

Pseudopilin residue E5 is essential for recruitment by the type 2 secretion system assembly platform.

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- Pseudopilin residue E5 is essential for recruitment by the type 2 secretion system assembly
 platform
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- 27 Summary
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30 Type II secretion systems (T2SSs) promote secretion of folded proteins playing important 31 roles in nutrient acquisition, adaptation and virulence of Gram-negative bacteria. Protein 32 secretion is associated with the assembly of type 4 pilus (T4P)-like fibres called pseudopili. 33 Initially membrane embedded, pseudopilin and T4 pilin subunits share conserved 34 transmembrane segments containing an invariant Glu residue at the 5th position, E5. 35 Mutations of E5 in major T4 pilins and in PulG, the major pseudopilin of the Klebsiella T2SS 36 abolish fibre assembly and function. Among the four minor pseudopilins, only PulH required 37 E5 for secretion of pullulanase, the substrate of the Pul T2SS. Mass-spectrometry analysis of pili resulting from the co-assembly of PulG^{E5A} variant and PulG^{WT} ruled out an E5 role in pilin 38 39 processing and N-methylation. A bacterial two-hybrid analysis revealed interactions of the full-length pseudopilins PulG and PulH with the PulJ-Pull-PulK priming complex and with the 40 assembly factors PulM and PulF. Remarkably, PulG^{E5A} and PulH^{E5A} variants were defective 41 42 in interaction with PuIM, but not with PuIF and co-purification experiments confirmed the E5-43 dependent interaction between native PulM and PulG. These results reveal the role of E5 in 44 a recruitment step critical for assembly of the functional T2SS, likely relevant to T4P 45 assembly systems.

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49 Introduction

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51 Bacterial interactions with their environment involve the transport of proteins and other 52 macromolecules across their cell envelope. In Gram-negative bacteria, several sophisticated 53 molecular machines facilitate protein secretion across two bilayer membranes (Costa et al., 54 2015). Among these, the type II secretion system (T2SS) promotes specific transport of 55 folded proteins across the outer membrane once they have reached the periplasmic 56 compartment though the Sec or the Tat systems (Voulhoux et al., 2001; Korotkov et al., 57 2012; Nivaskumar and Francetic, 2014). T2SSs are structurally and functionally similar to 58 type IV pilus (T4P) assembly systems (Berry and Pelicic, 2014). Unlike type IV pili, which 59 extend beyond the bacterial surface to promote attachment and motility, the T2SS 60 pseudopilus fibres are thought to be restricted to the periplasm. However, plate-cultured 61 bacteria that overexpress T2SS-encoding genes present extended pseudopili (hereafter 62 referred to as T2SS pili) on the cell surface, a feature that has facilitated their biochemical 63 and structural characterisation (Sauvonnet et al., 2000; Durand et al., 2003; Kohler et al., 64 2004). T2SS pili are helical homo-polymers composed of one major pseudopilin subunit 65 called PulG in K. oxytoca (Sauvonnet et al., 2000; Kohler et al., 2004). Mechanistic insights 66 gained from their structure function analysis have revealed that interactions between neighbouring pilus subunits P and P⁺¹ tightly correlate with fibre assembly and function in 67 68 protein secretion (Campos et al., 2010). The fact that these essential contacts take place in 69 the membrane implies a one-start helix rotational assembly mode for these and other type 4 70 pilus-like fibres (Nivaskumar et al., 2014).

Pseudopilus biogenesis is a multistep process that begins with co-translational insertion of pseudopilins in the inner membrane (IM) *via* the SRP and the Sec machinery (Arts *et al.*, 2007; Francetic *et al.*, 2007). Pseudopilins are made as precursors anchored in the IM through a positively charged N-terminal peptide that is removed prior to assembly by an inner membrane protease called the prepilin peptidase (Nunn and Lory, 1991). Following

cleavage of this peptide after a conserved Gly residue at the base of the transmembrane
segment, the N-terminal domain of the prepilin peptidase transfers a methyl group to the new
N-terminal residue of the mature pilin (Strom *et al.*, 1993; Aly *et al.*, 2013).

79 In addition to the highly abundant major pseudopilins that build the helical homo-80 polymeric fibre, T2SSs contain four minor pseudopilins, essential for function. Purified 81 globular domains of three of these minor subunits, GspJ, GspJ and GspK from 82 enterotoxigenic Escherichia coli form a quasi-helical complex, which is predicted to cap the 83 pseudopilus tip (Korotkov and Hol, 2008). Functional, biochemical and molecular dynamics 84 analysis of their full-length equivalents in the Pul T2SS showed that Pull, PulJ and PulK self-85 assemble in the IM in the absence of other T2SS factors (Cisneros et al., 2012a). The PulJ-86 Pull-PulK complex promotes initiation of pseudopilus assembly and presumably provides a 87 template for subsequent incorporation of PulH and PulG subunits. Mutants lacking Pull, PulJ 88 or PulK, but not PulH, assemble fewer PulG pili and are completely defective for secretion of 89 the specific Pul T2SS substrate, pullulanase (PulA) (Cisneros et al., 2012a). Biochemical 90 studies of the Pseudomonas aeruginosa Xcp T2SS show that the periplasmic domain of the 91 PulH homologue XcpU forms a complex with other minor pseudopilins in vitro by binding to 92 the PulJ homologue XcpW (Yanez et al., 2008; Korotkov and Hol, 2008; Douzi et al., 2009). 93 In addition, the major pseudopilin XpsG of Xanthomonas campestris T2SS and the minor 94 subunit XpsH, a PulH homologue, interact directly in vivo (Hu et al., 2002). Based on these 95 studies, PulH homologues are thought to provide a link between the priming complex and the 96 major pseudopilins (Yanez et al., 2008; Korotkov and Hol, 2008; Douzi et al., 2009; Cisneros 97 et al., 2012a).

