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Molecular dissection of the interface between the Type VI secretion TssM cytoplasmic domain and the TssG baseplate component.

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Running head: T6SS baseplate-TssM interactions

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ABSTRACT

The type VI secretion system (T6SS) is a multi-protein complex that catalyses toxin secretion through the bacterial cell envelope of various Gram-negative bacteria including important human pathogens. This machine uses a bacteriophage-like contractile tail to puncture the prey cell and inject harmful toxins. The T6SS tail comprises an inner tube capped by the cell-puncturing spike and wrapped by the contractile sheath. This structure is built on an assembly platform, the baseplate, which is anchored to the bacterial cell envelope by the TssJLM membrane complex. This membrane complex serves both as tail docking station and channel for the passage of the inner tube. The TssM trans-membrane protein is a key component of the membrane complex as it connects the inner and outer membranes. In this study, we define the TssM topology, highlighting a large but poorly studied 35 kDa-cytoplasmic domain, TssM_{Cyto}, located between two trans-membrane segments. Protein-protein interaction assays further show that TssM_{Cyto} oligomerizes and makes contacts with several baseplate components. Using computer predictions we delineate two sub-domains in TssM_{Cyto}, including a nucleotide tri-phosphatase (NTPase) domain followed by a 110-amino-acid extension. Finally, site-directed mutagenesis coupled to functional assays reveal the contribution of these sub-domains and of conserved motifs to the interaction with T6SS partners and to the function of the secretion apparatus.

Keywords: protein transport, protein secretion, Type VI secretion, bacterial competition, membrane complex.

INTRODUCTION

The Type VI secretion system (T6SS) is a versatile multi-protein secretory machine that is implicated in both inter-bacterial competition and anti-eukaryotic host activities. The T6SS delivers a broad arsenal of toxins with peptidoglycan, phospholipid or DNA hydrolysis activities, or that induce cytoskeleton re-arrangements directly into the target cell.¹⁻⁴

For toxin delivery, the T6SS uses a contractile mechanism that is comparable to that of *Myoviridae* phages or R-pyocins.⁵⁻¹⁰ This machine is composed of 13 core subunits, categorized in three sub-complexes^{8,10-12}: a cytoplasmic tubular structure built on an assembly platform – or baseplate (BP) – that is evolutionarily, structurally and functionally related to bacteriophage contractile tails^{5,13-15}, and anchored to the cell envelope by a membrane complex (MC).¹⁶

The T6SS tail is composed of an inner tube made of stacked Hcp hexameric rings and wrapped into a sheath-like structure, formed by the polymerization of TssB-TssC heterodimeric complexes, and that is assembled in an extended conformation.^{14,17-19} Indeed, the assembly of the tail can be followed by time-lapse microscopy: fluorescent-labelled sheath components assemble a ~ 600 nm-long tubular structure in tens of seconds, that then contracts in a few millisecond.^{14,20} The contraction of the sheath coincides with bacterial prey lysis, suggesting that, similarly to phages, sheath contraction propels the inner tube towards the target cell, allowing delivery of toxin effectors).^{8,11,20,21} The assembly of the tube and the sheath is coordinated by TssA, a protein that controls the elongation of the tail at the distal end and that maintains the sheath under the extended conformation.²² The inner tube is tipped by a spike constituted of a trimer of the VgrG protein, which is proposed to puncture the target cell membrane.^{13,23} The VgrG trimer is also part of the BP that is used as an assembly platform for the tail. Recently, the T6SS BP composition has been revealed. In addition to VgrG, it is composed of the TssE, -F and -G subunits, the homologues of the phage T4 gp25, gp6 and gp7 proteins, respectively, as well as of TssK a protein of unknown function with limited homologies to phage T4 gp8 or gp10 proteins that has been proposed to be a connector to the membrane complex.^{15,24-27} This MC is composed of the two TssL and TssM inner membrane (IM) proteins and of the TssJ outer membrane (OM) lipoprotein.²⁸⁻³² TssL and TssM interact in the IM whereas the C-terminal periplasmic domain of TssM contacts the TssJ lipoprotein close to the OM.^{16,29,30,33-35} The MC serves as a docking station for the BP

and the tail, but has also been proposed to serve as channel for the passage of the inner tube during sheath contraction.¹⁶ In the recent years, the assembly pathway of the T6SS has been well defined. T6SS biogenesis progresses from the outer membrane to the cytoplasm. It starts with the positioning of the TssJ lipoprotein and the successive recruitments of TssM and TssL.¹⁶ Recruitment of TssA then positions the baseplate complex onto the MC and primes the polymerization of the tail tube/sheath.^{15,22,36} This ordered assembly pathway requires tight contacts between the different subunits. Indeed, docking of the BP onto the MC requires multiple contacts including interactions of TssE and TssK with TssL, and of TssG and TssK with TssM.^{15,24} TssM is therefore a key component as it mediates contact with the OM TssJ lipoprotein, as well as with cytoplasmic BC subunits.

Here, we show that the enteroaggregative *Escherichia coli* TssM protein is a polytopic membrane protein, inserted into the inner membrane by three trans-membrane helices (TMH). The C-terminal portion of TssM is in the periplasm and interacts with TssJ.³⁴ TMH2 and TMH3 delimitates a ~ 35 kDa cytoplasmic domain, TssM_{Cyto}, which is conserved among TssM homologues. Computer analyses show that TssM_{Cyto} is constituted of two sub-domains: a sub-domain with a nucleotide tri-phosphatase (NTPase)-like domain followed by an extension. Indeed, TssM has been previously shown to bind and hydrolyse NTPs.³⁷ However, the role of the NTP-binding motif and its functional implication during T6SS activity is still a matter of debate.^{29,33} The extension comprises a eukaryotic DPY-30-like dimerization motif. We show that the NTPase-like domain mediates interaction with TssK whereas the extension is necessary and sufficient for TssM_{Cyto} oligomerization and interaction with TssG. Site-directed mutagenesis of conserved motifs within the extension revealed their contribution for TssM_{Cyto} oligomerization, TssM_{Cyto}-TssG interaction and for proper assembly of the T6SS. Our results thus provide details on the molecular interface between the T6SS membrane and baseplate complexes.

RESULTS

TssM is a polytopic IM protein

The TssM protein encoded within the enteroaggregative *E. coli sci-1* gene cluster (EC042_4539; Genbank accession (GI): **284924260**) is a large protein of 1129 amino acids.

Based on hydrophobicity plots, most widely used computer tools predict TssM as an inner membrane protein with three trans-membrane helices (TMH) (Fig. 1A). Indeed, fractionation experiments showed that TssM co-fractionates with membrane proteins (data not shown). To experimentally define the TssM topology and determine the TMH boundaries we performed a cysteine accessibility assay using the substituted cysteine accessibility method (SCAM, Ref 38). This assay relies on the ability of 3-(*N*-maleimidylpropionyl) biocytin (MPB), a sulfhydryl reagent to cross the outer membrane but not the inner membrane of Gram-negative bacteria including EAEC.^{30,31} TssM possesses nine native cysteine residues, with one (C727) predicted to locate in the periplasm. Hence, the WT TssM protein is labelled by MPB *in vivo* (Fig. 1B). In agreement with the computer predictions, a TssM protein in which the cysteine at position 727 is substituted to serine (C727S) was not labelled with MPB (Fig. 1B). These data suggest that C727 is located in the periplasm whereas all other 8 cysteine residues locate in the cytoplasm or are buried into the structure of the protein and then inaccessible to MPB. We then introduced cysteine substitutions in the C727S TssM variant at various positions along the protein (at positions 37, 67, 352 and 386) (Fig. 1A). All these mutated proteins were produced at similar levels (Fig. 1B) and were able to complement the effect of the *tssM* mutant in an Hcp secretion assay (data not shown). The A37C and S386C variants were biotinylated with MPB suggesting that the A37 and S386 residues locate in the periplasm (Fig. 1B). By contrast, the V67C and S352C variants were not labelled indicating that the V67 and S352 residues locate in the cytoplasm (Fig. 1B). All together, the data of the cysteine accessibility defined the topology of TssM: TssM is constituted of three TMH, with the N-terminus in the cytoplasm and the C-terminus in the periplasm. TssM spans the IM through two TMH oriented in-to-out (TMH1, residues 13-29; TMH3, residues 360-382) and one TMH oriented out-to-in (TMH2, residues 44-62) (Fig. 1C). TMH2 and TMH3 thus delimitate a ~ 35 kDa domain located in the cytoplasm, called hereafter TssM_{Cyto}.

The cytoplasmic domain of TssM oligomerizes and interacts with components of the T6SS membrane and baseplate complexes

The topology of TssM indicates the existence of two soluble domains, one in the periplasm (TssM_{Peri}, amino-acids 383-1129) and one residing into the cytoplasm (TssM_{Cyto}, amino-acids 63-359). The T6SS being a multi-protein complex, such large protein domains might be necessary for interacting with other T6SS components. Indeed, we and other have previously demonstrated that TssM_{Peri} interacts with the TssJ outer membrane lipoprotein in

Edwardsiella tarda and enteroaggregative *E. coli*.^{33,34} By contrast, little is known regarding the cytoplasmic domain of TssM. To gain further insights onto TssM_{Cyto} partners, we used TssM_{Cyto} as bait for an *in vivo* systematic bacterial two-hybrid assay. TssM_{Cyto} was fused to the T18 domain of the *Bordetella* adenylate cyclase and all the other T6SS proteins - or soluble domains - were fused either at their N- or C-terminus of the T25 domain. The results presented in Fig. 2 show that TssM_{Cyto} interacts with itself and with TssK whatever the constructions used. In addition, TssM_{Cyto} interacts with TssG and with the cytoplasmic domain of TssL (TssL_{Cyto}) when fused at the N-terminus of T25. In conclusion, TssM_{Cyto} is capable of oligomerization and interacts with components of the T6SS membrane (TssL_{Cyto}) and baseplate (TssK and TssG) complexes. These results are in agreement with previously published bacterial two-hybrid screens and co-immune precipitations that identified TssM-TssK, TssM-TssL and TssM-TssG interactions.^{15,24,29,30}

Sub-domain architecture of TssM_{Cyto}

Pfam, Blast and HHPred analyses suggest that EAEC TssM_{Cyto} is organized as a NTPase domain (TssM_{Cyto}/NTP; amino-acids 62-248; Pfam accession: **PF06858**) followed by a C-terminal extension (TssM_{Cyto}/Ct; amino-acids 254-360) (Fig. 3; Supp. Fig. S1-S2). However, despite the fact that the overall NTPase domain is conserved among the TssM_{Cyto} homologs, the sequence alignment shows that the NTP binding and hydrolysis motifs (Walkers A & B and NTP specific motif) are not well conserved (Supp. Fig. S1). In agreement with this observation, an evolutionary analysis of TssM_{Cyto} NTPase domains shows that they categorize into two sub-groups: while the TssM proteins encoded within the *Pseudomonas aeruginosa* H1, *Agrobacterium tumefaciens* and *Edwardsiella tarda* T6SS gene clusters carry a complete NTPase domain, a number of TssM, including that of EAEC, *Serratia* and *Citrobacter* possess a NTPase domain amputated of hydrolysis motifs (Supp. Fig. S1 and S2).

Our attempts to produce and purify the EAEC TssM_{Cyto} domain or the TssM_C NTPase sub-domain in order to gain structural information were unsuccessful, as the different constructs used were all insoluble. Consequently, we sought to construct homology-based models of both TssM_{Cyto} sub-domains using a bioinformatic approach. The TssM_{Cyto}/NTP structure was predicted using HHpred.³⁹ The program confirmed that the EAEC TssM_{Cyto}/NTP protein resembles the solved structure of various GTP-hydrolysing proteins (Supp. Fig. S3). The X-ray structure of the *Burkholderia thailandensis* EngB GTP-binding protein (PDB ID: **4DHE**)⁴⁰ was subsequently used as template to build a homology model of the EAEC

TssM_{Cyto/NTP} domain. Figure 3 shows that TssM_{C/NTP} adopts a compact fold consisting of a four-stranded parallel β -sheet with one side being in contact with three α -helices. The TssM_{Cyto/NTP} domain belongs to the α/β class, harbouring an incomplete Rossmann fold, a motif associated with nucleotide-binding proteins.⁴¹ The EAEC TssM_{Cyto/NTP} architecture is typical of P-loop nucleotide triphosphate hydrolases. The predicted structure of EAEC TssM_{Cyto/NTP} diverges however from the classical $\alpha\beta\alpha$ sandwich architecture as the second α -helix is not strictly sandwiching the β -strand. As expected, the large loops that bear the binding and hydrolysis motifs are absent in the EAEC TssM_{Cyto/NTP}, by contrast to EngB or the homology model of the *A. tumefaciens* TssM_{Cyto/NTP} domain (Supp. Fig. S4). The TssM_{Cyto/Ct} homology model was constructed using the Swiss-Model server based on the X-ray structure of the C-terminal domain of human DPY-30-like protein, a component of the eukaryotic histone methyltransferase complex (PDB ID: **3G36**)⁴² as template (Supp. Fig. S3). The TssM_{C/Ct} structure was confidently modelled from residue Q254 to N289 (Fig. 3). This fragment encompasses the two-helix 40-amino-acid Dpy-30 motif (Pfam accession: **PF05186**) found in DPY-30 proteins and involved in DPY-30 dimerization⁴².

