Identification and biochemical characterization of the CHLAMYDIA TRACHOMATIS type III secretion chaperone, SLC1, and its role in the translocation of the invasion-associated effector TARP.

Amanda J. Brinkworth

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IDENTIFICATION AND BIOCHEMICAL CHARACTERIZATION OF THE
CHLAMYDIA TRACHOMATIS TYPE III SECRETION CHAPERONE, SLC1,
AND ITS ROLE IN THE TRANSLOCATION OF THE INVASION-ASSOCIATED EFFECTOR TARP

By
Amanda J. Brinkworth

A Dissertation
submitted to the Faculty of the
Graduate School of the University of Louisville
in Partial Fulfillment of the Requirements
for the Degree of

Doctor of Philosophy
in Microbiology

Department of Microbiology and Immunology
University of Louisville
Louisville, Kentucky, USA

May 2011
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TARP

By

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A Dissertation Approved on

January 27, 2010

by the following Dissertation Committee:

________________________________________
Michah Worley

________________________________________
Yousef Abu-Kwaik

________________________________________
Richard D. Miller
DEDICATION

I would like to dedicate this dissertation to

my loving parents

Pam and Greg

and

my amazing husband

Jeff,

Without your love and support I never would have been able to accomplish my goals.
ACKNOWLEDGEMENTS

I would like to express my most sincere gratitude to my mentor Dr. Rey Carabeo, who has consistently gone above and beyond my expectations to help me in both science and life. I would also like to acknowledge my co-mentor Dr. James Graham, whose guidance and generosity have been pivotal to my work. My committee members Dr. Richard Miller, Dr. Yousef Abu-Kwaik, and Dr. Michah Worley all provided essential discussions and suggestions throughout my thesis, and I want to specifically thank you for all of your time and efforts. I want to gratefully acknowledge our collaborator, Dr. Richard Hayward at University College London for his invaluable advice and efforts toward all of the gel filtration chromatography work in this dissertation.

Furthermore, I want to specifically thank the Imperial College London masters and undergraduate students Antonio Tedim Pedrosa, Sevanna Shahbazian, and Katarzyna Roguska, who worked with me to complete the bacterial 2-hybrid and translocation studies. Their hard-work and enthusiasm was greatly appreciated. I would like to thank all of the former and current lab members: Jennifer, Illya, Chris, Scot, Tristan, Sophie, Denise, and David for their kindness, patience, and supportiveness throughout my thesis.

I specifically want to thank Sophie Nicod for being such a lovely person who always was there for me whenever lab or life became chaotic. Gunnar Schroeder, Alex Wong, Diana Munera, and Robert Fagan from the Centre for Molecular Microbiology and Infection at Imperial College London and Chris Price at University of Louisville were all excellent resources and were very helpful during my thesis. I especially want to acknowledge my best friend, Meena Vanchinathan, whose sense of humor and companionship has been essential to my graduate career. Finally I would like to acknowledge my wonderful and loving husband Jeff, who can never be thanked enough for his endless patience and support.
ABSTRACT
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ASSOCIATED EFFECTOR TARP
Amanda Brinkworth
May 2011

Chlamydia trachomatis is an obligate intracellular pathogen that utilizes a type III secretion system to enter mammalian cells and establish an intracellular niche. TARP, the translocated actin recruitment protein, is a chlamydial invasion protein known to be type III secreted by the metabolically inert elementary body upon docking to the mammalian cell surface. Because immediate secretion of TARP into host cells is necessary for entry, I hypothesized that a chlamydial chaperone binds to TARP and facilitates its translocation through the type III secretion apparatus. Most effector-binding type III secretion chaperones are small (14-18 kDa), have an acidic pI, and share a specific secondary structure of alternating alpha-helices (α) and beta-sheets (β). Typically, type III secretion chaperones dimerize and interact with their effectors as a complex of two molecules of chaperone to one effector molecule. Only 3 Chlamydia trachomatis proteins have been identified in EB’s that are predicted to be putative chlamydial type III effector chaperones. These are CT043, CT663, and CT088, which I have designated as Slc1, Slc2, and Scc1, respectively. These chaperones were tested for their interaction with the N-terminal 200 amino acids of TARP (HIS6-TARP1-200) by co-
immunoprecipitation. HIS$_6$-TARP$^{1-200}$ interacted specifically with Slc1, but not Scc1 or Slc2. This interaction was enhanced by coexpression of the recombinant proteins. To confirm this interaction and rule out the possibility of Slc1 heterodimerization enhancing the interaction with TARP, I employed a 2-hybrid system to test for TARP: chaperone and chaperone:chaperone interactions. I confirmed the specific interaction between Cya18-TARP$^{1-200}$ and Cya23-Slc1. I was also able to detect Slc1 interaction with itself as well as confirm a few other previously described chaperone-chaperone interactions.

Analysis by crosslinking and gel filtration chromatography indicated that Slc1 forms a stable dimer in solution. Complexes of the Slc1 chaperone dimer with TARP in a 2:1 stoichiometry were detected following purification from co-expressing bacteria, but not following addition of singly purified species. Expression of beta-lactamase fused to TARP$^{1-200}$ by the heterologous system Yersinia enterocolitica allowed for secretion of TARP into type-III inducing media (low calcium). Furthermore I was able to detect Slc1-dependent translocation of TARP into HeLa cells via the heterologous type III secretion system of Y. enterocolitica, and also by the SPI-2 system of Salmonella enterica serovar typhimurium.
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CHAPTER I

INTRODUCTION

Epidemiology

A myriad of diseases are caused by pathogenic Chlamydia species that can efficiently invade human mucosal epithelial cells. Genital infections of the superficial mucosal epithelium in the endocervix or urethra by serovars D-K can result in cervicitis, urethritis, endometritis, salpingitis, and proctitis. LGV1-3 of C. trachomatis are responsible for invasive genital infections that result in lymphogranuloma venereum, and serovars A-C are responsible for ocular infections that can result in blinding trachoma. Lung infections with C. pneumoniae or C. psittaci can result in mild upper respiratory or life-threatening pneumonias, respectively. Chlamydial infections are also endemic to koalas, guinea pigs, mice, sheep, and cattle.

Chlamydia is the most common bacterial sexually transmitted disease in the United States. Most Chlamydia infections are asymptomatic, allowing the bacteria to persist in the genital tract until it is cleared naturally by the host immune system. In a small fraction of infected females, long-term complications such as pelvic inflammatory disease (PID), ectopic pregnancy, and tubal infertility occur[1]. An extensive survey of Chlamydia genital infections in 2007 was reported by the Center for Disease Control (CDC) and they estimated over 1.1 million infections in the U.S. Their most convincing findings were from the screening of the National Job Training applicants that reported
infection rates of 13.2% of females and 7.2% of males aged 16-24 year olds were positive for *C. trachomatis* infections[2]. Furthermore, the World Health Organization has estimated over 90 million cases of *C. trachomatis* infection worldwide[3], with serovars D, E, and F being the most prevalent[4, 5].

*C. trachomatis* is also the causative agent of blinding trachoma that accounts for 16% of blindness worldwide. Ocular infection mainly affects young children (0-4 yrs old) and is cleared without complications within 2-4 weeks. However, familial transition to adults can result in serious scarring of the tarsal conjunctiva, inversion of the eyelid, and corneal scarring due to scratching by the in-turned eyelashes. Without treatment, continued irritation by chlamydial antigens will lead to corneal opacity and eventually blindness[6]. Trachoma is mainly a disease endemic to poverty-stricken villages with little access to clean water and medicine.

The World Health Organization currently funds a campaign to eradicate blinding trachoma by 2020 that utilizes the SAFE strategy (Surgery, Antibiotic treatment, Facial cleanliness, and Environmental improvement)[7]. Some advanced ocular infections require eyelid surgery in addition to antibiotic treatment to prevent blindness. Treatment with tetracycline derivatives (doxycycline) or macrolides (erythromycin, azithromycin) is sufficient to clear up most *C. trachomatis* infections. Mass treatment with antibiotics has been proposed to eradicate trachoma within specific communities, but evidence of pneumococcal resistance to these antibiotics has emerged as a major barrier to mass treatment strategies[8].

*C. pneumoniae* is responsible for 10% of mild upper respiratory infections. It has been estimated that 80% of people will have had *C. pneumoniae* infection in their
lifetime[9]. *C. pneumoniae* has also been found in the joints and in atherosclerotic plaques, associating the bacterium with chronic inflammatory diseases as well[10].

**Chlamydia pathogenesis**

The ability of *Chlamydia trachomatis* to cause damage to human tissues is a consequence of the host immune response to infection. Clearance of *C. trachomatis* genital infections involves a combination of innate and cellular immune responses[11, 12] The housekeeping gene, heat-shock protein 70 (Hsp70), from *Chlamydia* species is recognized by host cell toll-like receptors TLR2 in cooperation with CD14 to induce secretion of the pro-inflammatory cytokine IL-8 (interleukin-8)[13]. Initial infection of human epithelial cells results in the release of pro-inflammatory cytokines such as IL-1β, IL-6, GM-CSF (granulocyte-macrophage colony stimulating factor) and TNF (tumor necrosis factor) that attract monocytes, natural killer cells and neutrophils, as well as in releasing tissue-damaging matrix-metalloproteases (MMPs)[1, 13]. These recruited cells further release damaging MMP’s, elastase, and secrete IFN-γ (interferon-gamma) to drive a Th1-like CD4+ T-cell response. Further mediators of cellular immunity (CD8+ T-cells, B cells, and plasma cells) are recruited to the site of infection and secrete IFN-γ which activates IDO (indoleamine-2,3-dioxygenase) and subsequently blocks synthesis of tryptophan in RBs [1, 12].

Treatment of RB’s with IFN-γ, Penicillin, or nutrient or iron starvation in cell culture induces a “persistent” non-dividing, but transcriptionally active state. Under these conditions the RBs become enlarged while persisting as an “abberant” RB that can survive inside the host cell for extended periods of time. The fact that these RB’s can be
recovered from their persistent state and continue their developmental cycle upon removal of the persistence-inducing stimuli hints at possible role for developmental persistence in long-term “silent” chlamydial infections[14].

Decreased organismal load and decreased duration of repeat Chlamydia trachomatis genital infections indicate the induction of adaptive immunity [9, 15, 16]. Interestingly, repeat infections are common and more likely to result in long-term complications indicating that any acquired immunity is incomplete [17]. Host factors such as HLA-DR alleles and IL-10 promoter polymorphisms are also associated with clearance of the pathogen by influencing the ability of the infected persons to mount an effective Th1 immune-response[18].

In a minority of infections, the host immune response induces fibrosis and scarring of the fallopian tubes that are responsible for tubal infertility and ectopic pregnancy[1, 11]. However, it is still unclear which immune cells are responsible for infection-associated morbidity. Several studies have identified reduced IFN-γ production by CD4+ T-cells and Peripheral Blood Mononuclear Cells (PBMC’s) as an indicator of tissue damage and long-term sequelae.[19-21].

The challenge of designing a vaccine for immunization against Chlamydia trachomatis would be to initiate a bacteria-clearing Th1-response while avoiding a tissue-destructive inflammatory response in individuals with previous Chlamydia infections[11, 16]. In fact, a dead-whole cell vaccine against ocular Chlamydia trachomatis infections was administered in the 1960’s, which was effective in some individuals, while damaging to those who were re-infected[16]. Furthermore, an ideal vaccine would integrate
chlamydial antigens that would elicit an effective immune response against all pathogenic Chlamydia species.

*Adaptation of pathogenic Chlamydia to their mammalian hosts*

Comparison of ribosomal RNA from Chlamydiae with that of other modern prokaryotes indicates a divergence of Chlamydiae from Proteobacteria and other prokaryotes nearly 2 billion years ago. Pathogenic and environmental species of the order Chlamydiae are obligate intracellular prokaryotes that split about 700 million years ago. Environmental species reside within unicellular eukaryotes such as free-living amoeba and are ubiquitous in nature. Differences between the genome of the recently characterized *Acanthameoba* environmental strain UI25 and those of modern pathogenic species have provided insight into the adaptation of pathogenic family Chlamydiaceae to their niche in the mammalian cell.

The genome of the UWE25 strain is more than twice the size (~2.4 million) of those of the pathogenic strains (~1 million bp), while a core set of 711 CDS were conserved among these strains. Adaptation of the pathogenic Chlamydia to mammalian hosts involved a drastic reduction (~2000 CDS to 1000 CDS) in genes, many of which encoded proteins involved in biosynthetic and metabolic pathways. Pathogenic species are auxotrophic for most amino acids and nucleotides and compete with their host cells for them. Other adaptations unique to pathogenic Chlamydia include genes encode possible bacterial adhesins: the major outer membrane protein, MOMP (ompA) and the polymorphic outer membrane proteins (Pmps). These additions to the pathogenic...
chlamydial surface likely play a role in attachment to specific host cell types and/or immune evasion[22].

However, the cysteine-rich outer membrane proteins OmcA and OmcB that make up Chlamydia’s unique disulfide-bonded outer membrane are present in UI25. Unlike other gram-negative bacteria, Chlamydia do not synthesize a peptidoglycan layer on their surface for structure, instead highly crosslinked protein complexes of OmcA, OmcB, and MOMP provide stability and protect EB’s from environmental stress[23]. Conserved genes between UI25 and modern pathogenic Chlamydia indicate the importance of their gene products for maintaining intracellular survival. In addition to the outer membrane proteins (OmcAB), these conserved genes encode some inclusion membrane proteins (Incs), components of the type III secretion system (T3SS), as well as some type III secretion effectors.

**Type III Secretion**

Type III Secretion Systems are syringe-like molecular machines that are common in pathogenic and non-pathogenic gram-negative bacteria and transport specific bacterial proteins into eukaryotic cells. Pathogenic bacteria often utilize type III secretion systems deliver effectors that hijack cellular functions such as reorganizing the host cytoskeleton, modulating immune signaling or apoptotic factors. A more detailed discussion of type III effector functions will be discussed later. A functional type three secretion system consists of structural components that make up the inner and outer membrane base, the needle shaft, tip, and translocator proteins[24]. It also requires cytoplasmic chaperones...
and a membrane bound ATPase to effectively stabilize and deliver effector proteins, respectively.

These components of Type III Secretion Systems remain conserved across gram-negative species and also share close structural homology with flagellar components. Such structural conservation has allowed for the quick identification of putative chlamydial type III secretion proteins *in silico* analysis based on conserved structural and genetic domains[25-31]. Many groups have identified putative *Chlamydia* TTSS components by these approaches, but because of *Chlamydia* species’ genetic intractability, it has remained a challenge to confirm the function of these proteins. Most of our current knowledge of the structure of the chlamydial type III secretion apparatus comes from the discovery of *in vitro* protein-protein interactions between chlamydial type III components.[27, 32-41], while the detection of new chlamydial effectors has involved expression and secretion of recombinant chlamydial proteins in heterologous type III secretion systems of other bacteria[38, 40-43]. The predicted structure of the Chlamydia type III secretion apparatus and its components is visualized in Figure 1.

Assembly of the chlamydial type III apparatus begins with sec-dependent secretion of CdsC, an outer membrane lipoprotein with homology to secretins[44] followed by secretion of CdsJ into the inner membrane[45], and then CdsD into the integral membrane to join the former components into a scaffold that spans two bacterial membranes[31]. CdsD homologs in other bacteria function as scaffolding proteins to link CdsC and other integral membrane components. CdsD has been shown to co-immunoprecipitate with other integral membrane proteins as expected[36], but unique features such as forkhead association domains (FHA) and *in vitro* phosphorylation are
puzzling as to the actual role of CdsD in type III secretion[46, 47]. Homology with inner-membrane components from *Yersinia* species indicate that CdsJ provides a scaffold for the inner membrane components CdsR/S/T/U and V[46]. CdsS and CdsT interact with the C-ring component CdsQ, while the CdsU homolog in *Yersinia* YscU gets cleaved to allow secretion[48]. CdsD has been shown to interact with the cytoplasmic ATPase CdsN and with CdsL, a possible negative regulator of ATPase activity[35]. A complex network of interactions has been shown to occur *in vitro* between CdsQ, CdsD, CdsN, and CdsL [35, 49]. Upon assembly of the inner membrane and C-ring components, the machinery becomes secretion competent and secretes its needle subunits CdsF[27], possibly with the help of the putative molecular ruler protein CdsP[50]. Lastly secretion of the tip protein, identified as CT584 in *C. trachomatis*, completes the latent type III secretion apparatus and is likely to be responsible for activating secretion upon host cell contact[34]. CopN interacts with the ATPase CdsN and prevents secretion of any effectors[41, 51]
Figure 1. The Type III Secretion System of *Chlamydia*. The chlamydial *cds* components *C*-*V* correspond to their *ysc* homologues in *Yersinia*.
Interaction of the chlamydial EB with the host cell is sensed by the tip protein, which in turn relays an unknown signal to induce CopN secretion. Unblocked, the type III apparatus secretes pairs of translocator proteins (CopB/CopD or CopB2/CopD2) to the end of the needle such that they can form a pore in the host cell membrane[37, 38, 46]. After secretion of CopN and the translocator proteins, the ATPase is free to interact with effectors and ensure their translocation into the host cell.

