, β -, γ - and ϵ -Proteobacteria. Other phyla in which autotransporters have also been identified are *Chlamydiae*, genera *Chlamydia* (*C*.) and *Chlamydophila* (*Cp.*) (Henderson and Lam, 2001) and Fusobacteria (Desvaux et al., 2005). The chlamydial polymorphic membrane proteins or pmps contain the necessary Sec signal sequence for export through the inner membrane and can be divided into at least six subtypes based on phylogenetic analysis (Henderson and Lam, 2001). Although specific roles in pathogenesis have been described for some pmps [reviewed in Tan et al. (2006)], a role in the avoidance of the host immune response as already earlier proposed by Tanzer et al. (2001), is highly probable.

Whether or not the autotransporter pathway is energydependent is highly debated. Neither ATP nor GTP is present in the periplasm, so hydrolysis of these molecules cannot provide the energy for translocation of the passenger domain through the outer membrane and no evidence exists for a proton motive force across the outer membrane. It has been suggested that release of the auto-chaperone or folding of the β -barrel or passenger domain might provide the required energy [overview in Henderson et al. (2004)].

The autotransporter pathway differs from the two-partner secretion pathway in the fact that, instead of being produced as a single polypeptide, the passenger domain and the pore forming β -domain are translated as two separate proteins, referred to as TpsA (also called the exoprotein) and TpsB (also called transporter domain) family members, respectively. The genes encoding these two proteins are usually organized in an operon. Specific secretion signals are present in a highly conserved region of the secretion domain of TpsA. In addition to its channel function, TpsB probably also serves as a specific receptor of the TpsA signal peptide (Jacob-Dubuisson et al., 2001). Characteristic examples of autotransporters and the two-partner secretion pathway respectively are the IgA1 proteases of Neisseria spp. (Pohlner et al., 1987) and the 220 kDa filamentous hemaglutinin of Bordetella spp. (Jacob-Dubuisson et al., 2001).

The <u>AT-2</u> pathway is very similar to the classical autotransporters, but contains a remarkably short carboxy terminal translocator domain. Hence, three AT-2 polypeptides are required to form a functional β -barrel pore, which also functions as an adhesin of various bacteria (e.g. Hia of *Haemophilus influenza*, YadA of *Yersinia* (Y.) *enterocolitica* and UspA of *Moraxella catharralis*) (Scott-Tucker and Henderson, 2009).

Type VI secretion systems

Although only named Type VI secretion system (T6SS) in 2006 (Pukatzki et al., 2006), indications on the presence of T6SSs date from a decade earlier, with Hcp (haemolysin co-regulated protein) from *Vibrio cholerae* shown to be secreted without cleavage of a signal peptide. In the years that followed, similar mechanisms were described in *Rhizobium leguminosarum*, *Pseudomonas aeruginosa* and *L. pneumophila*. Das and Chaudhuri named the encoding gene clusters IAHP, for

'IcmF-associated homologous proteins', because they contain a gene encoding a homolog of the IcmF protein associated with Type IVB secretion in L. pneumophila (Das and Chaudhuri, 2003). This gene cluster is conserved among Gram-negative bacteria but the majority of proteins encoded have no homology to T4SSs or any other known secretion system components (Bingle et al., 2008). So far, in silico analysis has proven the existence of one or two T6SS clusters in more than 100 bacterial species, but a few species, including Burkholderia spp., can contain up to six clusters. Whether these multiple systems represent a redundancy for virulence or specificities for particular niches or hosts is still unclear. Core components conserved among all T6SSs include the T4SS IcmF- and IcmH-like proteins, a putative lipoprotein, the CIvP AAA+ ATPase (possibly providing the energy) and the Hcp and VarG (valine-alvcine repeats) proteins (Cascales, 2008). The presence of a functional T6SS in distinct bacterial isolates is often associated with the ability to induce host diseases or disorders. In the past three years, several functions have been attributed to translocated T6SS effector proteins, including adhesion to and invasion of the host cell and promoting intracellular growth within macrophages or survival and persistence within the host. These and other subversions of host cell functions (e.g. signaling cascades. inflammatorv responses. intracellular trafficking,...) are elegantly summarized in Cascales (2008).

Chaperone/usher pathway

The chaperone/usher (C/U) pathway is implicated in the assembly and secretion of a superfamily of virulenceassociated surface structures in Gram-negative bacteria (Thanassi et al., 1998). Only two proteins are required to mediate transport through the outer membrane: a periplasmic chaperone protein and an integral outer membrane protein termed usher. Typical structures assembled on the outer membrane through the C/U pathway are rigid, rod-like fibres, termed pili or fimbriae, but amorphous capsular and thin fibrillar structures have also been observed (Thanassi et al., 2009). C/U pathways are subdivided into six major phylogenetic groups, termed α -, β -, γ -, κ -, π -, and σ -fimbriae (Nuccio and Baumler, 2007). The prototypes of the C/U pathway are the adhesive Type 1 and P pili of uropathogenic E. coli (Abraham et al., 1988; Marklund et al., 1992). These pili are composite structures built from multiple subunit proteins, which are first exported to the periplasmic space via the Sec system (Table 1). In the periplasm, the chaperones bind to the subunits through a conserved carboxy terminal motif, allowing cleavage of the amino terminal Sec signal peptide. This way, correct folding is ensured and premature interactions between the pili subunits are prevented. The chaperone-subunit complexes are then targeted to a dimeric usher in the outer membrane, each containing a central β-barrel channel domain. Interaction of a first complex with one of both ushers, induces dissociation of the chaperone and opening of the usher channel (2-3 nm) allowing the subunit to pass. A second chaperone-subunit complex binds to the second usher in a similar way, and both subunits interact at the bacterial cell surface. Repetition of the whole process (which is energy-independent) allows extension of the pilus and twisting of the pili rod into its final helical conformation (Thanassi et al., 2005; 2009).

An alternative C/U pathway exists, exhibiting the same function as the classical C/U pathway, but displaying low sequence homology. It is found primarily in enterotoxigenic *E. coli* (ETEC) strains (Li et al., 2007).

Type III secretion system

Introduction

Unlike Type I and Type II secretion systems, secreting active proteins in the bacterial supernatant, Type III secretion systems (T3SSs) predominantly translocate proteins to the cytoplasm of eukaryotic host cells (Hueck, 1998). Factors inducing protein secretion include contact with the surface of the target cell ('contact dependent secretion', cds) and environmental factors such as a shift in temperature to 37 °C, a change in extracellular pH or iron concentration and depletion of calcium ions to millimolar concentrations. An overview of these phenomena can be found in Ramamurthi and Schneewind (2002).

The T3SS is by far the most complex of protein secretion systems known today: it is composed of approximately 25 proteins, predominantly localized in the cell envelope, spanning both the bacterial inner and outer membrane as well as the plasma membrane of the eukaryotic cell when actively translocating effector proteins (Plano et al., 2001). Most inner membrane structures are homologous to components of the 'flagellar basal body' in both Gram-negative and Gram-positive bacteria (Kubori et al., 1998; Macnab, 1999). The secretin SctC, localized in the outer membrane shows homology to PuID, the outer membrane secreton of the T2SS (Hueck, 1998). Genes coding for the structural components and translocated effectors are mostly clustered in a single pathogenicity island (PAI) on a plasmid or within the bacterial chromosome, leading to the assumption that these clusters are inherited as a single unit (Plano et al., 2001). Although the structural components of T3SSs are generally conserved, the actual arsenal of bacterial proteins that they deliver (collectively known as 'effectors' or T3SEs) is unique to each system (Galan and Wolf-Watz, 2006). Most of these effector proteins require cytoplasmic chaperones to protect the secreted substrates from premature interactions with the other components of the T3SS (Hueck, 1998). The whole process is energized by a cytoplasmic, inner membrane associated ATPase (Akeda and Galan, 2004).

Type III protein translocation is independent of the Secpathway and no periplasmic intermediate is generated through amino terminal processing, similar to Type I and Type VI secretion. However, several structural components of the T3SS do harbor the typical amino terminal Sec signal peptide, indicating that the Secpathway is probably involved in T3SS assembly (Hueck, 1998).

