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23 Abstract

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25 Bacteria have evolved macromolecular machineries that secrete effectors and toxins to survive and 26 thrive in diverse environments. The type VI secretion system (T6SS) is a contractile machine that is 27 related to Myoviridae phages. The T6SS is composed of a baseplate that contains a spike onto which 28 an inner tube is built, surrounded by a contractile sheath. Unlike phages that are released to and act 29 in the extracellular medium, the T6SS is an intracellular machine inserted in the bacterial membranes 30 by a trans-envelope complex. This membrane complex (MC) comprises three proteins: TssJ, TssL and 31 TssM. We previously reported the low-resolution negative stain electron microscopy structure of the 32 enteroaggregative Escherichia coli MC and proposed a rotational 5-fold symmetry with a 33 TssJ:TssL:TssM stoichiometry of 2:2:2. Here, cryo-electron tomography analysis of the T6SS MC 34 confirmed the 5-fold symmetry in situ and identified the regions of the structure that insert into the 35 bacterial membranes. A high resolution model obtained by single particle cryo-electron microscopy 36 reveals its global architecture and highlights new features: five additional copies of TssJ, yielding a 37 TssJ:TssL:TssM stoichiometry of 3:2:2, a 11-residue loop in TssM, protruding inside the lumen of the

38 MC and constituting a functionally important periplasmic gate, and hinge regions. Based on these 39 data, we revisit the model on the mechanism of action of the MC during T6SS assembly and function. 40

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42 Introduction:

43 In a competitive environment, the ability to communicate and outcompete neighbours 44 provides bacteria with key advantages to survive. The type VI secretion system (T6SS) is a 45 macromolecular complex involved in the release of toxins that disrupt essential functions in 46 competitor cells (Russell et al, 2014). The T6SS is associated with increased survival and 47 pathogenicity in bacteria expressing it (Zhao et al, 2018). The T6SS is composed of 13-14 core 48 proteins (Boyer et al, 2009), usually encoded in the same locus in the genome (Mougous et al, 2006). 49 The T6SS assembles a molecular spring-loaded dagger, which punctures the target cell to secrete 50 fully folded effector proteins into neighbouring bacteria (Russell et al, 2011) or eukaryotic hosts 51 (Barker et al, 2009). The full assembly consists of the trans-envelope TssJLM membrane complex 52 (MC) (Durand et al, 2015) that tethers the TssKFGE-VgrG baseplate (Brunet et al, 2015)(No Title), 53 onto which the tail polymerize. This tail comprises the inner tube made of stacks of Hcp hexamers 54 wrapped by the TssBC sheath proteins that polymerize in a helical conformation (Brunet et al, 2014; 55 Clemens et al, 2015; Kudryashev et al, 2015; Chang et al, 2017) and tipped by the spike VgrG which 56 can be sharpened by the PAAR protein (Renault et al, 2018; Shneider et al, 2013). Effectors are either 57 associated within the Hcp lumen, or directly or indirectly bound to the VgrG or PAAR spike 58 (Unterweger et al, 2017; Silverman et al, 2013; Shneider et al, 2013; Flaugnatti et al, 2016a; Quentin 59 et al, 2018). Upon contact with a neighbouring cell, unknown signals trigger the contraction of the 60 sheath causing the tube and spike to pierce the membranes and secrete the effectors (Basler et al, 61 2012).

62 While the baseplate, tube and sheath proteins are conserved among contractile injection 63 systems, the MC is specific to the T6SS. TssJ is an outer membrane lipoprotein (Aschtgen et al, 64 2008a) that positions first at the site of assembly, and then recruits TssM and TssL (Durand et al, 65 2015). TssM and TssL are two inner membrane proteins that share homology with two accessory 66 subunits associated with T4SSb, IcmF and IcmH/DotU (Aschtgen et al, 2012; Ma et al, 2009; Durand 67 et al, 2012; Logger et al, 2016). Not only does the MC anchor the baseplate to the inner membrane, 68 but it would also serves as a channel to allow the passage of the tail tube/spike and maintain the 69 integrity of the attacking cell during the translocation of the inner tube (Durand et al, 2015). The 70 different subunits and the MC have been extensively biochemically characterised and several crystal 71 structures of the components are available from various bacterial species (Felisberto-Rodrigues et al, 72 2011; Durand et al, 2012; Rao et al, 2011; Robb et al, 2012; Chang & Kim, 2015; Durand et al, 2015). 73 We previously reported the negative-stain electron microscopy (EM) structure of the TssJLM complex 74 from enteroaggregative Escherichia coli (EAEC). We determined that 10 TssJ lipoproteins are bound 75 to 10 TssM proteins, forming two concentric rings of pillars and arches that span the periplasm. The 76 arches were shown to link to a flexible base composed of the N-terminal part of TssM and 10 copies 77 of TssL (Durand et al, 2015). This study also revealed that the EAEC TssJLM complex assembles into a 78 5-fold rotationally symmetric trans-envelope structure. However, the symmetry mismatch between 79 the 5-fold symmetry of the MC and 6-fold symmetry of the baseplate (Nazarov et al, 2018) questions 80 whether the purified TssJLM MC reflects the in vivo situation. In addition, although most of the 81 available crystal structures can be fitted into this EM structure, we currently lack molecular details on 82 the whole complex, such as the precise location of the membranes, of the trans-membrane helices, 83 and the potential presence of a periplasmic channel.

84 Here, we first present the *in situ* cryo-electron tomography (cryo-ET) structure of the EAEC 85 TssJLM MC. These data confirm the 5-fold symmetry of the complex *in vivo*, and provide information 86 on the location of the inner and outer membranes. We then present the single particle (SPA) cryo-87 electron microscopy (cryo-EM) structure of the MC that describes the intricate molecular 88 architecture of the periplasmic portion of the complex. This high-resolution cryo-EM structure 89 reveals the presence of five additional copies of TssJ at the tip of the complex, the presence of a 90 periplasmic gate that closes a periplasmic channel. Finally, we demonstrate that the periplasmic gate 91 and the additional TssJ are required for T6SS-dependent activity in vivo.