Specific motifs are involved in TssM_{Cyto} oligomerization and interaction with TssG.

The structural organization of TssM_{Cyto} prompted us to investigate the contribution of the two sub-domains to the TssM_{Cyto} interactions. The interaction network of TssM_{Cyto/NTP} and TssM_{Cyto/Ct} was assessed by bacterial two-hybrid. As shown in Figure 4A, TssM_{Cyto/NTP} interacts with TssK, whereas TssM_{Cyto/Ct} mediates oligomerization and interactions with TssG and TssL_{Cyto}. Interestingly the interaction network of the isolated sub-domains, TssM_{Cyto/NTP} and TssM_{Cyto/Ct}, recapitulates the interaction network of TssM_{Cyto} (Fig. 4A), supporting the hypothesis of two independently-folded domains. The TssM_{Cyto/Ct}-TssG and TssM_{Cyto/NTP}-TssK interactions were further confirmed by co-immuno-precipitation experiments: TssG was co-precipitated with TssM_{Cyto} and TssM_{Cyto/Ct} (Fig. 4B, upper panel). As previously shown²⁴, TssK did not interact with TssM_{Cyto} by co-immuno-precipitation. However, our results suggest that it is prevented by the extension as the TssM_{Cyto/NTP} domain alone interacts with TssK (Fig. 4B, lower panel).

Interestingly, the TssM_{Cyto/Ct} sub-domain interacts with several partners from the membrane and baseplate sub-complexes. We therefore questioned whether these different interactions involve the same recognition motif or different binding epitopes on TssM_{Cyto}. The

sequence alignment of TssM_{Cyto} homologues emphasized two well-conserved regions, F278-E284 and L309-S315 (Supp. Fig. S1). Region F278-E294 is specifically conserved in TssM_{Cyto} lacking functional NTPase domain, whereas the L309-S315 motif is conserved among all TssM_{Cyto} (Supp. Fig. S1). Using site-directed mutagenesis, we engineered TssM_{Cyto} variants in which these motifs were targeted. Although these two motifs do not appear to be involved in TssM_{Cyto}-TssL_{Cyto} interactions, substitutions within the L309-S315 motif specifically abolished the TssM_{Cyto}-TssG interaction (Fig. 5A and 5B). We also noted that substitutions within the F278-E284 motif impacted TssM_{Cyto} oligomerization. However, although the L279W and L282W/A283W mutations prevented interaction with TssM_{Cyto} in the bacterial two-hybrid assay (Fig. 5A), only the L279W mutations had a strong effect on multimerization in the co-immunoprecipitation assay (Fig. 5B). It is worthy to note that the TssM_{Cyto/Ct} F278-E284 residues correspond to the dimerization motif in Dumpy-30 (DPY-30) proteins (Supp. Fig. S4B).

TssM_{Cyto} oligomerization and interaction with the TssG baseplate subunit are critical for T6SS function.

The mutations that specifically affect TssM_{Cyto} oligomerization and TssM_{Cyto}-TssG complex formation were tested for their repercussion on T6SS function. EAEC Sci-1 T6SS function could be monitored by measuring its antibacterial activity.⁴³ The four TssM substitutions were introduced within the native, chromosomal *tssM* gene. Fig. 6A shows that all four mutated strains were defective in T6SS-dependent killing of prey bacterial cells. We therefore conclude that TssM_{Cyto} oligomerization and interaction with TssG are required for proper function of the Type VI secretion apparatus.

T6SS biogenesis starts with the assembly of the membrane complex and is followed by (i) the recruitment of the baseplate and (ii) tail polymerization. We therefore sought to define which stage of T6SS biogenesis is impacted by these mutations. We first tested the effects of these mutations on T6SS sheath assembly by following the dynamics of a chromosomally-encoded TssB-sfGFP fusion using fluorescence microscopy (Fig. 6B). All the substitutions severely affected T6SS sheath assembly as indicated by the decrease in the number of sheath per bacterial cell (Fig. 6B). While ~ 25% of the wild-type cells assembled sheath structures, TssBC sheath assembled on rare occasions in cells carrying mutations affecting TssM_{Cyto} oligomerization (3-4 % of cells with sheath structures). The effect of the

TssM_{Cyto}-TssG disruption was even more drastic as sheath assembly was observed in ~ 1% of the cells.

Second, we tested the effect of these mutations on recruitment of the baseplate. By following the dynamics of a sfGFP-TssF fusion, a recent study concluded that the membrane complex recruits and stabilizes the T6SS baseplate.¹⁵ We therefore investigated whether the mutations affecting oligomerization of the cytoplasmic loop of TssM and contacts between this loop and the TssG baseplate component impact baseplate assembly, stability and recruitment. As TssG fusions to sfGFP were previously shown to be non functional¹⁵, we introduced the TssM point mutations in a strain producing the chromosomal and functional sfGFP-TssF fusion, a TssG protein partner, and we monitored the formation and stability of baseplate foci by fluorescence microscopy (Fig. 6C). All four mutations significantly decreased the number of cells with sfGFP-TssF foci and the average number of foci per cell (1 cluster in 15-25% of the mutated cells compared to 1-3 foci in ~ 65% for WT cells) (Fig. 6C and 6D). Taken together, these data demonstrate that mutations that affect TssM_{Cyto} oligomerization and TssM_{Cyto}-TssG complex formation abolish T6SS sheath formation and function by impacting T6SS baseplate assembly and stability.

DISCUSSION

In this manuscript, we report the characterization of the cytoplasmic domain of the T6SS membrane core complex protein TssM from EAEC. We showed that TssM_{Cyto} comprises two sub-domains, a domain resembling NTPases but lacking nucleotide binding and hydrolysis motifs, followed by a ~ 110-amino-acid extension. Protein-protein interaction studies revealed that this extension mediates TssM_{Cyto} oligomerization and interaction with the TssG baseplate subunit. We finally defined specific motifs involved in these interactions and reported that these interactions are critical for the assembly of a functional T6SS. Models summarizing the findings reported in this study are depicted in Fig. 7.

We first defined the boundaries of the TssM trans-membrane segments using cysteine accessibility experiments. We determined that TssM is constituted of three TMH. The TssM N-terminus locates in the cytoplasm and is followed by a trans-membrane hairpin, a cytoplasmic domain, and the third TMH, TMH3. Finally, the ~ 750-amino-acid C-terminal

domain locates in the periplasm (Fig. 1C). This topology is similar to the topology of the *Agrobacterium tumefaciens* TssM protein previously defined using translational reporter fusions.²⁹ Computer analyses of TssM proteins encoded within well-studied T6SS gene cluster showed that this topology is likely shared between all the homologues with the notable exception of the *P. aeruginosa* H1-T6SS TssM protein that is predicted to have a single TMH corresponding to TMH3 (Supp. Fig. S5).

The topology experiments also defined that TssM TMH2 and TMH3 delimit a 35-kDa cytoplasmic domain, TssM_{Cyto}. Our data showed that TssM_{Cyto} oligomerizes and interacts with TssL_{Cyto}. The TssM_{Cyto}-TssM_{Cyto} and TssM_{Cyto}-TssL_{Cyto} interactions have been reported in the *Agrobacterium tumefaciens* T6SS²⁹. The conservation of TssM_{Cyto} oligomerization and interaction with TssL_{Cyto} between *A. tumefaciens* and EAEC suggest that these contacts are important for T6SS function. Indeed, mutation of L279, a residue that participates to TssM_{Cyto} oligomerization, severely impacts T6SS function in EAEC. The low-resolution of the recently published electron microscopy (EM) map of the 5-fold symmetry TssJLM MC does not allow to use docking simulations to precisely locate the TssM cytoplasmic domain and therefore to provide insight onto its oligomeric state in the complex. However, stoichiometry analyses and reconstruction of the MC suggested it is constituted of 5 dimers of TssJLM heterotrimers.¹⁶ These information suggest that, similarly to the TssL cytoplasmic domain³², TssM_{Cyto} dimerizes. The biogenesis of the MC may therefore start with the formation of dimers of heterotrimers that will then symmetrize. Based on the TssM_{Cyto} interaction with the TssL cytoplasmic domain, it has been proposed that these two domains form the large base of the T6SS MC.¹⁶ This cytoplasmic base corresponds to the docking site for the TssEFGK-VgrG baseplate complex.^{15,16,22} Indeed, our bacterial two-hybrid and co-immune precipitation experiments confirmed that TssM_{Cyto} interacts with two baseplate components, TssK and TssG (Fig. 7).

Bio-informatic analyses predict that TssM_{Cyto} comprises a N-terminal NTPase domain and a C-terminal DPY-30-like domain named TssM_{Cyto/NTP} and TssM_{Cyto/Ct}, respectively. However, although the EAEC TssM_{Cyto/NTP} domain belongs to the NTPase fold, it misses the specific binding and hydrolysis motifs found in functional NTPases, such as the Walker A and B motifs. Sequence alignment of TssM_{Cyto/NTP} domains from various bacterial species revealed that they categorize in two sub-families. While a number of TssM_{Cyto/NTP} domains do not carry these motifs, other TssM possess Walker A and B signatures, including that of *A.*

tumefaciens, *Edwardsiella tarda* or *P. aeruginosa* H1-T6SS (Suppl. Fig. S2). Indeed, the detergent-solubilized *A. tumefaciens* TssM protein exhibits ATPase activity.³⁷ However, mutations of these motifs in *A. tumefaciens* and *E. tarda* did not have the same impact on the function of the T6SS. Production of the K124A TssM variant in *Edwardsiella* was fully functional as shown by T6SS-dependent Hcp, VgrG and EvpP release.³³ By contrast, ATP binding and hydrolysis regulate conformational changes within the periplasmic domain of the *A. tumefaciens* TssM protein and the *Agrobacterium* TssM K145A variant is unable to restore Hcp release.^{29,37} Therefore, TssM_{C/NTP} domains come in different flavours: active NTPase domains (e.g., *A. tumefaciens*), inactive NTPase domains (e.g., *E. tarda*) and NTPase fold lacking the functional motifs (e.g., EAEC). Interestingly, our protein-protein interaction studies showed that TssM_{C/NTP} contacts TssK. In the case of TssM with functional NTP domains, it would be interesting to test whether the presence of TssK influences NTP binding and hydrolysis.

The TssM_{Cyto} C-terminal extension shares structural homologies with the dimerization motif of DPY-30, a subunit of the histone methyltransferase complex in eukaryotic cells. This two- α -helix motif forms an antiparallel bundle at the dimer interface, which is mediated by extensive hydrophobic and van der Waals interactions (Suppl. Fig. S4).⁴² Indeed, this DPY-30-like hairpin and notably the conserved L279 residues are involved in TssM_{Cyto} oligomerization. In addition to its role in TssM_{Cyto} oligomerization, this sub-domain is also required for proper interaction with the cytoplasmic domain of TssL, TssL_{Cyto}, and with TssG, one of the components of the T6SS baseplate (Fig. 7). Whereas we have not identified in this study the residues of TssM_{Cyto/Ct} mediating the interaction with TssL_{Cyto}, a conserved hydrophobic sequence (L-A-G-I-V-F-S in EAEC) is required for TssM_{Cyto}-TssG complex formation. Taken together, our results show that this relatively small sub-domain is responsible for several interactions. Interestingly, a similar case has been reported for DPY-30, which is a partner of several complexes involved in the regulation of chromatin and nucleosome organization.⁴⁴ One may hypothesize that TssM_{Cyto/Ct} uses its DPY-30-like domain to interact sequentially with its different protein partners. Purification of the TssM_{Cyto}-TssL_{Cyto}-TssG complex or high-resolution structure of the different binary complexes involving TssM_{Cyto/Ct} will shed light on the dynamic nature of these interactions.