The T3SS is not necessarily a virulence hallmark

Since it was first described in *Yersinia* less than twenty years ago (Michiels et al., 1990), T3SSs to translocate specialized proteins to the cytoplasm of a eukaryotic host cell or the lumen of an intracellular organelle have been identified in numerous Gram-negative bacteria,

especially Proteobacteria, but so far, none were found in ε-Proteobacteria. The following list aims to give an overview but is far from complete, as with the increasing number of bacterial genomes being sequenced, more bacterial species with T3SSs will be identified. The T3SSs are best characterized in bacteria infecting mammals, e.g. Yersinia spp., Salmonella spp., Chlamydia and Chlamydophila spp., Shigella spp., Pseudomonas aeruginosa, enterohemorrhagic (EHEC) and enteropathogenic (EPEC) E. coli, Bordetella spp., Burkholderia spp., Edwarsiella tarda, Citrobacter rodentium and Chromobacterium violaceum (Hueck, 1998; Pallen et al., 2003a; Brito et al., 2004; Cornelis, 2006). Pathogens infecting fish, such as Aeromonas salmonicida subsp. salmonicida, Aeromonas hydrophila, Vibrio parahaemolyticus and Yersinia ruckeri also deploy T3SSs as a virulence determinant (Burr et al., 2003; Gunasena et al., 2003; Makino et al., 2003a; Pallen et al., 2003a), as is also the case for plant pathogens Xanthomonas spp., Erwinia spp., Pseudomonas spp. and Ralstonia solanacearum (Hueck, 1998; Pallen et al., 2003a). However, not only pathogens harbor a T3SS: Rhizobium spp. (soil bacteria, nitrogen fixation in plants), Sodalis glossinidius (tse-tse fly Glossina spp. symbiont) and SZPE (Sitophilus zeamais primary endosymbiont, in grain weevils) and other endosymbionts also possess a T3SS (Dale et al., 2001; 2002; Marie et al., 2001). Genome sequencing of the Protochlamydia (P.) amoebophila UWE25 symbiont of amoeba revealed genes encoding a complete T3SS as well as effector proteins (Horn et al., 2004) as is the case for commensals, such as non-pathogenic E. coli strains (Pallen et al., 2003a) or mutualists, e.g. Photorhabdus luminescens in entomophagous nematodes (Cornelis, 2006).

Role of multiple T3SSs in the same cell

In some bacteria, multiple T3SS encoding gene clusters are present. In Salmonella (S.) enterica, PAIs SPI-1 and SPI-2 code for two different T3SSs, inv/spa and spi/ssa, respectively (Hensel et al., 1995) and both systems depend on a common post-translational isomerase (Pallen et al., 2005). Likewise, three Yersinia-species (Y. pestis, Y. pseudotuberculosis and Y. enterocolitica) harbor, apart from a well-characterized plasmid encoded Ysc-Yop system, a chromosomally encoded T3SS (Ysa, SPI-2 like), which although degenerate is important in Yersinia virulence (Haller et al., 2000). Moreover, crosstalk and functional overlap between both systems have been demonstrated (Young and Young, 2002). In the case of Vibrio parahaemolyticus, a ubiquitous chromosome-1 T3SS is implicated in cytotoxicity, whereas the chromosome-2 encoded system, only present in strains pathogenic to humans, mediates enterotoxicity (Park et al., 2004). Most commensal and pathogenic E. coli strains also encode two T3SSs, the second being SPI-1 like (Makino et al., 2003b; Ren et al., 2004). Burkholderia pseudomallei and related species can harbor up to three T3SSs: two systems, TTS1 and TTS2, are homologous to T3SSs of plant pathogens and a third system (TTS3 or Bsa) is similar to SPI-1 of S. enterica. Fascinatingly, the TTS1, which is similar to T3SSs found in plant pathogens, appears to be restricted to *B. pseudomallei* strains that are pathogenic to humans (Rainbow et al., 2002; Stevens et al., 2002).

As a relationship between *B. pseudomallei* and plants has already been proposed (Mahfouz et al., 2006), its TTS1 may allow this human pathogen to infect plants as well.

A possible reason for the presence of multiple T3SSs is that the different systems function at different stages of the infection. This has already been established for *Salmonella*, where the major function of the SPI-1 T3SS is to enable invasion of epithelial cells in the early stage of infection, while the SPI-2 T3SS is required to allow replication inside macrophages. Subsequently, the bacteria can spread throughout the body and induce a systemic infection. Thus, the two T3SSs, each with its specific set of regulators and effectors, are needed during different periods of infection [reviewed in Schlumberger and Hardt (2006)].

A similar phenomenon occurs in *Chlamydiaceae*. Instead of possessing two separate T3SSs, only the components that are built into the plasma membrane of the eukaryotic cell (translocators CopB and CopD) and their specific chaperone are encoded twice by the genome. It is believed that CopB1 and CopD1 are required for translocation of effectors mediating internalization of the pathogen, while CopB2 and CopD2 are built into the inclusion membrane, to allow transport of effectors reprogramming the host cell functions to maintain intracellular growth (Ouellette et al., 2005; Hefty and Stephens, 2007).

Different families of T3SSs

As already mentioned in the introduction, structural components and translocated effectors are often clustered on elements with a high plasticity (plasmids or PAIs). Phylogenetic analysis on their amino acid sequences reveals the existence of at least seven different clades or families of related T3SSs: Ysc-Yop, SPI-1 (Inv-Mxi-Spa), SPI-2 (Ssa-Esc), Hrc1 (Hrp1) and Hrc2 (Hrp2) in plant pathogens, *Chlamydiales* (Sct) and *Rhizobiales* (Gophna et al., 2003; Pallen et al., 2005; Troisfontaines and Cornelis, 2005), each with their specific nomenclature (Table 2). Carrying out the analysis on different constituents of the T3SS generates the same tree, whatever the component used for computing (Gophna et al., 2003) and (Troisfontaines and Cornelis, 2005).

This indicates that all of the genes encoding T3SSs have evolved as large, intact genetic blocks. Within each of the seven families, genetic organization of the loci is identical (syntheny). The most striking feature of this evolutionary tree is that it is completely different from the bacterial evolutionary tree based on 16S rRNA (Gupta, 2000), implying that Type III secretion has been distributed among bacteria through horizontal gene transfer. For bacteria with multiple T3SSs, the corresponding proteins in each T3SS are xenologs, indicating that the presence of two T3SSs in a species did not result from a gene duplication event inside the species but rather is the consequence of two successive horizontal gene transfers (Troisfontaines and Cornelis, 2005).

So far, most described horizontal transfer of T3SS encoding gene clusters occurred among species sharing similar ecological niches (e.g. between plant pathogens or animal pathogens, but not from animal to plant pathogens), again indicating that physical proximity between donor and acceptor bacteria is indispensable for horizontal gene transfer events.

Translocation of effectors - models

The global architecture of the structural components in the bacterial membranes is well conserved among the different families of T3SSs. The basal structure consists of a cylindrical structure spanning the inner and outer membrane, associated with a C-ring like structure (homologous to the flagellum MS ring) and an ATPase. Depending on the families, a hollow stiff needle, a filament or a pilus extends from the basal structure, and consists of 100-150 molecules arranged according to a helical polymerization pattern (Blocker et al., 2008). Its length varies from 40 to 80 nm in animal pathogens, up to several µm in the plant pathogen Pseudomonas syringae allowing to protrude the thick plant cell wall (Cornelis, 2006). Upon contact with a target cell, a hetero-oligomeric translocation pore is built into the host cell membrane. Generally, three translocators make up the pore (e.g. YopB, YopD and LcrV in Yersinia spp.; IpaB, IpaC and IpaD in Shigella spp.), only the third (LcrV, IpaD) of which is hydrophilic, serving as a scaffold for the other, hydrophobic, translocators (Cornelis, 2006). In plant pathogens, only one T3SS translocon family protein been identified so has far. HrpF/NopX/PopF, exhibiting only limited homology to the YopB family from animal-pathogenic bacteria (Meyer et al., 2006). However, the HrpF homologs from Xanthomonas oryzae pv. oryzae and Pseudomonas syringae pv. tomato are not essential for bacterial pathogenicity, suggesting that additional proteins are involved (Buttner and Bonas, 2006).

The internal central channel of the whole system is approximately 2-3 nm in diameter, which is too small to allow folded proteins to pass through the tube. Therefore, effector proteins need to be unfolded prior to translocation (see further).

For animal-infecting bacteria, mechanisms of how T3SEs interact with the host cell are largely unknown, unlike for plant pathogens, whose primary role is to suppress the basal defence response. Described mechanisms include suppression of defence signaling, salicylic-acid dependent and cell wall based defences, as well as alteration of ethylene production. However, they may also induce a strong defence response, if the host possesses a specific resistance protein corresponding to a particular avirulence (Avr) protein (McCann and Guttman, 2008).

Substrate recognition – In search of the bar code

Whether or not T3SEs contain a conserved signal sequence has long been debated and research has predominantly focused on Yop effectors of *Yersinia* spp. It was first proposed that the secretion signal was an encrypted non-cleaved amino terminal sequence in the protein as these amino termini showed no significant amino acid homology. Relatively radical changes in amino acid sequence did not affect secretion, suggesting that the secretion signal of T3SEs was conformational rather than sequence based (Michiels and Cornelis, 1991). Anderson and Schneewind (1997) postulated that the signal was embodied within the 5' region of the mRNA coding for the substrate and that secretion was coupled to translation.

Table 2: Frequently used nomenclature for the conserved structural proteins of the T3SS across the different families and their homologs in the flagellum. Legend: Chlam.: Chlamydiales; Rhiz.: Rhizobiales; TL: translocon component; IM: inner membrane; IM: outer membrane; CP: cytoplasmic; MA: membrane associated; HCM: host cell membrane. Adapted from Hueck (1998), Meyer et al. (2006), Blocker et al. (2008) and Mueller et al. (2008).