Assembly of PulG fibres requires a functional complex called the assembly platform (AP) composed of the IM proteins PulC, PulF, PulL and PulM (Py *et al.*, 2001; Possot *et al.*, 2000). The cytoplasmic domain of PulL binds the hexameric ATPase PulE that provides energy for pseudopilus assembly (Abendroth *et al.*, 2005; Camberg *et al.*, 2007). Biochemical and structural analyses of AP components in T2SSs have provided molecular insights into their soluble domains and hints about their interactions and organization

104 (Korotkov *et al.*, 2011; Abendroth *et al.*, 2004b; Abendroth *et al.*, 2004a; Sandkvist *et al.*, 105 1995; Johnson *et al.*, 2007; Abendroth *et al.*, 2005). This information has led to the current 106 T2SS model, where the IM platform promotes pseudopilus assembly within a cage-like 107 compartment delimited by the secretin channel formed by a GspD dodecamer in the outer 108 membrane and the GspD-interacting partner GspC in the IM (Gsp being a generic term to 109 designate T2SS components) (McLaughlin *et al.*, 2012).

110 The hydrophobic segments of major (pseudo)pilins in bacterial T4P and T2SSs are 111 highly conserved and rich in branched chain amino acid residues. In addition to the 112 conserved Gly residue preceding the prepilin peptidase cleavage site, mature subunits 113 contain an invariant Glu residue at position 5 (E5), which is located near the cytoplasmic side 114 of the transmembrane segment and is essential for function (Strom and Lory, 1991; Aas et 115 al., 2007; Pugsley, 1993). In the T4P systems, residue E5 has been implicated in prepilin 116 processing (Pasloske and Paranchych, 1988) or N-methylation (Strom and Lory, 1991; Aas 117 et al., 2007). Based on structural studies of T4P, it has also been proposed that E5 is 118 involved in docking of pilins to the growing fibre during assembly, by forming electrostatic 119 contacts with the N-terminal amine of the last pilin subunit incorporated in the pilus (Parge et 120 al., 1995; Craig et al., 2006).

121 In contrast, the E5V substitution in PulG from K. oxytoca T2SS did not affect 122 processing and N-methylation (Pugsley, 1993). Analyses of the PulG pilus structure and 123 assembly show that E5 is not implicated in docking, but participates in fibre stabilisation. 124 More precisely, once the protomer P has been extracted from the membrane, its E5 residue interacts with two lysine residues (K28 and K35) of protomer P⁻³ (Campos et al., 2010; 125 126 Nivaskumar et al., 2014). However, the Ala substitutions of these Glu and Lys residues lead to strikingly different phenotypes: while $pulG^{K28A/K35A}$ mutants are fully functional in protein 127 secretion, *pulG^{E5A}* mutants are completely defective. This and other data suggest an 128 129 additional role for E5 in a step preceding the docking step in the pilus assembly pathway 130 (Nivaskumar et al., 2014). With the exception of the interactions between EpsG and the AP

131 component EpsL in Vibrio cholerae T2SS that have been revealed by chemical cross-linking 132 (Gray et al., 2011), little is known about the early events of the assembly process. To shed 133 some light on these steps, we sought to characterise interactions between minor and major 134 pseudopilins and to identify AP components that directly interact with pseudopilins using the 135 K. oxytoca T2SS as a model system. By combining bacterial two-hybrid and co-purification 136 approaches with mass spectrometry and functional analyses, we identified several 137 interacting partners of the K. oxytoca major pseudopilin PulG and revealed a novel role of the 138 conserved pseudopilin residue E5, which holds important clues to the assembly mechanism 139 of the T4P fibre family.

- 140
- 141 **Results**
- 142

143 Mass spectrometry analysis of PulG and PulG^{E5A} co-assembled into mixed pili

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145 As mentioned above, earlier studies showed that residue E5 is not required for PulG 146 processing or N-methylation (Pugsley, 1993), while studies of T4P suggest the opposite 147 (Strom and Lory, 1991; Aas et al., 2007). The assembly-defective pilin variants with E5 148 residue substitutions have been shown to co-assemble into pili in the presence of wild type 149 pilins (Pasloske, et al., 1989; Aas et al., 2007). However, while N-terminal sequencing of 150 Neisseria gonorrhoeae T4P made of PilE wild type and E5L variants revealed under-151 methylation of the pilins, it was not clear whether this under-methylation was specific to PilE^{E5L} (Aas *et al.*, 2007). To clarify the role of E5 in PulG, we used a similar approach and 152 tested whether the assembly-defective PulG^{E5A} variant can be co-assembled into pili with 153 154 PulG^{WT}. Therefore, we co-expressed the full set of genes encoding the Pul T2SS (on plasmid pCHAP8185, which includes PulG^{WT}) with the gene encoding assembly-defective PulG^{E5A} 155 156 variant (on plasmid pCHAP7790, Table 1). The surface proteins were isolated from these 157 bacteria and analysed by SDS-PAGE and Coomassie Blue staining (Experimental 158 Procedures and Figure 1A). A prominent band of approximately 14.5 kDa, the molecular

159 mass of PulG, was present in the sheared fraction (Figure 1A, lane 1). This band was not present in samples from the negative control strain producing PulG^{E5A} and the Pul T2SS 160 161 lacking PulG (Δ G, Figure 1A, lane 2). The intensity of this band was higher in the strain 162 where both plasmids carried the wild type copy of the *pulG* gene (Figure 1A, lane 3), 163 compared to the strain carrying a $\Delta pulG$ version of T2SS complemented with wild type pulG(Figure 1A, lane 4). To test whether PulG^{E5A} was co-assembled with PulG^{WT}, the gel slices 164 corresponding to the 14.5-kDa band were excised from lanes 1 (strain producing PulG^{WT} and 165 PulG^{E5A}) and 4 (producing PulG^{WT}) and analysed for protein content using top-down mass 166 167 spectrometry (analysis of intact proteins without any digestion) (Gault et al., 2014; Gault et al., 2015). For PulG^{WT} sample (Figure 1B), two major peaks were found at 14607.56 and 168 169 14621.54 Da, corresponding to the molecular mass of PulG (theoretical Mr: 14607.51 Da) and methylated PulG (theoretical Mr: 14621.53 Da). For the PulG^{WT} + PulG^{E5A} sample 170 (Figure 1C), two supplementary peaks corresponding to PulG^{E5A} (experimental: 14549.49 171 Da; theoretical: 14549.50 Da) and methylated PulG^{E5A} (experimental: 14563.53 Da; 172 173 theoretical: 14563.52 Da) were observed.