92 Results

93 Structure of the T6SS MC within the cell envelope

94 To observe the T6SS MC in its native cellular environment, we performed cryo-ET on bacterial 95 cells (Weiss et al, 2017). With the aim to have a sufficient number of particles to obtain an in situ 96 structure by subtomogram averaging, we imaged E. coli BL21(DE3) cells in which TssJLM were 97 heterologously overexpressed. This strain does not possess T6SS genes, thereby preventing any 98 crosstalk or protein-protein interactions between TssJLM and other natively present T6SS proteins. 99 Since *E. coli* is too thick to be directly imaged by cryo-ET, we pursued three different approaches to 100 tackle sample thickness. The first approach consisted of engineering a minicell-producing skinny 101 strain (Farley et al, 2016) of E. coli BL21(DE3), and thereby generating a minicell strain that is 102 compatible with the T7-based expression system. Although this strain produced minicells as small as 103 450 nm in diameter (Fig EV1A), their size still affected the contrast to an extent that did not allow for 104 sub-tomogram averaging. Nevertheless, characteristic inverted Y-shaped particles (side views, Fig

105 EV1B) and star-shaped particles (top views, Fig EV1C) could occasionally be observed in the 106 periplasm of these minicells. In a second approach, tomograms were recorded of cells that were 107 partially lysed and exhibited a high contrast (Fig EV1D, E). These cells, which have previously been 108 described as "ghost cells" due to their transparent appearance, still had mostly intact membranes 109 and conserved their cytoplasmic macromolecular complexes such as ribosomes (Fu et al, 2014). In a 110 third approach, we used a state-of-the-art cryo-sample thinning method called focused ion beam 111 (FIB) milling to thin *E. coli* BL21(DE3) cells in which TssJLM were heterologously overexpressed (Fig 112 EV1F, G). FIB milling allows to etch through a lawn of bacterial cells and thin them down to under 200 113 nm (Medeiros et al, 2018a, 2018b; Marko et al, 2007). This approach was more native, as it was 114 performed on intact rod-shaped wild-type BL21(DE3) cells. These tomograms had a high contrast and 115 provided great detail. To confirm the relevance of these observations, we carried out control 116 experiments in the native T6SS⁺ EAEC 17-2 strain. TssJLM particles were also readily observed by 117 cryo-ET, both when heterologously overproduced (Fig EV1H, note that the particles could 118 occasionally be found detached from the outer membrane: Fig EV1J) as well as under native 119 conditions, *i.e.* in wild-type EAEC 17-2 cells (Fig EV1I).

120 Sub-volumes of star-shaped (top views) and inverted Y-shaped (side views) particles were 121 manually picked and subsequently aligned and averaged. The resulting average was similar to the in 122 vitro T6SS MC structure published previously (Durand et al, 2015), with a tip and a core, made of 5 123 pairs of pillars forming a narrow central channel, that splits into 10 arches (Fig 1A-E). Importantly, the 124 5-fold symmetry was evident without applying symmetry (Fig 1A'). As it is the case for the TssJLM MC 125 solved by EM (Durand et al, 2015), the periplasmic core and the arches were well resolved, whereas 126 the inner membrane embedded base and the outer membrane-embedded cap were poorly resolved. 127 The Fourier shell correlation (FSC) curve indicated a resolution of 25 Å at a coefficient of 0.5 (Fig EV2 128 A). The prevalence of top views indicated that the average might be affected by a missing wedge (Fig. 129 EV2 B.

130 After aligning and averaging a set of sub-volumes, it can be useful to place the isosurface of 131 the average back in the original tomographic volume to analyze the positions and orientations of the 132 individual aligned particles. In this way, we obtained a clear view of the location of the MC within the 133 cell envelope (Fig 2A, Movie 1 and Fig EV2 C-E, Movie 2). The position of the membranes with respect 134 to the TssJLM highlighted that the tip was embedded in the outer membrane without crossing it, 135 while the MC was anchored in the inner membrane at the lower part of the arches (Fig 2B). In some 136 cases, densities could be seen spanning the inner membrane and extending into the cytoplasm (Fig 137 2C). In what could be an overexpression artefact, TssJLM particles were also found in cytoplasmic 138 membrane invaginations (Fig 2A). Altogether, these data confirmed the 5-fold symmetry of the T6SS 139 TssJLM MC *in situ* and provided further insights into the position of the inner and outer membranes.

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141 Structural analysis of the TssJLM complex by cryo-EM

We previously reported the negative stain EM structure of the EAEC TssJLM MC (Durand *et al*, 2015). Here, we used the same purification procedure and used SPA cryo-EM to obtain the cryo-EM structure of the 1.7 MDa T6SS MC at 4.9 Å overall resolution. With a 5-fold symmetry imposed, the local resolution ranged between 4.1 Å and 21.7 Å (Figs 3A and EV3A-E). The angular distribution of the particles was good and allowed us to reach a high resolution (Fig EV3 C, F).

The 5-fold symmetry is clearly visible in the 2D classes top views and the volume slices of the reconstruction, which retain a 5-pointed star shape (Fig EV3C,G). When no symmetry was applied during the reconstruction, the overall resolution decreased to 7.5 Å but the pentameric nature of the complex was maintained, with only one of the 10 arches displaying weaker density than the others causing the drop in resolution (Fig EV3H, I). This could indicate partial assembly of the complex *in vivo* or disassembly during its purification.

153 The overall structure resembles that obtained by negative stain (Fig EV3J(Durand et al, 2015)). 154 The architecture of the complex comprises a tip connected to a core region that extends to a base, 155 through a double ring of pillars with arches in proximity to the inner membrane (Fig 3A-B). Several 156 notable features are already evident from the cryo-EM density map of the full complex (Fig 3A). First, 157 the tip region and the base are disordered, and appear to be filled by random densities. Second, 158 cross-sections of the full complex show that the channel, across which tube/spike transport might 159 occur, is closed by a gate at the intersection between arches and pillars, above the inner membrane 160 (Fig 3B-E).

161 To better characterise the flexibility of the base of the complex, we collected tomograms on 162 the same frozen EM grids that were used to collect the SPA dataset (Fig EV4A-C). The base of 163 individual complexes appeared as very heterogeneous, with single arches often pointing in opposite 164 directions, or on the contrary several arches clumping together (Fig EV4A). On the other hand, the 165 core was rigid and resembled a 5-branched star. Moreover, about 20% of the particles possessed 166 only 3 or 4 out of 5 branches (Fig EV4B), indicating that partial assemblies could be stable. The 167 particles were lying in a thin layer of ice (25 nm) and were found at different orientations (Fig EV4C1-168 2).

169

170 The structure of the base

To try and overcome the inherent flexibility of the complex and better discern different features of the base, we performed a density subtraction of the tip, core and arches followed by a

173 focused refinement of the base with and without symmetry applied. We thus obtained 2D classes 174 and a 3D structure of the base at 17-Å resolution when a C5 symmetry was applied (Figs 3F and 175 EV4E-F). When observing the cross section of the base in the full complex, as indicated by arrows in 176 Fig3B, and in the subtracted structure (Fig EV4E), two 110 Å-wide linear densities are clearly visible, 177 separated by 40 Å. This double layer of density is consistent with the density diagram of a lipid 178 bilayer with the head groups being the most dense at a distance of ~4 nm from each other and fits 179 well a lipid bilayer composed of PE, obtained using the CHARMM-GUI (Jo et al, 2007) (Fig EV4E). We 180 conclude that the inner membrane sub-domain of the T6SS MC is filled by a lipid bilayer.