Sct	Ysc-Yop	SPI-1	SPI-2	Hrc1	Hrc2	Chlam.	Rhiz.	Flagella	Subcellular localization
SctV	YscV	InvA MxiA	SsaV	HrcV Hrpl	HrcV HrpO	SctV CdsV	PcrD	FlhA	IM
	LcrD					LcrD			
SctW	YscW	InvE	1		/	SctW	1	1	Secreted
	YopN LcrE	MxiC		HrpJ		CopN LcrE			
SctN	YscN	InvC Spal	SsaN	HrcN	HrcN HrpE	SctN CdsN	HrcN	Flil	CP, MA
SctO	YscO	Invl SpaM	SsaO	HrpO	1	SctO CdsO	ORF7	1	?
SctP	YscP	InvJ	SsaP		1	SctP	1	1	Secreted
		SpaN		HrpP		CdsP			
SctQ	YscQ	InvK	SsaQ	HrcQ	HrcQ	SctQ	HrcQ	FliN	IM
		SpaO		A&B	HrpQ	CdsQ		FliM	
		·						FliY	
SctR	YscR	InvL	SsaR	HrcR	HrcR	SctR	HrcR	FliP	IM
		SpaP		HrpW	HrpT	CdsR			
SctS	YscS		SsaS	HrcS	HrcS	SctS	HrcS	FliQ	IM
		SpaQ		HrpO	HrpU	CdsS			
SctT	YscT	InvN	SsaT	HrcT	HrcT	SctT	HrcT	FliR	IM
		SpaR		HrpX	HrpC	CdsT			
SctU	YscU		SsaU	HrcU	HrcU	SctU	HrcU	FlhB	IM
		SpaS		HrpY	HrpN	CdsU			
SctC	YscC	InvG	SsaC	HrcC	HrcC	SctC	NolW	1	OM
		MxiD	SpiA	HrpH	HrpA	CdsC			
SctD	YscD		SsaD			SctD	1	FliG	IM
			SpiB	HrpQ	HrpW	CdsD			
		PrgH			HrpD5				
SctF	YscF		SsaG	1	1	SctF	1	1	Needle
		MxiH				CdsF			
		Prgl							
Sctl	Yscl		1		1	/	1	1	CP?
				HrpB					
	LcrO	PrgJ							
SctJ	YscJ		SsaJ	HrcJ	HrcJ	SctJ	NoIT	FliF	Link between IM and OM
	YlpB	MxiJ			Hrpl	CdsJ			
		PrgK							
SctK	YscK	/	1		1	/	1	1	CP?
				HrpD					
SctL	YscL	1	SsaK	HrpE	HrcL	SctL CdsL	NolV	FliH	CP
TL	YopB	SipB	SseC		PopF	CopB1/2	/	1	HCM
		IpaB		HrpK	HrpF				
					NopX				
TL	YopD	SipC IpaC	SseD	1	1	CopD1/2	1	/	НСМ
TL	LcrV	SipD IpaD	SseB	1	1	LcrV	1	/	НСМ

Further research indicated that a physical property, namely amphipathicity, of the amino terminus is critical for secretion, although at least some secondary signal might reside in the mRNA (Lloyd et al., 2001). More recent research indicates the presence of a chaperonebinding domain (CBD) located within the first 100 amino acids of the effector protein, downstream from a short, uncleaved, amino terminal export signal (Birtalan et al., 2002).

A more detailed discussion on this topic can be found in Sorg and Cornelis (2009). In *Chlamydiaceae*, the bilobed

hydrophobic domain of the inclusion membrane proteins (Incs) was proven indicative for secretion by a T3SS (Subtil et al., 2001). Research by the same group indicated that, for *Chlamydiaceae* as well, a signal must be present within the 14 amino terminal amino acids of effector proteins, although no conserved sequence pattern could be detected (Subtil et al., 2005). Based on these observations a prediction tool for T3SEs was developed: EffectorT3, which is available online at <u>www.chlamydiaeDB.org</u>. Tests across different taxa indicate that a common mechanism is not restricted to a certain phylum, and that the ability to be transported by the T3SS seems to have been re-invented by different families of effectors several times (Arnold et al., 2009).

Specific T3SS chaperones

Generally spoken, chaperones assist in folding or unfoldina and assembly or disassembly macromolecular structures, such as proteins or protein complexes. Based on structural and functional analyses. at least five classes of specific T3SS chaperones can be distinguished. Chaperones of Class I assist the effector proteins, while Class II chaperones interact with the translocator proteins. Class III groups the chaperones of the related bacterial flagellar apparatus. Class IV is represented by only one member, CesA (chaperone for E. coli secretion A), accompanying the EspA filament protein of enteropathogenic E. coli (Yip et al., 2005). Yersinia YscE and its Pseudomonas homolog PscE, with their co-chaperones YscG and PscG, are classified as Class V (Wilharm et al., 2007). Class I and Class II chaperones are well characterized and will be discussed in further detail.

Class I chaperones

As mentioned above, Class I chaperones specifically interact with the T3SEs. A distinction is made whether only a single substrate is bound (Class IA, which is the case for the majority of them), or whether multiple substrate proteins can be served (Class IB, the so-called 'promiscuous chaperones'). Common features include their small size, acidic pl, their dimeric nature and being encoded adjacent to their corresponding cognate effector gene (Pallen et al., 2005). Although exhibiting a very low sequence homology (maximum 20% when aligning two different chaperone sequences), they do share a conserved fold. Moreover, T3SEs wrap their CBD around chaperone homodimers to hind hydrophobic regions on the surface of the chaperones. The common fold may therefore reflect a shared effector recognition pattern (Wilharm et al., 2007). Functions attributed to the chaperones include acting as a 3D export signal, establishing a temporal export hierarchy and storage of the effector proteins in an exportcompetent state prior to export. In addition, their presence could direct the effector protein to the ATPase, which unfolds exported proteins in an ATP-dependent manner and subsequently detaches the chaperone from its substrate, a function similar to that done by the bacterial AAA⁺ ATPase disassembly machineries (Akeda and Galan, 2005; Sorg and Cornelis, 2009).

Class II chaperones

Class II chaperones protect bacteria from the toxic properties exerted by the hydrophobic translocator proteins (Menard et al., 1994; Wattiau et al., 1994; Neyt and Cornelis, 1999). Examples are IpgC of *Shigella flexneri*, LcrH (SycD) of *Yersinia* spp., and Scc2/3 of *Chlamydiaceae*. All Class II chaperones share the same structure of three tandem tetratricopeptide repeats (TPRs), consisting of 34 amino acids per repeat and involved in protein-protein interactions (Pallen et al., 2003b). Similar repeats can be found in host cell chaperones Hsp70 and Hsp90. Darwin and Miller showed that unbound SicA of *S. typhimurium* interacts with the transcriptional regulator InvF, to activate

transcription from the effector gene promoters (Darwin and Miller, 2000). A similar transcriptional regulation has been described for IpgC in conjunction with MxiE (Mavris et al., 2002), while LcrH is presumed to be involved in mRNA degradation (Anderson et al., 2002). The higher concentration of free chaperones upon assembly of the translocon in the host cell membrane could then inform the bacterium that the pore is made and the T3SS is fully functional (Page and Parsot, 2002). Fascinatingly, the plant-associated Hrc1 and Hrc2 T3SSs do not seem to employ TPR chaperones (Pallen et al., 2005).

How these translocator chaperones exactly bind to their substrates is still a matter of debate. Is the CBD a unique region or not? Do they form a complex with both translocators or are both substrates bound independently in the cytoplasm? An overview on these topics and the stability of translocator-chaperone complexes can be found in Sorg and Cornelis (2009).

T3SS in Chlamydiales

History and first evidence of a chlamydial T3SS

Long before the first visualization of a T3SS in Salmonella (Kubori et al., 1998), Matsumoto and coworkers already described the presence of rosette-like structures and projections on the surface of both elementary bodies (EBs) and reticulate bodies (RBs) of different chlamydial species (Matsumoto, 1973; 1982b; Matsumoto et al., 1976). These hexagonally arranged patches, delineating a zone of close contact between the RB and inclusion membrane were long thought to facilitate the uptake of nutrients from the host cell cytoplasm, an idea designated as the "soup-through-astraw" hypothesis (Stephens, 1992). The first indication for the existence of a chlamydial T3SS came from Hsia et al. (1997) who identified a locus of seven, nonoverlapping open reading frames in Cp. caviae GPIC, the first four of which encode proteins with significant homology to T3SS proteins in other species. Shortly thereafter the same group postulated that the envelope spanning components of the chlamydial surface projections represent the chlamydial T3SS. The observation that this type of 'contact dependent secretion' is present in contact zones, protruding both IM and OM as well as the host cell membrane is consistent with a channel mediating the translocation of proteins from the bacterial cytoplasm into the host cell cytosol (Bavoil and Hsia, 1998). Subsequent genomic analysis revealed the presence of a complete set of T3S encoding genes in all Chlamydiaceae species sequenced so far (Stephens et al., 1998; Read et al., 2000; Shirai et al., 2000; Kalman et al., 1999; Azuma et al., 2006; Read et al., 2003; Thomson et al., 2005) as is the case for the environmental chlamydiae P. amoebophila UWE25 (Horn et al., 2004) and Simkania negevensis (Peters et al., 2007, Meyers et al., unpublished results).