To confirm the N-terminal methylation and presence of an alanine residue (A) instead of a Glu (E) at the 5th position in the sequence, we performed High Energy Collision Dissociation (HCD) fragmentation of the (14⁺) ions of both methylated PulG and methylated PulG^{E5A}. The MS/MS spectra are shown in Figure 2. The presence of series of b-type ions in both cases confirms without any ambiguity both the methylation of the N-terminus and the identity of the 5th amino acid.

These results show that PulG^{E5A} can assemble into pili, provided it is co-produced with PulG^{WT}. This suggests that PulG^{E5A} likely co-assembles with native PulG into mixed pili. The presence of methylated and non-methylated forms of both PulG variants in the pili shows further that methylation of pseudopilins is not required for assembly.

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185 Differential requirements of residue E5 for the major and minor pseudopilin function

187 With the exception of the PulK subunit, which caps the priming complex, all minor 188 pseudopilins in T2SS have the conserved E5 residue. Given its strong conservation among 189 (pseudo)pilins, we wondered whether E5 was also essential for minor pseudopilin function. 190 To address this question, we introduced E5A substitutions in minor subunits PuIH, PuII and 191 PulJ and assessed their ability to support PulA secretion in conditions of low, chromosome-192 like levels of *pul* gene expression. The results of three independent secretion assays are 193 shown in Figures 3A and supplementary Figure S1, and their quantification is shown in Figure 3B. Under these conditions, the PulG^{E5A} variant was as defective for PulA secretion as 194 195 the negative control strain lacking PulG (Figure 3A, lanes 1-6), as shown previously 196 (Nivaskumar et al., 2014), The pulH deletion mutant supported a low level of PulA secretion 197 (Figure 3A, lanes 11,12) confirming that PulH is required for efficient secretion in 198 physiological expression conditions (Cisneros et al., 2012a). Surprisingly, PulA secretion in the presence of the PulH^{E5A} variant was nearly abolished, suggesting a dominant negative 199 200 effect of the *pulH^{E5A}* allele (Figure 3A, lanes 7-12). In contrast, E5A substitutions in minor 201 pseudopilins Pull and PulJ did not affect their function (Figure 3A, lanes 13-18 and 19-24). Replacing the Met-5 with Glu in variant PulK^{M5E} had no effect on PulA secretion (Figure 3A, 202 203 lanes 25-30), confirming previous results obtained in conditions of *pul* gene overexpression 204 (Vignon et al., 2003). Thus, the requirement for E5 appears to be specific to PulG and PulH 205 pseudopilins, which are not part of the priming tip complex and have been implicated in the 206 downstream steps of pseudopilus assembly (Cisneros et al., 2012a).

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208 Pseudopilin interaction network

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The PulJ-Pull-PulK complex presumably provides a template for PulH incorporation (Cisneros *et al.*, 2012a), as suggested by the formation of the quaternary complex comprising all minor pseudopilin periplasmic domains in the *P. aeruginosa* Xcp T2SS (Yanez *et al.*, 2008; Korotkov and Hol, 2008; Douzi *et al.*, 2009). However, this complex did not interact with the globular domain of the major pseudopilin XcpT *in vitro* (Douzi *et al.*, 2009),

215 leaving the question of the mechanism that couples the initiation and elongation stages of 216 assembly open. Based on the findings of Hu and collaborators (Hu et al., 2002), it has been 217 proposed that the minor pseudopilin quaternary complex and the major pseudopilin interact 218 instead though their hydrophobic segments (Douzi et al., 2009). To test this possibility and to 219 characterise the pseudopilin interaction network, we employed the bacterial two-hybrid 220 (BAC2H) approach that allows one to study interactions between full-length, membrane-221 embedded proteins (Karimova et al., 1998). Using this method we had previously 222 demonstrated interactions of Pull with PulJ and PulK (Cisneros et al., 2012a). In the present 223 study, we fused the T18 and T25 fragments of adenylyl cyclase CyaA from Bordetella 224 pertussis to the N-terminus of all five full-length mature pseudopilins and performed BAC2H 225 assays (Experimental Procedures). Using T18-PulG as bait, we identified strong interactions 226 with two minor pseudopilins, PulH and PulJ (Figure 4). When T18-PulH was used as bait, a 227 specific interaction was observed only with T25-PulG. Although both T18-PulH and T25-PulH 228 chimera consistently showed a tendency to interact with PulJ, these interactions were not 229 statistically significant despite low P values (0.106 and 0.201, respectively) (Figure 4A and 230 Supplementary Dataset 1). The T18-Pull chimera showed specific interactions with PulJ and 231 PulK (Figure 4), in agreement with previous studies in several T2SSs (Cisneros et al., 2012a; 232 Korotkov and Hol, 2008; Douzi et al., 2009). Conversely, T18-PulJ interacted with Pull 233 (Figure 4A). In addition, T18-Pull, T18-PulJ and T18-PulK showed a weak interaction with 234 PulG. Of note, only PulG formed homo-dimers in the BAC2H assay, consistent with the 235 unique ability of major pseudopilins to form long homo-polymers (Durand et al., 2005). The 236 results of these analyses, summarised schematically in Figure 4B, reveal multiple novel 237 contacts between PulG and minor pseudopilins. Overall, the data are compatible with the 238 formation of complexes containing all five pseudopilin species.