It must be noted however that several of the secreted components and effectors require or utilize molecular chaperones to be efficiently secreted. Bacterial type III secretion chaperones have been organized into classes based on the type of substrates with which they interact. Class I chaperones mediate the translocation of type III effectors and are typically small (13-16 kDa), have an acidic pI (4-5), and have a tendency to dimerize. CT043, CT088 (also called Scc1), and CT663 have been predicted to be *C. trachomatis* class I chaperones based on their predicted secondary structure[28, 29, 52], but to date their interaction with effector proteins has not been confirmed. A recent yeast-2-hybrid screen identified an interaction between CT663 and CT088, indicating formation of a chaperone heterodimer[49]. The same screen also identified a new class Ib chaperone that was capable of interacting with multiple effectors as well with the C-ring component CdsQ which was described as the multi-cargo secretion chaperone, Mesc. Characterization of Mesc by crosslinking also indicated a homodimeric molecular structure. In addition, CT663 has been shown to interact with region 4 of the sigma-66 subunit of *Chlamydia* and the β-subunit of *E. coli* RNA polymerase in a fashion typical of anti-sigma factors[53].
The translocator proteins are secreted by Class II chaperones which have tetratricopeptide repeat (TPR) regions. In *Chlamydia trachomatis* the TPR-containing chaperones Scc2 and Scc3 are transcribed in operons with their sets of translocators, CopB1/CopD1 and CopB2/CopD2, respectively. Scc2 has been demonstrated to interact with both CopB1 and CopD1, but unexpectedly Scc3 has been shown to interact with CopN, the type III secretion-negative regulator. A third TPR-containing protein, CT274, has been detected in *Chlamydia* spp. and its small size and acidic pI indicate that it may have chaperone function. Novel interactions between CT274 and the hypothetical proteins CT668 and CT166 indicate possible *Chlamydia*-specific type III related proteins.

Type III chaperones that are responsible for the stability or secretion or type III components other than effectors or translocators have been classified as Class III chaperones[46]. The needle subunit CdsF and molecular ruler protein CdsP have been demonstrated to interact with chaperone products from their respective operons, *cdsEFG* and *cdsOP*. A yeast-3-hybrid assay indicated that CdsE and CdsG form a heterodimer that can interact with CdsF[27]. Multiple studies have identified an interaction between CdsP and its putative chaperone CdsO[33, 54]. Furthermore, CdsO was shown to form homodimers indicative of type III chaperones. Predicted and experimentally confirmed type III secretion chaperones and their substrates are summarized in Table 1.

Many roles have been reported for the interaction of Class I type III secretion chaperones with their cognate substrates. The most popular role for a Class I chaperone has been to bind approximately the 30-130 N-terminal amino acids of its cognate effector and to unfold the adjacent N-terminal secretion signal. Some chaperones have also been demonstrated to increase the solubility of their substrates or prevent their degradation[55-
58]. Directly mediating the interaction between the effector and ATPase has also been reported[59, 60]. Less common functions have included blocking secretion-inhibitory domains or enzymatic domains of effectors[61, 62].

Many pathogenic bacteria such as *Salmonella, E.coli, Shigella, Pseudomonas,* and *Yersinia* species utilize type III secretion systems to export virulence factors into target cells[63]. In these bacteria, the type III secretion-related genes are mostly located in pathogenicity islands or on specific virulence plasmids. However most type III genes in *Chlamydia* are spread out among ten operons while a few other type III-related genes are completely separate in the genome. In fact this disjointed type III structure is conserved among environmental *Chlamydia isolates*[64]. Approximately 10% of the *Chlamydia trachomatis* serovar D genome is annotated to be type III secretion-related. Considering the small size of the genome it is likely that these are all functional and necessary for productive infection.

In addition to type III secretion components, *Chlamydia* also contains an incomplete set of flagellar genes. Recent studies have found novel interactions between chlamydial flagellar and type III components. The flagellar ATPase FliI has been shown to have ATP-hydrolyzing activity and interacts *in vitro* with CdsL and CopN. Furthermore the flagellar homolog (FlhA) to the inner membrane protein CdsV can interact with CdsU, CdsQ, and CdsL[32]. These findings taken together with the fact that *Chlamydia* is non-motile indicate that the interaction of flagellar components with the type III secretion apparatus may be important in regulating its unique intracellular lifestyle[32, 46].
<table>
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<th>Chaperones</th>
<th>Interacting partners</th>
<th>Evidence</th>
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<td>Slc2 (CT663)</td>
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<td>CT161 (apparatus?)</td>
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<td>CdsF (needle subunit)</td>
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</table>
Type III secretion and Chlamydia’s developmental cycle

One common feature of all Chlamydia species is their biphasic developmental life cycle that imposes upon them an obligate intracellular growth restriction. A small non-infectious but metabolically inactive EB, or elementary body, attaches to a eukaryotic cell membrane by reversible electrostatic-interactions with unknown host receptors. A second irreversible interaction occurs and entry into the host cell is guaranteed by the translocation of effectors by a type III secretion apparatus. Pre-formed in the EB and pre-loaded with virulence factors, the chlamydial type III secretion apparatus translocates its EB-associated effectors into the host cytosol upon contact.

Because of their complex life cycle, it has been challenging for researchers to stably introduce nucleic acids into the bacterium. It is not yet possible to make knockouts, express recombinant proteins or perform small interfering Ribonucleic Acid (siRNA) knockdowns in Chlamydia. For this reason, much of the characterization of chlamydial proteins has focused on their interactions in host cells. Thus the roles of chlamydial type III secreted proteins or effectors in the developmental cycle of Chlamydia have been a major focus of research.

One particular EB-associated effector, TARP, the Translocated Actin Recruitment Phosphoprotein is detected in the host cytosol minutes after infection and is involved in hijacking host signaling molecules to the effect of inducing endocytosis of the attached EB[42, 66-73]. Close contact with the eukaryotic cell surface allows interaction of the EB’s disulfide-crosslinked outer membrane with host protein disulfide isomerase (PDI) such that the chlamydial outer membrane proteins become reduced and its uptake can occur. Immediately upon uptake into its endocytic vesicle, the EB’s DNA starts to
decondense and it begins to differentiate into a large metabolically active RB or reticulate body. A small metabolite interrupts the interaction of the chlamydial histone-like proteins Hc1 and Hc2 with chromatin and the subsequent decondensation of DNA allows for transcription of early genes.

Expression of inclusion membrane proteins, IncD-G, by RBs is detected within 2 hours of infection[74-76]. Highly divergent in sequence and function, these proteins share large bi-lobal hydrophobic domains of 40-60 amino acids in length that ensure their localization. Secretion and outer membrane localization of these Incs is pivotal to forming its unique intracellular niche, the “inclusion,” that diverges from the customary endocytic pathway. IncA, IncG and the Inc CT229 have been demonstrated to be type III secretion substrates by heterologous bacteria, while the mechanism of secretion of other Incs is unknown. Characterization of IncA revealed a eukaryotic SNARE-like domain that may mediate interaction with host vesicles. Similarly IncA has been shown to mediate homotypic fusion of inclusion vesicles[75]. Another host mechanism that *Chlamydia* mimics to influence the trafficking of host vesicles is the recruitment of Rab GTPases. The Inc CT229 has been shown to interact with Rab4[77], and the *C. pneumoniae* inc Cpn0585 can interact with Rabs 1, 10 and 11[78]. The Rab6-associating protein Bicaudal-1 has been found localized to the inclusion as well, possibly by the action of an inc protein[74]. Recruitment of Rab-GTPases to the inclusion can direct vesicles from the golgi, multivesicular bodies, and recycling pathways to the inclusion where essential nutrients, amino acids and specialized lipids can be acquired. Another characteristic of the early inclusion is its recruitment of dynein and subsequent migration toward the host microtubule organizing center (MTOC)[79].
A major mechanism of survival of the parasitophorous inclusion is to delay the host cell cycle and preventing apoptosis. The C. trachomatis IncG proteins have been shown to sequester the host protein 14-3-3 and prevent phosphorylation of the pro-apoptosis protein BAD [80, 81]. Cleavage of the cell cycle regulator cyclin B1 was also detected as a consequence of chlamydial infection [82]. The secreted protein CPAF (Chlamydia protease-like secretion factor) has been shown to have several roles during infection including degrading BH3-like apoptosis proteins [83].

Other effectors have been shown to be involved in suppressing host inflammatory signaling. ChlaDub1 and CT441 are able to interrupt NFkB-signalling by deubiquitinating IκBα and by degradation of RelA, respectively [84, 85]. Sequestering of Act1 by the C. pneumoniae effector CP0236 prevents it interaction with IL-17 and subsequent NFkB-activation [86]. CPAF has also been implicated in the degradation of host factors to prevent proper MHC and lipid antigen presentation [75].

As the inclusion becomes modified additional type III secretion needles are assembled and a second set of effectors is expressed. Expression of mid-cycle genes (6-16 hours p.i.) allows for intermediate metabolism such that RBs can continue to grow and multiply by binary fission. Although Chlamydia do not have a peptidoglycan layer, they are affected by treatment with Penicillin- indicating a possible role for peptidoglycan during division. This theory is corroborated by the transcription of peptidoglycan synthesis genes prior to cell division and the lack of an FtsZ homolog in Chlamydia species [23]. Possibly by a contact-dependent mechanism, newly divided RBs detach from the inclusion membrane and differentiate back to small infectious EBs. It has been proposed that as RBs divide and the inclusion size increases, individual RBs have less
surface contact with the inclusion membrane and detachment of the RB occurs, possibly by retraction of type III secretion apparatuses[75, 87, 88]. While microscopic observations and mathematical models support the contact-dependent model of detachment, the mechanism connecting detachment and differentiation remains elusive.

Two distinct mechanisms of EB exit have been identified. Ordered permeabilization of the inclusion membrane, nucleus, and host cell membrane has been demonstrated to release EB’s during cysteine protease-induced cell lysis. Extrusion, or a slow pinching off of the chlamydial inclusion, has also been observed at nearly the same frequency as cell lysis during C. trachomatis infection[89]. Future research to define any differences in infectivity, transmission, and pathogenesis by EB’s released by lysis versus extrusion will be highly anticipated.

Before secondary differentiation, however, proteins important for invasion, such as bacterial adhesins, components of the type III apparatus and invasion-associated effectors must be expressed and preloaded for immediate translocation into the host cell. Gene expression of type III components have been detected from 8 hours post-infection onward. However, specific subsets of type III-related genes have been shown to be temporally regulated[90-93], indicating functions at different points in the developmental cycle. Evidence of transcription of type III-related genes and their temporal classes are compiled in Table 2.
### Table 2. Expression of type III genes during the chlamydial developmental cycle.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Temporal class [39, 91-93]</th>
<th>Type III function</th>
</tr>
</thead>
<tbody>
<tr>
<td>lncG</td>
<td>E</td>
<td>Effector</td>
</tr>
<tr>
<td>CT229 (Inc)</td>
<td>E</td>
<td>Effector</td>
</tr>
<tr>
<td>lncA</td>
<td>M</td>
<td>Effector</td>
</tr>
<tr>
<td>TARP</td>
<td>M, VL</td>
<td>Effector</td>
</tr>
<tr>
<td>S1c1</td>
<td>M</td>
<td>Chaperone (I)</td>
</tr>
<tr>
<td>Scc1</td>
<td>C</td>
<td>Chaperone (I)</td>
</tr>
<tr>
<td>S1c2</td>
<td>M</td>
<td>Chaperone (I)</td>
</tr>
<tr>
<td>FliF (J)</td>
<td>M</td>
<td>Apparatus</td>
</tr>
<tr>
<td>FliN (Q)</td>
<td>L</td>
<td>Apparatus</td>
</tr>
<tr>
<td>FliA</td>
<td>M</td>
<td>Sigma 28</td>
</tr>
<tr>
<td>FliA (V)</td>
<td>M</td>
<td>Apparatus</td>
</tr>
<tr>
<td>FlhA (N)</td>
<td>L</td>
<td>ATPase</td>
</tr>
<tr>
<td>CdsC</td>
<td>L</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsE</td>
<td>M</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsL</td>
<td>M</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsJ</td>
<td>M</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsR</td>
<td>M, C</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsS</td>
<td>M, C</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsT</td>
<td>L, C</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsU</td>
<td>M, C</td>
<td>Apparatus</td>
</tr>
<tr>
<td>CdsV</td>
<td>M, L</td>
<td>Tip/Cap</td>
</tr>
<tr>
<td>CT274</td>
<td>M</td>
<td>Chaperone (II)</td>
</tr>
<tr>
<td>Scc2</td>
<td>L</td>
<td>Chaperone (II)</td>
</tr>
<tr>
<td>Scc3</td>
<td>M</td>
<td>Chaperone (II)</td>
</tr>
<tr>
<td>CopB</td>
<td>L</td>
<td>Translocator</td>
</tr>
<tr>
<td>CopD</td>
<td>L</td>
<td>Translocator</td>
</tr>
<tr>
<td>CopB2</td>
<td>M</td>
<td>Translocator</td>
</tr>
<tr>
<td>CopD2</td>
<td>M</td>
<td>Translocator</td>
</tr>
<tr>
<td>CopN</td>
<td>M</td>
<td>Neg. Regulator</td>
</tr>
</tbody>
</table>

1. E= early, M=Mid-cycle, L=Late-cycle, VL= Very Late C=constitutively active.
2. Type-III apparatus homolog to Flagellar proteins is in parenthesis.
Attachment

The initial association of *Chlamydia* elementary bodies to the epithelial cell surface has been shown to involve a heparin-sensitive reversible electrostatic interaction with host cells. A plethora of data has been accumulated regarding the role of heparan-sulfate like glycosaminoglycans (GAGs) in this attachment step. Early studies to determine possible bacterial adhesins demonstrated that infection can be limited by pretreatment of the bacteria or host cells with heparan-sulfate[94-101]. Coating of *Chlamydia trachomatis* and *pneumonaie* EBs with heparan-sulfate specific monoclonal antibody prevents infection[97]. Treatment of host cells or EBs with heparinase or with chemically-modified heparin-derivatives also prevented infection of *C. trachomatis* serovar E and L2 strains[99, 101]. Several groups have attempted to determine the role of host cell heparan sulfate in attachment by infecting heparan-sulfate deficient mutant CHO cell lines. A few of these studies have indicated a reduction in attached EBs to these cell lines, while a more recent study has indicated that host heparan-sulfate is not necessary for attachment.

Substantial evidence implicates the chlamydial antigen MOMP (major outer membrane protein) in the initial binding of host cells. It was shown that trypsin-mediated cleavage of the variable regions of MOMP inhibited *C. trachomatis* serovar B, but not L2 infection[102, 103]. Furthermore, *in vitro* treatment of host cells with MOMP-specific antibodies severely abrogates *C. trachomatis* EB binding and infection. EB-surface associated MOMP is linked to a glucosamine containing sulphated-polysaccharide that seems to be important for binding mammalian cells[96, 104-107]. N-glycanase cleavage of this polysaccharide prevents infection by *C. trachomatis* strains[105, 106, 108]. This
MOMP-associated GAG has likely complicated the interpretation of the heparin, heparan-sulfate, and heparinase experiments mentioned earlier.

A second temperature-dependent binding step was discovered by infection of a mutant CHO cell line with *Chlamydia trachomatis* L2 strain[109]. Attachment of EBs to wild-type CHO cells at 4°C could be reversed by addition of heparin, but attachment at 37°C could not. Isolation of a mutant cell line that could be competed for binding to EBs by heparin at 37°C indicated a loss of the second heparin-resistant, temperature-dependent receptor.

Many different host ligands have been proposed as this second receptor that remains irreversibly bound to EBs. Mannose-6-Phosphate treatment of epithelial cells can inhibit attachment and entry of *C. pneumoniae*, indicating a possible role for Mannose-6-Phosphate receptor/Insulin Growth Factor 2 as a receptor[110]. Knockdown of host cell Platelet-derived growth factor receptor-β (PDGFRβ) with siRNA and treatment with PDGRβ-monoclonal antibody reduce EB binding[111]. PDGRβ has also been indicated as an important player in *Chlamydia pneumoniae* infection by a recent systems biology study[112]. A recent study of the role of the cellular cystic fibrosis transmembrane conductance regulator (CFTR) during *C. trachomatis* infection describes CFTR-dependent uptake of EBs and an increase in EB internalization by CFTR-ectopically expressing cells[113].