Genetic organization

Chlamydiales are the only non-Proteobacteria that harbor a T3SS (Troisfontaines and Cornelis, 2005). In most Proteobacteria, genes encoding the structural components of the T3SS are clustered within a single chromosomal PAI or on a virulence-associated plasmid,

with sizes up to 200 kbp, flanked by vestiges of recent integration events, e.g. direct repeats or insertion sequences (IS) (Winstanley and Hart, 2001) and within the different T3SS families, gene order within the loci is well conserved (see above). By contrast, genes coding for the chlamydial T3SS are located in at least four clusters dispersed throughout the genome, predominantly coding for structural components, specific chaperones for effector proteins, and two translocator systems to transport secreted proteins across the host cell membrane, at least in *Chlamydiaceae*. The presence and location of effector encoding genes, but not structural genes differs between the species (Figure 2).

	Cluster 1			Cluster 2		Cluster 3		Cluster 4
С. trachomatis ст	088 090 115 117 119 228 2 089 091 116 118 229	32 442 456 529 5 541 5	59 561 563 562 564	570 572 569 571	576 579610 6	64 666 669 671 673 674 670 672	847 858	860 861 863
С. muridarum тсо	363 366 366 391 393 396 499 5 364 366 392 394 500	03 726 741 816 8 504 828	48 850 852 851 853	859 861 858 860	865 868 900 0 867	35 037 040 042 044 045 041 043	235 248	250 252 251 253
Cp. pneumoniae	0325 0323 0292 0187 0324 0322 0291 0186	1022 1020 1021 1019 1004	28 0826 0824 0825 0823	0817 0815 0818 0816	0811 0808 0761 07 0809	12 0707 0705 0703 0702 0710 0706 0704	0648 0585 0661	0572 0355 0556
Cp. felis	0550 0548 0517 0479 0459 036 0549 0547 0516 0458	59 0277 0275 0276 0274 0271 000 0256 0274	79 0077 0075 0076 0074	0068 0066 0069 0067	0062 0059 0017 05	0970 0968 0966 0965 0973 0969 0967	0914 0851 0926	0837 0820 0574
Cp. caviae ccao	0457 0459 0460 0530 0549 0550	0739 0741 0740 0742 0745 093	35 0937 0939 0938 0940	0946 0948 0945 0947	0952 0955 0996 00 0954	030 0035 0037 0039 0040 0032 0036 0038	0092 0078	0170 0186
Cp. abortus _{CAB}	443 445 476 516 535 444 446 477 556	706 708 707 709 725	004 906 908 907 909	915 917 914 916	921 924 966 0 923	1 033 036 038 040 041 037 039	092 080	167 182
	Cluster 1	Cluster 4		Cluster 2		Cluster 3		
	scc1 sctW sctV sctU	•	sctJ sctL sctR sctS sctT	gspG gspF gspE gspD	scc2 copB1 copD1cadd	sctD sctN ca037 pkn5 sctC sctF chap sctQ		1kb
Chlamydia	incDEFG incA	incBC crpA tarp cap1 mip					cpaf	copD2 scc3 copB2 lcrV
Chlamydophila	incCB inc?A <u>ca530 ca5</u>	lcrV copB2 cpaf 30 scc3 copD2					mip incA2 cap1	tarp <u>ca530</u> crpA

Figure 2: Comparison of the genetic organization of the T3SS genes of six different species of the *Chlamydiaceae*. Below each species, genes are numbered with the locus tag of the GenBank genome files (NC_000117, NC_002620, NC_000922, NC_007899, NC_003361 and NC_004552, respectively). Genetic organization in *Cp. psittaci* is presumed to be highly similar to that of *Cp. abortus* as *Cp. psittaci* strains resemble what might have been recent ancestors of *Cp. abortus* (Van Loock et al., 2003). Positions of corresponding homologs are indicated by color codes. Note that structural proteins [unified nomenclature after Hueck (1998)] appear in the same order in both genera, while (putative) effector proteins do not. The distance within the clusters is in scale, the distance between the clusters is not.

The first structural cluster of T3SS genes (*sctU*, *sctV*, *sctW* and *scc1*) was discovered in *Cp. caviae* (Hsia et al., 1997) (

Figure **2**, cluster 1). The *incDEFG* effector cluster (Scidmore-Carlson et al., 1999) is presumably specific for *Chlamydia* since psi-blast searches (Hsia et al., 1997) did only reveal homologs in *C. muridarum* and not in *Chlamydophilae*. The same is true for the CT228-229 cluster in *C. trachomatis* and its homologs in *C. muridarum*. *IncA* is immediately adjacent to this cluster in *Chlamydia*. The *incBC* cluster which is found 125 kb downstream of *incA* in *C. trachomatis*, is flipped to *incCB* and lies in front of it in *Chlamydophila*. The second structural cluster contains genes coding for SctJLRST as well as Scc2 and YopB/YopD homologs (translocator components CopB/CopD). A third structural

cluster contains *sctD*, *sctN*, *sctQ* and *sctC* as well as the Type III secretion (T3S) effector genes *ca037* (*sctP*) (Subtil et al., 2005), *pkn5* and a gene coding for a putative SctN chaperone (*chap*). The *tarp* gene is found before the second and third cluster in *Chlamydia*, together with effector encoding genes *crpA*, *cap1* and *mip*. In *Chlamydophila*, the same genes are present on the complementary strand downstream of the third cluster. For *scc3*, *copB2*, *copD2* and *lcrV* it is vice versa, as is for CT847 and its homologs.

In *P. amoebophila* UWE25 genes for only one translocon component (CopB, PC1385) and five putative translocon chaperones (PC1384, PC1386, PC0787, PC1525 and PC1110) are present, while no translocon components or associated chaperones are encoded within the *Simkania negevensis* genome (Peters et al., 2007) and

most effector encoding genes identified so far in *Chlamydiaceae* are absent from the environmental chlamydiae.

Genes coding for Type III secretion are usually low in GC content (40-45%) when compared to the rest of the genome (Hueck, 1998; Subtil et al., 2000). In *Chlamydiaceae* however, the GC content of the chlamydial T3S apparatus equals the GC content of the rest of the genome (approximately 40%) and no remnants of recent gene integration seem to flank the different T3SS gene clusters (Subtil et al., 2000).

Both the genetic organization in PAIs and the low GC content suggest that T3S genes originated in a chlamydial ancestor and were subsequently transferred to the Proteobacteria by means of horizontal gene transfer. This is corroborated by the fact that when constructing phylogenetic trees for some T3SS components, the chlamydial homologs are often placed at the bottom of the tree (Kim, 2001; Gophna et al., 2003). Moreover, the presence of T3S genes in environmental chlamydiae suggests that T3S was likely to be present in this lineage more than 700 million years ago, around the time that symbiotic and pathogenic chlamydiae diverged (Horn et al., 2004). The apparent absence of a T3SS in ε -Proteobacteria is probably due to subsequent gene loss.

The presence of four *gsp* genes in cluster 2 is not remarkable since assembly of the T3SS is probably dependent on the Sec-pathway (see above).

Composition of the chlamydial T3SS

In this section we will further focus on the composition and regulation of the chlamydial T3SS. As its closest phylogenetic relative is that of *Y. pestis* (Peters et al., 2007), many assumptions on the chlamydial T3SS are based on information on assembly and regulation of the Ysc-Yop T3SS in *Yersinia*.

Towards a unified nomenclature

In the past years, genes encoding structural components and chaperones of the chlamydial T3SS have been designated Ysc (Yersinia secretion), Lcr (low calcium response) or Syc (specific Yersinia chaperone) after the Yersinia T3SS. With T3SSs found in such a wide range of Proteobacteria, nomenclature of T3SS related genes and proteins became difficult to follow, as each protein is known by many different names according to species, first discovered function, etc. In a 'tour de force' to facilitate interpretation of T3S related research, a unified nomenclature was suggested by Hueck, who used the abbreviation Sct (secretion and cellular translocation), followed by a specific suffix, e.g. SctR, for the conserved structural core of the syringe, giving only chaperones and effectors in its load species specific names (Hueck, 1998). Another nomenclature system, this time specific for chlamydiae was proposed by Fields et al., who chose to use the original cds (contact dependent secretion) designation of Hsia et al. (1997) with the letter suffixes of the unified system (Fields et al., 2003), except for CopN (SctW), where the original designation of Hsia was kept (Hsia et al., 1997).

As chlamydiae have recently been suggested to possess the most ancient non-flagellar T3SS (Pallen et al., 2005; Troisfontaines and Cornelis, 2005), the unified Sct nomenclature as proposed by Hueck for the conserved structural proteins of the T3SS is used throughout this review. Chaperones and effectors were given their species specific names.