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240 Pseudopilin interactions with IM assembly platform components

242 Previous functional studies suggest that the minor pseudopilin complex primes fibre 243 assembly and that it might activate the ATPase PulE that is associated with the assembly 244 platform (AP) complex (Cisneros et al., 2012a). The AP is essential for T2SS function and 245 contains multiple copies of IM proteins PuIC, PuIF, PuIL and PuIM (Py et al., 2001; Possot et 246 al., 2000). To understand the physical and functional link between AP and the pseudopilin 247 complex and to elucidate molecular details of the initiation and elongation steps of assembly, 248 we assessed interactions between the pseudopilins and membrane-embedded AP 249 components by using the BAC2H system. Therefore, we fused the T18- and T25- CyaA 250 fragments to the N-terminal ends of full-length AP components PuIC, PuIF, PuIL and PuIM 251 (Experimental Procedures). In the BAC2H assays, both T18-PulG and T25-PulG interacted 252 specifically with the corresponding PulM- and PulF-CyaA chimera, but not with PulC-253 chimera. T18-PulG interacted very weakly with the T25-PulL, but the reciprocal pair did not 254 interact (Figure 5A), which was surprising, given that their homologues in V. cholerae, EpsL 255 and EpsG, interact in a cross-linking study (Gray et al., 2011). We confirmed that this was 256 not due to non-functional PulL chimera, since the latter did interact with corresponding PulM 257 hybrids (Figure 5A, black bars on the right), as expected based on previous studies (Possot 258 et al., 2000; Buddelmeijer et al., 2006). The results, summarized in Figure 5B, revealed 259 strong and specific interactions of PulG with the PulM and PulF components of the assembly 260 platform, suggesting their direct role in pseudopilus elongation.

Among the minor pseudopilins, PulH also interacted with PulF and PulM, although these interactions seem weaker than those observed between PulG and PulF or PulM, based on beta-galactosidase activity levels (Figure 5A). A weak "one-way" interaction was also detected between T18-PulK and T25-PulM chimera. In addition, PulF interacted weakly but significantly with Pull, PulJ and PulK in one out of two protein hybrid pairs. The high betagalactosidase activity of strains producing T18-PulF and T25-PulF chimera in the BAC2H system (Figure 5A) shows a strong propensity of PulF for dimer formation.

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269 The PulG^{E5A} and PulH^{E5A} variants are specifically defective in the interaction with PulM

271 Given the suggested role of E5A in an early step of the assembly process (Campos et 272 al., 2010; Nivaskumar et al., 2014), we asked whether the E5A mutation affects interactions of PulG with one of the assembly factors. In BAC2H assay the PulG^{E5A} variant showed equal 273 274 or stronger interaction signal with PuIF, compared to native PuIG, as indicated by the high beta-galactosidase activities of T18-PulG^{E5A} or T25-PulG^{E5A} hybrids (Figure 6A). In contrast, 275 the same PulG^{E5A} hybrids showed a significantly decreased interaction with PulM (Figure 276 277 6A). These results suggest that mutation E5A specifically alters the interface between PulG 278 and PulM. Likewise, while the E5A substitution in PulH did not affect the PulH-PulF 279 interaction, it reduced the interaction with PulM, supporting a role of E5 in PulM binding 280 (Figure 6B).

281 The PulG-PulM interaction data were further validated with native, full-length proteins. 282 We analysed PulG interactions with PulM and the effect of the PulG E5A substitution using a co-purification approach. PulG-His₆, PulG^{E5A}-His₆ or PulG^{WT} were co-produced with PulM in 283 284 the absence of other T2SS components (Experimental Procedures). Membrane proteins 285 were extracted from these strains with Triton-X-100 and separated on the Ni-NTA columns. 286 PulM was co-eluted with PulG-His₆ (Figure 7A, lanes 6-11). Despite the higher levels of the PulG^{E5A}-His₆ variant produced and retained on the column, the quantity of co-purified PulM 287 288 was significantly reduced (Figure 7A, lanes 17-22). Quantification of the relative amounts of 289 PulG and PulM shows that the amount of co-eluted PulM was reduced nearly ten-fold, 290 relative to PulG-His₆ (Figure 7B). As shown in Figure 7A, PulG and PulM were present in 291 very low amounts in the elution fractions when membrane proteins were extracted from the 292 control strain producing untagged PulG (lanes 28-33), indicating very low nonspecific binding 293 to the Ni-NTA resin.

The results of these affinity co-purification experiments were in good agreement with those of the BAC2H analysis, revealing specific binding of PulM to PulG *in vivo*. They confirm that the E5A substitution in PulG, which abolishes pseudopilus assembly and protein secretion, specifically affected the interaction between PulG and PulM.

299 Discussion

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301 Archaeal pili and flagella, as well as bacterial T2SS, T4P and competence systems use 302 similar machineries to assemble helical fibres from membrane-anchored (pseudo)pilin 303 subunits, suggesting a common mechanism. The globular (pseudo)pilin domains, exposed 304 on the fibre surface, show variability in size and sequence, presumably optimized to promote 305 specific functions (Berry and Pelicic, 2014; Nivaskumar and Francetic, 2014; Shahapure et 306 al., 2014). In contrast, their hydrophobic segments are highly conserved, which might reflect 307 constraints imposed by their biogenesis and assembly pathway (Campos et al., 2013). For 308 example, high hydrophobicity is key for efficient targeting to the SRP and Sec pathways 309 during integral membrane protein biogenesis (Francetic et al., 2007; Arts et al., 2007) and 310 additional constraints might be linked to specific interactions with the prepilin peptidase or 311 with fibre assembly factors. Here we used the Klebsiella T2SS as a model to study 312 pseudopilus assembly, focusing on the most conserved (pseudo)pilin residue E5 playing an 313 essential role in this process.