The interactions between TssM_{Cyto} and TssK and TssG, two components of the baseplate complex, might be important to recruit or to stabilize the BC at the cytoplasmic base

of the TssJLM MC. In addition to this structural role, it is likely that these interactions are required for proper function of the baseplate. As shown for other contractile structures such as bacteriophages, the baseplate serves as assembly platform for the tail, but is also responsible for initiating sheath contraction.^{26,45-47} The TssM_{Cyto}-TssG interaction might be therefore important to regulate baseplate assembly, recruitment or sheath assembly and/or contraction. Indeed, point mutations disrupting the TssM_{Cyto}-TssG interaction destabilize the baseplate complex and prevent elongation of the tail sheath. It is interesting to note that TssM undergoes structural transitions³⁷ and one may suggest that TssM conformational changes might be transduced to the baseplate via the TssM_{Cyto}/NTP-TssK and/or TssM_{Cyto}/Ct-TssG interactions leading to sheath assembly. As TssM has a large periplasmic domain and a short extension that lies outside of the cell¹⁶, it is a strong candidate to sense modifications of the cell envelope such as an attack by neighbouring cells, or contact with a prey, and to transmit the information to the baseplate complex. Further experiments will provide insights on how sheath assembly and/or contraction is regulated.

Our results also pointed that mutations disrupting TssM_{Cyto} oligomerization and TssM_{Cyto}-TssG complex formation affect T6SS-dependent activities as they abolish TssBC-sheath assembly and inhibit bacterial prey killing. The drastic effect of these mutations makes these interactions attractive targets for the rationale design of drugs that, by binding to TssM_{Cyto}, would hamper the T6SS activity and the delivery of harmful toxins. This approach has been successfully achieved in the case of the *Brucella* Type IV secretion VirB8 inner membrane subunit for which specific inhibitors of its dimerization were identified by a high-throughput bacterial two-hybrid screen and were further shown to inhibit *Brucella* infection of macrophages.^{48,49} This example emphasises the importance of understanding protein-protein interactions in bacterial secretion systems with the ultimate goal to target specific interactions with small molecule inhibitors.

MATERIALS and METHODS

Bacterial strains, media, growth conditions and chemicals. Strains used in this study are listed in Supplementary Table S1. *Escherichia coli* K-12 DH5 α , BTH101 and W3110 were used for cloning procedures, bacterial two-hybrid and co-immune precipitation, respectively. The *E. coli* K-12 W3110 strain carrying the pUA66-*rrnB* plasmid (*gfp* under control of the constitutive *rrnB* ribosomal

promoter, specifying strong and constitutive fluorescence, and kanamycin resistance)⁵⁰ was used as prey in antibacterial competition experiments. Enteroaggregative *E. coli* strain 17-2, and its $\Delta tssM$, *tssB-GFP* and *GFP-tssF* derivatives^{15,24,30} were used in this study. Chromosomal fluorescent reporter insertions were obtained using the modified one-step inactivation procedure⁵¹ using the red recombinase expressed from pKOBEG⁵² as previously described²⁸ using *pgfp*-KD4 as template for Polymerase Chain Reaction (PCR) amplification. Briefly, the sfGFP-coding sequence and the kanamycin cassette were amplified from the *pgfp*-KD4 vector¹⁵ with oligonucleotides carrying 50-nucleotide extensions homologous to regions adjacent to the site of insertion. The PCR product was column purified (PCR and Gel Clean up kit, Promega) and electroporated. Kanamycin resistant clones were recovered and the insertion of the kanamycin cassette at the targeted site was verified by PCR. Kanamycin cassettes were then excised using pCP20.⁵¹ *tssM* point mutations were engineered at the native locus on the chromosome by allelic replacement using the pKO3 suicide vector.^{53,54} Briefly, 17-2 *tssB-GFP* or *GFP-tssF* cells were transformed with a pKO3 plasmid in which a fragment of the *tssM* gene carrying the point mutations has been cloned (see below). Insertion of the plasmid into the chromosome was selected on chloramphenicol plates at 42°C. Plasmid sequences removal was then selected on 5% sucrose plates without antibiotic and *tssM* point mutation recombinant strains were screened by PCR and confirmed by DNA sequencing (Eurofins,MWG). Unless specified, cells were grown in Luria broth (LB) or in *sci-1* inducing medium (SIM: M9 minimal medium, glycerol 0.2%, vitamin B1 1 µg/mL, casaminoacids 100 µg/mL, LB 10%, supplemented or not with bactoagar 1.5%)⁵⁵ at 37°C with shaking. Plasmids were maintained by addition of ampicillin (100 µg/mL), kanamycin (50 µg/mL) or chloramphenicol (30 µg/mL). Gene expression from pASK-IBA37 and pBAD vectors was induced by the addition 0.1 µg/mL of anhydrotetracyclin (AHT, IBA Technology) and 0.02% of L-arabinose (Sigma-Aldrich), respectively.

Plasmid construction. Plasmids used in this study are listed in Supplementary Table S1. Polymerase Chain Reactions (PCR) were performed using a Biometra thermocycler using the Q5 high fidelity DNA polymerase (New England Biolabs). Restriction enzymes were purchased from New England Biolabs and used according the manufacturer's instructions. Custom oligonucleotides, listed in Supplementary Table S1, were synthesized by Sigma Aldrich. Enteroaggregative *E. coli* 17-2 chromosomal DNA was used as a template for all PCRs. *E. coli* strain DH5α was used for cloning procedures. With the exception of the pKO3-'*tssM*' vector, plasmids have been constructed by restriction-free cloning⁵⁶ as previously described³⁰. Briefly, genes of interest were amplified with oligonucleotides introducing extensions annealing to the target vector. The double-stranded product of the first PCR was then been used as oligonucleotides for a second PCR using the target vector as template. pKO3-'*tssM*' has been constructed by the restriction-ligation procedure. A *Bam*HI-*Sa*II PCR product corresponding to a fragment of the *tssM* gene (nucleotides 216-1569) was ligated into pKO3 digested by the same enzymes using T4 DNA ligase (New England Biolabs). Substitutions into

pIBA37-FLAG-TssM, pTssM_{Cyto}-T18, pT18-TssM_{Cyto} and pKO3-'*tssM*' have been introduced by site-directed mutagenesis using complementary pairs of oligonucleotides and the Pfu Turbo high fidelity polymerase (Agilent Technologies). All constructs have been verified by restriction analysis and DNA sequencing (Eurofins, MWG).

Antibacterial assay. The antibacterial competition growth assay was performed as described.⁴³ The wild-type *E. coli* strain W3110 bearing the GFP⁺ kanamycin-resistant pUA66-*rrnB* plasmid⁵⁰ was used as prey in the competition assay. The kanamycin-resistant pUA66-*rrnB* plasmid provides a strong constitutive green fluorescent (GFP⁺) phenotype. Attacker and prey cells were grown for 16 hours in SIM, then diluted 100-fold in SIM. Once the culture reached an OD₆₀₀ = 0,8, cells were harvested and resuspended to an OD_{600nm} of 10 in SIM. Attacker and prey cells were mixed to a 4:1 ratio and 20-μL drops of the mixture were spotted in triplicate onto a pre-warmed dry SIM agar plate. After incubation for 4 hours at 37°C, the bacterial spots were resuspended in LB and bacterial suspensions were normalized to an OD₆₀₀ of 0.5. For enumeration of viable prey cells, bacterial suspensions were serially diluted and spotted onto selective LB agar plates supplemented with kanamycin (for the *E. coli* prey cells). The experiments were done in triplicate, with identical results, and we report here the results of a representative experiment.

Substituted cysteine accessibility method (SCAM). Cysteine accessibility experiments were carried out as described^{57,58} with modifications.^{30,31} A 40-mL culture of strain Δ*tssM* producing the TssM or cysteine-substituted TssM derivatives was induced for *tssM* gene expression with 0.02% AHT for 2 hours. Cells were harvested, resuspended in buffer A (100 mM HEPES (pH 7.5), 250 mM sucrose, 25 mM MgCl₂, 0.1 mM KCl) to a final OD₆₀₀ of 12 in 500 μL of buffer A. MPB (Molecular Probes) was added to a final concentration of 100 μM (from a 20 mM stock freshly dissolved in DMSO) and the cells were incubated for 30 min at 25°C. β-Mercaptoethanol (20 mM final concentration) was added to quench the biotinylation reaction, and cells were washed twice in buffer A, and resuspended in buffer A containing *N*-ethyl maleimide (final concentration 5 mM) to block all free sulfhydryl residues. After incubation 20 min at 25°C, cells were disrupted by four passages at the French press at 800 psi. Membranes recovered by ultracentrifugation 40 min at 100 000 × *g* were resuspended in 1 mL of buffer B (10 mM Tris (pH 8.0), 100 mM NaCl, 1% (w/v) Triton X-100, protease inhibitor cocktail (Complete, Roche)). After incubation on a wheel for 2 hours, unsolubilized material was removed by centrifugation 15 min at 20, 000 × *g*, and solubilized proteins were subjected to immune precipitations using anti-FLAG M2 affinity gel (Sigma-Aldrich). After 3 hours of incubation on a wheel, the beads were washed twice with 1 mL buffer B, and once with buffer C (10 mM Tris (pH 8.0), 100 mM NaCl, 0.1% (w/v) Triton X-100). Beads were air-dried, resuspended in Laemmli buffer and subjected to SDS-PAGE and immunodetection with anti-FLAG antibodies, and streptavidin coupled to alkaline phosphatase.

Bacterial two-hybrid assay. The adenylate cyclase-based bacterial two-hybrid technique⁵⁹ was used as previously published.^{24,60} Briefly, pairs of proteins to be tested were fused to the isolated T18 and T25 catalytic domains of the *Bordetella* adenylyl cyclase. After transformation of the two plasmids producing the fusion proteins into the reporter BTH101 strain, plates were incubated at 30°C for 48 hours. Three independent colonies for each transformation were inoculated into 600 µL of LB medium supplemented with ampicillin, kanamycin and IPTG (0.5 mM). After overnight growth at 30°C, 10 µL of each culture were dropped onto LB supplemented with 40 µg/mL⁻¹ bromo-chloro-indolyl-β-D-galactopyranoside (X-Gal) and incubated for 16 hours at 30°C. The experiments were done at least in triplicate and a representative result is shown.

Co-immune precipitation. 100 mL of W3110 cells producing proteins of interest were grown to an OD₆₀₀ of 0.4 and the expression of the cloned genes were induced with AHT or L-arabinose for 45 min. The cells were harvested, and the pellets were resuspended in Tris-HCl 20 mM pH 8.0, NaCl 100mM, sucrose 30%, EDTA 1mM, lysozyme 100 µg/mL, DNase 100 µg/mL, RNase 100 µg/mL supplemented with protease inhibitors (Complete, Roche) to an OD₆₀₀ of 80 and incubated on ice for 20 min. Cells were lysed by three passages at the French Press (800 psi) and lysates were clarified by centrifugation at 20,000 × g for 20 min. Supernatants were used for co-immune precipitation using anti-FLAG M2 affinity gel (Sigma-Aldrich). After 3 hours of incubation, the beads were washed three times with 1 mL of Tris-HCl 20 mM pH 8.0, NaCl 100 mM, sucrose 15%, resuspended in 25 µL of Laemmli loading buffer, boiled for 10 min and subjected to SDS-PAGE and immunodetection analyses.

Fluorescence microscopy and statistical analyses. Overnight cultures of entero-aggregative *E. coli* 17-2 derivatives strains were diluted 1:100 in SIM medium and grown for 6 hours to an OD₆₀₀ ~ 1.0 to maximize expression of the *sci-1* T6SS gene cluster.⁵⁵ Cells were washed in phosphate buffered saline (PBS), resuspended in PBS to an OD₆₀₀ ~ 50 and spotted on a thin pad of 1.5% agarose in PBS and covered with a cover slip. Microscopy recording and digital image processing have been performed as previously described.^{15,16,18,20,24} The Z project (average intensity) plugin has been used to merge and flatten all Z planes. Microscopy analyses were performed at least six times, each with technical triplicate, and a representative experiment is shown. The number of sheath per number of cells and sfGFP-TssF foci was measured manually.

Computer analyses. Trans-membrane helix predictions were made using HMMTop⁶¹, TMHMM⁶², TMpred⁶³ and PHDhtm⁶⁴. Secondary structure predictions were made using the Psipred server (<http://bioinf.cs.ucl.ac.uk/psipred/>). Structural predictions and homology modelling of the tri-dimensional structure of TssM_{Cyto/NTP} and TssM_{Cyto/Ct} were performed using HHpred³⁹ or Swiss-Model⁶⁵, respectively. Figures were made using Chimera⁶⁶. Amino-acid sequences were aligned with

T-COFFEE⁶⁷ and phylogenetic analyses were performed with phylogeny.fr⁶⁸.