Model of the chlamydial T3SS

Depending on the species, 20 to 30 genes have been predicted or proven to encode T3SS structural proteins and chaperones. Homologous to the related flagellar system, components are classified into 1) the exporter, enveloping the basal body, 2) the basal body itself in the inner membrane, 3) the secreton piercing the outer membrane 4) the needle and 5) the translocon, assembling in the eukaryotic membrane at the top of the needle. One of the key players is the cytoplasmic inner membrane associated ATPase SctN. to energize the apparatus. Structural components assemble to form a molecular syringe for direct translocation of effector proteins into the eukaryotic host cell, and although still uncovered, the Sec-pathway is thought to be involved. Additionally, the specific chlamydia chaperones Scc1 and Scc2/3 (homologs of SycE and SycD/LcrH, respectively) assist translocation, while the regulatory protein SctW (or chlamydial outer protein CopN) is predicted to function similar to a cork on a flask. Chlamydiaceae actively modify inclusion membranes with a set of T3S effector proteins, known as inclusion membrane proteins or Incs (Subtil et al., 2001: Fields and Hackstadt, 2002; Rockey et al., 2002).

The degree of homology between T3S proteins from different organisms varies: from highly conserved (especially in the exporter and the basal body) to quite divergent (chaperones, translocon components) and even inapparent or absent (effector proteins). This decrease in homology corresponds to the organization of the T3SS, the molecular equivalent of a syringe, from the inside to the outside, the outside being subjected to a larger evolutionary pressure. At least 11 proteins have homologs in all currently described T3SSs: SctC, D, J, L, N, Q, R, S, T, U, V (Cornelis and Van Gijsegem, 2000). What follows is a general description of how the chlamydial T3SS might be composed (Figure 3), from the inside of the syringe to the top of its needle. The closer we get to the top of the needle, the more difficult it becomes to assign general functions to the components.

Exporter

The exporter consists of several proteins located in the inner membrane at its cytoplasmic face, surrounding the basal body and mediating active transport to the periplasmic space. It is composed of two families of proteins, on the one hand SctR, S and T and on the other hand SctU, V and N, which uses energy from ATP to export proteins across the IM (Aizawa, 2001).

SctR. The SctR protein is homologous to the Yersinia secretion protein R (YscR). In Yersinia this component may have a role in the negative pathway regulation of the calcium controlled Yop expression. SctR is an integral inner membrane protein paralogous to the basal body associated FliP of the flagellar apparatus (Aizawa, 2001). Models show a weakly conserved 70 amino acid hydrophilic domain and four predicted transmembrane helices with periplasmic amino and carboxy termini (Hueck, 1998) or three transmembrane domains with a cytoplasmic amino terminus and a periplasmic carboxy



Figure 3: Type III secretion system in *Chlamydiaceae*. On the left side, an inactive T3SS is shown, while the right side depicts the T3SS after activation. Prior to contact with the host cell, only LrcV is supposed to be present at the tip of the needle and secretion is regulated by SctW. Translocon component CopD is bound to its chaperone, as is the case for effector proteins. Upon host cell recognition, Cops are built into the eukaryotic membrane on top of an LcrV scaffold, thus allowing translocation of unfolded, unchaperoned effectors to the host cell cytoplasm. Legend: IM, bacterial inner membrane; OM, bacterial outer membrane. Refer to the text for details.

terminus (Ghosh, 2004). In *Cp. psittaci*, SctR is predicted to have five transmembrane segments (Beeckman et al., 2008).

SctS. SctS has two predicted transmembrane domains (Hueck, 1998) with probably cytoplasmic termini. It shows homology to FliQ of the flagellar system (Aizawa, 2001).

SctT. Similar to SctS and SctR and paralogous to flagellar FliR (Aizawa, 2001), SctT is an integral membrane protein, this time with six predicted transmembrane domains and the amino and carboxy termini presumably in the periplasm (Hueck, 1998).

SctU. SctU is homologous to YscU, an inner membrane protein with four transmembrane segments anchoring a large cytoplasmic carboxy terminal domain to the inner membrane (Allaoui et al., 1994). Its paralog in flagella is FlhB. Cleavage of *Yersinia* YscU is needed to acquire a conformation allowing recognition and subsequent membrane insertion of the translocon components (Sorg et al., 2007). In *Chlamydiaceae*, SctU was first discovered in *Cp. caviae* and was originally called Cds1

(Hsia et al., 1997). In *C. trachomatis*, RT-PCR analysis showed *sctU* expression from 12 h onwards (Fields et al., 2003) while *sctU* expression was not detected in *Cp. pneumoniae* (Slepenkin et al., 2003), suggesting variable importance of SctU in chlamydiae.

SctV. SctV is predicted to form a central proteinconducting channel across the inner membrane, based on its homology to Yersinia LcrD/YscV. These proteins are predicted to have at least six amino terminal transmembrane segments that anchor a large cytoplasmic carboxy terminal domain into the inner membrane (Plano et al., 1991). In *Chlamydiaceae*, SctV was originally detected in *Cp. caviae* and called Cds2 (Hsia et al., 1997).

SctN. This protein belongs to the family of cytoplasmic ATPases and demonstrates extensive homology with the ATP synthase F1 beta subunit (Woestyn et al., 1994). Following oligomerization into a dodecamer (Pozidis et al., 2003; Muller et al., 2006), it uses ATP to energize the T3SS or at least a part of it. Its subcellular location is cytoplasmic, but it is connected to the inner membrane,

possibly through SctV, although binding is probably not required for proper function. Three highly conserved regions are found: ATP/GTP binding domains A and B also called walker boxes A and B (Walker et al., 1982) and a Mg²⁺ binding motif (Yoshida et al., 1982). *Cp. pneumoniae* SctN interacts with the basal body components, SctD, Q and L as well as the regulator protein SctW (Stone et al., 2008) and is also believed to play a role in the delivery of effector proteins through ATP-dependent unfolding of proteins and releasing the chaperone from its substrate prior to translocation (Akeda and Galan, 2005; Sorg and Cornelis, 2009).

Basal body

The basal body of the T3SS is located in the bacterial cytoplasm and is necessary for correct anchoring of the needle complex. The *sctQ*, *R*, *S*, *T*, *U*, *V* and *J* genes all show significant homology to the genes of the flagellar system, indicating that their corresponding proteins are localized in or at least associated with the inner membrane. Both *sctD* and *sctL* show weaker homologies to their flagellar counterparts.

SctD. The 3' region of the *C. trachomatis CT664* gene has homology to *yscD* but is not consistently reported as a possible component of the chlamydial T3SS (Kim, 2001). SctD resembles the *Salmonella* PrgH required for anchoring of the *Salmonella* needle complex (Kubori et al., 1998). Two forkhead-associated (FHA) putative nuclear signaling domains of approximately 55 to 75 amino acids are present in the amino terminal region, and pull down assays suggest that SctD interacts with *Cp. pneumoniae* SctL and SctQ, probably through its carboxy terminus (Johnson et al., 2008).

SctJ. Most members of the SctJ family are lipoproteins, as they carry a characteristic lipid attachment site motif LxGC at the amino terminus as well as a hydrophobic stretch of 20 to 30 amino acids at the carboxy terminal end. They represent the morphogenic equivalent of the flagellar rotor FliF (MS ring) and probably bridge the inner and outer membranes (Allaoui et al., 1992; Hueck, 1998).

SctQ. Proteins belonging to the SctQ family are peripheral cytoplasmic proteins with a strongly conserved carboxy terminus and a variable amino terminus. Homology to flagellar proteins FliM and FliN has been reported (Aizawa, 2001; Ghosh, 2004) and functionally they are thought to link the conserved structural components to variable species-specific components as recombinant *Yersinia* YscQ provides a link between YscK and YscL (Jackson and Plano, 2000). In *Cp. pneumoniae*, cytoplasmic SctQ may play a role in shuttling chaperone/effector complexes to the ATPase SctN (Stone et al., 2008), apart from its interaction with SctW (R. Toor et al., unpublished data). Furthermore, SctQ contains a "surface presentation of antigens" (SpoA) domain and is thought to be incorporated in the chlamydial T3SS as tetramer (Johnson et al., 2008).

SctL. Members of the YscL family are mainly hydrophilic. Although little is known about their function, they probably act in a similar way to SctQ, as yeast three-hybrid studies showed that recombinant YscL linked YscN and YscQ, suggesting that a complex composed of YscN, YscQ, YscK and YscL is involved in the assembly and/or function of the *Y. pestis* Type III secretion apparatus (Jackson and Plano, 2000). In

Cp. pneumoniae, SctL is shown to form dimers (Stone et al., 2008).

Secreton

The secreton is defined as the collection of outer membrane proteins involved in the transport of effectors and translocon components across the outer membrane. SctC. SctC is clearly an outer membrane protein (Plano and Straley, 1995), forming a channel absolutely necessary for T3S. SctC has no paralogs in the flagellar system but belongs to the large GspD family of bacterial and phage proteins involved in transport of large molecules across the outer membrane (see above). Secretins form stable multimeric ring-like structures (Genin and Boucher, 1994) of around 600 kDa with an external and internal diameter of 200 and 50 nm. respectively. Interestingly, their ultrastructure shows similarity to the rosette-like structures described by Matsumoto (Matsumoto, 1973; 1982b; Matsumoto et al., 1976).