314 Although the E5 residue is involved in pseudopilus stabilisation via long-range 315 interactions, genetic data suggest it plays a critical role at a step preceding pseudopilin 316 docking. Using two complementary mass spectrometry methods we demonstrated unambiguously that PulG^{E5A} co-assembles with PulG^{WT} into mixed PulG pili containing the 317 methylated and non-methylated forms of PulG and PulG^{E5A}. This confirms previous findings 318 319 that E5A substitution does not prevent N-methylation of PulG (Pugsley, 1993) and shows 320 that methylation is not essential for pseudopilus assembly. This is in agreement with 321 assembly of T4P in the methylase-deficient *pilD* mutants of *P. aeruginosa* (Pepe and Lory, 322 1998) and with the small amount of non-methylated PulG detected in T2SS pilus 323 preparations (Köhler et al., 2004). Compared to PulG^{WT}, the degree of N-methylated PulG^{E5A} 324 was lower, which might be a consequence of *pulG* overexpression increasing the load on the methyl-transferase function of the prepilin peptidase PulO. Of note, the proportion of PulG^{E5A} 325

326 in the mixed pili was substantially lower than 50%, which could be explained by the 327 dependence on PuG^{WT} for assembly, as well as by the predicted impact of the E5A substitution on fibre stability, due to the loss of long-range interactions with K28^{P-3} and K35^{P-3} 328 (Nivaskumar et al., 2014). We propose that co-assembly of PulG^{E5A} relies on its ability to 329 330 form heterodimers with PulG^{WT} via the previously characterised electrostatic contacts of 331 conserved residues E44 and D48 of the incoming subunit (P⁻¹) with R87 and R88 of the P 332 subunit incorporated in the PulG pilus (Campos et al., 2010; Nivaskumar et al., 2014). This association with PulG^{WT} in the membrane would bypass the defect of PulG^{E5A} possibly 333 334 related to its targeting to the assembly site, as discussed below.

335 Pseudopilus assembly requires the formation of the PulJ-PulK priming complex, 336 essential for protein secretion by the T2SS (Cisneros et al., 2012a). While our previous 337 studies demonstrated the correlation between the formation of this complex and pseudopilus 338 assembly activation (Cisneros et al., 2012a; Cisneros et al., 2012b), the molecular events 339 linking the two processes have not been identified. To understand this link, we employed the 340 BAC2H approach and characterised the network of interactions between full-length 341 membrane-embedded pseudopilins and between individual pseudopilins and the AP 342 components. The results confirmed the Pull-PulJ and Pull-PulK interactions observed in vitro 343 in T2SSs from Vibrio (Yanez et al., 2008), E. coli (Korotkov and Hol, 2008) and P. 344 aeruginosa (Douzi et al., 2009). In the BAC2H assay, the full-length PulH showed only a 345 weak tendency to interact with PulJ, although the periplasmic domains of their Pseudomonas 346 homologues interacted in vitro (Douzi et al., 2009). In addition, contrary to the latter study, we 347 identified direct contacts of PulG with PulH, which are in agreement with the results obtained 348 in Xanthomonas T2SS studies (Hu et al., 2002). In the P. aeruginosa T4P, the BAC2H 349 analysis revealed very similar interactions between FimU, the PulH structural homologue, 350 and the major pilin PilA, showing that these contacts are conserved in both systems (Nguyen 351 et al., 2015). Morover, in the same study, FimU also failed to interact with FimW, the PulJ 352 equivalent in this T4P system. Finally, we also identified novel direct contacts of PulG with 353 PulJ that are in good agreement with the proposed structural models wherein GspH, but also

354 GspG, could be docked directly to GspJ at the base of the tip complex (Korotkov and Hol, 355 2008). An interaction between PulG and PulJ might bypass the requirement for PulH, 356 explaining the minor effect of *pulH* deletion on PulG pilus assembly (Cisneros *et al.*, 2012a) 357 and on PulA secretion (Sauvonnet et al., 2000) in conditions of pul gene overexpression. 358 However, when the T2SS components are not overproduced, this requirement becomes 359 more critical, consistent with the strongly defective PulA secretion in the absence of PulH 360 (Figure 3). In addition, all minor pseudopilin subunits showed a weak tendency to interact 361 with PulG in the BAC2H assay, which might explain the capacity of Pull-PulJ and Pull-PulK 362 minor pseudopilin pairs to partially restore initiation of PulG assembly in *ApulHIJK* mutants 363 (Cisneros et al., 2012a).

364 How do pseudopilins connect to the AP and with the associated ATPase, essential for 365 fibre elongation? The results of our BAC2H analysis show that PulH and PulG, but not the 366 other minor pseudopilins, interact specifically with two AP components, PulM and PulF, 367 which might provide a basis for the recruitment of the initiation complex to the assembly site. 368 We demonstrated that PulG and PulH interact with the conserved AP component PulF, 369 presumably via TM segments of the two partners, given the absence of prominent 370 periplasmic segments in PulF. In view of their high conservation, PulF and its equivalents in 371 T4P assembly systems might be the main factors imposing restraints on the pilin primary 372 sequence. The sequence conservation of pseudopilin TM segments is consistent with weak 373 but significant binding of all minor pseudopilins to PulF in the BAC2H assay (Figure 5). 374 Although we cannot exclude the possibility that these weak interactions anchor the PulJ-I-K 375 priming complex to PuIF, we favour the model wherein the quaternary minor pseudopilin 376 complex binds to PuIF via PuIH. In addition, PuIF showed a strong tendency to form dimers 377 in this assay, consistent with the crystallography data (Abendroth et al., 2009) and with the 378 observed oligomerisation of its homologue PilC from Thermus thermophilus (Karuppiah et 379 al., 2010). In the recent study of the Myxococcus xanthus T4P architecture by cryo-electron 380 tomography, the PulF homologue PilC also appears to form a dimer (Chang et al., 2016). 381 During fibre elongation, PuIF dimers could act as membrane scaffolds or docking sites for the