Other bacterial adhesins besides MOMP that have been implicated in host receptor binding include LPS, the outer membrane complex protein OmcB, and the polymorphic membrane proteins Pmp6, Pmp20, and Pmp21. OmcB from *C. trachomatis* serovars LGV1 and E show GAG-dependent and independent binding to host cells, respectively, indicating that it may be involved in either the primary or secondary
attachment step[114]. *C. pneumoniae* Pmp proteins contain tetrapeptide repeat motifs that mediate binding to epithelial cells, and a recent study indicates that the presence of multiple Pmps have an additive effect on attachment to host cells[115].

A complex role for Protein Disulfide Isomerase (PDI) in attachment was also discovered. Reduced attachment of *C. trachomatis* to a CHO cell line with a mutated-leader sequence in PDI could be restored by complementation with the full length PDI[116]. This requirement of PDI for attachment has been confirmed in all other pathogenic *Chlamydia*. However, it is suspected that PDI does not behave as an actual receptor but instead structurally associates with another host receptor to initiate attachment[117].

While enzymatic activity of host PDI is not necessary for attachment, it is an absolute requirement for chlamydial entry into epithelial cells. Thus, as the chlamydial EB is brought in close vicinity of the eukaryotic membrane by electrostatic interactions, another secondary host receptor in coordination with PDI irreversibly binds and reduces the crosslinked-outer membrane proteins and initiates uptake events such as the activation of the type III secretion system. Tight-binding of the type III secretion apparatus to the host cell membrane could indicate that the type III needle and tip proteins as possible secondary receptors, though further research in this area is needed to corroborate or disprove the role of the type III secretion apparatus in attachment.

Despite years of research dedicated to the search for host and chlamydial receptors that mediate attachment, the use of a variety of host cell types and chlamydial species has made it difficult to determine a general mechanism for attachment. A compendium of this work can be found in *Tables 3 and 4*. 
<table>
<thead>
<tr>
<th>Adhesin/Receptor</th>
<th>Serovars</th>
<th>Evidence</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MOMP</strong></td>
<td>L2, Cps</td>
<td>1. A 32 kDa EB protein bound Hela cell extracts and heparin.</td>
<td>[118, 119]</td>
</tr>
<tr>
<td></td>
<td>J, L2</td>
<td>2. Infection could be neutralized with mAB against 31 kDa protein.</td>
<td>[119]</td>
</tr>
<tr>
<td></td>
<td>B, L2</td>
<td>3. Trypsin-mediated cleavage of VD IV prevented attachment.</td>
<td>[102]</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>4. mAB against VD II and IV blocked electrostatic interactions.</td>
<td>[103, 104]</td>
</tr>
<tr>
<td></td>
<td>C. trach</td>
<td>5. Glycan cleaved from 32 kDa protein binds Hela cells.</td>
<td>[105]</td>
</tr>
<tr>
<td></td>
<td>L2</td>
<td>6. MOMP carbohydrate mapped to reveal N-linked high mannose oligosaccharide.</td>
<td>[106]</td>
</tr>
<tr>
<td></td>
<td>C. trach</td>
<td>7. Recombinant MOMP competes with EBs for binding Helas and had reduced binding to heparin sulfate synthesis-deficient CHO cells.</td>
<td>[120]</td>
</tr>
<tr>
<td><strong>Heparin Sulfate-like GAGS</strong></td>
<td>C. trach</td>
<td>1. Isolation of a heparin-sulfate-like GAG from EBs recovered from GAG-deficient CHO cells. Another study was unable to isolate the chlamydial GAG from EBs cultured in a separate HS-deficient CHO cell line.</td>
<td>[121, 122]</td>
</tr>
<tr>
<td></td>
<td>L2</td>
<td>2. GAG-on intracellular RBs visualized and cleaved by lyase.</td>
<td>[97]</td>
</tr>
<tr>
<td></td>
<td>LGV,E</td>
<td>3. Heparin-coated microspheres were taken up similarly to EBs.</td>
<td>[98]</td>
</tr>
<tr>
<td></td>
<td>Ctr, Cpn</td>
<td>4. Desulfation of L2 or E EBs by sodium chlorate treatment did not reduce infectivity.</td>
<td>[123]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. Treatment of C.pneumo strains and C.tr E EBs with heparin derivatives or heparinase inhibit infection</td>
<td>[101]</td>
</tr>
<tr>
<td><strong>OmcB</strong></td>
<td>LGV, E</td>
<td>1. OmcB was identified as binding heparin. The Binding region was mapped to a 20-mer peptide whose addition could out-compete binding by OmcB.</td>
<td>[124]</td>
</tr>
<tr>
<td></td>
<td>Cpneumo., L2, E</td>
<td>2. Addition of recombinant OmcB from LGV1 but not E prevents EBs from binding host cells. Attachment of LGV1 rOmcB but not E was abrogated in the presence of heparin.</td>
<td>[114]</td>
</tr>
<tr>
<td></td>
<td>Ctr, Cpn, Cps</td>
<td>3. Expression of C.pneumo OmcB on Yeast was sufficient for binding to epithelial cells and addition of heparin prevented this binding. Addition of OmcB recombinant</td>
<td>[125]</td>
</tr>
</tbody>
</table>
protein or monoclonal antibody prevents C. pneumo EB attachment to host cell.

1. Chalmydia LPS immunoprecipitates with CFTR.

1. Recombinant expression of Pmp6, 20, or 21 on yeast cells mediates adhesion to epithelial cells. Treatment of host cells with recombinant Pmps or mAb against these proteins prevents C pneumo infection. Binding can be attributed to the tetrapeptide motifs in these proteins.

LPS  E, LGV  protein or monoclonal antibody prevents C. pneumo EB attachment to host cell. 1. Chalmydia LPS immunoprecipitates with CFTR. [113]
Pmps  Cpn, L1 E  1. Recombinant expression of Pmp6, 20, or 21 on yeast cells mediates adhesion to epithelial cells. Treatment of host cells with recombinant Pmps or mAb against these proteins prevents C pneumo infection. Binding can be attributed to the tetrapeptide motifs in these proteins. [115]

<table>
<thead>
<tr>
<th>Receptor</th>
<th>Cell types</th>
<th>Evidence</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDGFRβ</td>
<td>S2, Hela</td>
<td>1. Treatment of host cells with PDGFRβ siRNA or mAb reduces C. trachomatis L2 binding.</td>
<td>[111]</td>
</tr>
<tr>
<td>CFTR</td>
<td>Hela</td>
<td>2. Uptake of C. trachomatis EBs is blocked by CFTR mAbs and by treatment with CFTR inhibitors. Also there was a reduction in EB uptake by CFTR knockout mice. CFTR can also be co-immunoprecipitated with LPS.</td>
<td>[113]</td>
</tr>
<tr>
<td>M6P receptor</td>
<td>Hela</td>
<td>1. Mannose-binding protein interacts with MOMP and prevents interacation of C. trachomatis with host cells.</td>
<td>[126]</td>
</tr>
<tr>
<td></td>
<td>HMEC-1</td>
<td>2. Addition of M6PR or M6P analog prevents attachment of C. pneumo but not C. trachomatis strains to endothelial cells.</td>
<td>[110]</td>
</tr>
<tr>
<td>PDI</td>
<td>CHO-6</td>
<td>1. The CHO-6 cell line is mutant for PDII and is resistant to attachment that could be recovered with FL PDII.</td>
<td>[116]</td>
</tr>
<tr>
<td></td>
<td>CHO-6, Hela</td>
<td>2. siRNA knockdown of PDII prevents infection. <em>Chlamydia</em> does not bind directly to PDII-gpi.</td>
<td>[117]</td>
</tr>
<tr>
<td>Receptor</td>
<td>Cell types</td>
<td>Evidence</td>
<td>Ref</td>
</tr>
<tr>
<td>------------------</td>
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<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Heparin-sulfate</td>
<td>Hela</td>
<td>1. 32 kDa and 18 kDa EB surface proteins are prevented from binding host cell by heparin.</td>
<td>[118]</td>
</tr>
<tr>
<td></td>
<td>Hela, CHO761</td>
<td>2. Infection inhibited by adding heparin-sulfate receptor or heparitinase.</td>
<td>[121, 127]</td>
</tr>
<tr>
<td></td>
<td>L929, Hela</td>
<td>3. Heparitinase treatment of LGV and trachoma EBs prevented attachment.</td>
<td>[127]</td>
</tr>
<tr>
<td></td>
<td>Hela, L929</td>
<td>4. Heparin inhibition of LGV and trachoma biovar infections was rescued by competition with a sulphated decasaccharide.</td>
<td>[94]</td>
</tr>
<tr>
<td></td>
<td>CHO</td>
<td>5. A heparin sulfate synthesis-deficient CHO cell line had reduced EB infectivity. A separate HS-deficient CHO cell line was resistant LGV but not E infection.</td>
<td>[120]</td>
</tr>
<tr>
<td></td>
<td>L929, Hela</td>
<td>6. Attachment of LGV EBs to was completely inhibited by pretreatment with heparin or heparitinase.</td>
<td>[107, 122]</td>
</tr>
<tr>
<td></td>
<td>CHO</td>
<td>7. Attachment of Trachoma biovar EBs was only partially inhibited by heparin or heparitinase pretreatment.</td>
<td>[107, 128]</td>
</tr>
<tr>
<td></td>
<td>Hela, McCoy</td>
<td>8. Serovar E attachment not inhibited by heparin or heparan sulfate, but L2 was completely inhibited.</td>
<td>[128]</td>
</tr>
<tr>
<td></td>
<td>Hela, McCoy</td>
<td>9. Heparitinase blocked C. psittaci EB attachment at 4° and 37° while heparin blocked attachment at 4°.</td>
<td>[95]</td>
</tr>
<tr>
<td></td>
<td>Hela</td>
<td>10. Inhibition of C. trachomatis infectivity by sulphated polymers or polysaccharides in vitro (cell culture) but not in vivo (mouse model)</td>
<td>[96]</td>
</tr>
<tr>
<td></td>
<td>Hela, CHO</td>
<td>11. Heparin sulfate mAb binds to C. trach and C. pn EB surface and neutralizes infectivity.</td>
<td>[97]</td>
</tr>
<tr>
<td></td>
<td>McCoy, Hela</td>
<td>12. Heparitinase treatment of host cells prevented L2 infection and the HS-synthesis deficient CHO cell line was less sensitive to infection.</td>
<td>[99]</td>
</tr>
<tr>
<td></td>
<td>CHO</td>
<td>13. Desulfation of host cells by sodium chlorate inhibited L2 but not E infection.</td>
<td>[123]</td>
</tr>
<tr>
<td></td>
<td>HL</td>
<td>14. The HS mutant cell line CHO-18.4 was infected equally by L2 and D and was comparable to infection of WT CHO cells.</td>
<td>[100]</td>
</tr>
<tr>
<td></td>
<td>CHO</td>
<td>15. Treatment of host cells with heparitinase and heparin derivatives prevented infection by C. pn strains and C. tr E.</td>
<td>[101]</td>
</tr>
<tr>
<td></td>
<td>CHO</td>
<td>16. OmcB-expressing yeast from Cpn and C. tr-L1 were unable to bind heparitinase treated or GAG-deficient CHO cells.</td>
<td>[125]</td>
</tr>
</tbody>
</table>
Entry

Following attachment of EBs, uptake by the endocytic pathway is induced by chlamydial factors. Uptake of chlamydial EBs by epithelial cells is associated with the recruitment of actin and tyrosine phosphorylation of proteins at the site of entry for C.trachomatis[42, 73, 129, 130]. This actin recruitment is dependent on the activation of the small Rho GTPase Rac in C.trachomatis infection, while both Cdc42 and Rac activation are important for C.caviae entry[72, 131]. A tyrosine-phosphorylated protein associated with actin recruitment has been identified to be the chlamydial type III translocated protein TARP. C.trachomatis LGV TARP contains six tyrosine-rich repeat domains that can be tyrosine phosphorylated by several host kinases containing the Src-homology 2–domain (SH2). The SH2-containing kinases Src, Yes, Fyn, Syk, and Abl are all able to phosphorylate tyrosine residues of C.trachomatis TARP[66, 68, 132]. The observation that individual knockdown of any one of these kinases does not completely abrogate tyrosine phosphorylation reinforces the importance of TARP phosphorylation for efficient manipulation of host signaling cascades and actin remodeling. Type III effector-induced actin polymerization and cytoskeletal rearrangement has been extensively studied in other bacteria pathogens. Enteropathogenic E.coli stimulates actin-pedestal formation through the type III translocation of Tir (translocated intimin receptor) into host cells where it is tyrosine-phosphorylated and associated with Arp2/3 recruitment[133, 134]. The SPI-1 (Salmonella Pathogenicity Island-1) type III secretion system induces bacterial uptake by translocation of effectors that induce membrane ruffling and Rho GTPase activation[135].
Tyrosine phosphorylated residues in *C. trachomatis* TARP are able to recruit the guanine nucleotide exchange factors Sos-1 and Vav-2[67, 68]. A complex of Sos-1, Eps8, and Abi-1 or Vav-2 with PIP3 (phospho-inositol 3,4,5-P3) have been shown to activate Rac in a phosphotyrosine dependent manner[67]. Rac-dependent recruitment of actin has been shown to involve Abi-1 and WAVE-2 dependent formation of the actin-nucleating Arp 2/3 complex[69]. While *C. trachomatis* L2 and D TARP s contain 6 and 3 tyrosine-rich repeats (TRR), respectively, *C. caviae, C. pneumoniae,* and *C. psittaci* do not contain TRRs. All chlamydial TARP s do contain proline-rich domains that induce TARP aggregation and C-terminal WH2-like domains that are capable of nucleating actin [70]. It has been proposed that the WH2 domains on TARP may contribute to the further polymerization of the short actin filaments already induced by the Arp2/3 complex[69].

A recent protein microarray indicated the interaction of an SH2-containing adaptor protein SHC1 with TARP derived phosphopeptides. Infection of SHC1 knockdown cells sensitized *Chlamydia* to TNF-induced apoptosis, and detection of SHC1-regulated genes included those involved in apoptosis and growth[136]. Taken together, previous findings clearly indicate that TARP plays a critical role in the entry and survival of *Chlamydia trachomatis* in host cells, and thus its translocation through the bacterial type III secretion apparatus into mammalian cells must be absolutely efficient to ensure survival of the organism.

Besides TARP, the EB-associated type III effectors CT166 and CT694 have been reported to be associated with entry-related signaling. As a homolog to the clostridial glucosylating toxins, the mono-glucosylation and subsequent deactivation of Rac by CT166 has been demonstrated in Hela cells. Since this effector is translocated into host
cells shortly after infection it is likely that it provides a balance to the actin remodeling induced by TARP during *C. trachomatis* entry[137]. The association of CT694 with the host protein AHNAK to alter stress fibers may also be important for invasion.[138]

Tyrosine-phosphorylation of the host protein Ezrin has also been discovered as a direct result of *C. trachomatis* infection of epithelial cells[139]. Like TARP, the tyrosine-phosphorylation of Ezrin upon infection led to its association with actin, specifically in microvilli.[140] The specific kinase(s) involved in Ezrin tyrosine phosphorylation during *Chlamydia* infection have not been determined, but previous studies have demonstrated that this event can be a direct result of host receptor binding and activation[141].

Chlamydial attachment to and subsequent phosphorylation of PDGFRβ results in phosphorylation of WAVE2, Cortactin, and the guanine-exchange factor Vav2[111]. Phosphorylation and subsequent activation of these molecules is associated with actin-remodeling events. It is clear that multiple and redundant signaling pathways are hijacked by *Chlamydia* species to ensure efficient uptake of the organism. Upon cell contact, immediate secretion of chlamydial factors such as TARP into the host cytosol is critical for initiating actin-cytoskeletal remodeling and subsequent endocytosis of the EB.
HYPOTHESIS

I hypothesize that a chlamydial factor is responsible for the efficient association of TARP with the type III secretion apparatus and its ensuing translocation into the host cell cytosol. I propose that the *Chlamydia trachomatis* Class I type III secretion chaperone S1c1 interacts with TARP and is responsible for its efficient translocation through the type III apparatus.
CHAPTER II.

MATERIALS AND METHODS

Bacterial strains and cell culture.