Miscellaneous. SDS-polyacrylamide gel electrophoresis was performed using standard protocols. For immunostaining, proteins were transferred onto nitrocellulose membranes, and immunoblots were probed with primary antibodies, and goat secondary antibodies coupled to alkaline phosphatase, and developed in alkaline buffer in presence of 5-bromo-4-chloro-3-indolylphosphate and nitroblue tetrazolium. The anti-FLAG (M2 clone, Sigma-Aldrich), anti-VSV-G (clone P5D4, Sigma-Aldrich) monoclonal antibodies, the alkaline phosphatase-conjugated streptavidin (Pierce) and alkaline phosphatase-conjugated goat anti-mouse secondary antibodies (Beckman Coulter) have been purchased as indicated and used as recommended by the manufacturer.

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LEGEND TO FIGURES

Figure 1. TssM is a polytopic inner membrane protein.

A. The trans-membrane helices predicted using the algorithms listed on the left are represented by black rectangles. The EAEC Sci-1 TssM natural cysteine residues, as well as the cysteine substitutions engineered in this study are indicated. Filled circles indicate cysteine residues labelled with the 3-(*N*-maleimidylpropionyl) biocytin (MPB) probe, whereas open circles indicate unlabelled cysteine residues. Arrowheads indicate trans-membrane segments determined experimentally. **B.** Accessibility of cysteine residues. Whole EAEC $\Delta tssM$ cells producing wild-type TssM (WT) or the indicated mutant proteins were labelled with the MPB probe, lysed, solubilized, and TssM and mutant proteins were immunoprecipitated with anti-FLAG-coupled beads. The precipitated material was subjected to

SDS-PAGE and Western blot analysis using anti-FLAG antibody (TssM detection, upper panel) and streptavidin coupled to alkaline phosphatase (MPB-labelled TssM detection, lower panel). Molecular weight markers are indicated on the left. **C.** Topology model for the EAEC TssM protein at the inner membrane. The localizations of the labelled and unlabelled cysteine residues are indicated by filled and open circles respectively. The cytoplasmic (TssM_{Cyto}) and periplasmic (TssM_{Peri}) domains are highlighted in green and orange respectively. The three trans-membrane segments identified by the accessibility studies are shown, with their membrane boundaries (in blue).

Figure 2. TssM_{Cyto} interaction network identified by bacterial two-hybrid analysis.

BTH101 reporter cells carrying pairs of plasmids producing the indicated T6SS proteins fused to the T18 or T25 domain of the *Bordetella* adenylate cyclase were spotted on X-Gal-IPTG indicator LB agar plates. Only the cytoplasmic (Cyto) or periplasmic (Peri) domains were used for membrane-anchored proteins. Controls include T18 and T25 fusions to TolB and Pal, two proteins that interact but unrelated to the T6SS.

Figure 3. Structural architecture of TssM_{Cyto}.

A. The cytoplasmic domain of TssM, TssM_{Cyto}, delimited by the trans-membrane helices 2 and 3 (TMH2 and TMH3 respectively) could be partitioned into an NTPase-like domain (NTP, blue) and a C-terminal extension (Ct, green). TssM_{Cyto/NTP} was modelled using HHpred based on the X-ray structure of the *Burkholderia thailandensis* EngB GTP-binding protein (PDB ID: **4DHE**). TssM_{Cyto/Ct} was modelled using SwissModel based on the X-ray structure of the C-terminal domain of the human DPY-30-like protein, a component of the histone methyltransferase complex (PDB ID: **3G36**). All images were made with Chimera.⁶⁶

Figure 4. The two TssM_{Cyto} sub-domains mediate interaction with two baseplate components.

A. Bacterial two-hybrid assay. BTH101 reporter cells carrying pairs of plasmids producing the indicated T6SS proteins fused to the T18 or T25 domain of the *Bordetella* adenylate cyclase were spotted on X-Gal-IPTG indicator LB agar plates. Only the cytoplasmic (c) or periplasmic (p) domains were used for membrane-anchored proteins. Controls include T18 and T25 fusions to TolB and Pal, two proteins that interact but unrelated to the T6SS. **B.** Co-immunoprecipitation assay. Soluble lysates from 5×10^{10} *E. coli* K12 W3110 cells producing FLAG-tagged TssM_{Cyto} (M_{Cyto_FL}), FLAG-tagged TssM_{Cyto/NTP} (NTP_{FL}) or FLAG-tagged TssM_{Cyto/Ct} (Ct_{FL}) and VSV-G-tagged TssG (TssG_V) or TssK (TssK_V) proteins were subjected to immunoprecipitation with anti-FLAG-coupled beads. The total soluble (Tot) and the immunoprecipitated (IP) material were separated by 12.5% acrylamide SDS-PAGE and immunodetected with anti-VSVG (TssG_V or TssK_V) monoclonal antibodies. Molecular weight markers (in kDa) are indicated on the left. The asterisk on the lower panel indicates a degradation product from the TssK protein.

Figure 5. Specific conserved motifs mediate TssM_{Cyto} dimerization and interaction with TssG

A. Bacterial two-hybrid assay. BTH101 reporter cells carrying pairs of plasmids producing the indicated T6SS proteins fused to the T18 or T25 domain of the *Bordetella* adenylate cyclase were spotted on X-Gal-IPTG indicator LB agar plates. Only the cytoplasmic (c) or periplasmic (p) domains were used for membrane-anchored proteins. Controls include T18 and T25 fusions to TolB and Pal, two proteins that interact but unrelated to the T6SS. **B.** Co-immunoprecipitation assay. Soluble lysates from 5×10^{10} *E. coli* K12 W3110 cells producing FLAG-tagged TssM_C WT (M_{Cyto_FL}) or mutants and VSV-G-tagged TssG (TssG_V) or TssM_{Cyto} (TssM_{Cyto_V}) proteins were subjected to immunoprecipitation with anti-FLAG-coupled beads. The total soluble (Tot) and the immunoprecipitated (IP) material were separated by 12.5% acrylamide SDS-PAGE and immunodetected with anti-FLAG (lower panels) and anti-VSV-G (upper panels) monoclonal antibodies. Molecular weight markers (in kDa) are indicated on the left.

Figure 6. TssM_{Cyto} oligomerization and interaction with the TssG baseplate are essential for T6SS function, sheath assembly and baseplate stability

A. Anti-bacterial assay. *E. coli* K-12 prey cells (W3110 *gfp*⁺, kan^R) were mixed with the indicated attacker cells, spotted onto Sci-1 inducing medium (SIM) agar plates and incubated for 4 hours at 37°C. The number of recovered *E. coli* prey cells is indicated in the graph (in log10 of colony-forming unit (cfu)). The circles indicate values from three independent assays, and the average is indicated by the bar. **B.** Image recordings of the TssB-sfGFP fusion protein in the indicated cells. Statistical analyses (*n*=number of sheath/ cell) are indicated under each strain. The number of cells studied per strain (*n*) is 150. The scale bare is 2 μm. **C.** Image recordings of the sfGFP-TssF fusion protein in the indicated cells. The scale bare is 2 μm. **D.** Statistical analyses of sfGFP-TssF in the indicated strains. Shown are box-and-whisker plots of the measured number of sfGFP-TssF foci per cell for each strain with the lower and upper boundaries of the boxes corresponding the 25% and 75% percentiles respectively. The black bold horizontal bar represents the median values for each strain and the whiskers represent the 10% and 90% percentiles. Outliers are shown as open circle. *n* indicates the number of cells analyzed per strain.

Figure 7. Schematic representation of the TssM_{Cyto} interaction network. **A.** Schematic representation of the TssJLM membrane complex (MC) and its interactions with the TssKEFG-VgrG baseplate complex (BC). The TssM_{Cyto/NTP} and TssM_{Cyto/Ct} and the TssL cytoplasmic domain (TssL_{Cyto}) are shown in blue, green and orange respectively. The interactions defined in this study are indicated by red arrows. Interactions determined previously²⁴ or in the accompanying article⁶⁹ are shown in blue dashed arrows. **B.** Schematic representation of a TssM_{Cyto} dimer (TssM_{Cyto/NTP} are shown in blue,

TssM_{Cyto/Ct} are shown in green). The cartoon highlights the TssM_{Cyto/Ct}-TssM_{Cyto/Ct} interface and the interaction with TssK, TssG and the cytoplasmic domain of TssL.

LEGEND TO SUPPLEMENTARY FIGURES

Supplementary Figure S1. Sequence alignment of the cytoplasmic domain from TssM homologues. T-COFFEE sequence alignment of the cytoplasmic domain of TssM of (from top to bottom) entero-aggregative *Escherichia coli* Sci-1 (EAEC 17-2), *Yersinia pseudotuberculosis* (IP31758) T6SS-5, *Klebsiella pneumoniae*, *Serratia* sp. (strain M24T3), *Citrobacter rodentium* (sp. A1), *Proteus penneri*, *Myxococcus xanthus* (DK1622), *Yersinia pseudotuberculosis* (IP31758)T6SS-3, *Salmonella enterica* Typhimurium (LT2), *Agrobacterium tumefaciens* (C58), *Pseudomonas aeruginosa* H1-T6SS (PA01), *Aeromonas hydrophila* (ATCC7966) and *Edwardsiella tarda*. Local homologies are indicated by a colour code (bad to good from blue to red). The Walker and GTP motifs usually associated with NTPases are indicated, as well as the motifs required for TssM_{Cyto} oligomerization and interaction with TssG. The numbering corresponds to the sequence of the EAEC Sci-1 TssM protein. The figure has been prepared with T-COFFEE.⁶⁷

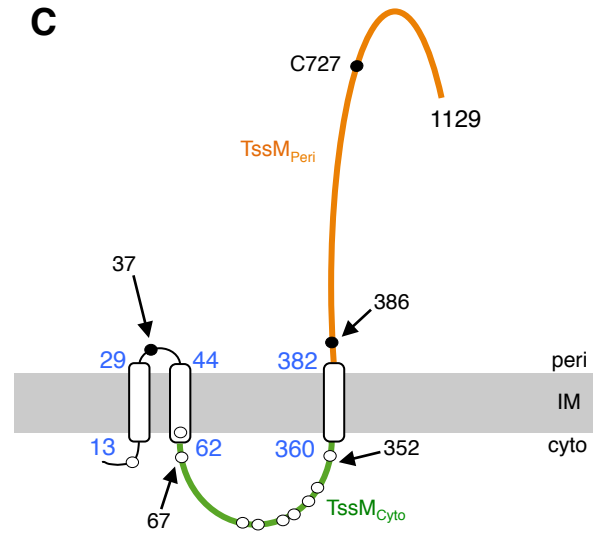
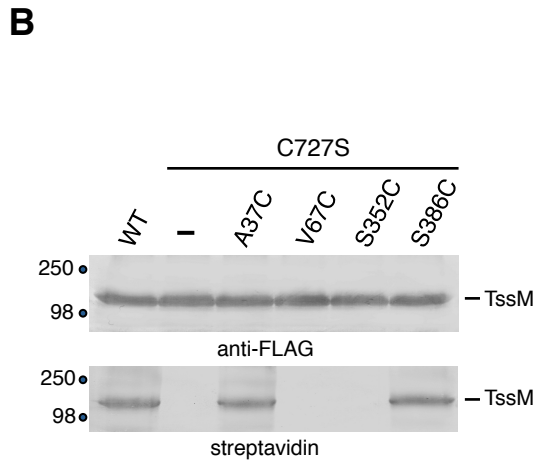
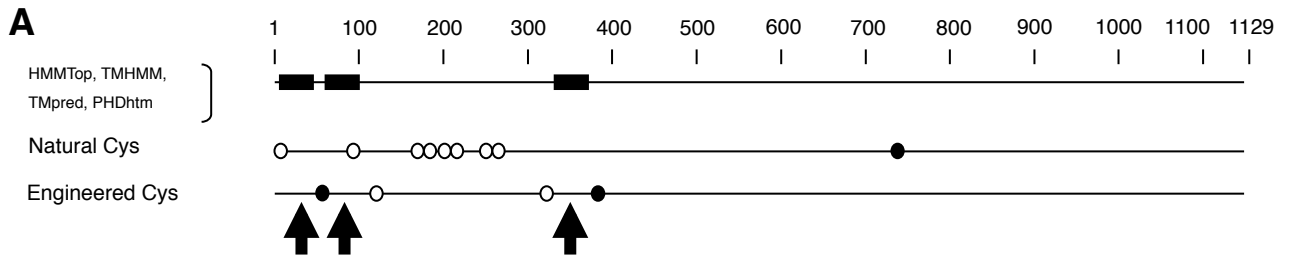
Supplementary Figure S2. Phylogenetic tree of TssM_{Cyto} homologs. Phylogenetic analysis of the TssM_{Cyto} sequences shown in Supplementary Fig. S1. The TssM cytoplasmic loops categorize into two phylogenetic clusters, one with complete NTPase domains (orange) and one with NTPase domains but with degenerated Walker motifs (blue). The figure has been prepared with phylogeny.fr.⁶⁸