SctW. This gene was first discovered in Y. enterolitica and originally named LcrE, Ysc4b or YopN (Viitanen et al., 1990). The chlamydial homolog SctW (CopN) was discovered in Cp. caviae (Hsia et al., 1997) where it might control the polarized contact-dependent secretion of other Cops, and thus seems be involved in a mechanism for regulating the activity of the T3SS (Cheng et al., 2001), Fields and Hackstadt (2000) showed C. trachomatis SctW to be associated with the inclusion membrane and to be T3SS secreted by a heterologous Yersinia system. Homologs of this protein return in all T3SSs, are mainly surface associated (Forsberg et al., 1991) and at some stages associated with the basal body or the secreton. Regarding its function, there is controversy whether SctW should be regarded as an outer membrane protein of the secreton, or rather be classified as an effector. Expression of SctW might provoke detachment of C. trachomatis RBs from the inclusion membrane, shutting off the T3SS as a consequence (see further).

Needle

The needle is the hollow conduit through which the T3SS proteins travel to cross the membranes and the peptidoglycan layer, which is not present in chlamydiae (Hatch, 1996; Moulder, 1993).

SctF. In *Yersinia*, the needle consists of approximately 150 YscF subunits, on top of which up to five LcrV molecules are attached (Broz et al., 2007). Recently, the *C. trachomatis* needle subunit SctF was identified as CT666 using a bioinformatics approach. The protein is concentrated in the outer membrane of EBs and is surface exposed as a component of an extracellular needle-like projection. During infection, SctF is localized in the inclusion membrane adjacent to membrane-associated RBs (Betts et al., 2008).

SctP. A gene encoding an YscP homolog, the ruler protein acting as a molecular ruler controlling the length of the needle (Journet et al., 2003), has not yet been identified in chlamydiae. However, based on sequence homology and its location in the genome, Cpn0705 of *Cp. pneumoniae* could be a ruler protein, but no further studies have been completed with this protein (Stone et al., 2008). Its homolog in *Cp. caviae* CCA00037 was previously shown to be secreted by *Shigella flexneri*, as

well as the homologous protein from *C. trachomatis*, CT671 (Subtil et al., 2005).

Translocon

Upon contact with a target cell, a hetero-oligomeric translocation pore is built into the eukaryotic host cell membrane, consisting of T3SS released proteins assisting translocation into the host cell. In Yersinia, the translocon pore consists of YopB and YopD, on top of an assembly platform of three to five LcrV molecules. Among species, translocator sequences are not conserved, but there are structural similarities in transmembrane segments and coiled-coil regions [reviewed in Mueller et al. (2008)]. In C. trachomatis and Cp. pneumoniae, proteins without signal sequence and with size and hydrophobicity profiles reminiscent to the ones of YopB and YopD were identified as the chlamvdial translocators (Subtil et al.. 2000). Remarkably, in each species two homologs were found: CT579/Cpn0808 or copD and CT860/Cpn1019 or copD2 as homolog for yopD and CT578/Cpn0809 or copB and CT861/Cpn1020 or copB2 in close proximity to the sycD chaperone homologs CT576/Cpn0811 (scc2 or lcrH1) and CT862/Cpn1021 (scc3 or IcrH2), respectively.

CopB. CopB1 was the first chlamydial homolog of YopB and it functions as a putative translocator component of the chlamydial T3SS, consistent with its presence in the *C. trachomatis* inclusion membrane. On the contrary, CopB2 was demonstrated in the host cell cytosol (Fields et al., 2005).

CopD. CopD1 and CopD2 are the chlamydial YopD homologs, with CopD1 being translocated through the heterologous SPI-2 T3SS of *S. enterica* sv. *typhimurium* (Ho and Starnbach, 2005).

LcrV. The needle culminates in a tip complex consisting of multiple copies of a protein of the LcrV family, also known as adaptor proteins (Blocker et al., 2008). In addition to its essential role in the insertion of the translocators into the host cell membrane, the *Yersinia* LcrV pentamer on top of the T3SS needle is involved in host cell sensing and causes immunosuppression by triggering IL-10 release by macrophages [reviewed in Hamad and Nilles (2007)]. Until now, no LcrV homolog had been identified in chlamydiae (Stone et al., 2008). However, all currently sequenced *Chlamydiaceae* contain at least one LcrV encoding gene in their genome, adjacent to the *scc3/copB2/D2* gene cluster (cluster 4 in

Figure **2**, author contribution). Whether the corresponding protein is functional is elusive and requires further attention.

Chaperones

Type III secretion chaperones are specific, associating with one or multiple secretion substrates, typically encoded on an adjacent gene. Their function includes stabilizing their target, maintaining a secretion competent state or serving as a secretion pilot. In *Yersinia*, they are also involved in establishing a secretion hierarchy and participate in feedback regulatory cascades [overview in Fields and Hackstadt (2006)]. The chlamydial genomes contain at least nine predicted T3S chaperones, at least seven of which contain predicted amphipathic helices. These will now be discussed in further detail. **Scc1**. Scc1 is the <u>Specific chlamydia chaperone 1</u> discovered by Hsia et al. (1997), CT088 in *C. trachomatis*. Scc1 shares limited sequence similarity to SycE of *Yersinia spp.*, a rather small protein of about 15 kDa carrying a carboxy terminal amphipathic alpha helix. SycE family members are acidic (pl 4,4 – 5,2) and bind their secreted or translocated partner, which is YopE in the case of SycE (Cornelis, 2000). Moreover Scc1 contains a CesT domain, homologous to the CesT chaperone of the *E. coli* translocated Tir T3SE protein (Delahay et al., 2002).

CT043 and CT663. As for Scc1, both predicted chaperones contain a CesT domain, but their respective chlamydial substrates have not yet been uncovered.

Scc2/3. Scc2 and Scc3 are homologous to the LrcH chaperone of the Yersinia translocators, chaperoning CopB1/D1 and CopB2/D2, respectively, in chlamydial T3SSs. As with the other members of the LcrH/SycD family, both Sccs possess three tandem TPRs implicated in protein-protein interactions (Pallen et al., 2003b). In addition, both chaperones associate with YopD in a heterologous Yersinia T3SS and coprecipitate with CopB from *C. trachomatis* extracts (Fields et al., 2005). Surprisingly, *Cp. pneumoniae* Scc3, adjacent to CopB2 and CopD2, was found to interact with SctW in a two-hybrid system, suggesting to function as a chaperone for SctW as well (Slepenkin et al., 2005).

CT274. With its typical TPR domain, CT274 and its chlamydial homologs are classified in the group of Class II chaperones, and might therefore be involved in transcriptional regulation of T3S gene expression (see above).

CT665 and CT667. These two recently discovered chaperones are involved in regulation of SctF in *C. trachomatis*. Originally named CdsE (SctE) and CdsG (SctG), both proteins interact with each other and a SctF-CdsG interaction was only detectable in the presence of CdsE (Betts et al., 2008).

CT670/SctO. Proteins of this family have various sizes and exhibit little sequence similarity. In *Yersinia*, YscO is a mobile protein required for high-level expression and secretion of LcrV and Yops (Payne and Straley, 1998). Its counterpart in the flagellar system is FliJ, a cytoplasmic protein occupying a chaperone function (Payne and Straley, 1998; Minamino et al., 2000). This is in accordance with the presumed chaperone function of the homologous *Cp. pneumoniae* Cpn0706 protein (Stone et al., 2008), the encoding gene of which is localized at the predicted location of the *sctO* gene, in addition to its predicted coiled-coil conformation (unpublished result).