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182 An additional TssJ is present in the full complex

183 To focus on the best-resolved region of the cryo-EM map, the base was subtracted from the 184 tip, a 2D classification and a masked 3D refinement was performed to obtain the structure of the 185 core at 4.5 Å (Figs 3G and EV5A-B), with a local resolution ranging from 4 Å to >10 Å (Fig EV 5C). The 186 known crystallographic structure of the C-terminus of TssM (aa. 869-1129) bound to TssJ (PDB 4Y7O) 187 could be easily fitted in the outer and inner pillars, with a correlation of 0.8505 and 0.565 188 respectively, leaving an extra density (Fig EV5D-E). Interestingly, an extra TssJ subunit, TssJ', which 189 was not observed in the low resolution complex (Durand et al, 2015) fits in this extra density with a 190 correlation of 0.879 (EV5E-F). We conclude that the T6SS MC comprises 15 TssJ proteins, 3 TssJs for 2 191 TssM (Fig 4A). No additional residues were visible for the N- and C-termini of TssJ (1-21 and 151-155), 192 disordered in both the crystal and the cryo-EM structures. After placing the extra TssJ, we refined the 193 TssM-TssJ crystal structure and the newly fitted TssJ copy against the cryo-EM map to obtain the 194 structure of each TssM and TssJ monomer within the whole MC.

195 TssJ' binds to the MC through previously unknown interfaces. If we consider the TssJ.i, TssJ.o 196 and the TssJ' subunits (Figs 4A-C and EV6A-B) with TssJ.i and TssJ.o being in contact with TssM.i and 197 TssM.o respectively, they sit in the same position as in the crystal structure and in the outer and 198 inner pillars (Fig 4B). By contrast, TssJ' binds to TssM.o and TssJ.i strongly (Fig 4C, Table EV1). In 199 particular, the interaction of TssJ' with TssJ.i is specifically strong, as their contact is mediated not 200 only by hydrogen bonds but also by salt bridges (R31 with E34, and D97 with R33; see Fig EV6C). TssJ' 201 binds to TssM.i via hydrogen bonds only and this interaction is comparable to that between TssJ.i and 202 TssJ.o with TssM.i and TssM.o respectively (surface of interaction 573 $Å^2$ and -2.6 kcal/mol Δ G) (Fig 203 4B-C, Table EV1).

204 TssJ' is required for MC assembly and T6SS activity

205 To gain further information on the function and in vivo relevance of TssJ', mutations that 206 specifically impact the TssJ'-TssJ.i interface were engineered onto the chromosome, at the native 207 locus. Two residues, Arg-31 and Asp-97 (Fig EV6C), were targeted as they form a salt bridge with E34 208 and R33 in TssJ.a, respectively (Table EV 1 and Fig EV6C). The R31E, D97A and D97K substitutions 209 were then tested for their ability to outcompete a fluorescent E. coli competitor strain. Although the 210 R31E and D97A did not significantly impact T6SS antibacterial activity, the D97K mutation abolished 211 proper function of the T6SS (Fig 4D). The assembly and stability of the T6SS MC was then assessed by 212 fluorescence microscopy using a chromosomally-encoded and functional fusion protein between 213 TssM and the super-folder GFP (sfGFPTssM, Durand et al, 2015) (Fig 4E). As previously observed 214 (Durand *et al*, 2015), the sfGFPTssM strain forms stable foci. By contrast, cells producing the TssJ D97K 215 variant form small and unstable fluorescent sfGFPTssM foci (Fig. 4E). Time-lapse fluorescence 216 microscopy recordings showed that about 90% of the foci observed in the $_{stGFP}$ TssM strain (n = 50) are 217 stable over the 600-sec recording time, in agreement with the previous observation that the EAEC 218 T6SS MC is stable and serves for several contraction/elongation cycles (Durand et al., 2015). By 219 contrast, with a mean lifetime of ~ 107 sec (n = 50) the $_{sfGFP}$ TssM fluorescent foci observed in cells 220 producing the TssJ D97K variant are drastically less stable (Fig EV 7A). We then tested whether the 221 TssJ D97K variant promotes sheath assembly, using chromosomally-encoded functional sfGFPTssB 222 fusion. Fig. 4F shows that, contrarily to the wild-type parental cells in which dynamic sheaths can be 223 observed, no sheath polymerization occurs in presence of TssJ-D97K (Fig. 4F and EV Fig.7B). 224 Altogether, these results demonstrate that TssJ'-TssJ.i interface is required for the stability of the 225 T6SS MC, sheath formation and T6SS antibacterial activity.

226

227 A flexible hinge within the TssM periplasmic domain

228 We were able to confidently build de novo the periplasmic domain of TssM including its N-229 terminal fragment (residues 579 to 869) that was missing in the crystal structure (Durand et al, 230 2015). This was done in Coot (Emsley et al, 2010) by using Phyre2 secondary structure predictions 231 (Kelley et al, 2015) and RaptorX evolutionary covariance interaction predictions (Källberg et al, 2012) 232 as validation tools (Fig 5A and EV8A-C). The cryo-EM structure of the TssM periplasmic domain 233 slightly differs from the X-ray structure (Fig EV9A). From the C-terminus, helix 869-891 extends to 234 amino-acid 841 with a slight kink at residue Pro-870. The remaining N-terminal fragment forms an α -235 helical domain comprising 8 helices that snake back and forth to the inner membrane (Fig 5A). The 236 region closest to the membrane was too flexible to be resolved and for an atomic model to be built 237 (Fig EV8B). The cryo-EM pseudoatomic model of the fully assembled TssM-TssJ complex shows it

238 forms a bell shape composed of two rings of pillars that twist around each other (Fig 5A). Within each 239 asymmetric unit, two copies of TssM are present, named TssM.o in the external pillar and TssM.i in 240 the internal pillar. The inner and the outer pillar TssM proteins are superimposable with the 241 exception of a 23° kink located at residue 867 (Fig EV9B). These two TssM subunits interact front-to-242 back (Fig EV9C) with an area of 1168 $Å^2$, a binding energy of -9.9 kcal/mol, and a ΔG of -5.2 kcal/mol. 243 Each TssM.i also interacts with two adjacent TssM.i within the inner TssM ring at an angle of 76°, 244 with an area of 1529 $Å^2$, a binding energy of -12 kcal/mol and a ΔG of -5.32 kcal/mol (Figs EV6A and 245 EV9D). Finally, TssM.o⁺¹ also interacts with TssM.i and the two are oriented at 68° from one another 246 (Figs EV6A and EV9E).

247 A poorly-defined density that sits in the core region between TssM.i and TssM.o⁺¹, was 248 attributed to the C-terminus of TssM. We built a small loop that terminates into a helix. (Fig EV9F). 249 This region is disordered in the outer pillar monomer (TssM.o). Additionally, the resolution of the 250 pillars gets worse towards the basal side, and no secondary structure could be identified when we 251 tried to build *de novo* the atomic model of TssM. Despite this high degree of flexibility, we produced 252 a model of the periplasmic region between amino acids 390 and 550 based on a RaptorX contact 253 predictions. This model fits well (correlations of 0.819 and 0.827) into the remaining densities (Fig. 254 EV9G). The EM density shows that while they are separated at the level of the arches, the inner and 255 the outer pillar re-join at the level of the inner membrane (Fig EV9G).