Chemically competent Top10 and BL21star E. coli strains from Invitrogen were used for plasmid propagation and protein expression, respectively. The adenylate cyclase (cya) deficient DHM1 E. coli strains were grown at 30°C on indicator plates (LB + 40 μg/ml X-gal + 1mM IPTG or McConkey + 1% maltose) for 48 hours or in Luria-Bertani broth (LB) at 37°C. All other E. coli strains were grown at 37°C in LB broth or agar plates. The Yersinia enterocolitica WT strain MRS40 or ΔYscN E40(pMSl41) strain were grown in LB or Brain-Heart-Infusion (BHI) broth at 26°C. Media was supplemented with 100 μg/ml carbenicillin (carb), 60 μg/ml kanamycin (kan), 12.5 μg/ml tetracycline (tet), or 35 μg/ml Nalidixic Acid, when appropriate. The wild-type EPEC serotype O127:H6 strain E234/69 and the cesT knockout strain were obtained from Gadi Frankel’s lab and grown in LB broth. The Salmonella enterica typhimurium strain 12023s was obtained from David Holden’s lab and grown in LB broth. Hela cells were cultivated at 37°C with 5% CO2 in DMEM supplemented with 10% FBS and 2 mM L-glutamine.

Plasmid construction.
Slc1, Sccl, Msc, Slc2, and the N-terminal 600 nucleotides of TARP were amplified from *Chlamydia trachomatis* genomic DNA (gDNA). Cest and the N-terminal 600 nt of Tir from enteropathogenic *E. coli* gDNA, and YopH from *Y. enterocolitica* MRS40 gDNA by the polymerase chain reaction (PCR) using the primers listed in Table 5.

Cloning into pET101, pET200, pBAD-TOPO and pENTR-D/SD-TOPO utilized the endogenous topoisomerase activity of Invitrogen directional cloning vectors. The PCR fragments CACC-CT043-FLAG-stop (and CACC-CT663-FLAG-stop were cloned into pET101/D-TOPO, resulting in the plasmids pET101-Slc1-FLAG and pET101-Slc2-FLAG, respectively. The PCR fragment CACC-TARP1-200 was cloned into pET200/D-TOPO, resulting in the plasmid pET200-TARP1-200. All of the constructed plasmids were transformed into Top10 cells, recovered in SOC medium for one hour at 37° C, and plated on LB agar supplemented with carb, kan or both. The resulting colonies were restreaked and grown overnight. Presence of the correct plasmid inserts was confirmed by colony PCR and sequencing. The PCR fragments CACC-CT043 and CACC-CT088-FLAG were cloned into pENTR/SD/D-TOPO, resulting in the plasmids pENTR-Slc1 and pENTR-Scc1-FLAG, respectively. These new entry vectors were recombined with pET-56-DEST using LR recombinase II reaction mix (Invitrogen), transformed into Top10, and selected for on LB+carb agar overnight. The resulting colonies were restreaked on carb plates and grown overnight. The presence of the correct plasmids was determined by colony PCR and sequencing. Plasmid DNA was purified from the Top10 genetically modified strains by using Qiaprep Spin Miniprep Kit.(Qiagen). The BL21 star expression strain was singly or co-transformed with pET200-TARP200, pET101-Slc1, pET101-Slc2, and pET56-Scc1-FLAG. For the expression of chlamydial proteins in *Yersinia*, I digested
Slc2 and Scc1 from pet101-Slc2 and pet101-Scc1 with XbaI and Scal-HF (NEB). These digested inserts were then ligated into the XbaI and Scal sites of pBAD18 (American Tissue Culture Collection). Mcsc and Slc1 were cloned into the pBAD-TOPO vector. SLIM PCR was used to add a KpnI site downstream of TARp in the vector pet200-TARp1-200 to form pet200-TARp1-200-KpnI. This vector was then digested with NdeI and KpnI and the resulting NdeI-TARp1-200-KpnI fragment was cloned into these sites in pCX340. For co-expression in the bacterial 2-hybrid system, all of the PCR inserts were generated to have flanking XbaI and KpnI sites and were cloned into the same sites in pUT18C and/or pKT25 prior to co-electroporation into DHM1 E. coli.

**Protein expression, lysate preparation, and protein purification.**

Overnight cultures of BL21star strains were diluted 1:20 in LB + antibiotic, and grown to OD600 of 0.6. Expression of recombinant proteins was induced by addition of 0.5 mM isopropyl-β-D thiogalactopyranoside (IPTG) and incubated for 3 hours with shaking. Bacteria were centrifuged at 8000g for 20 minutes, and the pellet was resuspended in 1/10 culture volume of lysis buffer (50mM NaH2PO4, 300mM NaCl, 0.05% Triton, 0.1 mg/ml Dnase I, 1 mg/ml lysozyme, Halt Protease Inhibitor Cocktail (Pierce)). For preparation of lysates that were added to NiNta Magnetic Beads, 25 mM imidazole was added to the lysis buffer. Bacteria were incubated in lysis buffer for 30 minutes at room temperature, sonicated to shear DNA, and stored at -80°C. Purifications of Slc1-FLAG and His6- TARp1-200 for use in crosslinking and gel filtration experiments were performed using M2-Anti-FLAG agarose (Sigma) and Ni-Nta Agarose (Qiagen), respectively. Lysozyme was left out of the lysis buffer and the french press was used to
release soluble proteins from these lysates prior to clearing by centrifugation at 15,000 g for 30 minutes. Purification fractions were run on 12% SDS-PAGE gels and proteins were stained with a 0.2% Coomassie Blue R250 solution.

Coimmunoprecipitation assays and immunodetection.
Bacterial expression lysates were thawed and partially cleared by centrifugation at 4000 g for 15 minutes prior to addition to M2 Anti-FLAG Agarose (Sigma) or NiNta Magnetic Beads (Qiagen). 30 mM imidazole was added to 500 µl lysates prior to incubation with the magnetic beads on an end-over-end rotator overnight at 4°C. The flow-through fraction or supernatant was separated from the beads with a magnetic separator, and the beads were washed five times with NiNta Native Wash Buffer (50 mM Na₂HPO₄, 300 mM NaCl, 20 mM imidazole, 0.05% Triton X-20, pH 8.0). The first (W1) and fifth (FW) wash fractions were kept for analysis, and the native protein complexes were eluted from the beads by incubation with 100 µl NiNta Native Elution Buffer (50 mM Na₂HPO₄, 300 mM NaCl, 300 mM imidazole, 0.05% Triton X-20, pH 8.0) at 4°C for 4 hours. For each coimmunoprecipitation experiment with M2 Anti-FLAG agarose, 1 ml of bacterial lysate was added to 100 µl of packed agarose resin and incubated on an end-over-end rotator at 4°C overnight. The flow-through fractions were collected by centrifugation at 3000 g for 1 minute, and the resin was washed five times with tris-buffered saline (TBS, prior to elution of the native protein complexes by incubation with 100 µl of 200 ng/ml 3XFLAG peptide (Sigma) in TBS at 4°C for 4 hours-overnight. Immunoprecipitations of S-tagged proteins were done as per the S-tag ReK purification kit manual. (Novagen). Immunoprecipitation fractions were incubated 1:1 with Laemmli sample buffer (Biorad).
+ 0.5 % &beta;-mercaptoethanol at 95°C for 10 minutes prior to separation by SDS-PAGE. Proteins were transferred to polyvinylidene fluoride (PVDF) membrane and detected by Western blot probed with 6XHIS, DDDDK, and S-tag specific antibodies from Abcam (cat#: ab18588, ab21536, ab18184, ab49763, ab1187). HRP-conjugated goat-anti-mouse (Millipore 12-349) or HRP-goat-anti-rabbit (Pierce #185415) were used as secondary antibodies when necessary, followed by development with Millipore Immobilin Western Substrate.

&beta;-galactosidase assay.

A modified Miller assay was followed[142]. Briefly, co-expressing DHML strains were grown overnight in LB+antibiotic at 37°C followed by a 1:50 subculture and growth in fresh LB+antibiotic to an OD600 of 0.5-0.8. The OD600 of the culture was recorded and 20 ul of culture was added to 80 ul of permeabilization buffer (20 mM KCl, 2 mM MgSO₄, 0.8 mg/mL CTAB (hexadecyltrimethylammonium bromide), 0.4 mg/mL sodium deoxycholate, 5.4 µL/mL beta-mercaptoethanol) and incubated at 30°C for 30 minutes. The time was recorded as 600 ul of pre-warmed substrate solution (60 mM Na₂HPO₄, 40 mM NaH₂PO₄, 1 mg/mL o-nitrophenyl-β-D-Galactoside (ONPG), 2.7 µL/mL β-mercaptoethanol) was added to each permeabilized sample and incubated at 30 °C. Upon developing a yellow color, the samples were stopped with 700 ul of 1 M Sodium Carbonate (Na₂CO₃) and the stop time was noted.

Colocalization assay.
The open reading frames of Slc1 and Slc2 were cloned into pENTR/D-TOPO and recombined into pDest-N-mcherry by the LR recombinase reaction as described by the Gateway expression manual. (Invitrogen) Mcherry-Slc1, Mcherry-Slc2, or Mcherry gus were co-transfected with full length C. caviae GFP-TARP into Cos-7 cells with Fugene 6 Transfection Reagent as indicated in the Fugene 6 Technical Bulletin. (Roche) Transfected constructs were expressed for 24 hrs before the cells were fixed with 4% Paraformaldehyde on glass coverslips and analyzed with an Olympus Fluoview 500 Laser Scanning Microscope.

**Secretion Assay.**

Overnight cultures of Y.enterocolitica strains containing pcx340 and pBAD vectors were diluted to an OD600 of 0.2 in 5 ml of BHI + 2.5 mM CaCl2 (represses Yop production) or BHI + 0.4% glucose + 5 mM EGTA + 20 mM MgCl2 (induces Yop production) for 2 hours at 26° C. Expression of recombinant proteins were induced with the addition of 1 mM IPTG and 13 mM L-Arabinose, upon shifting to 37°C (type III secretion-inducing conditions). Secretion was induced for 4 hours prior to centrifugation at 3000g for 20 min. The top 9 ml of the supernatant were removed and respun at 12000g for 30 minutes at 4°C. This supernatant was further cleared with a 0.45 uM filter and proteins were precipitated with ice-cold 10% TCA. Precipitated proteins were pelleted, washed twice with ice-cold acetone, and dried prior to resuspension in a proportional volume of Lysis buffer B (0.1 M Tris-Hcl, 0.1 M NaH2PO4, 300 mM NaCl, 6 M Urea) to cell pellet. Lysates were treated 1:1 with Laemmli + 5% βME and heated to 95° C for 15 minutes prior to separation on a 12% SDS-PAGE gel. Protein concentration in the whole-cell
pellet was determined by bicinchoninic acid assay (BCA) and the protein in whole-cell pellets was normalized to 1 mg/ml in Lysis buffer B.

**Yersinia enterocolitica Translocation Assay.**

Hela cells were plated in black, flat, glass-bottomed 96-lll plates at a density of 1 X 104 cells per ll and allowed to grow for 24 hours prior to infection. One day prior to infection TEM-1 fusion protein (pcx340) and pBAD18-chaperone expressing *Y. enterocolitica* strains were inoculated into 3 ml of BHI broth + tet ± carb and grown at 26°C overnight. The following day, the cultures were diluted 1:20 in BHI + antibiotics and grown at 26°C for 1 hr. 1 mM IPTG and 13 mM L-Arabinose were added to the cultures and were grown for 2 more hours at 26°C. Monolayers were washed twice with HBSS, and infected with *Y. enterocolitica* at an MOI of 10. After 1 hr of infection at 37°C, the monolayer was washed twice and fresh DMEM was added. The infection progressed for 2 more hours at 37°C in 5% CO2. Infected cells were washed 3 times with HBSS and topped with 100ul of HBSS. The 1xCCF2/AM substrate was prepared as per the manufacturer’s manual (Invitrogen) and 20 ul was added to each ll. Infected cells were incubated at RT for 1.5 hrs in the dark prior to fluorescence detection at 450 and 520 nm with a fluorimeter (excitation at 410 nm).

**Enteropathogenic E.coli and Salmonella typhimurium translocation assays.**

Enteropathogenic E.coli or *Salmonella* typhimurium was inoculated into LB broth + tetracycline and chloramphenicol and allowed to grow at 37°C overnight. Cultures were diluted 1:33 in fresh LB broth + antibiotics and allowed to grow to mid-log phase at 37°C
C (shaking for *Salmonella*, no shaking for EPEC). Bacteria were pelleted, washed and resuspended in PBS and added to 80% confluent HeLa cell monolayers at a multiplicity of infection (MOI) of 50. For the EPEC and SPI-1 assays, the monolayers were infected for 2 hours prior to the washing and CCF2/AM steps (see above), while the *Salmonella* were allowed to infect cells for 9 hours in the SPI-2 assay at an MOI of 10.
CHAPTER III
IDENTIFYING INTERACTIONS BETWEEN TARP AND TYPE III CHAPERONES FROM CHLAMYDIA TRACHOMATIS

Since TARP translocation is a critical event that ensures the uptake and survival of *Chlamydia*, I sought to identify any type III secretion chaperones that may facilitate its translocation. From earlier *in silico* studies, I knew that three such type III class I (effector) chaperones were predicted in *C. trachomatis* genome. I manually aligned the predicted secondary structures of the open reading frames CT043, CT088, and CT663 with known class I effector chaperones from other pathogenic bacteria (Figure 2). In this figure I annotated the aforementioned chaperones as SIC1, Scc1, and SIC2. Previous studies had coined Scc1 (*Specific Chlamydial Chaperone 1*) for CT088[41]. Throughout this study I refer to CT043 and CT663 as SIC1 and SIC2 for SycE-Like Chaperone 1 and 2, respectively.

The predicted chlamydial chaperones shared the α-β-β-α-β-β secondary structure characteristic of type III effector chaperones. Interestingly, all three putative chaperones were reported to be present in the elementary body proteome, [143-146] suggesting that these chaperones may function in the translocation of EB-associated effectors. These findings convinced us to test these putative chlamydial chaperones for their ability to interact with the invasion-associated effector TARP.
Figure 2. Secondary structure alignment of type III chaperones.

Three *C. trachomatis* proteins share predicted secondary structural features characteristic of known type III export chaperones. Primary amino acid sequence alignment of known type III export chaperones CesT (EPEC), SycE (*Yersinia enterocolitica*) and SicP (*Salmonella enterica* serovar *typhimurium*) with the putative export chaperones Slc1, Slc2 and Scc1 of *C. trachomatis*. Secondary structure was predicted using Jpred and sequences aligned manually to according to the α-β-β-α-β-β-α signature characteristic of some type III export chaperones. Residues predicted to form α-helices and β-strands are highlighted dark grey and light grey, respectively.
Interaction of the N-terminal 200 amino acids of TARP with the chaperone Slc1.

Since chaperones typically bind their effectors within the N-terminal 30-150 amino acids, I chose to test the N-terminal 200 amino acids of TARP for binding by the three putative chlamydial chaperones Slc1, Slc2, and Sec1. I utilized two independent approaches – co-immunoprecipitation and the bacterial two-hybrid system to confirm a chaperone-binding partner for TARP. Both of these approaches allowed us to identify interactions between native recombinant proteins and have both been used previously to identify type III chaperone:effector interactions[38, 147]. Co-immunoprecipitation is a directed blunt-force approach for identifying in vitro interactions, while the bacterial two-hybrid assay allows for real-time interaction of expressed proteins and is conducive to quantitative analysis.

For bacterial expression in the co-immunoprecipitation assays, I cloned the N-terminal 200 amino acids of L2 TARP into pET200 (Figure 3) and expressed the recombinant N-terminus of TARP with an N-terminal 6XHIS tag (HIS<sub>6</sub>-TARP<sup>1-200</sup>). Each of the predicted chlamydial chaperones were cloned into pET101 (Figure 5) and expressed to have C-terminal FLAG tags. A stop codon was included after the FLAG tag to exclude the vector-encoded 6XHIS tag from the chaperone open reading frame.