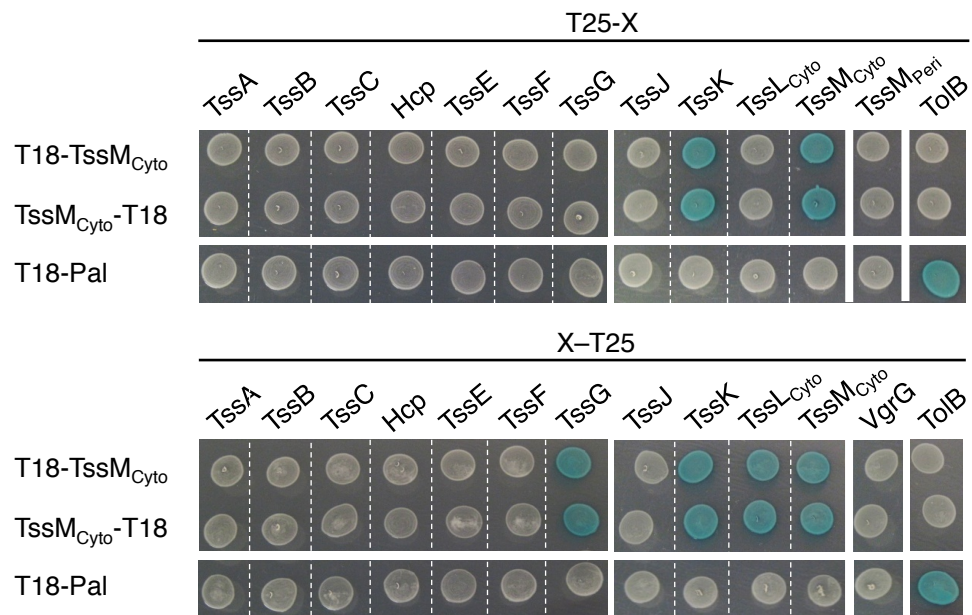
Supplementary Figure S3. Sequence homology of TssM_{Cyto} sub-domains. A. Sequence alignment of TssM_{Cyto/NTP} with *B. thailandensis* EngB. **B.** Sequence alignment of TssM_{Cyto/Ct} with DPY-30. Identical residues are coloured white and underlined in red while strongly similar residues are coloured red. The secondary structures of the EngB and DPY-30 proteins are indicated below the sequences, as well as the percentage of identity and similarity of these proteins with the corresponding TssM_{Cyto} sub-domains.

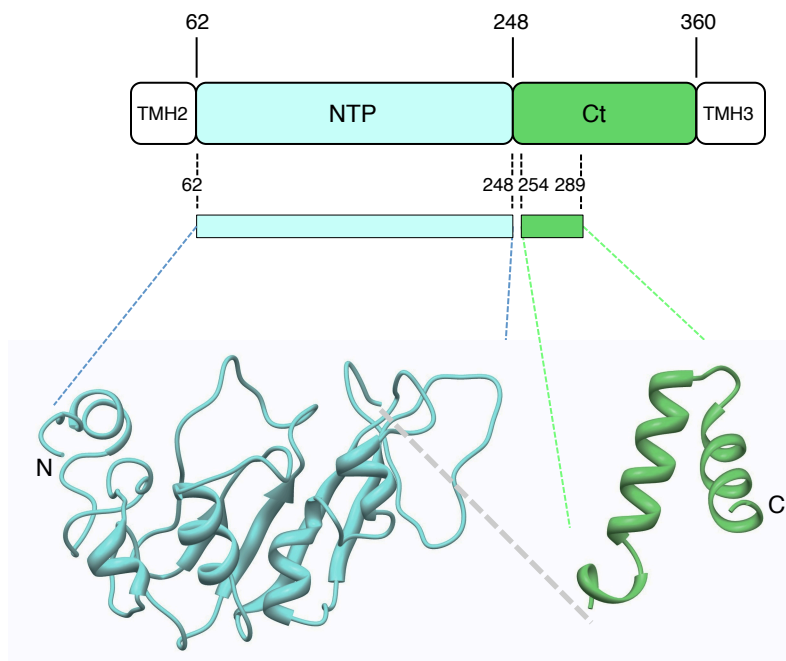
Supplementary Figure S4. Homology models of TssM_{Cyto} sub-domains. A. Homology models of the EAEC Sci-1 (blue, left panel) and *A. tumefaciens* (orange, right panel) TssM_{Cyto/NTP} domains based on the X-ray structure of the *Burkholderia thailandensis* EngB GTP-binding protein (PDB ID: 4DHE). The GTP-binding site of the *A. tumefaciens* TssM_{Cyto/NTP} is shown in sticks. The models have been generated with HHPred.³⁹ **B.** Homology model of the EAEC TssM_{Cyto/Ct} domain (green, left panel) based on the X-ray structure of the C-terminal domain of human DPY-30-like protein (PDB ID: 3G36). The residues involved in TssM_{Cyto/Ct} oligomerization are indicated by arrowheads. The model has been generated with Swiss-Model.⁶⁵ The structures of the DPY-30 C-terminal monomer and dimer

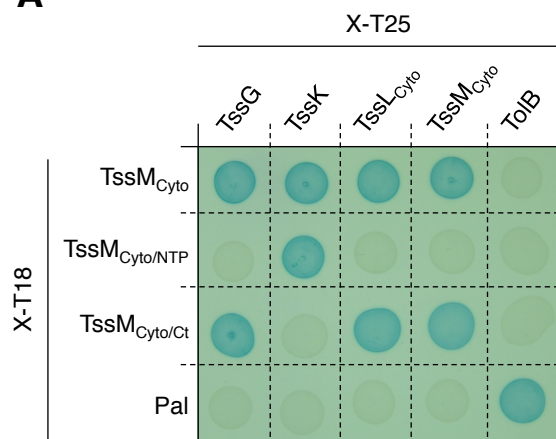
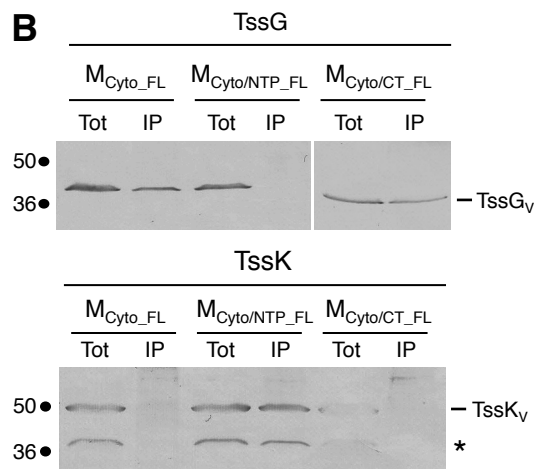
are shown in the middle and right panels respectively. The residues at the DPY-30 dimer interface are indicated by arrowheads. All images have been prepared with Chimera.⁶⁶

Supplementary Figure S5. Topologies of TssM protein homologues. Trans-membrane helix predictions of selected TssM proteins from the indicated strains. Trans-membrane helices are represented by black rectangles whereas TssM_{Cyto/NTP} and TssM_{Cyto/Ct} domains are indicated by blue and green lines respectively.