256 The periplasmic TssM gate

257 The inner pillars of TssM form a channel with a diameter that varies between 2.6 Å and > 20 Å 258 (Fig 5B-C). The site of constriction observed in the cryo-EM density (Figs 3B,E) corresponds to loop 259 776-786 in the atomic model of TssM (Fig 5D). Specifically, residues Gln-779 and Asn-780-781 260 maintain the constriction via polar interactions (Fig 5D). In the external pillar, the same loop interacts 261 with loop 600-625 on the neighbouring pillar, providing further stabilisation of the structure (Fig 262 EV10A). Conservation analysis of the sequence with related proteins predicted by the ConSurf server 263 (Celniker et al, 2013), indicated that the sequence of the loop 776-786 is poorly conserved amongst 264 species (Fig EV10B) although the presence of a loop at this position is a conserved feature. As 265 previously proposed (Durand et al, 2015), these data suggest that the purified TssJLM MC is in a 266 closed state. Large conformational changes, including modification of the constriction and movement 267 of the inner pillars are therefore required to allow the passage of the tube/spike during sheath 268 contraction.

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270 The TssM periplasmic gate is required for MC assembly and T6SS function

271 To test the function of the periplasmic gate, several mutations were engineered at the tssM 272 locus in the wild-type EAEC 17-2 strain and the function of the T6SS was assessed as previously 273 described (Fig 5E-G). To covalently stabilize the contacts between the internal pillars and thus 274 prevent MC opening, GIn779 and Asn780 were substituted with cysteines (Q779C-N780C). 275 Conversely, a constitutively open gate was created by deleting a large portion of the constriction 276 loop (Δ 777-783). Antibacterial competition assays showed that the Q779C-N780C variant loses the 277 ability to outcompete competitor cells, whereas the single control mutant Q779C did not (Figure 5E). 278 Deletion of the Δ 777-783 loop also impaired the T6SS antibacterial activity (Figure 5E). Fluorescence 279 microscopy recordings further show that sfGFPTssM Q779C-N780C and sfGFPTssM Δ 777-783 do not 280 assemble TssM foci since, in contrast to the parental strain, diffuse fluorescent pattern were 281 observed (Figs 5F and EV11). These results demonstrate that the MC is not properly assembled when 282 the integrity of the periplasmic gate is impacted. As expected, these mutant cells were not able to 283 assemble T6SS sheaths (Figs 5G and EV11).

284 Discussion

285 In this study, we report the *in situ* and *in vitro* structures of the T6SS TssJLM MC from EAEC. 286 The cryo-ET structure confirmed the 5-fold symmetry and general architecture in vivo while the high-287 resolution cryo-EM structure provided molecular details about the periplasmic portion of the 288 complex (Fig 1, Fig 3 and Movie 3). As previously defined, the pentameric propeller-like structure 289 composed of 10 pillars intertwined with each other was observed both in situ and from purified 290 material (Figs 1 and 3C-E). Cryo-ET analyses allowed to position both the inner and outer 291 membranes. As anticipated based on biochemical experiments showing that TssJ is a periplasmic 292 lipoprotein attached to the outer membrane by an acyl anchor (Aschtgen et al, 2008), the tip of the 293 complex is embedded in the outer membrane (Fig 2B). However, neither the predicted detergent cap 294 in the SPA cryo-EM structure nor the cryo-ET data, indicate that TssJLM breaches or crosses the outer 295 membrane (Fig 2, Fig 3A-B). Nevertheless, it has been shown that this region is extracellularly 296 exposed in wild-type EAEC cells (Durand et al, 2015). Although the cryo-EM structure demonstrates 297 that the C-terminal region of TssM locates in the periplasm, we propose that the recruitment of 298 specific T6SS components induces MC conformational changes and cell surface exposition of the 299 TssM C-terminus. While our previous study suggested that the inner membrane locates at the level 300 of the arches (Durand et al, 2015), the cryo-ET analyses revealed that the inner membrane surrounds 301 the base (Fig 2B). Moreover, some tomograms also revealed the most basal parts of the MC, which 302 cross the inner membrane into the cytoplasm (Fig 2A, C). The in situ cryo-ET structure presents an 303 apparent elongation of the tip region compared to the cryo-EM structure (Fig EV12A, B and Movie 3), which could correspond to an additional density associated to the outer membrane or which could alternatively be explained by the missing wedge. Nevertheless, the similarities between both structures allowed the atomic model of TssJ - TssM to be docked into the *in situ* average (Fig EV12C, D), whereas the cryo-EM structure could be placed in a cellular context (Fig EV12E and Movie 3).

308 The base of the complex in this higher resolution structure was not better resolved than in the 309 negative stain structure (Figs 3A, F, and EV3D). This base should comprise 10 copies of the TssM and 310 TssL cytoplasmic domains (Durand et al, 2015). TssL forms dimers (Zoued et al, 2018; Durand et al, 311 2012; Zoued et al, 2016) and the crystal structure of its cytoplasmic hook-like domain has been 312 reported from various species including EAEC, P. aeruginosa and V. cholerae (Durand et al, 2012; 313 Robb *et al*, 2012; Chang & Kim, 2015; Wang *et al*, 2018)(Durand *et al*, 2012; Robb *et al*, 2012; Chang 314 & Kim, 2015). The TssM cytoplasmic domain is comprised between the N-terminal transmembrane 315 hairpin and a third transmembrane helix (Ma et al, 2009; Logger et al, 2016). No high-resolution 316 structure of the TssM cytoplasmic domain is available, although a model has been built based on 317 homology with DPY-30 and NTPases (Logger et al, 2016). Unfortunately, due to the poor resolution of 318 the base, we did not succeed to confidently fit the TssL and TssM cytoplasmic domains in this 319 density. Additional assays to improve the resolution such as the use of nanodiscs or amphipols 320 proved to be unsuccessful (Fig EV13A). The flexibility of the TssJLM base, which did not allow for it to 321 be resolved, might be due to the absence of other T6SS cytoplasmic components such as the 322 baseplate. A similar observation was made for the type III secretion system, in which the presence of 323 the cytoplasmic sorting platform orders the IM components (Hu et al, 2017). One may hypothesize 324 that this flexibility is essential for the docking of the hexameric baseplate, and to accommodate the 325 five-fold to six-fold symmetry mismatch. One alternative hypothesis is that the disorder at the centre 326 of the base structure is caused by the presence of a lipid bilayer encircled by the TssM and TssL 327 proteins. In the *in vivo* situation, the MC assembles first, before the recruitment of the baseplate (Durand et al, 2015; Brunet et al, 2015), and hence, one can expect that a lipid bilayer at the 328 329 entrance of the TssJLM lumen would be present before baseplate docking to prevent the leakage of 330 solutes and proton-motive force.