Expression of recombinant TARP and chaperones such that they had different tags allowed us to purify interacting proteins using two different immobilization approaches: HIS<sub>6</sub>-TARP<sup>1-200</sup> with Ni-Nta magnetic beads or chaperone-FLAG with M2 Anti-FLAG agarose. The separate recombinant tags for TARP and chaperones were also important for immunodetection by Western blot analysis.
The open reading frames of the N-terminal 200 amino acids of TARP were amplified from *C. trachomatis* Serovar L2, *Mouse pneumonitis, C. pneumoniae*, and *C. caviae* by PCR. (Fig 3) Slc1, Scc1, and Slc2 were amplified from *C. trachomatis* serovar L2 genomic DNA by PCR. (Fig 5) The purified PCR products were cloned into Invitrogen’s directional cloning vectors (pET200 and pET101) by using the energy stored in a bond between the 3’ phosphate of the linear vector and *Vaccinia* virus topoisomerase I to join the 5’ CACC overhang from PCR products with the GTGG overhang of the linear vector. This allows for directional blunt-end annealing of the PCR products to the linear vectors without any digestion and ligation steps. Transformation of these cloned vectors into chemically competent Top 10 cells gave a high yield on antibiotic-selective plates, and colonies were checked for correct inserts by colony PCR. (Figures 4 and 6) Plasmid DNA from each positive clone was confirmed for the correct inserts by sequencing (Genomic Core-Baxter II, University of Louisville).
Figure 3. Amplification of TARP inserts for cloning into pET200.
(Top) Vector diagram of the pET200 directional topo cloning vector from Invitrogen.
(Bottom) PCR-amplified inserts of the N-terminal 200 amino acids of TARP from different Chlamydial species are displayed on a 1% agarose gel.
Fig 4. Colony PCR of transformants of pET200- TARp1-200.
Colony PCR was performed on kanamycin-selected transformants and the 1.2% agarose gels displayed here indicate the presence of the TARP200aa inserts for each Chlamydial species in the pET200-TARP1-200 vector.
Figure 5. Amplification of chaperone inserts for cloning into pET101.
(Top) Vector diagram of the pET101 reactional topo cloning vector from Invitrogen.
(Bottom) PCR-amplified inserts of the putative type III secretion chaperones from C. trachomatis serovar L2 are displayed on a 1% agarose gel.
Fig 6. Colony PCR of transformants of pET101-chaperones. Colony PCR was performed on kanamycin-selected transformants and the 1.2% agarose gels displayed here indicate the presence of *C. trachomatis* serovar L2 CT663, CT088, and CT043 inserts in the pET101 vector.
Cleared BL21 star *E. coli* lysates containing HIS₆-TARP¹⁻²⁰⁰, chaperone-FLAG, or a mixture of the 2 lysates were incubated with M2 Anti-FLAG agarose for 4 hours at 4°C to allow protein complexes to form and become immobilized on the agarose. After 5 wash steps to remove any non-specific proteins, I eluted the protein complexes with 250 mM 125 μg/ml 3XFLAG peptide. The specific retention of TARP by the chaperones was monitored in the flow-through, wash, and elution fractions by Western blot analysis (Figure 7).

The M2 Anti-FLAG beads retained HIS₆-TARP¹⁻²⁰⁰ in the presence of recombinant Slc1, but not Slc2 or Scc1 indicating a specific interaction with Slc1, but not the other chaperones. HIS₆-TARP¹⁻²⁰⁰ was present in all of the co-immunoprecipitation samples as evidenced by its presence in the flow-through fraction, therefore the lack of interaction with Slc2 and Scc1 was not due to an absence of TARP in the combined lysates. I can also eliminate excess protein forcing an interaction because the low level of Imidazole (20 mM) in the wash buffer in combination with the multiple wash steps should eliminate any non-specific interactions. One minor caveat of this approach is that mixing of separately expressed, prefolded proteins may not be ideal for studying their native interaction.
Figure 7. Pull-down of TARP by chlamydial chaperones from singly expressed lysates. Immunoblots of flow-through (FT), first wash (W1), final wash (FW), and elution (E) fractions following M2 Anti-FLAG co-immunoprecipitation of Slc1-FLAG, Slc2-FLAG or Scc1-FLAG in the presence or absence of His₆-TARP¹-²₀₀ from laboratory E.coli lysates following individual expression were detected using anti-FLAG and anti-His₆ antibodies, respectively.
To confirm the observed Slc1 binding to HIS$_6$-TARP$^{1-200}$, I performed the reverse co-immunoprecipitation reaction in which TARP was immobilized to Ni-Nta magnetic beads and monitored the different fractions for Slc1-FLAG content by Western blot (Figure 8). In this experiment, lysate from a BL21 star *E. coli* strain that co-expresses Slc1-FLAG and HIS$_6$-TARP$^{1-200}$ was incubated with Ni-Nta magnetic beads. Fractions were processed and detected as stated previously. As expected, I found that the Ni-Nta magnetic beads when co-expressed with HIS$_6$-TARP$^{1-200}$ retained high levels of recombinant Slc1, but none was retained in its absence.

In addition, I tested whether co-expression of Scc1 or Slc2 with TARP enhanced their interaction (data not shown) and I was unable to detect any Scc1 or Slc2 in the elution fraction, giving confirmation that TARP interacts specifically with the putative chaperone Slc1. I also observed a relatively higher level of TARP retained in the reverse co-immunoprecipitation experiment that utilized lysates from *E. coli* co-expressing both TARP and Slc1.
Figure 8. Pulldown of chaperones by TARP following coexpression

Immunoblots of flow-through (FT), final wash (FW), and elution (E) fractions following Ni-Nta immunoprecipitation of His<sub>6</sub>-TARP<sup>1-200</sup> from laboratory E. coli lysates following expression of TARP<sup>1-200</sup>-His<sub>6</sub> or Slc1-FLAG alone, or after co-expression as indicated. Slc1-FLAG and TARP<sup>1-200</sup>-His<sub>6</sub> were detected using anti-FLAG and anti-His<sub>6</sub>, respectively.
To confirm independently the Slc1-TARP interaction demonstrated by co-immunoprecipitation, the bacterial two-hybrid technique was used[148, 149]. The N-terminus of chlamydial TARP was fused to the C-terminus of the *Bordatella pertussis* adenylate cyclase (cya) 18-fragment (Cya18-TARP) in the vector pUT18C (Fig 9). Each of the chlamydial chaperones, including the recently described Mcsc, which has a chaperone-like activity, was cloned into the vector pKT25 (Figure 9) and expressed as C-terminal fusions to the 25 fragment of cya (Cya25-Slc1, Cya25-Scc1, Cya25-Mcsc, and Cya25-Slc2). The TARP and chaperone inserts were amplified by PCR and ligated into the XbaI and Kpnl sites of the pUT18C and pKT25 (Figures 10 and 11) vectors, respectively. The ligated vectors were transformed into BL21 star and transformants were checked for the correct inserts by colony PCR followed by DNA sequencing (Fig 10 and 11).
Figure 9. Diagram of the pUT18C and pKT25 β-lactamase fusion vectors. (Top) Diagram of the pUT18C vector. (Bottom) Diagram of the pKT25 vector. Colony PCRs of ampicillin-selected pUT18C-TARP00 transformants separated on a 1% Agarose gel.
Figure 10. Cloning of the N-terminus of TARP into pUT18C.
(Top) PCR of the TARP200aa insert with XbaI and KpnI sites run out on a 1.2% Agarose gel. (Bottom) Colony PCR of the ampicillin-selected pUT18C-TARP200aa transformants run out on a 1.2 % Agarose gel.
Figure 11. Confirmation of pKT25-chaperone transformants.
Kanamycin-selected pKT25 transformants were checked for their chlamydial chaperone inserts. Colony PCR for SICl (Top left), Msc (Top right), Scc1 (Bottom left) and CT663 (Bottom right) were separated on 1.2% agarose gels.
Figure 12. Diagram of the *E.coli* 2-hybrid system.
Co-expression of the T18 and T25 fusion proteins in adenylate cyclase-deficient (cya) *E.coli* yields two scenarios: 1. (Left) The fusion proteins do not interact and the cya toxin is not reconstituted. No cyclic-AMP (cAMP) is formed and the lactose and maltose operons are not transcribed. 2. (Right) The fusion proteins interact, bringing together the T18 and T25 fragments and reconstituting the toxin. Cya mediates the formation of cAMP, and it further interacts with the catabolite activator protein (CAP) to co-activate transcription of lactose and maltose operons.
These fusion vectors were then co-expressed in the cya deficient *E. coli* strain DHM1. In this bacterial 2-hybrid assay, interaction between fusion proteins results in the reconstitution of the adenylate cyclase (cya) enzymatic domain and catalyzes the conversion of ATP to (cAMP). Available cAMP, along with the catabolite activator protein (CAP) co-activates the maltose and lactose operons (Fig 12). Activation of the lactose operon leads to Beta-galactosidase (β-gal) production, which cleaves the colorimetric substrate X-gal to yield blue colonies on LB + x-gal + IPTG plates. IPTG was included in the plates to relieve repression of the endogenous lactose operon by lacI. Activation of the maltose operon in colonies growing on McConkey + 1% maltose plates leads to a color change of the phenol red in the agar to fuchsia in the presence of acidic maltose degradation products [148, 149]. This bacterial 2-hybrid system gives quick, reproducible results on indicator plates, and protein-protein interactions can be quantified in a β-galactosidase reporter assay. I was able to detect an interaction between Cya18-TARP and Cya25-Slc1 by blue and fuchsia colonies on X-gal and McConkey plates, respectively (Figure 13). Bacteria co-expressing Cya18-TARP and Cya25-Scc1, Cya25-Msc, or Cya25-Slc2 were similar to the negative control, and did not produce an obvious color change to blue or fuchsia on indicator media.
Figure 13. Bacterial 2-hybrid interactions between TARP and chaperones on indicator plates.
LB + X-gal + IPTG (left) or McConkey + maltose (right) agarose plates indicate interaction between co-expressed Cya15 or Cya18 fused chlamydial chaperones in the cya deficient *E. coli* strain DHM1. Upregulation of the lactose and maltose operons results from fusion-protein interaction and is indicated by blue or fuchsia bacteria. These results are representative of multiple trials.
The results obtained from colonies on indicator plates were confirmed and quantified by monitoring β-galactosidase activity as previously described. Briefly, I permeabilized co-expressing DHMI strains and measured the β-gal-mediated conversion of ONPG to the colorimetric molecule ONP. This measurement of β-gal activity was converted to Miller Units in order to take into account the cell number and length of time for color development in contrast to the indicator plate method that does not take bacterial cell numbers into account. The background level of interaction in the Cya_{18}-Empty/ Cya_{23}-Empty co-expressing strain was subtracted from the Miller Units for each sample.

Extracts from DHMI strains co-expressing Slc1 and TARP indicated 3-fold higher level of β-galactosidase production compared to the strain containing Slc2 and TARP (Figure 14). Both the Slc2/TARP and Scc1/TARP co-expressing DHMI extracts contained roughly 5-fold less β-galactosidase than the Slc1/TARP coexpressing strain. These results confirm the findings for a TARP-Slc1 interaction shown on indicator plates and in the co-immunoprecipitation assay.
Figure 14. $\beta$-galactosidase activity of TARP:chaperone bacterial 2-hybrid interactions.
The relative beta-galactosidase activity of *E.coli* DHM1 strains co-expressing Cya$^{18}$-TARP$^{1-200}$ and Cya$^{25}$-Sle1, Cya$^{25}$-Sle2, Cya$^{25}$-Sec1 or Cya$^{25}$-Msc. Activity is expressed as Miller Units after subtraction of the background Cya$^{25}$-Empty: Cya$^{18}$-Empty interaction. Data is representative of multiple trials.
Interaction of Slc1 with TARP from other Chlamydiae.

Slc1 is highly conserved between chlamydial species, but there is a complete lack of amino acid similarity between the N-termini of different chlamydial TARPS. Alignment of the secondary sequence of the N-terminal 200 amino acids of TARP from *C. trachomatis* L2, *C. muridarum* (MoPn), *C. caviae* (GPIC), and *C. pneumoniae* with the bioinformatics program ClustalW indicated conserved and semi-conserved features among the sequences (Figure 15). Several hydrophobic sites (in red) were spatially conserved between TARPs, and closer inspection reveals that these hydrophobic sites are flanked by conserved threonines and serines. However these features are conserved only through the N-terminal 100 amino acids, indicating a possible chaperone binding site within these TARPs. This alignment led us to believe that Slc1 from the L2 serovar could possibly interact with the other chlamydial TARPS.

I first chose to look at the localization of putative chlamydial chaperones to full length *Chlamydia caviae* (GPIC) TARP due to its ability to form aggregates in mammalian cells. Full length TARP contains C-terminal actin-nucleating domains that are prone to aggregation and forms a characteristic string-of-pearls pattern when over-expressed in mammalian cells. I was able to acquire the GPIC-GFP-TARP vector encoding a C-terminal GFP (green fluorescent protein) from the Hackstadt lab at Rocky Mountain Laboratories. The three putative chlamydial chaperones Slc1, Scc1, and CT663 were PCR-amplified from L2 genomic DNA and subcloned into the Invitrogen pENTR/D/TOPO entry vector (Figure 16). The attL1 and attL2 recombination sites on this vector allow for recombination into a number of Destination vectors with the homologous sites attR and attB. For mammalian expression with an N-terminally fused
Mcherry protein I recombined the chaperones from the entry vector into pDEST-N-Mcherry (Figure 16). This allowed us to look for colocalization of the red chaperones with green aggregates of GPIC TARP.

The vectors GPIC-GFP-TARP and Mcherry-gus, Mcherry-Slc1, or Mcherry-Slc2 were co-transfected into Cos-7 cells and the co-localization of recombinant proteins was visualized by confocal microscopy (Figure 17). While the host cytosolic protein, gus localized some with TARP aggregates, this is probably just due to bleed-through from the green channel into the red channel. Slc2 localized to the TARP aggregates to the same extent as gus indicating a negative result for interaction. However, Slc1 overwhelmingly colocalizes with TARP aggregates in this assay. Of the two putative chaperones, only Slc1 was specifically recruited to TARP, confirming it as the specific type III chaperone for TARP and indicating that Slc1 can interact with the N-terminus of TARP from multiple chlamydial species.
Figure 15. Amino acid alignment of the N-terminus of TARP from different pathogenic *Chlamydiae*.

The N-terminal 200 amino acids of TARP from *C. caviae* (GPIC), *C. pneumoniae* (CPn), *C. trachomatis* (L2), and *C. muridarum* (MoPn) were aligned with clustalW. Residues in red indicate regions of conserved hydrophobicity, while light blue indicates conserved residues.
Figure 16. Diagram of the recombination vectors used to express chaperones in mammalian cells.
Figure 17. Localization of Slc1 to full-length C. caviae TARP aggregates
100X magnified images from confocal microscopy reveal colocalization of GPIC TARP and Slc1 in cotransfected Cos-7 cells. Cells were co-transfected with GPIC-GFP-TARP and Mcherry-gus (host cytosolic protein), Mcherry-Slc1, or Mcherry-Slc2.
I continued to test the two other chlamydial TARPs for their ability to bind L2 Slc1. Interactions between *C. pneumoniae* and *C. muridarum* TARP with Slc1 were tested by co-immunoprecipitation with M2 Anti-FLAG agarose followed by immunoblotting. Addition of Slc1 and Cpn or Mopn TARP expression lysates allowed for their interaction which was confirmed by the presence of both of these TARP species in the Slc1-containing elution fractions (Figure 18).

**Figure 18. Pulldown of the N-terminus of *C. muridarum* and *C. pneumoniae* TARPs with Slc1.**

Immunoblots of the lysates (Lys), flow-through (FT), final wash (FW), and elution (E) fractions from M2 Anti-FLAG immunoprecipitated recombinant Slc1 and the N-terminal 200 amino acids of *C. muridarum* (MoPn) or *C. pneumoniae* (C.Pn) TARPs are probed with their respective antibodies.
CHAPTER IV
IDENTIFICATION AND CHARACTERIZATION OF CHLAMYDIAL CHAPERONE: CHAPERONE INTERACTIONS

Specific interaction between the chaperone Slc1 and TARP prompted us look deeper into the relationship of this chaperone: effector interaction. Translocation chaperones typically exist as either as a homodimer or a heterodimer prior to their interaction with their cognate effectors. Since our negative results for interaction between Slc2 or Scc1 and TARP only included possibly homodimers, I felt it necessary to see if the formation of Slc2-Slc1 or Scc1-Slc1 heterodimers could contribute to TARP binding. I chose to characterize all possible chlamydial chaperone dimers to reveal any interactions that could be important for binding with TARP and other chlamydial effectors.

Discovery of chlamydial chaperone: chaperone pairs

Therefore, I investigated the ability of Slc1 to form homodimers with itself or heterodimers with either Slc2 or Scc1 using both co-immunoprecipitation and bacterial two-hybrid techniques. As before, I are taking advantage of separate recombinant tags on the proteins to purify protein complexes and to detect the protein composition of these complexes by Western blot analysis. I first wanted to detect whether Slc1 can interact
with itself or with other chaperones so I employed M2 Anti-FLAG agarose and Anti-S-tag resin to purify chaperone dimers.