382 minor pseudopilin complex (or the nascent pilus) on one hand, and the incoming PulG 383 subunits on the other, to facilitate assembly. In the large-scale BAC2H interaction study of 384 T4P from Neisseria meningitidis, the major pilin PilE interacted with the PulF homologue 385 PilG, and both PilE and PilG formed homodimers, much like PulG and PulF (Georgiadou et 386 al., 2012). In the P. aeruginosa and Myxococcus xanthus T4P, the PulF homologue called 387 PilC interacts with the ATPase PilB (Takhar et al., 2013; Bischof et al., 2016), consistent with 388 its role in transduction of mechanical energy generated by the ATPase. In the E. coli bundle-389 forming pili (Bfp) the ATPase activity of BfpD is stimulated by specific regions of the PulF 390 homologue BpfE (Crowther et al., 2005).

391 Importantly, the BAC2H results revealed interactions of PulG and PulH with the PulM 392 component of the AP. Moreover, the direct interaction of native PulG and PulM was 393 confirmed by a complementary co-purification approach. So far, PulM has been implicated in 394 stabilization of PulL within the AP complex (Possot et al., 2000). While the results of the 395 BAC2H analysis confirm PulM interaction with PulL, widely observed and well-studied in 396 other T2SSs (Py et al., 2001; Johnson et al., 2007; Lallemand et al., 2013), the strong and 397 specific interactions with PulG and PulH reveal a novel role for PulM, possibly in pseudopilin 398 recruitment to the assembly complex. Furthermore, we have shown that E5A substitutions in 399 PulG and PulH interfere with their interactions with PulM in the absence of any other Pul 400 factors, which implies that, in both cases, the E5 residue is a key feature of the PulG-PulM 401 and PulH-PulM interfaces. These interaction defects caused by the E5A substitution likely account for the dramatic assembly defect of PulG^{E5A} variant. The trans-dominant effect of the 402 *pulH*^{E5A} allele might be due to the reduced ability of PulH^{E5A} to target the priming complex to 403 404 PulM, as discussed below. In contrast, residue E5 is dispensable for function of Pull and PulJ 405 subunits, although it might provide optimal packing with the N-termini of the distal 406 neighbouring pseudopilins in the complex, as proposed previously (Craig et al., 2003).

407 Our results show that PulG interacts strongly and specifically with PulM, but only very 408 weakly with PulL, which apparently contradicts the studies in *Vibrio*, showing that EpsG could 409 be cross-linked with EpsL (Gray *et al.*, 2011). This difference in interaction might reflect

410 differences between the Vibrio and Klebsiella T2SSs, or could be due to a difference in the 411 techniques used to detect G-L interactions. The interaction of EpsG with EpsL was observed 412 using chemical cross-linking, which might enhance and stabilize a weak or indirect 413 association (Gray et al., 2011). Here, we identified direct associations of PulG with PulM and 414 PulF in the absence of other T2SS components using BAC2H, and also, in the case of PulG 415 and PulM, by co-purification in the presence of nondenaturing detergents. Nevertheless, both 416 studies support a similar model wherein interactions of major pseudopilin with the L-M 417 complex result in pilin recruitment to the assembly site.

418 The architecture of the T2SS, proposed based on partial high- and low-resolution 419 structural data of individual components (McLaughlin et al., 2012) is very similar to that of the 420 Thermus thermophilus or Myxococcus xanthus T4P determined by cryo-electron tomography 421 (Gold et al., 2015; Chang et al., 2016). In all models, the secretin and the AP components 422 homologous to PuIC, PuIL and PuIM delimit a cage-like compartment that encloses PuIF 423 within and connects directly with the ATPase PulE at its base. While cryoEM tomography 424 shows that the pilus occupies the interior of this compartment, PilO (a PulM homologue) and 425 its interacting partner PilN (corresponding to the transmembrane and periplasmic domains of 426 PulL) were placed in the lower periplasmic ring surrounding the cavity (Chang et al., 2016).

427 Integrated with the previous data, the results of this study allow us to propose a 428 schematic working model of pseudopilus assembly initiation and elongation (Figure 8). In this 429 model, one of the initial steps, following pseudopilin maturation by the prepilin peptidase, 430 would be the assembly of the priming complex PulJ-PulI-PulK and its binding to PulH and 431 PulG (step 1). Consistent with the direct binding of PulH and PulG to PulF, and with PulF 432 self-interaction, this complex would associate with a dimer of PuIF (step 2). In step 3, the 433 pseudopilins and PuIF would bind to PuIM, outside or within the pre-assembled complex that 434 includes the secretin PuID channel in the OM bound to the IM components PuIC and PuIL. 435 An independent assembly of this latter complex is consistent with the results of cellular 436 localisation studies of T2SS components fused to fluorescent protein tags (Buddelmeijer et al., 2009; Lybarger et al., 2009), as well as with the T4P cryo-electron tomography data 437

(Chang et al., 2016). Importantly, association of PulM with PulF-pseudopilin complex 438 439 requires residue E5 of PulG and PulH (Figure 8, step 3). Results of the T4P study by cryo-440 tomography suggest that the ATPase is recruited to the assembly site in the final step of the 441 T4P complex biogenesis (Chang et al., 2016). A similar order of assembly, in which the 442 incorporation of the PulF-pseudopilin complex into the incomplete T2S machine would 443 precede the recruitment of the ATPase PulE, is possible in T2SS (steps 4 and 5). However, 444 there is an important difference between the two systems at the level of PulL, which 445 corresponds to a fusion between PilM and PilN of T4P, leaving open a possibility that 446 ATPase recruitment occurs independently of PulF.