331 The high-resolution structure of the EAEC TssJLM MC also revealed new interesting and 332 functional features. First, five additional TssJ subunits, called TssJ', were identified in the tip complex. 333 These TssJ' proteins interact with the TssJ proteins of the outer pillars (TssJ.i). Mutations that 334 interfere with TssJ'-TssJ.i interaction impair the functional integrity of the MC and hence inactivated 335 the T6SS (Fig 4D-F). Second, we identified a 11-amino-acid loop in TssM that protrudes from each 336 inner pillar to the centre of the channel, thus creating a constriction that is observed in the density 337 map (Fig 3D, E). Each loop is stabilized by the adjacent loop via Asn/Gln pairings. Such weak 338 interactions could be easily displaced by the VgrG/PAAR spike upon baseplate docking or during

339 firing. Our mutational analyses demonstrate that this constriction is important for TssJLM MC 340 formation and T6SS activity. Periplasmic constrictions are usual features of trans-envelope 341 complexes. The best characterized examples include the OM T2SS and T3SS secretins, and the CsgG 342 curli secretion channel who present one or two periplasmic gates to prevent leakage (Fig EV14A-B) 343 (Yan et al, 2017; Spagnuolo et al, 2010; Goyal et al, 2014). While we do not know the role of this 344 constriction in the T6SS, we propose that these loops may stabilize the MC in its closed conformation 345 during the resting state. The cryo-EM structure defined two hinge regions that exhibit a certain 346 degree of flexibility (Fig EV9B,G). These two hinges result in the formation of the two layers of pillars, 347 with the inner layer obstructing the channel. A large conformational change of these pillars is 348 therefore necessary to open the channel for the passage of the tube/spike complex. Interestingly, with an interaction surface of 1540 $Å^2$ (ΔG of 6.1 kcal/mol) the TssM inner pillars contacts are 349 350 considerably less stable than the contacts within the T2SS secretin (interaction surface of 5353.7 $Å^2$ 351 and ΔG of -52.4 kcal/mol). The displacement of the pillars could be controlled by the flexibility of the 352 hinge regions.

353 Based on these data, we propose a model in which the T6SS TssJLM MC is assembled in a 354 closed state. In this conformation, five pillars are oriented toward the centre of the complex to close 355 the complex at the outer membrane and hence to protect the cell from periplasmic leakage or from 356 the entry of toxic compounds. This conformation is further stabilized by the interactions of the TssM 357 protruding loops. The flexibility of the MC cytoplasmic base allows the proper positioning of the 358 TssKFGE-VgrG baseplate, and accommodates the five-to-six symmetry. The docking of the baseplate 359 positions the VgrG/PAAR spike in proximity to the inner membrane. Once in contact with the target 360 cell, a signal transmitted to the baseplate triggers the contraction of the sheath, allowing the passage 361 of the tube/spike complex through the MC. The hinge regions undergo a tectonic conformational 362 change that opens the channel and the tip complex. The MC then returns to the resting, closing state 363 allowing a new cycle to start (Figure 6).

364

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383

384 **Declaration of interests**

- 385
- 386 The authors declare no competing interests.
- 387

388

389 Figures legends

390

391 Figure 1. Subtomogram average of the TssJLM complex *in situ*

A-E. Isosurface of the subtomogram average (in pink) with an applied C5 symmetry and 0.69 nm tomographic slices (A-E) at the indicated heights. The average is shown in side view, whereas the slices represent perpendicular slices. The 5 pairs of pillars formed a narrow central channel and separated into 10 arches towards the base (base not visible at the used threshold). The division of the structure into subparts was adapted from (Durand *et al.*, 2015). Subvolumes were extracted from cryotomograms of ghost cells and FIB-milled intact *E. coli* BL21 cells heterologously expressing TssJLM.

- **A'**. Slice (0.69 nm) through the non-symmetrized average. Note that the C5 symmetry was visible.
- 400

401 Figure 2. Position of TssJLM in the cell envelope

402 **A.** Slice (9.7 nm) through a cryotomogram of a FIB-milled *E. coli* BL21 cell expressing TssJLM. The 403 average shown in Figure 1 was placed back at the positions and orientations of the individual 404 subvolumes that were used to generate the final average. The zoomed-in area highlights the position 405 of the TssJLM particles within the inner and the outer membrane. Some particles were found in 406 cytoplasmic membrane invaginations, as indicated by white arrows. Scale bar 100 nm.

407 B. Isosurface of the final average (orange) merged with the isosurface of a second average (with
408 higher threshold; grey mesh). The panel shows the positioning of TssJLM with respect to both inner
409 (IM) and outer membranes (OM). The distances corresponded to the widest and longest dimensions
410 of the TssJLM complex.

411 **C.** Cryotomographic slices (9.7 nm) showing side views of TssJLM. In these examples, the basal parts

412 of TssJLM (red arrows) could be seen extending into the cytoplasm. Scale bar 10 nm.

413

414 Figure 3. 3D structure of the TE complex of the T6SS

A. Autosharpened Cryo-EM density of the full TE complex composed of TssJ, TssL and TssM. The
 inner pillars are coloured in green and the outer pillars in blue. The unstructured tip and base are in
 gray, while the top of the core is in orange.

418 **B.** Vertical cross section of the cryo-EM density. Colouring is according to A. The position of the 419 slices for C-E are indicated. The density of the base appears to be composed of two leaflets, indicated

420 by 2 arrows.

421 **C-E.** Cross sections of A, sliced at positions indicated in B.

422 **F.** Subtracted, masked and unsharpened cryo-EM density of the mostly unstructured base

423 **G.** Locally sharpened cryo-EM density of the subtracted and masked core. Three different views are

424 shown. The colouring is according to A

425

426 Figure 4. The TssJ' monomer and its function

A. Ribbon diagram and locally sharpened surface representation (transparent= of the three TssJ
protomers in orange, labelled TssJ.a, TssJ.A and TssJ' for the internal, external and additional
monomer respectively. In light green and blue are the TssM.a and TssM.A protomers respectively.

430 **B.** Superimposition of the ribbon diagrams of TssM-TssJ heterodimer within the internal (green-431 orange) and the external (blue-orange) pillars.

432 **C.** Ribbon diagram of the TssM.A (external pillar) with the TssJ.A and the TssJ' (orange)

433 D. Antibacterial assay. Prey cells (Gfp+ kanR E. coli W3110) were mixed with the indicated attacker 434 cells, spotted onto Sci-1-inducing medium (SIM) agar plates and incubated for 4 h at 37°C. The image 435 of a representative bacterial competition spot is shown on the upper part. The relative fluorescent 436 level (in AU) and the number of recovered E. coli recipient cells are indicated in the lower graph (in 437 log10 of colony forming units (cfu)). The assays were performed from at least three independent 438 cultures, with technical triplicates and a representative technical triplicate is shown. The circles 439 indicate values from technical triplicate, and the average is indicated by the bar. E. Fluorescence 440 microscopy recordings showing sfGFPTssM foci in the parental (WT) and TssJ mutated strains (TssJ 441 D97K). TssM foci containing cells are indicated by arrowheads. Microscopy analyses were performed 442 independently three times, each in technical triplicate, and a representative experiment is shown. 443 Scale bars, 1 µm. F. Fluorescence microscopy recordings showing TssBsfGFP sheath in the parental 444 (WT) and TssJ mutated strains (TssJ D97K). Fluorescent sheath containing cells are indicated by 445 arrowheads. Microscopy analyses were performed independently three times, each in technical 446 triplicate, and a representative experiment is shown. Scale bars, 1 µm.