By co-immunoprecipitating Flag-tagged S1c1 (S1c1-F) with S-tagged S1c1 (S1c1-S) on M2 Anti-FLAG beads, I was able co-elute them from the beads and thus demonstrate that S1c1 can multimerize with itself (Figure 19). However I could not detect any co-elution of S1c2-F with S1c1-S following immunoprecipitation on M2 agarose, indicating a lack of interaction between the two proteins. Similarly, I was also able to detect an interaction between S1c1-S and S1c1-F by co-immunoprecipitation with S-Rek resin, but not between Scc1-S and S1c1-F (Figure 20). In both co-immunoprecipitation assays detection of the S1c1: S1c1 interaction was minimal. This is not surprising considering the fact that many known type III secretion chaperones in other bacteria preferentially form dimers in solution, leaving few monomers around for potential dimer formation in our assay.
Figure 19. Pulldown of FLAG-tagged Slc1 and Slc2 in the presence of S-tagged Slc1. Immunoblots of the FT (flow-through), W1 (first wash), FW (final wash), and E (elution) fractions following M2 Anti-FLAG co-immunoprecipitation of FLAG-tagged chaperones (Slc1-F or Slc2-F) in the presence or absence of S-tagged Slc1 (Slc1-S) from co-expressed BL21 *E. coli* lysates were detected with αFLAG or αS-tag antibodies, respectively.
Figure 20. Pulldown of Slc1 in the presence of Scc1.
Immunoblots of FT, FW, and E fractions following S-Rek resin immunoprecipitation of Scc1-S in the presence or absence of Slc1-F were detected with antibodies against their respective tags.
Therefore I continued to investigate these interactions by E.coli 2-hybrid analysis, testing all possible chaperones: chaperone interactions. I was able to detect interactions between Cya\textsubscript{18}-Sle\textsubscript{1}:Cya\textsubscript{25}-Sle\textsubscript{1}, Cya\textsubscript{18}-Mesc: Cya\textsubscript{25}-Mesc, and Cya\textsubscript{18}-Sle\textsubscript{2}:Cya\textsubscript{25}-Sccl by blue and fuchsia colonies on X-gal and McConkey plates, respectively (Figure 21). The Mesc:Mesc and Sle\textsubscript{2}:Sccl interactions had been previously identified in a recent yeast-two hybrid study [49]. I tested the expression of the Cya\textsubscript{18}-Sccl and Cya\textsubscript{25}-Sle\textsubscript{2} recombinant proteins to try to understand why the reciprocal 2-hybrid didn’t indicate interaction, and was able to detect expression of both by coomassie staining (data not shown). All other chlamydial chaperone: chaperone interactions were similar to the negative control, and did not produce an obvious color change to blue or fuchsia on indicator media.
Figure 21. Bacterial 2-hybrid interactions between chlamydial chaperones. LB + X-gal + IPTG (left) or McConkey + maltose (right) agarose plates indicate interaction between co-expressed Cya15 or Cya18 fused chlamydial chaperones in the cya deficient *E. coli* strain DHM1. Upregulation of the lactose and maltose operons results from fusion-protein interaction and is indicated by blue or fuchsia bacteria.
Quantitative analysis of the β-galactosidase activity of the Cya_{18}-chaperone / Cya_{25}-chaperone co-expressing strains indicated a very strong interaction between Cya_{18}-Mcsc and Cya_{18}-Mcsc, whereas the Slc1:Slc1 and Slc2:Scc1 interactions were stronger than the background chaperone: chaperone interactions (Figure 22). Other chaperone: chaperone interactions were higher than background levels but still lower than the interactions that had been previously confirmed on indicator plates. Taken together, these bacterial 2-hybrid data indicate a novel Slc1-Slc1 interaction and confirms the previous Mcsc-Mcsc and Slc2:Scc1 interactions that were discovered in a recent yeast 2-hybrid study. All together the co-immunoprecipitation and bacterial 2-hybrid results indicate that Slc1 forms homodimers, but not heterodimers. Given our previous findings that TARP does not interact with Scc1 or Scc2, it is likely that only Slc1 homodimers interact with TARP.
Figure 22. Quantitative analysis of bacterial 2-hybrid chaperone: chaperone interactions.

The relative beta-galactosidase activity of *E.coli* DHM1 strains co-expressing the recombinant Cya\textsuperscript{18}-chaperones and Cya\textsuperscript{25}-chaperones. Activity is expressed as Miller Units after subtraction of the background Cya\textsuperscript{25}-Empty: Cya\textsuperscript{18}-Empty interaction. Data is representative of multiple trials.
**Determination of the molecular composition of Scl homo-complexes**

I expect the apparent Scl1:Scl1 interactions shown by co-immunoprecipitation and bacterial 2-hybrid analysis was a result of the formation of homodimers, however the rare case of trimer and tetramer forms of type III chaperones have also been observed to interact with type III effectors in other bacteria. I chose to detect all native Scl1 complex forms by stabilizing their interaction with a crosslinker and separating the complexes on an SDS-PAGE gel. Since I wanted to crosslink M2-Anti-FLAG agarose purified Scl1 in PBS, I chose to use a hydrolysable crosslinker that could be reduced in the presence of β-mercaptoethanol (BME), but not in its absence. In addition I wanted to use a crosslinker that would react with the primary amines (mostly N-termini) of purified proteins. The compound DTSSP binds cysteines and primary amines at both ends and has been modified from its original non-hydrolyzable form, DSP, to be able to dissolve and react in solution. (Figure 23) Crosslinking of purified FLAG-Scl1 with the hydrolysable DTSSP reagent revealed the ability of this protein to form a dimer in the absence of the crosslinker; this dimeric complex was able to withstand denaturation in the SDS-PAGE gel (Figure 23). In fact, purified Scl1 that was reduced (BME) and heated to 90°C, but not crosslinked still showed faint dimerization at the expected ~40 kDa size. The addition a small concentration of crosslinker (0.25 mM) revealed a higher molecular weight Scl1 complex with a size (~60 kDa) consistent with trimeric species. Since non-crosslinked, non-reduced purified Scl1 readily migrated to a dimer-sized position on a non-reducing SDS-PAGE gel it is clear that Scl1 forms stable dimers.
Figure 23. Stabilizing Scl1 complexes by the crosslinker DTSSP.
Multimerization of Scl1. (Top) Chemical structure of the crosslinker 3, 3’-Dithiobis (sulfosuccinimidylpropionate (DTSSP). (Bottom) An immunoblot of β-mercaptoethanol (βME), mock-treated (-), or 0.25 mM DTSSP crosslinked (+) purified Scl1 was probed with αFLAG antibody following separation in 12% SDS-PAGE gel under non-reducing conditions.
While I was able to detect dimerization of SIC1 by crosslinking, the presence of a trimeric species required independent confirmation using gel filtration chromatography, which is a more gentle analytical technique that did not require crosslinking. Upon purification of SIC1-FLAG with M2 Anti-FLAG agarose, I loaded 50 µl of 100 ng/ml purified SIC1 into a Superdex 200 10/300 GL column in PBS. The column washes the protein through sepharose gel with additional PBS. Larger species get stuck in the gel sooner, and thus get eluted from the column first. Prior to elution of 500 µl fractions along the length of the gel, the relative abundance of the protein is detected at a wavelength of 280 nm.

The relative abundance of protein in each of the eluted fractions is shown on the y-axis, while the relative size of each species (peak) is indicated by the elution fractions on the x-axis. Each elution fraction was monitored for SIC1 content by Western blot. Higher molecular weight complexes are eluted earlier. Only one prominent peak was detected for the SIC1 complex and when compared to known molecular weight standards in PBS, it was determined to be ~40 kDa in size (Figure 24). SIC1 was detected in the TCA-precipitated elution fractions under the dimeric peak by immunoblotting. Crosslinking and gel filtration of purified SIC1 indicates that it preferentially forms homodimers, indicating that this is the form it likely takes when interacting with TARP.
Figure 24. Detection of native Slc1 species by gel filtration chromatography
The relative abundance of native M2-purified Slc1 complexes were detected and eluted in 500 ul fractions by gel filtration chromatography. An immunoblot of trichloracetic acid (TCA)-precipitated fractions was detected with αFLAG antibody.
CHAPTER V

FUNCTIONAL DETERMINATION OF SLC1 IN THE TRANSLOCATION OF TARP BY HETEROLOGOUS TYPE III SECRETION SYSTEMS

TARP: SIC1 complexes indicate the expected 2:1 stoichiometry of type III chaperone: effector interactions

I was able to detect a complex formed between SIC1 and TARP by detection of a shift in complex size between purified recombinant TARP and TARP co-expressed in the presence of SIC1 on a silver-stained native gel and on its corresponding 6XHis-probed immunoblot. Since native gels separate proteins based on their shape and size, it is difficult to extrapolate an approximate molecular weight of the complex (Figure 25).

To gain some insight into the composition of the complex formed by TARP and SIC1, a similar chromatography approach was used to analyze purified HIS6-TARP1-200 and the HIS6-TARP1-200/SIC1-FLAG complex. The chromatography conditions produced readily identifiable peaks for HIS6-TARP1-200 from SIC1-FLAG quite easily (Figure 26).

To determine the approximate size of the protein complex formed by TARP and SIC1, the Ni-Nta agarose eluates from lysates containing co-expressed HIS6-TARP1-200 and SIC1-FLAG were subjected to gel filtration chromatography. Ni-Nta purified protein

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from the lysate containing HIS₆-TARP₁⁻²₀₀ was used as a reference control. The Slc1 and TARP content of the co-expressed elution fractions was detected with antibodies against their respective tags by Western blot. Since the co-expressed complex was purified with Ni-Nta agarose, I expect the sample to contain both monomeric TARP and TARP in a complex with Slc1. Here I see that TARP-containing elution fractions correspond to a peak between 13.9 kDa and 29 kDa and a peak >75 kDa, while Slc1 is present only in the elution fractions corresponding to the >75 kDa peak. These sizes are in agreement with monomeric TARP and a 1:2 complex of TARP to Slc1. Taken together with our Slc1 dimerization data, the results indicate that a dimer of the Slc1 chaperone forms a complex with one molecule of HIS₆-TARP₁⁻²₀₀ when co-expressed. In order to detect the Slc1-TARP complex by chromatography, co-expression of Slc1 and TARP appeared to be necessary because the addition of pre-purified components failed to produce a similar >75 kDa-peak (Figure 27).
Figure 25. Native gel shift of TARP: Slc1 complex
Native purified protein complexes are shown on both silver stain and Western blot. All protein species from the purified fractions are displayed on the silver-stained Native PAGE gel on the left. TARP-containing complexes are detected in the identical purified fractions with α6XHIS antibody on the immunoblot on the right. Arrows indicate the complex formed under co-expressing conditions.
Separation and detection of co-expressed Slc1: TARP complexes. The relative abundances of native NiNta-purified TARP\textsuperscript{1-200} complexes alone or co-expressed with Slc1 were detected and eluted in 500 ul fractions (top) by gel filtration chromatography. Immunoblots of the TCA-precipitated elution fractions were detected with α6XHIS and αFLAG antibodies (bottom).

Figure 26. Detection of TARP: Slc1 native complex (coexpressed).
Figure 27. Detection of TARP: Slc1 native complex (Combined).
The separation and detection of M2-purified Slc1 (Red), Slc1 +TARP (Blue), or Ni-Nta-purified TARP (Green) was achieved by gel filtration chromatography. The peaks on the chromatogram correspond to the molecular weight standards (top). The relative abundances at A280 are indicated by the y-axis and the corresponding immunoblots of the 500μl fractions on the x-axis were TCA-precipitated and detected with α6XHIS and αFLAG antibodies.
**Translocation of TARP by heterologous type III secretion systems**

While the biochemical data suggests that Sic1 interacts with TARP in a fashion typical of type III chaperone: effector complexes, I desired to know what role Sic1 plays in TARP translocation by the chlamydial type III secretion apparatus. Unfortunately, *Chlamydia*’s lack of a genetically tractable system prevented us from directly testing Sic1’s role in TARP’s translocation from the chlamydial EB into mammalian epithelial cells. Fortunately, I was able to express our recombinant chlamydial proteins in the pathogenic bacteria *Yersinia enterocolitica*, *Enteropathogenic E. coli* (EPEC), and *Salmonella typhimurium* and detect TARP’s translocation through their heterologous type III secretion apparatuses into Hela cells.

To accomplish this, I took advantage of an existing translocation assay that involved expressing possible effectors in the pCX340 vector that C-terminally fuses them to enzymatically active β-lactamase, and the chaperones were cloned into an arabinose-inducible expression vector pBAD18-Cm (Figure 28). The chaperones were amplified from *C. trachomatis* L2 genomic DNA by PCR (Figure 29) and cloned into the KpnI and XbaI sites on the vector. A KpnI site was added downstream of the TARP$^{1-200}$ open reading frame in the pET200-TARP$^{1-200}$ vector by SLIM PCR (Figure 30), while TARP was cloned into the NdeI and KpnI sites of pCX340 (Figure 31).
Figure 28. Diagrams of the bacterial expression vectors pBAD18-Cm and pCX340.
Figure 29. Cloning chlamydial chaperones into the pBAD18-Cm vector
(Top) Insert PCRs of CT043 and Mcsc open-reading frames with flanking XbaI and Scal sites. (Bottom) Colony PCRs from chloramphenicol-selected pBAD18-chaperone transformants.
Figure 30. Adding a KpnI site to the pET200-TARP200 construct by SLIM PCR.
(Top) Site-directed Lipase-Independent Mutagenesis (SLIM) based PCR was performed to add a KpnI site downstream of the TARP200 open reading frame in the pET200-TARP200aa vector and the PCR product is displayed here on a 0.7% Agarose gel.
(Bottom) pET200-TARP200aa-KpnI plasmid DNA prepared from kanamycin-selected BI21star E.coli transformants were checked for the presence of a KpnI site by restriction digest as displayed here on a 0.7% Agarose gel.
Figure 31. Cloning TARP into the pCX340 vector.
(Top) The TARP200aa open reading frame was digested with KpnI and NdeI from the pET200-TARP200 vector and pCX340 was double digested as well prior to ligation as indicated here on the 1% Agarose gel. (Bottom) Colony PCR was performed on tetracycline-selected transformants and separated on a 1.2% Agarose gel.
Upon contact with mammalian cells, the bacteria will translocation their type III effectors into the host cell. Treatment of the HeLa cell monolayers with CCF2/AM, a coumarin molecule bound to hexane by a β-lactam ring, following infection allows for an especially innovative fluorescence assay that takes advantage of the fact that the intact substrate fluoresces at 520 nm and the cleavage product, coumarin, fluoresces at 450 nm (after excitation at 410 nm). The presence of TEM1-fused bacterial effectors in the host cell is reflected in the cleavage of the β-lactam ring to release the coumarin molecule, which I can measure its fluorescence at 450 nm. Thus I can detect fluorescence resonance energy transfer, (FRET) of the intact molecule in the green spectrum at 520 nm, while I can detect cleavage of the substrate as a loss of FRET in the blue spectrum at 450 nm. A diagram of the basis of this assay is shown in Figure 32.
In the bacterial cell, β-lactamase fused TARP is expressed from the pCX340-TARP vector. Two hours post-infection of HeLa cell monolayers, cultures are treated with CCF2/AM, a molecule that undergoes FRET (Fluorescence Resonance Energy Transfer) when excited at 410 nm and emits fluorescence in the green spectrum at 450 nm. Translocation of β-lactamase fused TARP by the bacterial Type III Secretion System (T3SS) into the eukaryotic cell and subsequent cleavage of the β-lactam ring portion of CCF2/AM results in a loss of FRET and an alternate emission of fluorescence in the blue spectrum at 450 nm.
The *Yersinia* *ysc* family type III secretion system is encoded on its virulence plasmid[63]. The type III secretion system from *Yersinia pseudotuberculosis* has been previously used to express and secrete chlamydial translocators and effectors, including TARP, into low-calcium media[38, 41, 150]. I was also able to detect TARP secretion into media by wild-type *Yersinia enterocolitica* similar to levels of secretion of the *Yersinia* effector, YopH (Figure 33).

However, *Yersinia* strains have not been tested for their ability to translocate chlamydial proteins into eukaryotic cells. I chose to utilize the type III secretion system of *Yersinia enterocolitica* to determine the role of Slc1 in the secretion and translocation of TARP. Based on previous findings, I expect that some TARP will be secreted and possibly translocated in the absence of its cognate chaperone[42]. However, our interaction studies indicate that the chaperone Slc1 binds TARP in a fashion typical of export chaperones convincing us that Slc1 will enhance TARP’s interaction with the type III secretion apparatus and thus increase the amount TARP secreted or translocated by the *Yersinia enterocolitica* type III secretion system.