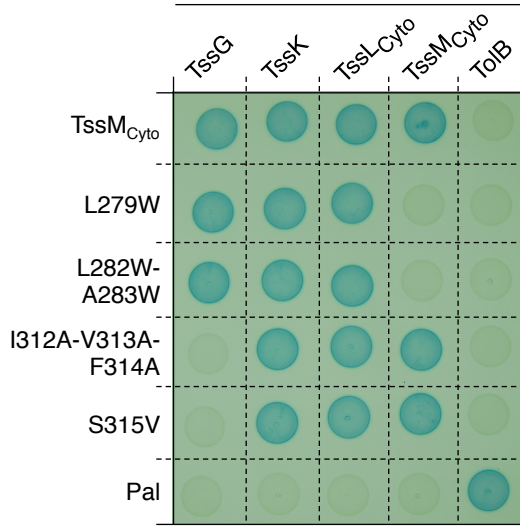


A**B**

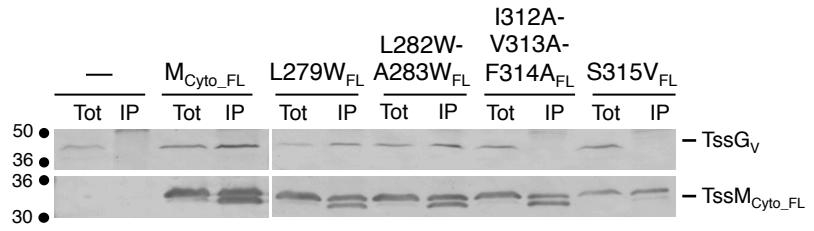
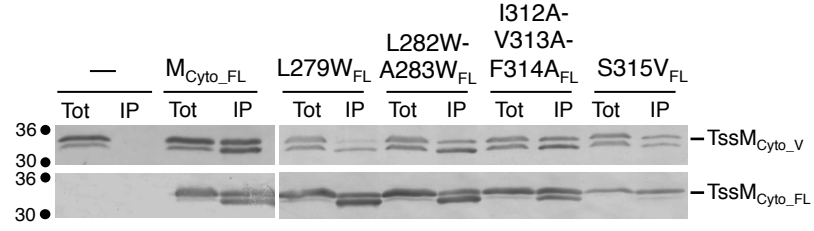
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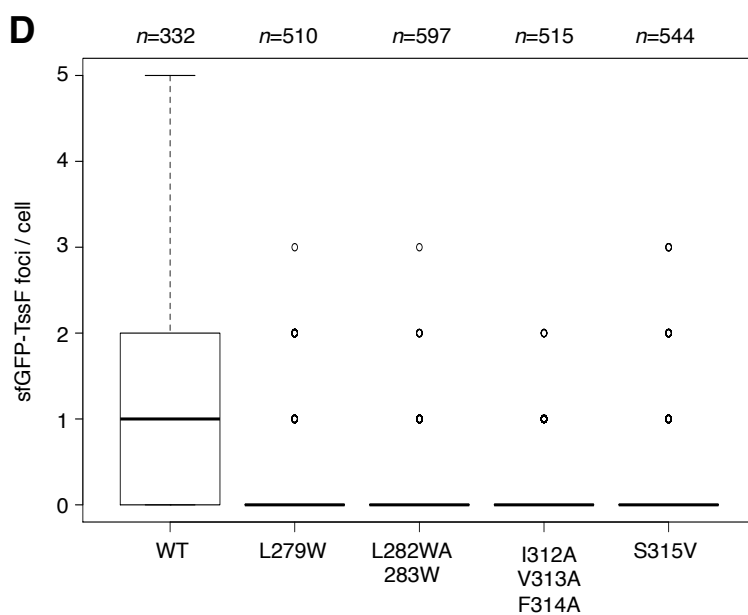
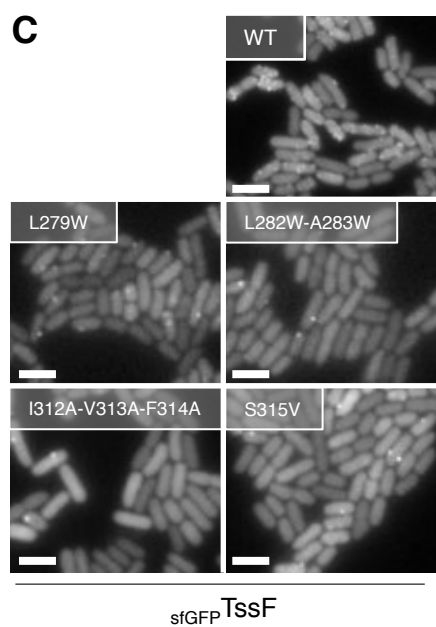
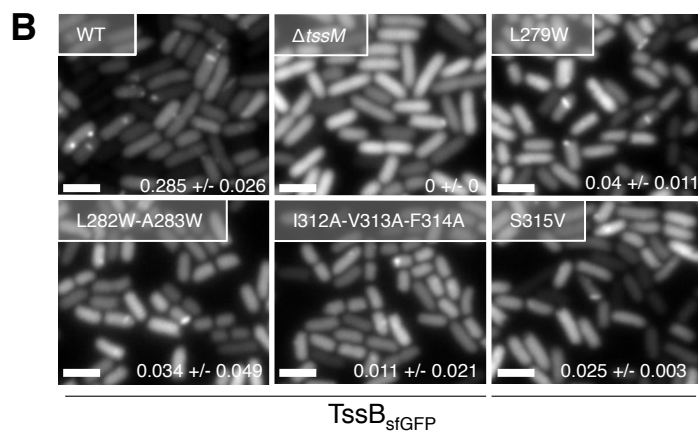
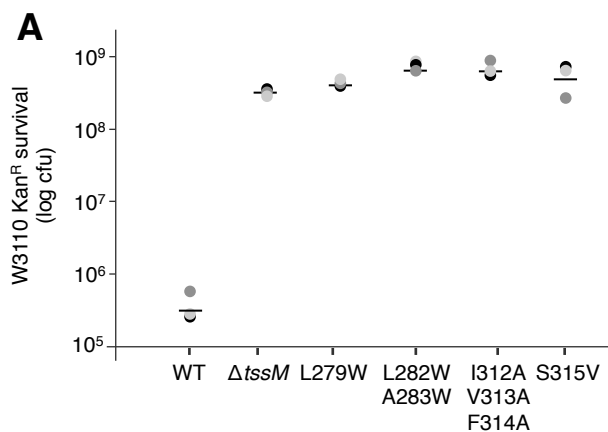
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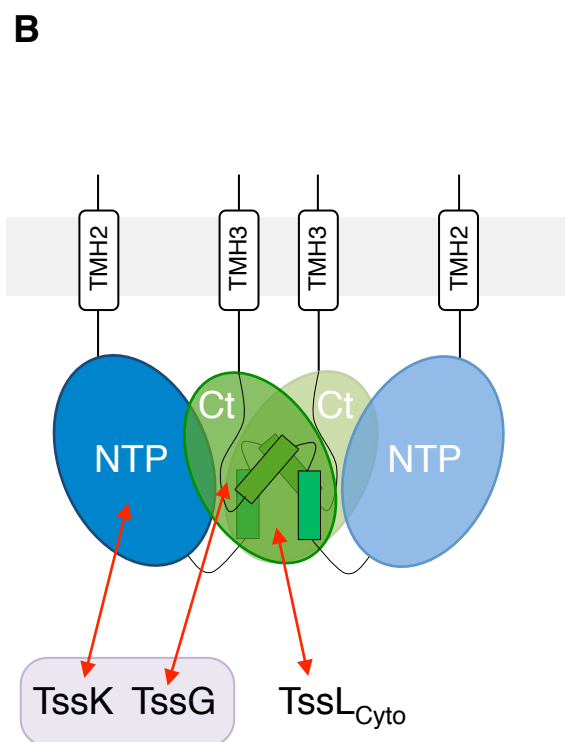
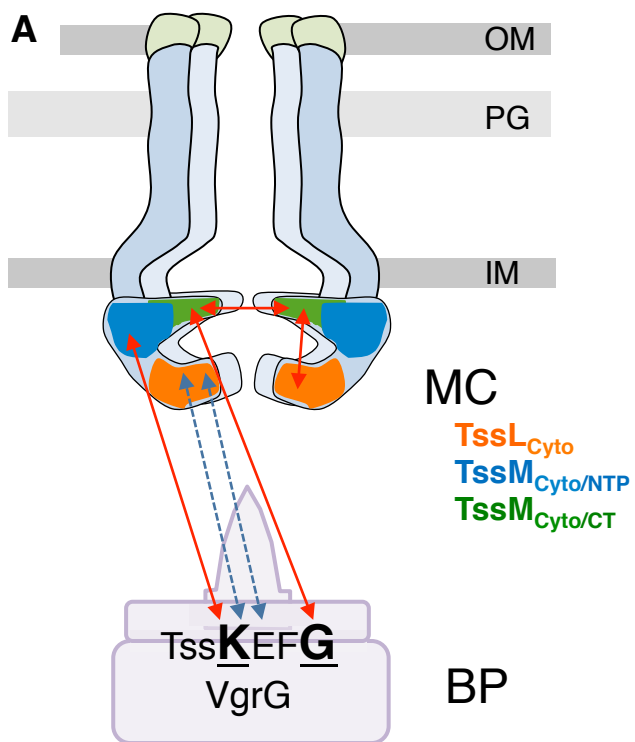
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**B**

TssG

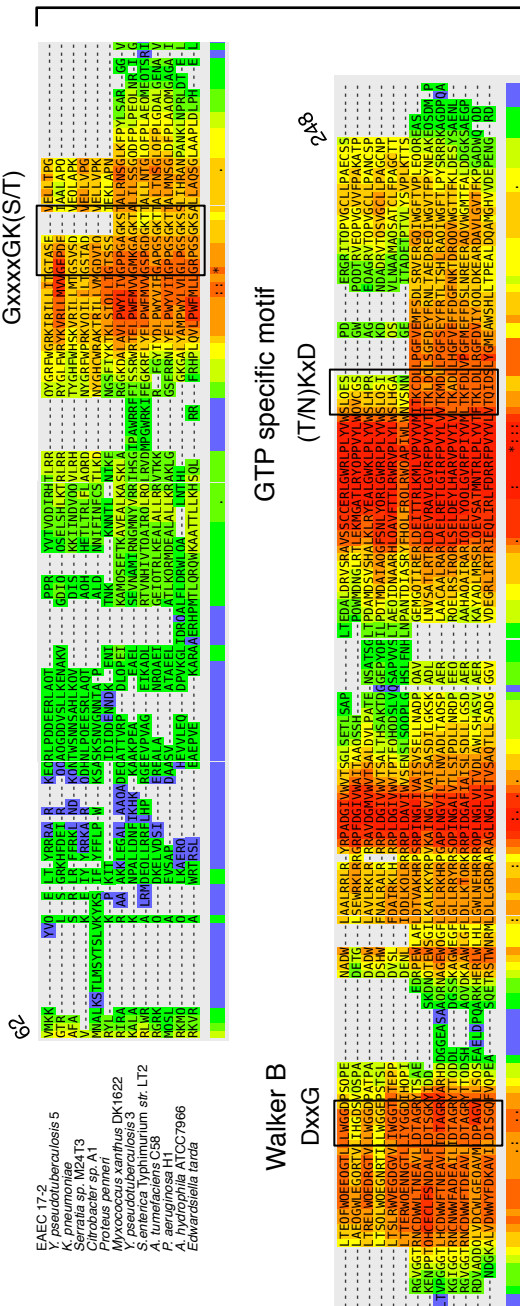
TssM_{Cyto}





Walker A
GxxxxGK(S/T)

NTPase
domain

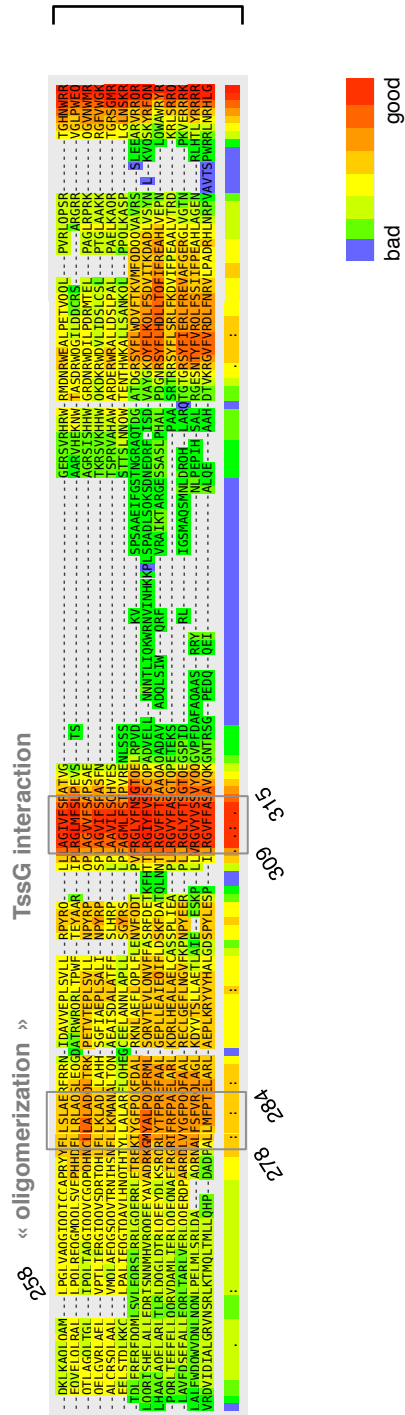


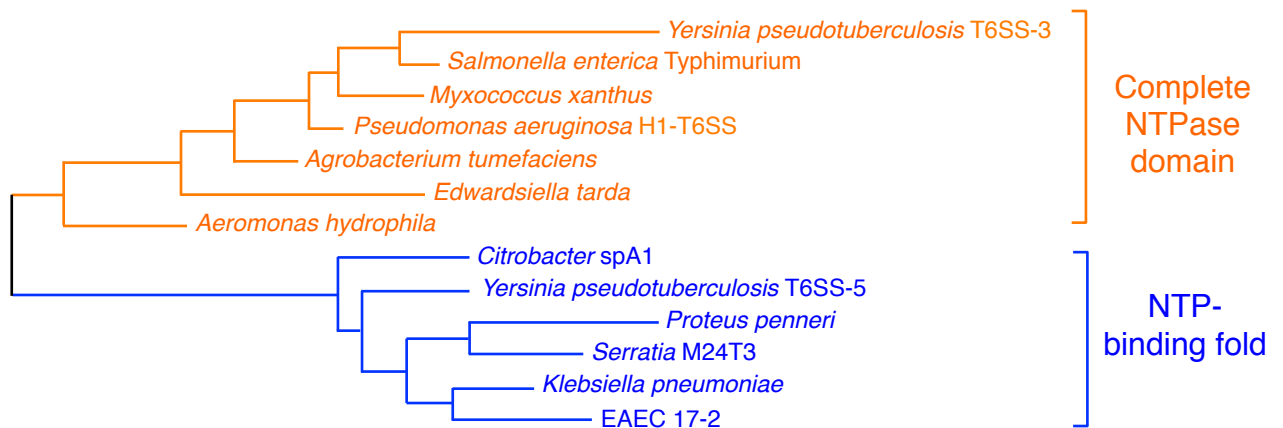
« oligomerization »