Effector proteins – chlamydial T3SS substrates

Type III secretion effector proteins are highly divergent, both in sequence and function between T3SS possessing organisms. For instance, *Yersinia* Yops mainly target the intracellular host cell signaling pathway to benefit their survival outside the cell and escape the innate immunity. On the other hand, *Salmonella* Sips (SPI-1 T3SEs) induce cell invasion, while Sses from the SPI-2 T3SS aid in proliferation inside macrophages (Hueck, 1998). Unlike in other pathogens with a T3SS, chlamydial effectors are scattered throughout the genome and in the past years, several efforts have been made to identify chlamydial T3SEs (Subtil et al., 2001; Fields et al., 2003; Slepenkin et al., 2003; Lugert et al., 2004; Clifton et al., 2005; Subtil et al., 2005; Jamison and Hackstadt, 2008; Kleba and Stephens, 2008), with the family of inclusion membrane proteins (Incs) being the best one characterized so far. Although the Incs exhibit a very low sequence similarity, a characteristic bilobed hydrophobic region of at least 50 amino acids is present in each of them. Biological functions for the Incs have been proposed in the past years. These will briefly be discussed hereafter, but for a more detailed description we refer to an excellent review by Kostryukova et al. (2008). IncA is a serine/threonine phosphoprotein, exposed on the cytoplasmic side of the developing inclusion, where it is phosphorylated by host cell kinases, both during chlamydial infection and endogenous expression of IncA in eukaryotic cells (Rockey et al., 1997). This protein has been extensively studied and is believed to be functionally involved in the homotypic fusion of C. trachomatis inclusions (Hackstadt et al., 1999). Decreasing the temperature to 32 °C is correlated with an impaired export of IncA to the inclusion membrane and delays the formation of a single inclusion (Fields et al., 2002). The biochemical properties of C. trachomatis and Cp. caviae IncA are conserved as both orthologs carry a eukaryotic SNARE domain (responsible for fusion of membrane structures in eukarvotes) and self-associate to form a tetrameric structure similar to the SNARE complex, possibly interacting with eukarvotic SNARES to hijack part of the cellular traffic (Delevoye et al., 2004). The adjacent Cpn0187 is also predicted to be an Inc as well as a methyltransferase, and is only present in Chlamydophila spp. (Kalman et al., 1999). IncD to G are expressed as an operon in the early stages of the developmental cycle, suggesting an important role in remodeling the nascent chlamydial inclusion. In immunofluorescence studies, patterns for these Incs resemble IncB staining: distinct patches of the inclusion membrane are stained, suggesting regions of close contact between the RB and the inclusion membrane (Scidmore-Carlson et al., 1999). C. trachomatis IncG is normally exposed on the cytoplasmic surface of the inclusion (Hackstadt et al., 1999) and was used as a bait to determine host proteins that interact with the inclusion. The eukaryotic interaction partner was proven to be 14-3-3β, a phosphoserinebinding protein involved in several signaling pathways as a scaffold for phosphotransfer. This specific interaction could be confirmed by the demonstration of two specific 14-3-3β binding sites at the carboxy terminal region of IncG, and colocalization studies in cell cultures infected with C. trachomatis serovars A, B, Ba, C, D, E, F, G, H, I, J, K and L1-3. In Cp. pneumoniae, Cp. caviae and C. muridarum infected cells no such interaction could be observed, which is not surprising given the absence of IncG in the first two species and the large sequence difference with C. muridarum IncG (Scidmore and Hackstadt, 2001). The inverse approach was followed to identifv possible interactions with eukarvotic RabGTPases localized in the inclusion membrane region. In a yeast-2-hybrid screening with Rab4 GTPase as a bait, only CT229 interacted. Furthermore, Rab4 was found to colocalize with CT229 on the inclusion membrane surface, an interaction requiring the presence of a GTP-binding domain (Rzomp et al., 2006). Similarly, Cpn0585 interacts with Rab1, Rab10 and Rab11 (Cortes

et al., 2007). CrpA, also known as CT442 in C. trachomatis, is one of the Incs predicted by the study of Bannantine et al. (2000). This protein was erroneously regarded as a cysteine rich protein in the chlamydial outer membrane (Clarke et al., 1988; Lambden et al., 1990) but contains only four Cys residues and has been shown to localize to the inclusion membrane by immunochemistry (Bannantine et al., 2000). CrpA is recognized by CD8⁺ T-cells, and may have access, together with other Incs, to the class I MHC-processing pathways of infected cells (Starnbach et al., 2003). Some proteins localized in the inclusion membrane lack the characteristic bilobed hydrophobic domain. Two of them, SctW and Cap1, are proven T3S effector proteins (Fields and Hackstadt, 2000; Subtil et al., 2005). SctW probably acts as a gating protein, preventing or allowing T3S proteins to be translocated across the bacterial membranes, while **Cap1** (Class I Accessible Protein-1, CT529) is another CD8⁺ T-cell antigen (Fling et al., 2001). This protein is expressed during the early stages of the C. trachomatis developmental cycle and by 8 h after infection, sufficient Cap1 protein is expressed to stimulate CD8⁺ T cells. Cap1 may therefore play an important role in early immune recognition of C. trachomatis- infected cells (Balsara et al., 2006). CT147 is transcribed early during the development of C. trachomatis and shows significant homology to EEA1 (mammalian Early Endosomal Antigen 1), involved in trafficking and fusion of mammalian endosomes. Although CT147 lacks homology with the region of EEA1 that has been shown to bind Rab5, it has maintained the carboxy terminal Zn-finger, suggesting CT147 could bind to endosomes but circumvent endosomal fusion, a novel strategy for chlamydial endosomes to avoid fusion with

lysosomes (Belland et al., 2003). The above mentioned results suggest that chlamydial Incs are capable of interfering with vesicle trafficking, guiding inclusion development, avoiding phagolysosomal fusion, acquiring nutrients and diverting important host signaling molecules from their designated pathways. However, there are distinct differences in how chlamydial species interact with intracellular membranes, so caution should be exercised when extrapolating findings to other chlamydial species.

Other secreted T3S effector proteins are summarized in Table 3. These include the Translocated actin recruiting protein Tarp, mediating host cell internalization through the recruitment of actin filaments to the site of chlamydial attachment (see further). CADD (Chlamydia protein associating with death domains) has been shown to be involved in modulation of host cell apoptosis via binding to the death domains of tumor necrosis factor family receptors. This novel redox protein toxin has limited sequence identity with coenzyme PQQ (pyrrolo-quinoline quinone) synthesis protein C (PqqC) family members, which are part of the six-step PQQ synthesis pathway in bacteria (Stenner-Liewen et al., 2002; Schwarzenbacher et al., 2004). As already mentioned above, SctP might be a ruler protein involved in determination of the needle length of the T3SS (Stone et al., 2008). Pkn5 was originally annotated as a putative serine/threonine kinase and postulated to be T3SS related when the genome of C. trachomatis was sequenced (Stephens et al., 1998).

Effector	Location	Gene number	Possible function
Tarp	Surface of EB, host cell cytosol	CT456/Cpn0572	Invasion, actin recruitment
CADD	Proximity of the inclusion	CT610/Cpn0761	Modulation of host cell apoptosis
SctP	Bacteria and host cell cytosol	CT671/Cpn0705	Putative T3SS needle ruler protein
Pkn5	Inclusion and host cell cytosol	CT673/Cpn0703	S/T kinase activity
Mip	Surface of EB, RB and IncM	CT541/Cpn0661	Induction of cytokine response
Hypothetical	Unknown	CT668/Cpn0708	Unknown
Hypothetical	Bacteria and host cell cytosol	CT694/ absent	Interaction with human AHNAK
Hypothetical	Unknown	CT847/Cpn1004	Interacts with human GCIP

Table 3: Overview of chlamydial T3S effector proteins. The corresponding gene number in *C. trachomatis* or *Cp. pneumoniae* is given. Legend: IncM, inclusion membrane

Since then, C. trachomatis Pkn5 was proven to be secreted by the heterologous Salmonella SPI-2 T3SS (Ho and Starnbach, 2005) as well as to be secreted into the inclusion membrane of Cp. pneumoniae by Herrmann et al. (2006). The lipoprotein Mip is homologous to the macrophage infectivity potentiator (Mip) gene of L. pneumophila (Lundemose et al., 1993a; Localization studies first 1993b). reported Cp. pneumoniae and Cp. caviae Mip to be surface exposed on EBs and RBs (Rockey et al., 1996; Montigiani et al., 2002). Later on, Herrmann et al. (2006) demonstrated that Cp. pneumoniae secretes Mip into the host cell cytoplasm. They concluded that the substrate proteins of Mip might be other chlamvdial proteins/virulence factors in the inclusion membrane or host cell proteins whose conformation might be altered by Mip peptidyl prolyl cis-trans isomerase activity. Genes encoding CT668 and its homologs in other chlamydial species are located in the third T3S gene cluster (

Figure 2), and at least for *C. trachomatis*, this protein has been shown to be Type III secreted (Sisko et al., 2006). No function or cellular localization for this protein has so far been identified. In contrast, CT694 is localized separate from bacteria only in the early phase of infection, and interacts with the repeat region of human AHNAK, presumably leading to Raf/MEK/ERK activation and modulation of the actin cytoskeleton. T3SS dependency was shown by heterologous translocation through the Y. pseudotuberculosis T3SS. As CT694 is only found in C. trachomatis, this protein could contribute in aspects unique compared to the other chlamydial species (Hower et al., 2009). CT874, secreted through the heterologous T3SS of Y. pseudotuberculosis, interacts in the host cell cytoplasm with a eukaryotic protein termed GCIP (Grap2 cyclin D-interacting protein) a multi-functional ubiquitously expressed protein that is degraded during chlamydial infection (Chellas-Gery et al., 2007).

Role of T3S in the chlamydial lifecycle

The T3SS plays an essential role throughout the chlamydial developmental cycle. In the following paragraphs, a short description is given of how chlamydiae employ their T3SS to modify the intravacuolar environment to their benefit at different stages in the life cycle.

Differential gene expression – resolving the 'chlamydial Type III paradox'

Upon irreversible attachment to the host cell, EBs directly translocate pre-existing effectors (e.g. Tarp) to the host cell cytosol (Clifton et al., 2005; Jewett et al.,

2006). In addition, expression and secretion of *Cp. pneumoniae* IncA, IncB and IncC (Lugert et al., 2004) and *C. trachomatis* IncC (Fields et al., 2003) occurs already within 2 h (early cycle) post infection. However, expression of T3S-specific Sct-genes is not detectable until 8 to 12 h post infection, implying that *de novo* synthesis of T3SSs only appears from mid-cycle onwards (Fields et al., 2003; Lugert et al., 2004).