Wild type and type III ATPase (YscN) knockout strains of *Y. enterocolitica* expressing TARP ± Slc1 were allowed to infect HeLa cell monolayer for 2 hours prior to CCF2/AM treatment. A loss of FRET (β-lactamase-mediated cleavage) to FRET (no cleavage) ratio was used to detect translocation. Translocation of the positive control, YopH was detected at a fluorescence ratio of ~0.8, and the negative control empty vector indicated at ratio of less than 0.1 (Figure 34). I was able to detect Slc1-mediated translocation of TARP as indicated by the ratio of 0.67 by co-expressing bacteria and 0.43 from TARP-only expressing bacteria. Because the YscN knockout strains were unable to translocate...
any effectors, I can be absolutely positive they are being translocated through the 

*Y. enterocolitica* type III secretion needle.

**Figure 33. Secretion of HIS₆-TARP¹⁻²⁰⁰ by the Yersinia enterocolitica type III secretion system.** Immunoblots of TCA-precipitated *Yersinia enterocolitica* culture supernatants (S) and whole cell pellets (P) under type III secretion inducing conditions (-Ca) or repressing conditions (+Ca) were probed with β-lactamase mAb. Secretion was tested by wild-type (WT) or type III secretion knockout (dYscN) *Yersinia enterocolitica* strains.
Figure 34. Translocation of TARP by *Yersinia enterocolitica*.
The fluorescence ratio of cleaved CCF2/AM substrate (450 nm) to uncleaved substrate (520 nm) indicates the relative translocation of β-lactamase fused effectors by the *ysc* type III secretion system of *Y. enterocolitica*. Wild-type (WT) and the Type III ATPase *yscN* knock-out (T3KO) strains bearing the β-lactamase fusion vector pcx340 with or without pBAD-S1c1 were allowed to infect HeLa cells prior to treatment with CCF2/AM substrate and fluorescence detection.
I next used the Enteropathogenic *Escherichia coli* (EPEC) to test the translocation of TARP. To date there has been no evidence of secretion of chlamydial proteins by the EPEC type III secretion of TARP. However, EPEC is known to translocate the type III effector Tir that has very similar function *in vivo* as TARP. They both induce massive cytoskeleton rearrangement to ensure close attachment and survival of the bacteria. Furthermore alignment of type III secretion chaperones has revealed that Tir’s cognate chaperone CesT has very similar secondary structure as Sic1. Therefore I thought it likely that the EPEC type III secretion could translocate TARP in the presence of its cognate chaperone Sic1.

EPEC attaches to epithelial cells and begins to translocate its effectors as soon as 30 minutes post-infection. I was able to detect translocation of the EPEC effector EspH as a fluorescence ratio of 1.5 after 1 hour of infection. Using the same infection conditions I was unable to detect any chaperone-dependent or -independent translocation of TARP. (Figure 35), and lengthening the duration of infection did not change this negative result (data not shown).
**Fig 35. Translocation of TARP by Enteropathogenic *E. coli***

The fluorescence ratio of cleaved CCF2/AM substrate (450 nm) to uncleaved substrate (520 nm) indicates the relative translocation of β-lactamase fused effectors by the type III secretion system of Enteropathogenic *E. coli* (EPEC). Wild-type (WT) and the Type III ATPase *escN* knock-out (T3KO) EPEC strains bearing the β-lactamase fusion vector pcx340 with or without pBAD-Slc1 were allowed to infect HeLa cells prior to treatment with CCF2/AM substrate and fluorescence detection.
Salmonella typhimurium contains two functional type III secretion systems that are temporally regulated and are responsible for translocating distinct sets of effectors into epithelial cells. The SPI-1 (Salmonella pathogenicity island-I) type III secretion system translocated effectors upon contact with host cells to induce uptake of Salmonella into endocytic vesicles. After complete engulfment of the bacterium, it switches to secretion of effectors involved in maintaining its Salmonella-containing vesicle (SCV) and in redirecting host cell vesicle trafficking to ensure its intracellular survival[63]. Because the SPI-1 T3SS is responsible for secreting proteins involved in invasion of eukaryotic cells, it is an attractive choice for testing TARPs translocation. However, the SPI-2 system resembles that of Chlamydia because of its ability to secrete from an intracellular vesicle indicating that it also is worth trying as a heterologous secretion system.

As indicated by the fluorescence ratios of infected monolayers, the Salmonella SPI-1 secretion system was unable to translocate TARP under any conditions (Figure 36), but the SPI-2 system seemed to minimally translocate TARP in a chaperone-independent manner, but achieved efficient translocation of TARP in the presence of its cognate chaperone Slc1 (Fig 34). However the high ratios for the β-lactamase only (pcx340-Empty) translocation casts some doubt on the reliability of this assay under SPI-2 conditions. Furthermore, this assay has remained untested for translocation during intracellular growth (8-12 hours post-infection); instead its use has been for detection of translocation of substrates immediately after attachment (1-3 hours post-infection).
Figure 36. Translocation of TARP by Salmonella typhimurium SPI-1 and SPI-2
The fluorescence ratio of cleaved CCF2/AM substrate (450 nm) to uncleaved substrate (520 nm) indicates the relative translocation of β-lactamase fused effectors by the SPI-1 and SPI-2 type III secretion systems of Salmonella typhimurium. Wild-type (WT) Salmonella strains bearing the β-lactamase fusion vector pcx340 with or without SIC1 were allowed to infect HeLa cells prior to treatment with CCF2/AM substrate and fluorescence detection. SPI-1 mediated translocation was detected 2 hours post-infection, while SPI-2 mediated translocation was detected 10 hrs post-infection.
The differences in TARP translocation by different heterologous type III secretion systems is likely due to the inability of the cytoplasmic components from other type III secretion systems to recognize the TARP-Slc1 complex. To gain insight into this possibility, I analyzed the relatedness of the chlamydial type III ATPases CdsN and FliI with the type III ATPases from *Salmonella*, *Yersinia*, and EPEC. The amino acid sequences of these paralogs were aligned with ClustalW and the resulting alignments were organized into unrooted dendograms (Figure 37). The CdsN tree indicates that *Yersinia* type III ATPase YscN and the SPI-2 ATPase SsaN are similar to the non-flagellar chlamydial ATPase, while the ATPases from the SPI-1 and EPEC are less closely related, supporting our findings that *Yersinia* type III system and the *Salmonella* SPI-2 system could both recognize the TARP: chaperone complex and subsequently translocate TARP into HeLa cells. Furthermore, phylogenetic analysis indicates that the chlamydial flagellar type III ATPase FliI is more similar to the EPEC and SPI-1 non-flagellar ATPases than to relevant ATPases in *Y. enterocolitica* and the *Salmonella* SPI-2 systems.
Figure 37. Phylogenetic analysis of multiple type III components from bacterial pathogens.
Unrooted phylograms were created from ClustalW-generated multiple sequence alignments of the *C. pneumoniae* (cpn) and *C. trachomatis* (ctr) ATPases CdsN (top) and Flil (bottom) with the relevant ATPases from *S. typhimurium* (stm), and Enteropathogenic *E. coli* (ecg). Orthologs were identified for comparison by the Kyoto Encyclopedia of Genes and Genomes (KEGG).
### TABLE 5. STRAINS AND PLASMIDS USED IN THIS STUDY

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pBAD18  Arabinose-inducible promoter  Guzman (1995)
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pBAD-CT663  pBAD18 with CT663-FLAG insert  This study

**TABLE 5. PRIMERS USED IN THIS STUDY**

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98
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DISCUSSION

Many intracellular bacterial pathogens use specialized secretion systems to translocate effectors into eukaryotic cells and in turn these effectors recruit, modify, or hijack cellular components to the benefit of the bacterium. Effectors of bacterial type III secretion systems have been reported to be involved in remodeling the eukaryotic cytoskeleton, altering the immune response, hijacking trafficking vesicles, disrupting cell-to-cell junctions, and causing cytotoxicity[63].

Survival and pathogenesis of Chlamydia spp. is entirely dependent on its ability to invade mammalian epithelial cells and develop a unique niche that is conducive to intracellular growth and replication. Translocation of effectors by a type III apparatus is a major strategy for Chlamydia entry, inclusion formation, and survival in eukaryotic cells[46, 84, 85]. Recruitment of host kinases and actin, in addition to activation of Rac and actin-polymerization aid in endocytosis, and are functions attributed to the type III invasion associated effector TARP[42, 66, 67, 69, 132]. Chlamydia trachomatis urogenital, lymphogranuloma, and ocular strains synthesize TARP homologs that contain three functional domains: tyrosine-rich repeats for recruitment of host kinases and subsequent Rac activation, a proline-rich aggregating domain, and actin-nucleating WH2-like domains.
Detected and tyrosine-phosphorylated in host cells within minutes after cell contact, TARP translocation is essential for the efficient uptake of *C. trachomatis* EBs. TARP homologs in *C. pneumoniae*, *C. caviae*, and *C. psittaci* do not contain tyrosine-rich repeat domains, but they are still able to invade host cells. In addition the WH2-like domains are sufficient to nucleate actin in host cells, which are also important for infection[70, 71]. This redundancy between TARP domains indicates another possible function for *C. trachomatis* TARP tyrosine-phosphorylation during chlamydial infection. It has been suggested that the tyrosine phosphorylation may help ensure specific uptake pathways that prevent phagolysosomal fusion. This theory has been supported by the detection of TARP localized to the inclusion throughout the developmental cycle[42].

Given TARP’s pivotal roles in entry and intracellular survival, I considered possible type III secretion chaperones that would increase the efficiency of TARP translocation into host cells. I determined that the chlamydial chaperone Slc1 specifically interacts with TARP and mediates its translocation by the heterologous type III secretion system of *Yersinia enterocolitica*. Identification of putative type III related genes in bioinformatics studies identified three putative chlamydial effector chaperones: Slc1, SLC2, and Scc1. Since Class I effector chaperones share the specific secondary structure $\alpha-\beta-\beta-\alpha-\beta$, putative chlamydial chaperones were readily identified by *in silico* approaches. Detection of these three proteins in elementary bodies indicated that any one of them could be involved in maintaining TARP in a secretion-competent state for immediate translocation into host cells.
In other gram-negative bacteria, previously identified type III Class I effector chaperones demonstrated binding of their effectors adjacent to their N-terminal secretion signal, usually within the N-terminal 30-130 amino acid range.

**Identification of the TARP’s cognate secretion chaperone Slc1**

I used a direct co-immunoprecipitation approach for identifying a chlamydial chaperone that could interact with the N-terminal 200 amino acids of *C. trachomatis* L2 TARP. I was specifically able to co-immunoprecipitate Slc1 with TARP in two independent co-IP approaches. Separate expression of Slc1 and TARP in *E. coli* prior to incubation of their lysates with M2 Anti-FLAG agarose yielded an interaction between Slc1 and TARP, but co-expression of these proteins prior to co-immunoprecipitation greatly increased their ability to interact. This indicates to us that Slc1 may mediate folding of TARP or prevent its pre-mature interaction with other *E. coli* components when expressed in the same bacterium. However, neither co-immunoprecipitation technique was able to demonstrate an interaction between TARP and the other putative chlamydial chaperones Slc2 or Scc1.

One other explanation for the increased interaction between Slc1 and TARP during co-expression would be that Slc1 prevents TARPs degradation or solubility. The cognate chaperone SycE in *Yersinia* helps to solubilize its cognate aggregate-proned effector YopE[151], and similarly the EPEC chaperone CesT prevents the degradation of the effector Tir[152]. Despite the described ability of many known type III chaperones to solubilize and stabilize their effectors, I was not able to detect any changes in solubility...
or stability of TARP when expressed in the presence or absence of its chaperone (data not shown).

One caveat of detecting chaperone: TARP interactions in *E. coli* lysates, is that an *E. coli* protein may be mediating the interaction detected between Slc1 and TARP. It is difficult to refute this claim, because attempts to find an interaction between pre-purified TARP and Slc1 were unsuccessful. Because this possible *E. coli* factor did not mediate an interaction between TARP and Ssc1 or Ssc2, which are predicted to have extremely similar secondary and tertiary structures as Ssc1, I propose that it is highly unlikely for an *E. coli* factor to be so specific for Slc1 and TARP.

Our co-immunoprecipitation findings were confirmed by the specific interaction detected between Slc1 and TARP in an adenylate-cyclase (cya) based bacterial 2-hybrid system[148]. Because the readout for interaction of cya-fused target and bait proteins in this system was the cAMP-mediated up-regulation of the lactose and maltose operons, I was able to detect interactions in a variety of assays. Both X-gal and McConkey plates indicated a range of protein-protein interactions from low-affinity (chaperone-effector) to high affinity (leucine-zipper proteins). While the indicator plates allowed for obvious qualitative results of chaperone-effector interactions, they were not able to differentiate between slow and fast growing strains. If allowed to grow long enough, even the negative control strains would begin to change color on these indicator plates, therefore I had concerns that any slow-growing strains would not indicate obvious interactions.

To alleviate this problem as well as to quantitatively compare protein-protein interactions, I employed the β-galactosidase assay which not only takes into account the amount and time of ONPG conversion, but also the total number of bacteria in each
sample. Compared to co-expression with other chaperones tested in this assay, DHM1
*E.coli* co-expressing Slc1 and TARP converted 2-3 times as much ONPG to the
colorimetric substrate ONP, confirming the bacterial 2-hybrid results on indicator plates.

**Characterizing all possible chlamydial chaperone-chaperone interactions**

The tendency of type III chaperones to dimerize prior to interacting with their
effectors, led us to detect whether Slc1 tends to form homodimers or heterodimers. From
co-immunoprecipitation assays I was able detect Slc1 interactions with itself, but not with
the other putative chaperones Slc2 or Scc1. Since I singly expressed the chaperones prior
to adding the lysates to the M2 Anti-FLAG agarose or S-rek resin, it is likely that Slc1
dimers were formed early, leaving very little monomeric Slc1 to interact with other
chaperones. Instead of testing co-expression in the co-IP assays, I thought it best to test
co-expressed chaperone-chaperone interactions in the bacterial 2-hybrid system. In this
system I was able to confirm previous chlamydial chaperone-chaperone interactions as
well as confirm our Slc1-Slc1 interaction. Obvious interactions on indicator plates
included Slc1-Slc1, Mesc-Mesc, and Slc2-Scc1. However, I was not able to detect the
reverse Cya18-Scc1/Cya25-Slc2 interaction indicating that the dimer formed may only
allow interaction of the 18 and 25 fragments when the dimer is formed in the Cya18-Slc2/
Cya25-Scc1 orientation and not the other way around. Furthermore, I was able detect
expression of both 18-Scc1 and 25-Slc2; consequently the lack of bidirectional
interaction was not due to the constructs being problematic.

Quantitative analysis of chaperone-chaperone interactions in the bacterial 2-
hybrid strains indicated the same positive interactions as on indicator plates, but there
also seemed to be a low level of interaction between most of the chaperones. The Slc1-Slc1 interaction was only 2-fold higher and the Slc2-Scc1 interaction was only slightly higher than most of the chaperone-chaperone strains. I speculate that *Chlamydia* maintains a low level of interaction between type III class I chaperones to enable the bacterium to quickly acquire new type III effectors. This is supported by the fact that *Candidata Protochlamydia ameobophilia* maintains SycE-like Class I chaperones in the same gene order as in pathogenic *Chlamydia*, however the chaperones themselves have completely diverged in amino acid sequence[22, 29]. In addition, recent studies have identified species-specific effectors among chlamydial pathogens[84], indicating a mechanism for the quick selection of new type III effectors for adaptation to new host cell types must exist. It has been noted in other bacteria that there is little conservation in sequence or structure of type III effectors, despite the requirement that all effectors must be adequately unfolded to ensure secretion through the ~4 nM diameter needle[29, 59, 153]. Thus type III chaperones may be necessary to partially unfold disordered regions in effectors so that they may effectively interact with the type III ATPase[59, 153].

Akeda *et al.* maintain that the ATPase actually does the majority of unfolding effectors in addition to providing active (ATP-dependent) separation of effectors from cognate chaperones[59]. Recent crystallizations of chaperone: effector complexes support this theory that chaperones only act to locally unfold or reorder the effector at the chaperone binding domain (CBD). Whether this localized unfolding allows for the unfolding of the N-terminal secretion signal or makes the secretion signal more accessible to the ATPase is not clear.
Besides being unfolded, secretion of type III effectors requires an N-terminal secretion signal that is possibly recognized by the ATPase. However, this signal is highly divergent and efforts to predict secretion substrates based on their N-terminal amino acid composition has remained challenging. Recent studies have used machine-learning approaches that focused on the N-terminal 25 amino acids of known type III effectors to predict the probability that a protein is in fact an effector[25, 30]. Predicted secretion signals were mostly disordered, largely hydrophobic, often with polar-hydrophobic-polar residue patterns, and were serine and threonine-rich. One such prediction program identified 86 putative C. trachomatis effectors, 24 of which had already been experimentally confirmed to be type III effectors[30].