TssG interaction

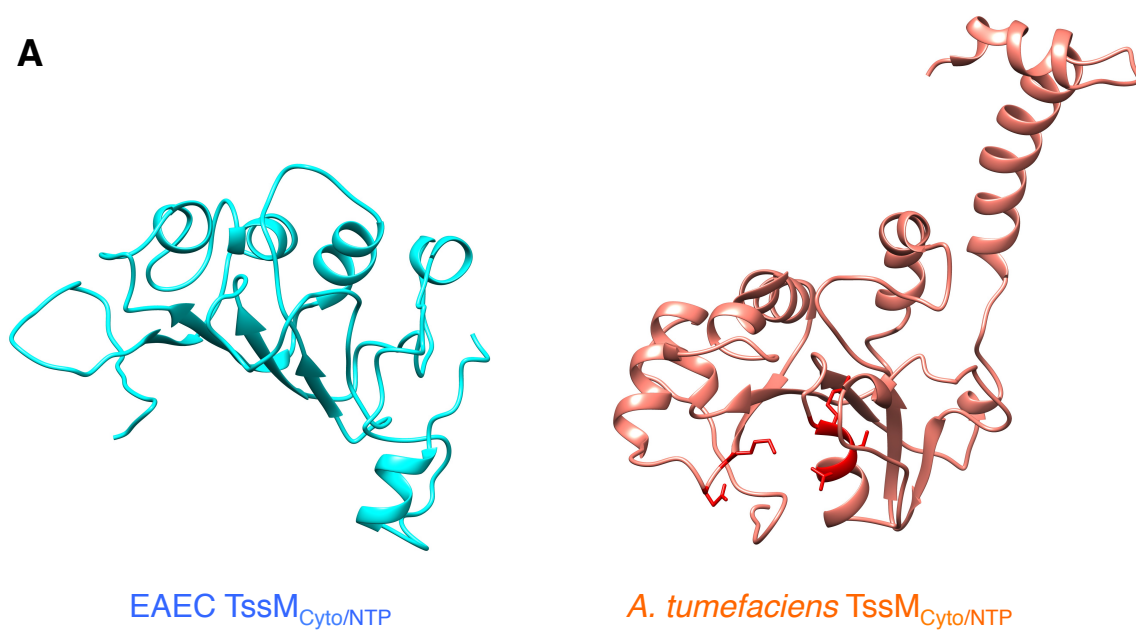
278 284 306 315

Cter
domain

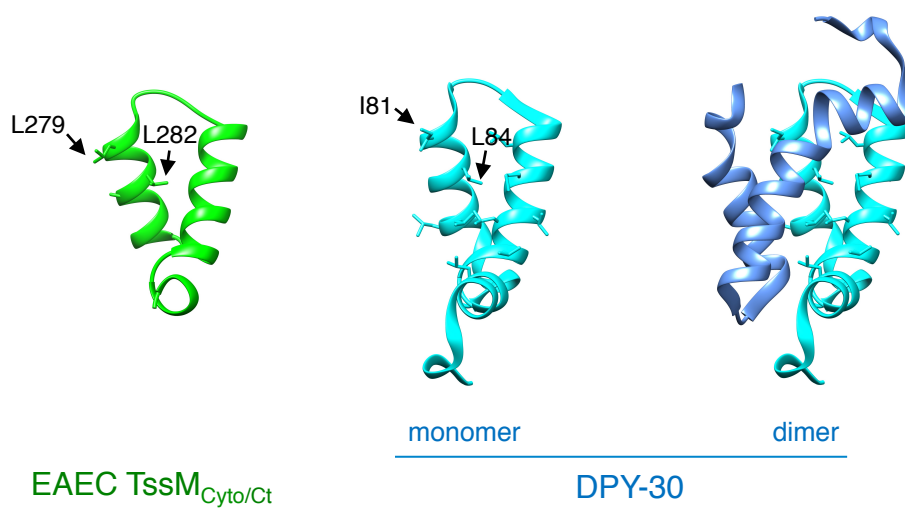


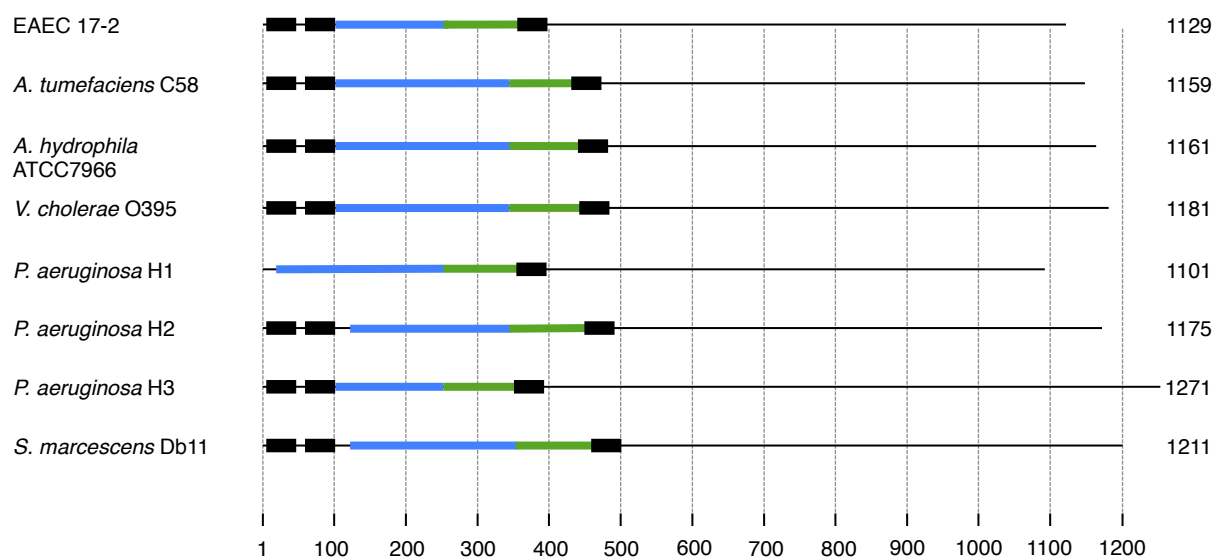


A



B





SUPPLEMENTAL DATA

**Molecular dissection of the interface between the Type VI secretion
TssM cytoplasmic domain and the TssG baseplate component.**

L. Logger, M.S. Aschtgen, M. Guérin, E. Cascales & E. Durand

Supplemental Table S1. Strains, plasmids and oligonucleotides used in this study.

Strains

Strains	Description and genotype	Source
<i>E. coli</i> K-12		
DH5 α	F-, Δ (<i>argF-lac</i>)U169, <i>phoA</i> , <i>supE44</i> , Δ (<i>lacZ</i>)M15, <i>relA</i> , <i>endA</i> , <i>thi</i> , <i>hsdR</i>	New England Biolabs
W3110	F-, lambda- IN(<i>rrnD-rrnE</i>)1 <i>rph</i> -1	Laboratory collection
BTH101	F-, <i>cya</i> -99, <i>araD</i> 139, <i>galE</i> 15, <i>galK</i> 16, <i>rpsL</i> 1 (<i>Str r</i>), <i>hsdR</i> 2, <i>mcrA</i> 1, <i>mcrB</i> 1.	Karimova <i>et al.</i> , 1998
Enteroaggregative <i>E. coli</i>		
17-2	WT enteroaggregative <i>Escherichia coli</i>	A. Darfeuille-Michaud
17-2 Δ <i>tssM</i>	17-2 deleted of the <i>tssM</i> gene of the <i>sciI</i> T6SS gene cluster	Aschtgen <i>et al.</i> , 2010
17-2 <i>tssB-gfp</i>	<i>gfp</i> inserted upstream the stop codon of <i>tssB</i> in 17-2	This study
17-2 Δ <i>tssM tssB-gfp</i> Ω Kan	<i>gfp</i> inserted upstream the stop codon of <i>tssB</i> in 17-2 Δ <i>tssM</i>	This study
17-2 <i>tssB-gfp tssM</i> -L279W	Chromosomal point mutation Leu279-to-Trp substitution of <i>tssM</i> in 17-2 <i>tssB-gfp</i>	This study
17-2 <i>tssB-gfp tssM</i> -L282W-A283W	Chromosomal point mutation Leu282 and Ala283-to-Trp substitutions of <i>tssM</i> in 17-2 <i>tssB-gfp</i>	This study
17-2 <i>tssB-gfp tssM</i> -I312A-V313A-F314A	Chromosomal point mutation Ile312 and Val313 and Phe314-to-Ala substitutions of <i>tssM</i> in 17-2 <i>tssB-gfp</i>	This study
17-2 <i>tssB-gfp tssM</i> -S315W	Chromosomal point mutation Ser315-to-Val substitution of <i>tssM</i> in 17-2 <i>tssB-gfp</i>	This study
17-2 <i>gfp-tssF</i>	<i>gfp</i> inserted downstream the start codon of <i>tssF</i> in 17-2	Brunet, Zoued <i>et al.</i> , 2015
17-2 Δ <i>tssM gfp-tssF</i> Ω Kan	<i>gfp</i> inserted downstream the start codon of <i>tssF</i> in 17-2 Δ <i>tssM</i>	This study
17-2 <i>gfp-tssF tssM</i> -L279W	Chromosomal point mutation Leu279-to-Trp substitution of <i>tssM</i> in 17-2 <i>gfp-tssF</i>	This study
17-2 <i>gfp-tssF tssM</i> -L282W-A283W	Chromosomal point mutation Leu282 and Ala283-to-Trp substitutions of <i>tssM</i> in 17-2 <i>gfp-tssF</i>	This study
17-2 <i>gfp-tssF tssM</i> -I312A-V313A-F314A	Chromosomal point mutation Ile312 and Val313 and Phe314-to-Ala substitutions of <i>tssM</i> in 17-2 <i>gfp-tssF</i>	This study
17-2 <i>gfp-tssF tssM</i> -S315W	Chromosomal point mutation Ser315 to Val substitution of <i>tssM</i> in 17-2 <i>gfp-tssF</i>	This study

Plasmids

Vectors	Description	Source
Vectors for chromosomal insertions		
pKD4	One-step gene inactivation vector, Kan ^R	Datsenko & Wanner, 2000
pgfp-KD4	superfolder- <i>gfp</i> cloned upstream the 3' –FRT site of pKD4. Used for chromosomal C-terminal GFP fusion	Brunet, Zoued <i>et al.</i> , 2015
pKOBEG	Recombination vector, phage λ <i>rec$\gamma$$\beta$</i> operon under the control of the pBAD promoter, Cm ^R	Chaverroche <i>et al.</i> , 2000
Expression vectors		
pUA66- <i>rrnB</i>	<i>P_{rrnB} ::gfpmut2</i> transcriptional fusion in pUA66	Zaslaver <i>et al.</i> , 2006
pASK-IBA37(+)	cloning vector, <i>Ptet</i> , fl origin, Amp ^R	IBA Technology
pASK-IBA37- _{FLAG} TssM	<i>sci-1 tssM</i> carrying N-terminal FLAG tag cloned into pASK-IBA37	Aschtgen <i>et al.</i> , 2010
pIBA37- _{FLAG} TssM-C727S	Introduction of the TssM Cys727-to-Ser substitution into pIBA37- _{FLAG} TssM	This study
pIBA37- _{FLAG} TssM-C727S-A37C	Introduction of the TssM Ala37-to-Cys substitution into pIBA37- _{FLAG} TssM-C727S	This study
pIBA37- _{FLAG} TssM-C727S-V67C	Introduction of the TssM Val67-to-Cys substitution into pIBA37- _{FLAG} TssM-C727S	This study
pIBA37- _{FLAG} TssM-C727S-S352C	Introduction of the TssM Ser352-to-Cys substitution into pIBA37- _{FLAG} TssM-C727S	This study
pIBA37- _{FLAG} TssM-C727S-S386C	Introduction of the TssM Ser386-to-Cys substitution into pIBA37- _{FLAG} TssM-C727S	This study
pIBA37-TssM _{Cyto} - _{FLAG}	<i>sci-1 tssM</i> residues 62-360 cloned into pASK-IBA37, C-terminal FLAG epitope	Zoued <i>et al.</i> , 2013
pIBA37-TssM _{Cyto} -NTP _{FLAG}	<i>sci-1 tssM</i> residues 62-273 cloned into pASK-IBA37, C-terminal FLAG epitope	This study
pIBA37-TssM _{Cyto} -Cter _{FLAG}	<i>sci-1 tssM</i> residues 274-360 cloned into pASK-IBA37, C-terminal FLAG epitope	This study
pIBA37-TssM _{Cyto} -L279W _{FLAG}	Introduction of the TssM Leu279-to-Trp substitution into pIBA37-TssM _{Cyto} _{FLAG}	This study
pIBA37-TssM _{Cyto} -L282W-A283W _{FLAG}	Introduction of the TssM Leu282 and Ala283-to-Trp substitutions into pIBA37-TssM _{Cyto} _{FLAG}	This study
pIBA37-TssM _{Cyto} -I312A-V313A-F314A _{FLAG}	Introduction of the TssM Ile312 and Val313 and Phe314-to-Ala substitutions into pIBA37-TssM _{Cyto} _{FLAG}	This study
pIBA37-TssM _{Cyto} -S315V _{FLAG}	Introduction of the TssM Ser 315-to-Val substitution into pIBA37-TssM _{Cyto} _{FLAG}	This study
pBAD33	cloning vector, P15A origin, <i>Para</i> , <i>araC</i> , Cm ^R	Guzman <i>et al.</i> , 1995
pBAD33-TssM _{Cyto} -VSV-G	<i>sci1 tssM_{Cyto}</i> cloned into pBAD33, C-terminal VSV-G epitope	This study
pBAD33-TssG _{VSV-G}	<i>sci1 tssG</i> cloned into pBAD33, C-terminal VSV-G epitope	Brunet, Zoued <i>et al.</i> , 2015
pBAD33-TssK _{VSV-G}	<i>sci1 tssK</i> cloned into pBAD33, C-terminal VSV-G epitope	Brunet, Zoued <i>et al.</i> , 2015