447

448 Figure 5. The TssM protein and the periplasmic gate

449 A. The high-resolution structure of MC complex in different orientations. TssJ is in orange, TssM is in450 green and blue according to their position as an internal or external pillar respectively.

451 B. The pore radius formed by the internal pillar of TssM protomers is highlighted as dots and mapped
452 using HOLE. The atomic model of TssM is coloured according to secondary structure (Helices in pink
453 and strands in brown)

454 **C.** A graph showing the pore radius calculated along the pore centre of the internal pillar of TssMs.

455 **D.** Ribbon diagram of the region surrounding the periplasmic gate, with the amino acids involved in

456 the formation of the gate in atom form. Gln779 and Asn780-781 from the internal pillars of TssM

457 (green) are labelled in one of the protomers

458 E. Antibacterial assay. Prey cells (Gfp+ kanR E. coli W3110) were mixed with the indicated attacker 459 cells, spotted onto Sci-1-inducing medium (SIM) agar plates and incubated for 4 h at 37°C. The image 460 of a representative bacterial competition spot is shown on the upper part. The relative fluorescent 461 level (in AU) and the number of recovered E. coli recipient cells are indicated in the lower graph (in 462 log10 of colony forming units (cfu)). The assays were performed from at least three independent 463 cultures, with technical triplicates and a representative technical triplicate is shown. The circles 464 indicate values from technical triplicate, and the average is indicated by the bar. F. Fluorescence 465 microscopy recordings showing sfGFPTssM foci in the parental (WT) and TssM mutated strains (TssM 466 Q779C/N780C, TssM Δ777-783). TssM foci containing cells are indicated by arrowheads. Microscopy 467 analyses were performed independently three times, each in technical triplicate, and a 468 representative experiment is shown. Scale bars, 1 µm. G. Fluorescence microscopy recordings 469 showing TssBsfGFP sheath in the parental (WT) and TssM mutated strains (TssM Q779C/N780C, TssM 470 Δ 777-783). Fluorescent sheath containing cells are indicated by arrowheads. Microscopy analyses 471 were performed independently three times, each in technical triplicate, and a representative 472 experiment is shown. Scale bars, 1 µm.

473

474 Figure 6. Summary of the Type 6 secretion system cycle of action

The T6SS assembly begins first with the recruitement of the membrane complex (MC) in its resting state (1). The MC recruits the baseplate (BP) and the tail tip complex (TTC) is assembled (2). The recently solved structures of the BP (No Title) and MC (this paper) in the membrane context are shown in the inset. A conformational change leads to the channel opening (3) and release of the toxin onto the VgrG spike by contraction of the TTC (4). Once the secretion has occurred, the TTC is recycled (5) and the MC can return to its resting state (6).

481

482

483 Materials and methods

484

485 Strains and media

486 Strains used in this study are listed in Table EV2. The E *E. coli* K-12 W3110 bearing the pUA66-*rrnB* 487 vector (Kan^R and GFP⁺, (Zaslaver *et al*, 2006)) was used as recipient for antibacterial competition 488 assays. Strains were routinely grown in lysogeny broth (LB) rich medium or in Sci-1-inducing medium 489 (SIM; M9 minimal medium, glycerol 0.2%, vitamin B1 1 μ g.mL⁻¹, casaminoacids 100 mg.mL⁻¹, LB 10%, 490 supplemented or not with bactoagar 1.5%) (Brunet *et al*, 2011) with shaking at 37°C.

491

492 Strain construction

493 tssM and tssJ point mutations were engineered at the native locus on the chromosome by allelic 494 replacement using the pKO3 suicide vector (Link et al, 1997) into the enteroaggregative E. coli 17-2 495 strain. Briefly, 17-2 WT strain was transformed with a pKO3 plasmid in which a fragment of the tssM 496 or tssJ gene carrying the point mutations has been cloned (see below). Insertion of the plasmid into 497 the chromosome was selected on chloramphenicol plates at 42°C. Plasmid sequences removal was 498 then selected on 5% sucrose plates without antibiotic and *tssM* point mutation recombinant strains 499 were screened by PCR and confirmed by DNA sequencing (Eurofins, MWG). Chromosomal fluorescent 500 reporter insertions into the enteroaggregative E. coli 17-2 strain mutated in tssM or tssJ was 501 achieved by using a modified one-step inactivation procedure (Datsenko & Wanner, 2000) as 502 previously described (Aschtgen et al, 2008b) using plasmid pKOBEG (Chaveroche et al, 2000). Briefly, 503 a kanamycin cassette was amplified from plasmid pKD4 using oligonucleotide pairs carrying 5' 50-504 nucleotide extensions homologous to regions adjacent to the gene to be deleted. After 505 electroporation of 600 ng of column-purified PCR product, kanamycin-resistant clones were selected 506 and verified by colony-PCR. The kanamycin cassette, inserted at the gene locus on the bacterial 507 chromosome, was then excised using plasmid pCP20, leaving an FRT scars. Gene deletions were 508 confirmed by colony-PCR and sequencing.

509

510 Fluorescence microscopy, image treatment and analyses

Fluorescence microscopy experiments were performed as described (Brunet *et al*, 2013; Zoued *et al*, 2013). Briefly, cells were grown overnight in LB medium and diluted to $A_{600nm} \sim 0.04$ in SIM. Exponentially growing cells ($A_{600nm} \sim 0.8-1$) were harvested, washed in phosphate-buffered saline buffer (PBS), resuspended in PBS to $A_{600nm} \sim 50$, spotted on a 1.5% agarose pad and covered with a cover slip. Fluorescence micrographs were captured using AxioImager M2 microscope (Zeiss) equipped with an OrcaR2 digital camera (Hamamatsu). For time lapse fluorescence microscopy, images were recorded with a Nikon Eclipse Ti microscope equipped with an Orcaflash 4.0 LT digital

518 camera (Hamamatsu) and a perfect focus system (PFS) to automatically maintain focus so that the 519 point of interest within a specimen is always kept in sharp focus at all times despite mechanical or 520 thermal perturbations. Fluorescence images were acquired with a minimal exposure time to reduce 521 bleaching and phototoxicity effects, typically 200 ms for TssB-sfGFP and 300 ms for sfGFP-TssM. For 522 image treatment, noise and background were reduced using the 'Subtract Background' (20 pixels 523 Rolling Ball) and Band plugins of imageJ (Rasband, 2012). The sfGFP foci were automatically detected 524 using the microbeJ plugin (Ducret et al, 2016). Floating bars representing the number of detected 525 foci for each strain were made using GraphPad (https://www.graphpad.com). Microscopy analyses 526 were performed at least three times, each in technical triplicate, and a representative experiment is 527 shown.