Considering the small number of possible chaperones, it is clear that not all of these effectors will require a chaperone for their translocation. Thus a major enigma in bacterial type III secretion is why some effectors require chaperones while others do not. In EPEC and Yersinia, a hierarchy of secretion is established in which effectors that have cognate chaperones are secreted preferentially to those without chaperones[62, 154, 155]. Since few chlamydial chaperone-effector interactions have been detected thus far, I have no indication as to whether such a hierarchy exists during infection, but it is easy to imagine how such a hierarchy could play a role in regulating Chlamydia's developmental cycle.

The recently discovered type III secretion chaperone Mcsc was not predicted to be a Class I chaperone by in silico analysis, but its association with effectors and ability to form homodimers in a yeast 2-hybrid screen[49] led us to test its ability to interact with TARP as well as its ability to form heterodimers in our bacterial 2-hybrid assays. I did
not detect any interaction between TARP and Mcsc, but the Mcsc-Mcsc co-expressing DHM1 strain had nearly 6 fold higher β-galactosidase activity than the Slc1-Slc1 interaction and as much as 10 fold higher than other background chaperone-chaperone interactions. Mcsc is an atypical Class Ib chaperone that has been shown to have multiple effectors and also interacts with several type III C-ring components, therefore its role in Chlamydia infection may differ from the typical effector chaperones and cannot be compared directly with other Class I chaperones.

Being associated with both type III secretion effectors and cytoplasmic components of the type III apparatus, the Mcsc dimer has been speculated to play a role in co-translational secretion of effectors[49]. No evidence has been presented thus far to suggest the coupling of effector translation and non-flagellar type III secretion in Chlamydia, nor in other bacteria. However, a model of flagellar secretion coupled with translation of the FlgM mRNA transcript has been proposed in Salmonella typhimurium[156]. In addition to Mcsc’s ability to interact with apparatus components, the Chlamydia non-flagellar type III components have also been shown to interact with flagellar components[157]. Thus it is possible that the Chlamydia type III secretion system is capable of unique mechanisms of regulating substrate translation and translocation.

**Oligomerization state of Slc1 and Slc1: TARP complexes**

Further characterization of the Slc1-Slc1 interaction was detected with cross-linking and gel filtration chromatography. While the cross-linking with DTSSP was only supposed to stabilize the already formed Slc1 homo-complexes, I contend that trimeric
Slc1 species were induced in the presence of the cross-linker. Purified Slc1 was highly concentrated in our samples and separate molecules of Slc1 were likely in close enough contact to become cross-linked. The most revealing aspect of the cross-linking assay was the finding that Slc1 homo-dimers could be detected on a non-reducing SDS-PAGE gel even without cross-linking treatment.

Gel filtration chromatography only confirmed our hunch that native Slc1 forms stable dimeric species in solution. A single native Slc1 complex migrated through the sepharose column to a position corresponding to a molecular weight of approximately 40 kDa, which is certainly indicative of homo-dimers. However, gel filtration chromatography is sensitive to both the size and molecular weight of the sample, so I cannot pinpoint an exact molecular weight with this assay. This approach was also useful for detecting the native Slc1: TARP complex. I expected to find a complex of a combined molecular weight of ~ 65 kDa (2 chaperone molecules to 1 TARP), but the Slc1 and TARP-containing elution fractions corresponded to a size >75 kDa. One reason for this larger size could be that the complex actually has a stoichiometry of 2:2, however this is unlikely considering the plethora of crystallization data supporting a 2:1 stoichiometry for type III chaperone: effector complexes. Instead I claim that the partially unfolded TARP in combination with the chaperone dimer has an extended size that caused the complex to elute earlier than expected from the sepharose column. The importance of this 2:1 stoichiometry in mediating the interaction with the type III secretion apparatus is not fully understood, but it is certainly an area worth investigating considering both Class II and Class III chaperones also form dimers to interact with their substrates[153]. The conservation of this chaperone dimer/substrate complex among
bacterial type III secretion systems has led to the conclusion that this complex forms a three-dimensional secretion signal that directs it to the type III apparatus[151].

A three-dimensional secretion signal that could trigger recognition by the type III ATPase has been proposed multiple times, but only a recent study was able to demonstrate the role of a chaperone-mediated translocation signal. The type III effector YopE from *Yersinia pseudotuberculosis* was dependent on a solvent exposed patch corresponding to the chaperone-binding domain for translocation, but interestingly these solvent exposed residues were not essential for chaperone binding[158]. This finding indicated that the type III chaperone orders the chaperone-binding domain of the effector such that a second secretion signal is exposed on the effector surface. Further work with type III chaperone-effector complexes that have been previously crystallized will indicate if the chaperone-mediated solvent-exposed patch is the universal three-dimensional translocation signal.

Considering that *Chlamydia* has two possible type III ATPases (CdsN and FliA) that are expressed during infection, it is probable that there is a difference in effector recognition between these ATPases[35]. Whether these ATPases (or possibly combinations of cytoplasmic type III components) would be able to differentiate between N-terminal secretion signals (S1) and/or chaperone-mediated translocation signals (S2) is yet to be determined. Thus future experiments testing the interaction of alternate chlamydial ATPases with chaperone-substrate complexes *in vitro* and in heterologous systems will be revealing as to the importance of different secretion signals in type III effector recognition, as well as clue us in to possible ATPase switching during actual infection. Speculation about ATPase switching may seem far-fetched when considering
the complete lack of precedence in other known bacterial type III secretion systems. However since the chlamydial type III secretion systems have evolved in isolation from the *Proteobacteria* for more than 2 billion years there will almost certainly be *Chlamydia*-specific type III interactions and functions that are not present in the other gram-negative type III secretion systems[22].

**Roles for chaperone-effector interactions in developmental cycle regulation**

As mediators of translocation type III chaperones may play an important role in the regulation of the EB to RB and/or RB to EB transitions that are necessary for establishing infection and spreading to surrounding tissues. Release of an effector, such as TARP, may allow for interaction of its cognate chaperone, Slc1, with another effector, whose translocation could trigger post-invasion events. Such an ordered process could conceivably be maintained by the specificity of the effectors’ chaperone binding domain for a specific chaperone dimer. CesT for instance, in EPEC, is known to act as chaperone for multiple effectors, having the highest affinity for the invasion-associated effector Tir, which is also the first of its cognate effectors to be detected in the host cell[159].

The apparent interaction of the chlamydial chaperone Slc2 (CT663) with both *E.coli* RNA polymerase (RNAP) and the chlamydial major sigma factor $\sigma^{66}$ indicates a possible regulatory role for type III chaperones[159]. It is tempting to speculate that the Slc2 interaction with $\sigma^{66}$ of the chlamydial RNAP holoenzyme allows for subsequent gene transcription by the alternate sigma factor $\sigma^{28}$, possibly initiating transcription of a set of genes involved in the RB to EB differentiation or maintaining the persistent RB. Both *CT663* and the $\sigma^{28}$-encoding gene *fliA* are mid-cycle genes and therefore their
expressed products are likely to play a role in late developmental cycle events[93]. The Slc2-Scc1 interaction detected in two-hybrid studies may sequester Slc2 and prevent it from acting as an anti-sigma factor during specific stages of infection. Determination of Scc1’s cognate effector or function during infection may provide insight into how and when Slc2 could act to regulate transcription. However, this area definitely needs to be further investigated to really determine the importance of Slc2 in gene regulation.

**Dependence of TARP’s translocation on Slc1 in heterologous systems**

While our biochemical assays only serve to indicate protein-protein interactions between putative chlamydial type III chaperones and between the chaperone Slc1 and the invasion-associated effector TARP, the findings beget many possible roles for these interactions during infection. I contend that Slc1’s main role is to achieve translocation of its cognate effector TARP and subsequently ensure the efficient uptake of *Chlamydia* by non-phagocytic eukaryotes. Using several different heterologous bacterial type III secretion systems I attempted to determine the chaperone requirement for TARP translocation. Translocation of β-lactamase fused effectors was reported by the cleavage of a lipophilic β-lactam ring-containing compound CCF2/AM. Because excitation of the cleaved and non-cleaved substrates at 410 nm produced emissions of alternate fluorescence wavelengths (510 nm and 450 nm, respectively), I was able to accurately measure TARP translocation into a population of HeLa cells by a ratio of fluorescence. This CCF2/AM based translocation assay has been widely used for detecting translocation of type III and type IV secreted effectors by *Shigella, Legionella, Salmonella, Yersinia, E.coli, and Pseudomonas*[160-163].
The reproducibility of this assay is based in the fact that total fluorescence is measured from a population of infected HeLa cells, so as long the multiplicities of infection are equal among compared samples, then the differences in substrate absorption and number of cells in each III are cancelled out when the fluorescence ratios are calculated. I was able to detect translocation of β-lactamase-fused TARP by *Yersinia enterocolitica* and the SPI-2 system of *Salmonella typhimurium*. Co-expression with its cognate chaperone significantly increased TARP translocation by both species.

However, I was unable to detect any TARP translocation by enterropathogenic *E.coli* or by the SPI-1 system of *S. typhimurium*, even in the presence of SICl. Secretion or translocation of chlamydial type III effectors had been previously reported with the SPI-1, SPI-2, and *Y. enterocolitica* type III systems, but not with EPEC. One possible reason for the lack of TARP translocation by EPEC could be that its ATPase recognizes a different N-terminal secretion signal. Alternatively, it could be that a chlamydial chaperone-mediated three-dimensional secretion signal cannot be recognized by the EPEC type III apparatus.

It has been previously shown that some chlamydial effectors are preferentially translocated by the heterologous SPI-1 or SPI-2 type III secretion systems[43]. Similarly, in our translocation assays I see a preference for translocation of TARP by the SPI-2 system. However, the fluorescence ratio had previously only been applied to measure the translocation of effectors within a few hours of infection, and the SPI-2 secretion system translocates effectors from its intracellular *Salmonella* containing vesicle between 6 and 10 hours post-infection[161]. I had to consider the possibility the some of the CCF2/AM substrate was able to access the non-translocated β-lactamase-fused effectors inside the
intracellular *Salmonella*, thus leading to a background level of cleaved substrate in this assay. SPI-2 translocation of β-lactamase fused effectors had been demonstrated previously but instead of calculating a fluorescence ratio, they observed blue and green fluorescence by live-cell microscopy[159]. Their microscopy results revealed a low level of conversion of the green to blue signal, but they were still able to obviously distinguish between cells in which a β-lactamase fused effector had been translocated (blue) from cells with no translocation (green). This background level of cleavage falsely skews the fluorescence ratio to indicate translocation in the (Empty) β-lactamase vector only negative control. However, I still believe that the elevated fluorescence ratio for TARP in the presence of Slc1 is significantly higher than this background, indicating actual chaperone-mediated translocation by the SPI-2 secretion system of *S. typhimurium*.

Secretion of TARP into media had been previously demonstrated by *Y. pseudotuberculosis*[42], and I was able to confirm *Yersinia*‘s ability to secrete TARP by *Y. enterocolitica*. However, TARP’s translocation into eukaryotic cells had not previously been tested, and even though TARP is secreted into media without a chaperone, secretion and translocation of effectors by *Yersinia* has been shown to have differential requirements for chaperones[62]. It has been shown that for *Yersinia* effectors, the N-terminal secretion signal is sufficient for secretion into low-calcium media, while translocation of effectors into host cells requires their cognate chaperones. This dogma is in agreement with our findings that TARP is secreted into media similarly to the positive control *Yersinia* effector YopH, but TARP requires Slc1 to be efficiently translocated into host cells.
I performed phylogenetic analysis of cytoplasmic type III flagellar and non-flagellar ATPases from *C. trachomatis* and *C. pneumoniae* in comparison with the SPI-1 and SPI-2 ATPases from *S. typhimurium*, EscN from EPEC, and YscN from *Y. enterocolitica* to assess whether the relatedness of ATPases could explain TARP translocation differences. The chlamydial non-flagellar ATPase CdsN was more closely related to YscN than InvC, SsaN, or EscN. This finding agrees with the fact that *Y. enterocolitica* was able to secrete and translocate TARP. The Slc1-dependence of TARP for efficient translocation of *Yersinia* indicates that the Slc1: TARP complex interacts with YscN in much the same way as it interacts with CdsN during chlamydial infection. Since TARP is capable of being translocated by the *Salmonella* SPI-2 type III secretion system but not by the SPI-1 system, it was interesting to note that the SPI-2 ATPase ssaN was more closely related to CdsN than the SPI-1 ATPase InvC. These phylogenetic comparisons of type III ATPases imply that the Slc1: TARP complex is also likely to be able to interact with chlamydial CdsN. The agreement between ATPase phylogeny and functional interaction with the effector TARP gives credibility to our heterologous translocation assays.

Taken together, these findings indicate a role for Slc1 in the translocation of TARP by the chlamydial type III secretion apparatus (Figure 38).
Figure 38. Model of Slc1-mediated translocation of TARP into eukaryotic cells. In the chlamydial EB, Slc1 associates with the N-terminus of TARP to reorganize its chaperone binding domain, which in turn is recognized by the type III secretion apparatus. Further unfolding by the ATPase allows TARP translocation through the type III secretion apparatus and into the eukaryotic cytosol where molecules of TARP form aggregates. Multiple domains on TARP are involved in actin polymerization that facilitates invasion of eukaryotic cells. TARP aggregates are nucleate actin at their WH2-like domains, while the tyrosine rich repeats become phosphorylated by host kinases. These phosphorylate repeats act as magnets for host signaling molecules and recruit further kinases guanine exchange factors that are involved in Rac activation and subsequent Arp2/3 formation.
CONCLUSIONS

This dissertation presents the first indication of a type III secretion chaperone mediating the translocation of a chlamydial effector into the cytosol of its target host cell. Importantly, this translocated effector, TARP, has been shown to be essential for ensuring the efficient invasion of non-phagocytic cells and subsequently enabling \textit{Chlamydia} to propagate. Not only does the requirement of the chaperone SIC1 in TARP translocation indicate a similar mechanism of effector translocation as detected in other type III-containing organisms, it also reveals the function of a previously uncharacterized protein, which is not an easy feat considering the genetically-intractable nature of \textit{Chlamydia spp.}

Biochemical characterization of the specific interaction between SIC1 and TARP indicate a chaperone-mediated translocation signal that is likely recognized by a type III ATPase. Detection of this translocation by the heterologous \textit{Salmonella} SPI-2 and \textit{Y. enterocolitica} type III secretion systems but not by enteropathogenic E.coli or the \textit{Salmonella} SPI-1 systems reveals a common mechanism for the chaperone-effector recognition by type III components. Comparison of the ATPases between these species indicates a role for the chlamydial ATPase CdsN in recognition of the SIC1: TARP complex. Further characterization the interaction of CdsN with this complex as well as other type III chaperone: substrate complexes may define a mechanism of differential
selection of type III substrates for translocation such as the recognition of three-dimensional chaperone-substrate translocation signals.

Recent work has demonstrated the inhibition of chlamydial growth and division by chemical inhibitors of type III secretion. The exact mechanism of this inhibition is unknown, and a likely target of these and future inhibition studies is the type III ATPase. Mapping the specific interactions that direct substrate recognition in the presence and absence of the type III chaperones will be pivotal to guiding the design of type III inhibitors for basic research and possibly for antibiotic use against several gram-negative pathogens.

The conservation of the type III secretion system among distantly related pathogenic and environmental Chlamydiaceae, all of which exhibit the biphasic intracellular lifestyle, indicates a role for the type III secretion system in maintaining this common lifestyle. Type III inhibitor studies will also provide valuable insight in the mechanisms directing the regulation of the Chlamydia’s unique biphasic developmental cycle. Substrate-switching, chaperone-mediated translocation hierarchies, interaction of type III components with transcription machinery, and sequestering of type III chaperones or substrates are all potential mechanisms that Chlamydia spp. may exploit to ensure efficient survival and transitioning between infectious (EB) and replicating (RB) forms.
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EDUCATION

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<tr>
<th>INSTITUTION AND LOCATION</th>
<th>DEGREE</th>
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RESEARCH PUBLICATIONS

Amanda J. Brinkworth, Antonio Tedim Pedrosa, Katarzyna Roguska, Sevanna Shahbazian, James E. Graham, Richard D. Hayward, Rey A. Carabeo. The type III chaperone Ssc1 mediates translocation of the C. trachomatis invasion-associated effector TARP. (Submitted to PLOS PATHOGENS, Feb 2011)