447 During pseudopilus elongation, PuIG, possibly in the form of dimers, would be 448 recruited to the assembly site via interaction with PuIM, through the critical residue E5 (step 449 6). PulG would dock to the available PulF "acceptor" site and bind to the membraneembedded PulG^{P+1} subunit incorporated in the pseudopilus. ATP hydrolysis could drive 450 451 rotation of the pilus via PulF to spool the protomer P into the growing fibre (step 7), following 452 the one-start helix assembly model that we proposed and described earlier (Campos et al., 453 2010; Nivaskumar et al., 2014). Fibre elongation would comprise cycles of targeting, docking 454 and spooling events (steps 6 to 9).

Based on this model, we propose that co-assembly of PulG^{E5A} relies on its ability to 455 form heterodimers with PulG^{WT}, which would promote targeting to the assembly site 456 457 (Campos et al., 2010; Nivaskumar et al., 2014). The trans-dominant effect of PulH^{E5A} variant 458 would be due to its association to pseudopilin-PulF complex but not to PulM. While providing 459 testable hypotheses, this model raises numerous questions, notably those related to the 460 mechanism of exoprotein substrate recruitment and transport. Whether a recruitment step 461 accompanies uptake of exoprotein substrate molecules, or whether there are mechanisms 462 that control substrate entry to the preformed assembly site, are only some of the questions 463 that need to be addressed by further investigation.

The interaction network of pseudopilin subunits and assembly platform proteins in the inner membrane described here will provide the basis for further biochemical and structure

466 function analysis of pseudopilus biogenesis and its mechanistic link with T2SS-mediated467 protein secretion.

468

469 **Experimental Procedures**

470

471 Bacterial strains and plasmids

472

The Escherichia coli strain DH5 α [F' lacl^Q Δ lacZM15 pro+ Tn10)] was used for recombinant 473 474 DNA experiments. Strains PAP7460 [\(\alpha\)(lac-argF)U169 araD139 relA1 rpsL150 \(\alpha\)malE444 475 malG501] and PAP5207 [Δ (lac-argF)U169 araD139 relA1 rpsL150 pcnB::Tn10 (F' lacl^Q) 476 $\Delta lacZM15 \text{ pro+})$ were used for pul gene expression studies. The Δcya strain DHT1 (Dautin 477 et al., 2000) was used for bacterial two-hybrid experiments. The bacteria were grown in LB 478 medium supplemented with antibiotics, as required: chloramphenicol (Cm) (25 μ g ml⁻¹), 479 ampicillin (Ap) (100 µg ml⁻¹) or kanamycin (Km) (15 or 25 µg ml⁻¹). The expression of the *pul* 480 genes was induced by addition of 0.4 % maltose. Isopropyl-β-D-thiogalactoside was added to 481 induce *lacZ* promoter-controlled gene expression in strain PAP5207.

482

483 Plasmid construction

484

485 Plasmids used in this study are listed in Table 1. Plasmid DNA purification, gel extraction and 486 PCR product purification were performed using appropriate Qiagen kits. Restriction 487 enzymes, DNA ligase and other molecular biology reagents were purchased from Fermentas 488 or New England Biolabs. The high-fidelity Pwo polymerase (Roche) was used for PCR 489 amplification and site directed mutagenesis using the modified Quick-change method. The 490 list of oligonucleotides (synthesized by Sigma Genosys) used for cloning or site-directed 491 mutagenesis is provided in Supplementary Table 1. To construct the BAC2H chimera, the pul 492 genes were PCR-amplified using corresponding primers, treated with DpnI and cloned into 493 pKT25 and pUT18c vectors using *Kpn*l and *Eco*RI enzymes. All plasmids were sequenced494 by GATC.

495

496 Bacterial two-hybrid and statistical analysis

497

498 Competent cells of strain DHT1 were co-transformed with pUT18C and pKT25 derivatives 499 and bacteria were grown for 48 h at 30°C on LB plates containing Ap and Km. Colonies were 500 picked at random and inoculated into 5 ml cultures in LB containing Km and Ap, grown 501 overnight and inoculated the next day into fresh medium containing 1 mM IPTG. Bacteria 502 were cultured to mid-log phase and β -galactosidase activity was measured as described 503 (Miller, 1972). At least 2 independent experiments were performed with 3 randomly picked 504 transformants. Mean values were presented by bar graphs, and error bars indicate standard 505 deviation. Microsoft Excel software was used for data processing and presentation. The 506 statistical analysis was performed using the non-parametric Kruskal-Wallis test, followed by 507 Dunn's post-test for multiple comparisons, using the Prism software.

508

509 Purification of PulG pili

510

511 PAP7460 bacteria were transformed with pCHAP8185 plasmid containing all pul T2SS 512 genes including *pulG^{WT}*, or its derivative pCHAP8184 lacking *pulG*, and with pSU18 or its derivatives pCHAP1205 (PulG^{WT}) or pCHAP7790 (PulG^{E5A}) carrying different *pulG* alleles 513 514 under control of the *lacZ* promoter. Bacteria were cultured on LB agar containing Ap, Cm and 515 0.4 % maltose. After incubating for 48 h at 30°C, bacteria were harvested and resuspended 516 in LB medium. Pili were sheared by vortex treatment and 20 passages through a 26-Gauge 517 needle. Two consecutive centrifugation steps at 16 000 g for 20 min were used to separate 518 bacteria and pili. The pilus fractions were further collected by ultracentrifugation in rotor Ti60 519 at 150 000 g and pili were resuspended in 20 mM HEPES for further analysis by SDS-PAGE 520 and mass spectrometry.