528

529 Interbacterial competition assay

530 The antibacterial growth competition assay was performed as previously described (Flaugnatti et al, 531 2016b). Wild-type E. coli K-12 strain W3110 bearing the pUA66-rrnB plasmid (conferring kanamycin 532 resistance and constitutive GFP fluorescence (qfp gene under the control of the ribosomal rrnB 533 promoter, (Gueguen & Cascales, 2013) was used as recipient. Attacker and recipient cells were 534 grown for 16 h in LB medium, diluted in SIM to allow maximal expression of the sci-1 gene cluster 535 (Brunet et al, 2011). Once the culture reached A_{600nm} ~ 0.8, cells were harvested and normalized to 536 A_{600nm} = 0.5 in SIM. Attacker and recipient cells were mixed to a 4:1 ratio and 15-µl drops of the 537 mixture were spotted in triplicate onto a pre-warmed dry SIM agar plate. After incubation for 4 h at 538 37°C, the bacterial spots were resuspended in LB and bacterial suspensions were normalized to 539 A_{600nm} = 0.5. For the enumeration of viable prey cells, bacterial suspensions were serially diluted and 540 spotted onto kanamycin LB plates. The assays were performed from at least three independent 541 cultures, with technical triplicates and a representative technical triplicate is shown.

542

543 **Protein preparation**:

The expression and purification of the TssJLM complex was carried out as previously described (Durand *et al*, 2015), with the exception that the cryo-EM grids were prepared immediately after the HisTrap Elution. For the amphipole-containing sample, the Strep-Trap elution was incubated with amphipoles A8-35 (Anatrace, USA) and subjected to gel filtration on a superpose 6 (GE Healthcare, UK) to remove residual detergent.

549

550 Cryo-EM grids preparation and data acquisition

551 C-flat $\[Martin et al, \]$ 552 C-flat $\[Martin et al, \]$ 2016). 3.5 $\[Martin et al, \$

same side for 2s in a Leica EM GP at 80% humidity and 4 °C, before being plunge frozen in liquid ethane (-184°C). Micrographs (Fig EV3B) at a nominal magnification of 120,000 X were collected in a Talos Arctica electron microscope equipped with a Falcon 3EC camera (Thermo Fisher, Waltham, MA, USA) in linear mode and with a pixel size of 1.24 Å. Dose-fractionated movie frames 20/micrograph were acquired for 1 s with a total electron flux of 120 e/Å/s. The defocus range chosen for the automatic collect was 0.7 to 2 μm, which resulted in an actual range between 0.4 to 5 μm.

For the amphipoles-containing MC collection, 3019 movies composed of 25 frames at a defocus range between 0.7 and $2\mu m$, were collected at 1.38Å pixel size with a 5s exposure time and 15 e/pix/s exposure rate at the Krios 2 at the Diamond eBIC facility.

562

563 Cryo-EM image processing

564 The 16,000 movies collected were aligned using MotionCor2, with dose weighting (6 $e^{-/A^2/frame}$) 565 and with 5X5 patches applied (Zheng et al, 2017). gCTF was used to estimate the CTF parameters 566 (Zhang, 2016) and low quality images were discarded. Relion 2.1 (Scheres, 2012) autopicked 227,527 567 particles and after several rounds of 2D classification in cryosparc (Punjani et al, 2017) and a 568 heterogeneous ab initio reconstruction (2 classes), 37,435 particles were converted using the script 569 csparc2star.py (Asarnow, 2016) and selected for a final 2D classification in relion 2.1 (Fig EV3C), of 570 which 36,828 particles were selected. An initial unmasked refinement using the ab initio model from 571 crysoparc, gave us a resolution of 7.6 Å with 5-fold applied symmetry and a soft mask of 450 Å. This 572 refined structure was used to do a movie refinement with all the frames and a polishing step with 573 RELION2.1. The final masked refinement of the full structure gave a final resolution of 4.9 Å with a C5 574 symmetry applied, and 7.9 Å with no symmetry applied (Fig EV3E, I), The disordered tip and base 575 were subtracted and a masked refinement around the core structure yielded a final resolution of 4.6 576 Å (Fig EV5B). The base focused refinement was also performed on subtracted particles, without the 577 tip and the core regions, to a resolution of 17Å (Fig EV4F). The resolution for all densities except the 578 base, was calculated by masked postprocessing according to the "gold standard" method using 0.143 579 as the FSC value cut-off, or 0.5 for the low resolution reconstruction (Rosenthal & Henderson, 2003) 580 and the local resolution of the core was calculated by relion 2.1 (Fig EV5 C).

581 For figures and to build *de novo* pseudoatomic models in Coot (Emsley *et al*, 2010), the cryo-EM 582 density was initially sharpened using phenix.autosharpen (Terwilliger *et al*, 2018) and later with 583 LocalDeblur (Ramírez-aportela *et al*, 2018). Fitting of density, correlation calculations, molecular 584 graphics and analyses were performed on UCSF Chimera (Pettersen *et al*, 2004). For the amphipoles 585 dataset, 2D classes were calculated from a total of 8637 particles (Fig EV13A).

586

587 Model building.

588 Model building proceeded by fitting the PDB 4Y7O (Durand et al, 2015) into the density 2 times for 589 each pillar, with the cross correlation being calculated using Chimera (Pettersen et al, 2004)(Fig 590 EV5D). The model was manually built in Coot (Emsley et al, 2010) using bulky sidechains and 591 secondary structure predictions (Kelley et al, 2015) as guides. The de novo structure was validated 592 and the register adjusted using residue contact prediction (Bouvier et al, 2018). The map of the co-593 evolution contacts was aligned with those of the built PDB using the MapAlign software (Ovchinnikov 594 et al, 2017). Where discrepancies were observed, the register was modified to fit the predicted 595 contact maps (Fig EV8). The model was then refined using one round of rosetta.refine (Wang et al, 596 2016) and phenix.real space refine (Afonine et al, 2018).

597

598 Validation of the data

- The model was validated as in the protocol in Refmac5 (Murshudov *et al*, 2011). The FSC map to model was calculated with the sharpened map (FSC_{sum}). The model was shaken by 0.5 Å and the FSC map to model was calculated with one Half map (FSC_{work}). This refined model was then used to calculate the FSC map to model with the other Half map (FSC_{free}) (Fig EV15A).
- The cross correlation between each amino acid in the model and map was also calculated with phenix.real_space_refine (Afonine *et al*, 2018) (Fig EV15B) and the Molprobity score (Chen *et al*, 2010). was obtained from the online server (Table EV5) Pore radius calculations were carried out using the HOLE (Smart et al, 1996) plugin in Coot and the protein interfaces were analysed with PISA (Krissinel & Henrick, 2007).
- 608

609 Strains, media and chemicals

- 610 The strains, plasmids and nucleotides used in this study are listed in Tables EV2 and 3. For the cryoET 611 studies, E. coli K-12 BL21(DE3) and enteroaggregative E. coli EAEC strain 17-2 were used for protein 612 overexpression before plunge freezing. Strains were routinely grown in LB-Miller or in Sci-1-inducing 613 medium (SIM; M9 minimal medium, glycerol 0.2%, vitamin B1 1 mg ml⁻¹, casaminoacids 100 mg ml⁻¹, 614 LB 10%, supplemented or not with bactoagar 1.5%) (Brunet et al, 2011) with shaking at 37°C. Plasmids were maintained by the addition of ampicillin (100 mg ml⁻¹ for E. coli K-12, 200 mg ml⁻¹ for 615 616 EAEC), kanamycin (50 mg ml⁻¹) or chloramphenicol (30 mg ml⁻¹). Expression of genes from pRSF (in 617 BL21) and pBAD33 (in EAEC) vectors was induced for 2-3h with 1 mM of isopropyl-b-D-thio-618 galactopyrannoside (IPTG) or 0.3% L-arabinose, respectively.
- 619