Bacterial Two-Hybrid vectors

pT18-FLAG	Bacterial two-hybrid vector, ColE1 origin, <i>Plac</i> , T18 fragment of <i>Bordetella pertussis</i> CyaA, Amp ^R	Battesti & Bouveret, 2008
pTssM _{Cyto} -T18	Cytoplasmic region of <i>tssM</i> (TssM ₆₂₋₃₆₀ fragment) cloned upstream T18 in pT18-FLAG	Zoued <i>et al.</i> , 2013
pT18-TssM _{Cyto}	Cytoplasmic region of <i>tssM</i> (TssM ₆₂₋₃₆₀ fragment) cloned downstream T18 in pT18-FLAG	Zoued <i>et al.</i> , 2013
pTssM _{Cyto} -NTP-T18	<i>sci-1 tssM</i> residues 62-273 cloned upstream T18 into pT18-FLAG	This study
pTssM _{Cyto} -Cter-T18	<i>sci-1 tssM</i> residues 274-360 cloned upstream T18 into pT18-FLAG	This study
pTssM _{Cyto} -L279W-T18	Introduction of the TssM _{Cyto} Leu279-to-Trp substitution into pTssM _{Cyto} -T18	This study
pTssM _{Cyto} -L282W-A283W-T18 This study	Introduction of the TssM _{Cyto} Leu282 and Ala283-to-Trp substitutions into pTssM _{Cyto} -T18	
pTssM _{Cyto} -I312A-V313A-F314A-T18	Introduction of the TssM _{Cyto} Ile312 and Val313 and Phe314-to-Ala substitutions into pTssM _{Cyto} -T18	This study
pTssM _{Cyto} -S315V-T18	Introduction of the TssM _{Cyto} Ser315-to-Val substitution into pTssM _{Cyto} -T18	This study
pT18-Pal	<i>pal</i> cloned downstream the T18 coding sequence in pT18-FLAG	Battesti & Bouveret, 2008
pT25-FLAG	Bacterial two-hybrid vector, P15A origin, <i>Plac</i> , T25 fragment of <i>Bordetella pertussis</i> CyaA, Kan ^R	Battesti & Bouveret, 2008
pT25-TssA	<i>tssA</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssA-T25	<i>tssA</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssB	<i>tssB</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssB-T25	<i>tssB</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssC	<i>tssC</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssC-T25	<i>tssC</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-Hcp	<i>hcp</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pHcp-T25	<i>hcp</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssE	<i>tssE</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssE-T25	<i>tssE</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssF	<i>tssF</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssF-T25	<i>tssF</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssG	<i>tssG</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssG-T25	<i>tssG</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-VgrG	<i>vgrG</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pVgrG-T25	<i>vgrG</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssJ _{sol}	<i>tssJ_{sol}</i> (TssJ ₂₋₁₅₂ fragment of the processed form) cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssJ _{sol} -T25	<i>tssJ_{sol}</i> (TssJ ₂₋₁₅₂ fragment of the processed form) cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssK	<i>tssK</i> cloned downstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssK-T25	<i>tssK</i> cloned upstream the T25 coding sequence in pT25-FLAG	Zoued <i>et al.</i> , 2013
pT25-TssL _{Cyto}	Cytoplasmic fragment of <i>tssL</i> (TssL ₁₋₁₈₄ fragment) cloned downstream the T25 coding sequence in pT25-FLAG	Durand <i>et al.</i> , 2012
pTssL _{Cyto} -T25	Cytoplasmic fragment of <i>tssL</i> (TssL ₁₋₁₈₄ fragment) cloned upstream the T25 coding sequence in pT25-FLAG	Durand <i>et al.</i> , 2012
pT25-TssM _{Cyto}	Cytoplasmic fragment of <i>tssM</i> (TssM ₆₂₋₃₆₀ fragment) cloned downstream T25 in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssM _{Cyto} -T25	Cytoplasmic fragment of <i>tssM</i> (TssM ₆₂₋₃₆₀ fragment) cloned upstream T25 in pT25-FLAG	Zoued <i>et al.</i> , 2013

pT25-TssM _{peri}	Periplasmic région of <i>tssM</i> (TssM ₃₈₆₋₁₁₂₉ fragment) cloned downstream T25 in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTssM _{peri} -T25	Periplasmic région of <i>tssM</i> (TssM ₃₈₆₋₁₁₂₉ fragment) cloned upstream T25 in pT25-FLAG	Zoued <i>et al.</i> , 2013
pTolB-T25	<i>tolB</i> cloned upstream the T25 coding sequence in pT25-FLAG	Battesti & Bouveret, 2008

Chromosomal mutagenesis vectors

pKO3	<i>sacB</i> , <i>repA</i> (pSC101 ^{tr}), M13 origin, Cm ^R	Link <i>et al.</i> , 1997
pKO3- <i>tssM</i>	Region of <i>tssM</i> cloned into the pKO3 vector	This study
pKO3- <i>tssM</i> -L279W	Introduction of the TssM Leu279-to-Trp substitution into pKO3- <i>tssM</i>	This study
pKO3- <i>tssM</i> -L282W-A283W	Introduction of the TssM Leu282 and Ala283-to-Trp substitutions into pKO3- <i>tssM</i>	This study
pKO3- <i>tssM</i> -I312A-V313A-F314A	Introduction of the TssM Ile312 and Val313 and Phe314-to-Ala substitutions into pKO3- <i>tssM</i>	This study
pKO3- <i>tssM</i> -S315W	Introduction of the TssM Ser315-to-Val substitution into pKO3- <i>tssM</i>	This study

Oligonucleotides

Name	Destination	Sequence (5' → 3')
For site-directed mutagenesis ^a		
M-C727S-5	pASK-IBA37- _{FLAG} TssM	GAATACGCTGGCGGTTTCAGGGATCCACTGGCCAGCCCCGGGAAG
M-C727S-3	pASK-IBA37- _{FLAG} TssM	CTTCCCGGGGCTGGCCAGTGGATCCCTGAACCGCCAGCGTATTC
M-A36C-5	pIBA37- _{FLAG} TssM-C727S	GATAACCCGATATGGAGCATACCTAGGGTGTGAAACACGCCGGGATCAAATAC
M-A36C-3	pIBA37- _{FLAG} TssM-C727S	GTATTTGATCCCGGCGTGTTCACACCCTAGGTATGCTCCATATCGGGTTATC
M-V67C-5	pIBA37- _{FLAG} TssM-C727S	CTGCCTGTGATGAAAAAATATTGTCAGGAAGTACATATCGAC
M-V67C-3	pIBA37- _{FLAG} TssM-C727S	GTCGATATGTCAGTTCCTGACAATATTTTTTCATCACAGGVAG
M-S352C-5	pIBA37- _{FLAG} TssM-C727S	AATTACCTGTCCGGCTACAGCCCTGCAGGACAGGTCATAACTGGCGCAG
M-S352C-3	pIBA37- _{FLAG} TssM-C727S	CTGCGCCAGTTATGACCTGTCCCTGCAGGGCTGTAGCCGGACAGGTAATTG
M-S386C-5	pIBA37- _{FLAG} TssM-C727S	GGTGGTTTCCTTTCTGGCAAATCGATGTCTGGTTGCTGAAGTCAG
M-S386C-3	pIBA37- _{FLAG} TssM-C727S	CTGTACTTCAGCAACCAGACATCGATTGCGCAGAAAGGAAACCACC
A-L279W	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	CCCTCGGTATTATTTTTTGGTTGTCGCTGGCAGAGC
B-L279W	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	ATAATACCGAGGGGACAGCATATCTGCTG
A-L282W-A283W	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	CCCTCGGTATTATTTTTTGTGTCGTGGTGGGAGCGATTTCAGACGAA
B- L282W-A283W	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	ATAATACCGAGGGGACAGCATATCTGCTG
A-I312A-V313A-F314A	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	GTTGTTGCTGGCGGGTGC GGCGGCGAGTCCGGCAACTGTGCG
B- I312A-V313A-F314A	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	CGCCAGCAACAACTGACGGTATGGACGGAG
A- S315V	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	GTTGTTGCTGGCGGGTATTGTTTTCTGCGCGGCAACTGTGCGGCG
B- S315V	pTssM _{Cyto} -T18, pIBA37-TssM _{Cyto_FL} or pKO3- <i>tssM</i>	CGCCAGCAACAACTGACGGTATGGACGGAG
For plasmid construction ^{b,c,d}		
5-pIBA37-TssM _{Cyto} -NTP _{FLAG}	insertion of <i>tssM</i> ₆₂₋₂₇₃ fragment into pASK-IBA37	<u>GACAAAAATCTAGAAATAATTTTGTTTAACTTTAAGAAGGAGATATACA</u> <u>AATGGTGATGAAAAAATATGTTTCAGGAAGTACATATC</u>
3-pIBA37-TssM _{Cyto} -NTP _{FLAG}	insertion of <i>tssM</i> ₆₂₋₂₇₃ fragment into pASK-IBA37	<u>GATGGTGATGGTGATGCGATCCTCTGCTAGCTTATTTATCATCGTCGTCTT</u> <u>TATAATCGGCACAGCATATCTGCTGTATCCC</u>
5- pIBA37-TssM _{Cyto} -CT _{FLAG}	insertion of <i>tssM</i> ₂₇₄₋₃₆₀ fragment into pASK-IBA37	<u>GACAAAAATCTAGAAATAATTTTGTTTAACTTTAAGAAGGAGATATACA</u> <u>AATGCCTCGGTATTATTTTTTGTGTCGCTGGC</u>
3- pIBA37-TssM _{Cyto} -CT _{FLAG}	insertion of <i>tssM</i> ₂₇₄₋₃₆₀ fragment into pASK-IBA37	<u>GATGGTGATGGTGATGCGATCCTCTGCTAGCTTACTTGTTCATCGTCGTCTT</u> <u>TATAATCTCTGCGCCAGTTATGACCTGTGCGGGA</u>

T25T18C-5-TssM _{Cyto} -NTP	insertion of <i>tssM</i> ₆₂₋₂₇₃ fragment downstream the T18 domain	<u>CGGATAACAATTTACACAGGAAACAGCTATGACCATGGTGAT</u> <u>GAAAAAATATGTTTCAGGAACTGACATATCG</u>
T18C-3- TssM _{Cyto} -NTP	insertion of <i>tssM</i> ₆₂₋₂₇₃ fragment downstream the T18 domain	<u>CCTCGCTGGCGGCTAAGCTTGGCGTAATGGCACAGCATATCTG</u> <u>CTGTATCCC</u>
T25T18C-5-TssM _{Cyto} -CT	insertion of <i>tssM</i> ₂₇₄₋₃₆₀ fragment downstream the T18 domain	<u>CGGATAACAATTTACACAGGAAACAGCTATGACCATGCCTCG</u> <u>GTATTATTTTTTGTGTCGCTGGC</u>
T18C-3- TssM _{Cyto} -CT	insertion of <i>tssM</i> ₂₇₄₋₃₆₀ fragment downstream the T18 domain	<u>CCTCGCTGGCGGCTAAGCTTGGCGTAATTCTGCGCCAGTTATGA</u> <u>CCTGTGCG</u>
5-pBAD-TssM _{Cyto}	insertion of <i>tssM</i> ₂₇₄₋₃₆₀ fragment into pBAD33	<u>CTCTCTACTGTTTCTCCATACCCGTTTTTTTTGGGCTAGCAGGAGGTATTAC</u> <u>ACCATGGTGATGAAAAAATATGTTTCAGGAACTGACATATCGAC</u>
3-pBAD-TssM _{Cyto} -VSVG	insertion of <i>tssM</i> ₆₂₋₃₆₀ fragment into pBAD33	<u>GGTCGACTCTAGAGGATCCCCGGGTACCTTATTTCTTAATCTATTCA</u> <u>ATATCTGTATATCTGCGCCAGTTATGACCTGTGCGGGA</u>
5-pKO3-BamHI-TssM _{Cyto}	insertion of <i>tssM</i> fragment into pKO3	<u>GACCGGATCCCGACGCAGGGCCCGTAAG</u>
3-pKO3-SalI-TssM _{Cyto}	insertion of <i>tssM</i> fragment into pKO3	<u>CCCGGTCTGACCTGTGGCCATTCCCGCATGAG</u>

For strain construction ^c

5- <i>tssB</i> - <i>gfp</i>	insertion of <i>gfp-mut2</i> at the 3' end of <i>tssB</i>	<u>CCGGCACTGAGTCAGACGCTGCGTGATGAACTGCGTGCACTGGTGCCGGA</u> <u>AAAGGCGGCAGCGCCGCGGAGGG</u>
3- <i>tssB</i> - <i>gfp</i>	insertion of <i>gfp-mut2</i> at the 3' end of <i>tssB</i>	<u>GCAACGTTCTTTCTTTCTGTACAGACATCAGCATTTTCTCTCGTAATCCG</u> <u>TTAAACATATGAATATCCTCCTTAGTTCCTATTCCGAAGTTCC</u>

^a Mutagenesized codon in **Bold**.

^b Sequence annealing on the target plasmid underlined.

^c FLAG or VSV-G epitope coding sequence *italicized*.

^d Restriction sites in **Bold**.

^e Sequences corresponding to the downstream and upstream regions of the gene to be inserted underlined