These observations immediately raised the question how chlamydiae can secrete and translocate their effector molecules early after infection, even before assembly of a functional T3SS. A possible solution for this 'chlamydial Type III paradox' came from studies by Vandahl et al. (2001), Fields et al. (2003) and Lugert et al. (2004), demonstrating the existence of the Type III secretion apparatus on both C. trachomatis and Cp. pneumoniae EBs. Proteome analysis of Cp. pneumoniae EBs by Vandahl et al. (2001) revealed that all components of the T3SS are present in EBs. Presence of CopD1 was highly abundant, its chaperone Scc2 being expressed late in the developmental cycle, as is the case for the majority of T3SS structural genes (Slepenkin et al., 2003). The presence of high amounts of CopD1 in purified EBs implied that it had been retained in the bacteria during the late phase of the developmental cycle. They suggested that CopD1 is preloaded in EBs, ready to be secreted by T3S upon contact with a host cell. This hypothesis could be further strengthened by the finding of the other translocon protein CopB1 in the cytoplasm of Cp. pneumoniae infected cells (Lugert et al., 2004). Additional and striking evidence was provided by Fields et al. (2003), demonstrating the existence of a T3S apparatus on EBs by matrix-assisted laser desorption ionization time-offlight (MALDI-TOF) and immunoblotting of purified EB extracts using polyclonal antibodies specific for the studied proteins. Based on their findings Fields et al. (2003) proposed a model in which T3SSs are present on EBs and mediate secretion of effectors such as Tarp and early Incs. Mid-cycle expression of T3S genes would then replenish the secretion apparatuses on RBs and serve as a source of secretion pores for subsequently formed EBs. This was also confirmed by Maurer et al. showing that genes coding for EB proteins were primarily connected with products in late, but not tardy (showing steadily increasing expression at the end of the cycle) expressed gene clusters (Maurer et al., 2007).

As mentioned above, *Chlamydiaceae* possess two sets of translocators, each with their specific chaperone, needed during different periods of infection and therefore expressed at different time points during the developmental cycle. In *C. trachomatis*, RT-PCR analysis showed *copB1*, *copD1* and *scc2* expression from 20 h post infection onwards (Fields et al., 2003), while expression from 24 h post infection onwards could be observed in *Cp. pneumoniae* for *scc2* (Slepenkin et al., 2003; Lugert et al., 2004). The second chlamydial translocon cluster (*scc3/copB2/copD2*) was expressed earlier: from 12 h post infection in *C. trachomatis*, and from the earliest studied time point on (1.5 - 2 h) in *Cp. pneumoniae* (Slepenkin et al., 2003; Lugert et al., 2004). This is in accordance with CopB2 and CopD2 built into the inclusion membrane to allow cellular reprogramming and CopB1 and CopD1 mediating translocation of proteins during the internalization phase (Ouellette et al., 2005; Hefty and Stephens, 2007).

Adhesion and entry

Adhesion of chlamydial EBs to the host cell is mediated by a number of possible adhesins such as MOMP (Su et al., 1990), Hsp70 (Raulston et al., 1993), OmcB (Ting et al., 1995; Moelleken and Hegemann, 2008) and pmpD (Wehrl et al., 2004; Crane et al., 2006) Analogous to what is described for *Shigella flexneri*, T3SS translocon components could be involved as well in the irreversible attachment of EBs, assuming that CopB1 binds the hyaluronan receptor CD44 in lipid rafts, thereby inducing complete assembly of the T3SS translocon in the eukaryotic membrane, followed by an interaction of the CopB1-D1-LcrV complex with a nearby $\alpha_5\beta_1$ integrin receptor on the host cell (Watarai et al., 1996; Skoudy et al., 2000).

Translocation of Tarp into the host cell within minutes following attachment induces actin recruitment and the formation of pedestal-like structures beneath the attached EB. At least two accompanying signaling cascades (Arp2/3 dependent or not) have already been described for *C. trachomatis* (Jewett et al., 2006; Lane et al., 2008), apart from other signaling pathways mediating EB internalization, most likely initiated after translocation of additional currently uncharacterized T3SEs.

In addition to microfilament dependent entry through the lipid raft signaling platforms, *Chlamydiaceae* also employ receptor mediated uptake by clathrin-coated pits. Possible ligands could include surface exposed T3SS components, e.g. the outer membrane secretin SctC or the needle protein SctF.

Inhibition of phagolysosomal fusion

During infection, *Chlamydiaceae* resist the host cell nonspecific defence system of phagolysosomal fusion. Inhibition of acidification of the inclusion is most likely accompanied by an active modification of the inclusion membrane (Scidmore et al., 2003). Possible candidates are chlamydial Incs involved in preventing fusion between the chlamydial inclusion and lysosomes (Hackstadt et al., 1997; Wyrick, 2000). In *Salmonella*, binding of SPI-2 T3SE SpiC with mammalian Hook3 disrupts Golgi morphology and alters lysosomal distribution, thus probably preventing phagosomelysosome fusion (Shotland et al., 2003; Bakowski et al., 2008). Likewise, chlamydiae might translocate unknown T3SEs to the host cell cytoplasm as well.

Role of Incs in the subversion of intracellular host cell traffic

Whilst avoiding fusion with lysosomes, the inclusions migrate to the microtubuli organizing centre - Golgi

region to fuse with sphingomyelin-containing exocytic vesicles of the host cell. In addition, glycerophospholipids and other lipids can be acquired through fusion with multivesicular bodies (Beatty, 2006) and lipid droplets (Cocchiaro et al., 2008). The *Chlamydiaceae* thus manage to intercept nutrients and constituent molecules from normal eukaryotic trafficking pathways, mimicking the phospholipid composition of the host cell in which the bacteria are grown (Hatch and McClarty, 1998).

Elucidation on the mechanisms that divert lipids to the vacuole came from the fact that the chlamydial inclusion interacts with RabGTPases, key regulators of host vesicle fusion. For example, the C. trachomatis specific CT229 Inc probably recruits Rab4 to the inclusion membrane surface, in an attempt to regulate the intracellular trafficking or fusogenicity with the chlamydial inclusion (Rzomp et al., 2006). Similarly, Cp. pneumoniae Inc Cpn0585 interacts with Rab1, Rab10 and Rab11 and may mediate their recruitment to the Cp. pneumoniae inclusion. The subsequent interactions are dependent on Rabs bound to GTP, suggesting that the inclusion membrane localized Rab proteins are functional (Rzomp et al., 2006; Cortes et al., 2007).

Another important player in the interception of vesicular trafficking might be IncA. In addition to its involvement in the homotypic fusion of *C. trachomatis* inclusions (Hackstadt et al., 1999), IncA eukaryotic SNARE-like motifs may represent a mimicry mechanism to interact with a subset of host SNAREs, leading to the selective recruitment of intracellular compartments around the inclusion (Delevoye et al., 2008).

Proliferation and engaging into late differentiation

As it was often observed that RB replication occurs while Chlamydiaceae are in close contact with the inclusion membrane, it was suggested that this juxtaposition could be needed for growth. This contact is thought to be mediated by T3SSs, most likely corresponding to the surface projections previously described by Matsumoto (1982a). Detachment of RBs from the inclusion membrane and coupled T3SS inactivation would constitute the signal for redifferentiation into EBs, the socalled 'contact-dependent development hypothesis' (Hackstadt et al., 1997; Bavoil and Hsia, 1998; Bavoil et al., 2000). Since then mathematical models linking the number of T3S projections and consecutive events during intracellular development in normal and persistent infections have been described (Wilson et al., 2006; Hoare et al., 2008). A schematic diagram is given in Figure 4.

Briefly, upon attachment of EBs, preloaded T3SSs translocate effector proteins into the host cell to promote internalization, survival and differentiation into RBs. Contact between the RBs and the inclusion membrane upregulates T3S gene-expression, leading to a rise in the number of projections per bacterium, and triggers binary fission of RBs. As the RBs replicate, the inclusion expands, reducing the contact area between RB and the inclusion membrane, and thus T3S activity. When the number of T3SSs in contact with the inclusion membrane drops below a certain threshold, physical detachment coincides with the upregulation of *sctW* expression, only expressed late in normal RBs.



Figure 4: Schematic overview of the contact-dependent chlamydial development hypothesis. Preloaded T3SSs of EBs are implicated in internalization and early survival, while neosynthesized T3SSs promote replication. Adapted from Wilson et al. (2006). IB: intermediate body. Refer to the text for details.

The question then becomes: 1) does overcrowding of the inclusion physically 'squeeze out' RBs to the lumen of the inclusion, thereby inducing SctW expression and downregulation of T3S or rather 2) does the low number of T3SSs per RB shut down T3S through SctW expression, thereby provoking physical detachment? In any case, progressive detachment of bacteria from the inclusion membranes coincides with the observed asynchronicity in the redifferentiation process from RBs to EBs (Wilson et al., 2006; Peters et al., 2007; Hoare et al., 2008).

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