620 **Preparation of frozen-hydrated specimens**

621 Plunge freezing was performed according to (Weiss *et al*, 2017). *E. coli* BL21 or EAEC cells were 622 concentrated by centrifugation to an OD_{600} of 3 - 20 and then mixed with protein A – 10 nm gold 623 conjugate (Cytodiagnostics Inc.). The higher concentrations of cells were used when preparing grids 624 for cryo-focused ion beam (cryo-FIB) milling to form "bacterial lawns" of several layers of bacteria on 625 top of each other. Bacterial lawns were found to be more amenable to cryo-FIB milling then 626 individual cells. A 3 μ L droplet of the sample was applied to a carbon-coated EM copper grid (R2/1, 627 Quantifoil) that had been previously glow-discharged for 90 s at -25 mA using a Pelco easiGlow[™] 628 (Ted Pella, Inc.). The grid was plunge-frozen in liquid ethane-propane (37 %/63 %) using a Mark IV 629 Vitrobot (Thermo Fisher Scientific). The forceps were mounted in the Vitrobot (27°C, humidity 95%) 630 and the grid was blotted from both sides or only from the backside by installing a Teflon sheet 631 (instead of a filter paper) on the front blotting pad. Grids were stored in liquid nitrogen.

632

633 Cryo-focused ion beam milling

634 Cryo-focused ion beam (cryo-FIB) milling was used to prepare samples of plunge-frozen cells that 635 could then be imaged by electron cryotomography (Marko et al, 2007). Our cryo-FIB milling workflow 636 has been detailed in (Medeiros et al, 2018b). Frozen grids with lawns of E. coli BL21 cells 637 overexpressing TssJLM were clipped into modified Autogrids provided by J. Plitzko or a commercial 638 prototype provided by Thermo Fisher. We then transferred the grids into the liquid nitrogen bath of 639 a loading station (Leica Microsystems) and clamped them onto a "40° pre-tilted TEM grid holder" 640 (Leica Microsystems). The holder with grids was shuttled from the loading station to the dual beam instrument using the VCT100 transfer system (Leica Microsystems). The holder was mounted on a 641 642 custom-built cryo-stage in a Helios NanoLab600i dual beam FIB/SEM instrument (FEI). The stage 643 temperature was maintained below -154°C during loading, milling and unloading procedures. Grid 644 quality was checked by scanning EM (SEM) imaging (5 kV, 21 pA). The samples were then coated with 645 a Platinum (Pt) precursor gas using the Gas Injector System. We adapted a "cold deposition" 646 technique that was published previously (Hayles et al, 2007) (needle distance to target of 8 mm, 647 temperature of the precursor gas of 27 °C, and open valve time of 5 s). Lamellae were milled in 648 several steps. We first targeted two rectangular regions to generate a lamella with ~2 µm thickness 649 with the ion beam set to 30 kV and ~400 pA. The current of the ion beam was then gradually reduced 650 until the lamella reached a nominal thickness of 150-400 nm (ion beam set to ~25 pA). Up to 6 651 lamellae were milled per grid. After documentation of the lamellae by SEM imaging, the holder was 652 brought back to the loading station using the VCT100 transfer system. The grids were unloaded and 653 stored in liquid nitrogen.

654

655 Electron cryomicroscopy and electron cryotomography

656 *E. coli* BL21 and EAEC cells (overexpressing TssJLM where indicated), cryo-FIB-processed *E. coli* BL21 657 cells overexpressing TssJLM, and purified TssJLM samples were examined by electron 658 cryotomography (cryoET). Images were recorded on a Tecnai Polara TEM (Thermo Fisher Scientific) 659 equipped with post-column GIF 2002 imaging filter and K2 Summit direct electron detector (Gatan), 660 or on a Titan Krios TEM (Thermo Fisher Scientific) equipped with a Quantum LS imaging filter and K2 661 Summit (Gatan). Both microscopes were operated at 300kV and the imaging filters with a 20 eV slit 662 width. The pixel size at the specimen level ranged from 4.93 Å to 4.05 Å. The latter pixel-sized was 663 used for the sub-tomogram average. Tilt series covered an angular range from -60° to +60° with 2° 664 (lamellae, sheath preparations) increments and -10 to -6 μ m defocus, or in focus (0 μ m defocus) 665 when the data was collected on the Titan Krios with a Volta phase plate (Thermo Fisher Scientific) 666 (Danev & Baumeister, 2016). The total dose of a tilt series was 60-100 e /Å². Tilt series and 2D 667 projection images were acquired automatically using UCSF Tomo (Zheng et al, 2007) on the Tecnai 668 Polara and SerialEM (Mastronarde, 2005) on the Titan Krios. Three-dimensional reconstructions and 669 segmentations were generated using the IMOD program suite (Kremer et al, 2005).

670

671 Sub-tomogram averaging

672 Tomograms used for subtomogram averaging were not CTF-corrected, as most of the particles 673 were extracted from tomograms collected in focus with the Volta phase plate. Individual particles 674 were identified visually in tomograms as 5-branched stars shapes in top and bottom views and as 675 inverted-Y shapes in side views and their longitudinal axes were manually modelled with open 676 contours in 3dmod (Kremer et al, 1996). The manual particle picking and first round of sub-677 tomogram averaging were performed with the PEET software package on tomograms that were 678 binned by 4 (1k reconstructions). Model points, the initial motive list, and the particle rotation axes 679 were generated using the stalkInit program from the PEET package (Nicastro et al, 2006). This 680 approach allowed the definition of each structure's longitudinal axis as the particle y-axis. 28474 681 individual particles extracted from cryotomograms of E. coli BL21 ghost and FIB-milled cells were 682 averaged using PEET with a box size of 44 pixels in x and z, and 72 pixels in y for the final step on data 683 binned by 2 (2k reconstruction, final pixel size 8.1 Å). A random particle was chosen as a first 684 reference. Missing wedge compensation was activated. The final motive lists obtained after this 685 initial average performed on tomograms that were binned by 4 were then translated and used to 686 perform a new round of sub-tomogram averaging on tomograms that were binned by 2 (2k 687 reconstructions). From individual particles and after analysing the resulting average, C_5 symmetry 688 was imposed. The Fourier shell correlation curves were calculated in PEET to estimate resolution. A 689 cylindrical mask centred on the structure's longitudinal axis was applied to the volumes using 690 imodmop (IMOD package) in order to mask neighbouring structures and the membranes during 691 averaging. 3dmod (IMOD package) and UCSF Chimera (Pettersen et al, 2004) were used for

- 692 visualization of the averages. 3dmod was used for generating all the movies, except for the morph
- and the atomic model visualization in Movie 3 that were generated in UCSF Chimera.
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