522 Mass spectrometry

523

524 All samples were desalted by C₄ ZipTip® (Millipore) and eluted directly into a 10 µL spray 525 solution of methanol:water:formic acid (75:25:3). Between 2 and 6 µL were introduced into 526 an Orbitrap Velos mass spectrometer, equipped with ETD module (Thermo Fisher Scientific, 527 Bremen, Germany) using a TriVersa NanoMate® (Advion) in positive ion mode. The spray 528 voltage was set to 1.2-1.6 kV and back-pressure to 0.3-0.4 psi. A full set of automated 529 positive ion calibrations was performed immediately prior to mass measurement. The 530 transfer capillary temperature was lowered to 100°C, sheath and auxiliary gasses switched 531 off and source transfer parameters optimised using the auto tune feature. Helium was used 532 as the collision gas in the linear ion trap. The FT automatic gain control was set at 1×10^6 for 533 MS experiments. Spectra were acquired in the FTMS in full profile mode with between 10 534 microscans over several minutes, with averaging on and set to max, and a resolution of 535 60,000 at 400 m/z. The final few spectra were then averaged using Qualbrowser in Thermo 536 Xcalibur 2.1 and deconvoluted using Xtract to produce zero charge mass spectra.

537 For MS/MS experiments, the FT automatic gain control was set at 2x10⁵. Ions corresponding 538 to the isotopic distribution of a single charge state (14⁺) were selected with the largest 539 possible window to avoid overlap with neighbouring species but minimize signal loss. HCD 540 was performed at 27 eV and spectra were acquired in the FTMS in full profile mode at a 541 resolution of 60,000 at 400 m/z, with 10 microscans and with averaging on and set to the 542 maximum value. The final few spectra were then averaged using Qualbrowser in Thermo 543 Xcalibur 2.1 and deconvoluted using Xtract to produce singly charged MS/MS spectra. 544 MS/MS spectra were interpreted manually.

545

546 SDS-PAGE and immuno-detection

548 Proteins from bacterial extracts were separated by electrophoresis on 9 % or 10 % 549 polyacrylamide gels and transferred onto nitrocellulose membranes (ECL, Amersham) using 550 a semi-dry blotting apparatus. Membranes were blocked with 5% milk in TBST (10 mM Tris, 551 150 mM NaCl, 0.05% Tween 20) and incubated with polyclonal antisera raised against 552 purified PuIA, PuIG and MalE-PuIM (diluted 1:1000 in TBST-5% milk), followed by four 10-553 min washes and incubation in horseradish peroxidase-coupled anti-rabbit antibody (1:40,000; 554 Amersham). Membranes were developed by enhanced chemiluminescence ECL2 (Thermo) 555 or Western Lightning Plus ECL (PerkinElmer) and recorded using the Typhoon phosphor-556 imager (GE) or LAS 4000 imager (Fujifilm). ImageJ software (Abramoff et al., 2004) was 557 used to quantify the density of bands.

558

559 PulA secretion assay

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561 PAP5207 bacteria were transformed with pCHAP8185 plasmid derivatives containing pul 562 genes with corresponding pseudopilin gene knockouts, and a pSU18 empty vector or its 563 derivatives containing the missing wild type or mutant pseudopilin gene (Table 1). Bacteria 564 were grown in LB containing Ap and Cm in the presence of 0.4% maltose, 1/10 volume of M9 565 salts and 1 mM IPTG to early stationary phase (OD_{600nm} >2). Cultures were normalized to 566 OD_{600nm} = 1 and bacteria were pelleted by centrifugation for 10 min at 16000 g at 4°C. 567 Bacteria were resuspended in SDS-sample buffer at final concentration of 1 OD_{600nm} per ml. 568 Supernatant fractions were centrifuged again for 10 min and mixed with equal volume of 2x 569 SDS-sample buffer. Samples corresponding to 0.05 OD_{600nm} from each fraction were 570 separated by SDS-PAGE on 9% Tris-Tricine gels. Proteins were transferred to nitrocellulose 571 membranes (ECL Amersham) by semi-dry electro-transfer. Polyclonal antisera were used for 572 immuno-detection of PulA.

573

574 Co-purification of PulG-bound proteins

Bacteria of strain PAP7460 producing PulG-His₆, PulG^{E5A}-His₆ or PulG^{WT} variants encoded 576 by pCHAP1362, pCHAP7785 and pCHAP8658, respectively and PulM (from plasmid 577 578 pCHAP2393) were cultured at 30°C in LB medium containing Cm and Ap to $OD_{600nm} = 1$. 579 Bacteria were collected by centrifugation and broken by sonication in cold TBS (20 mM Tris-HCl, pH 7.4, 150 mM NaCl) with 100 µg.ml⁻¹ of lysozyme. The lysate was cleared by 580 581 centrifugation to remove unbroken debris. Membranes were collected by ultracentrifugation 582 at 186000 x g and resuspended in cold TBS, followed by solubilisation with 2% Triton X-100. 583 Ultracentrifugation was repeated at 150000 g to remove non-solubilised membranes. Ni-NTA 584 resin beads were washed with ~10 volumes of TBS and the solubilised membrane fraction 585 was incubated with Ni-NTA resin at 4°C. The flow-through was collected, followed by seven 586 washes with two column volumes of TBS supplemented with 20 mM imidazole. Proteins 587 were eluted with TBS containing 300 mM imidazole.

588

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590

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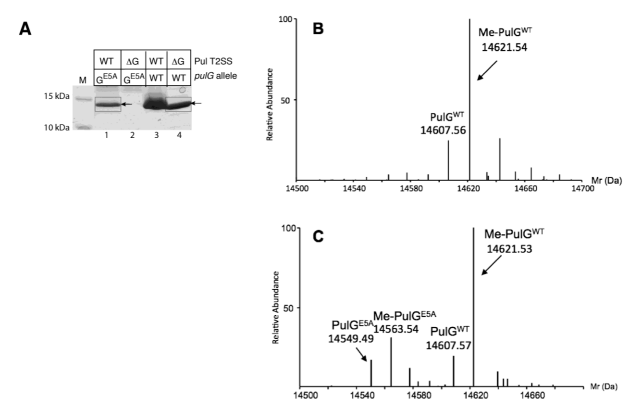
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 System of *Vibrio vulnificus. J. Mol. Biol.* **375**: 471–486.
- 786

790 Table 1. Plasmids used in this study

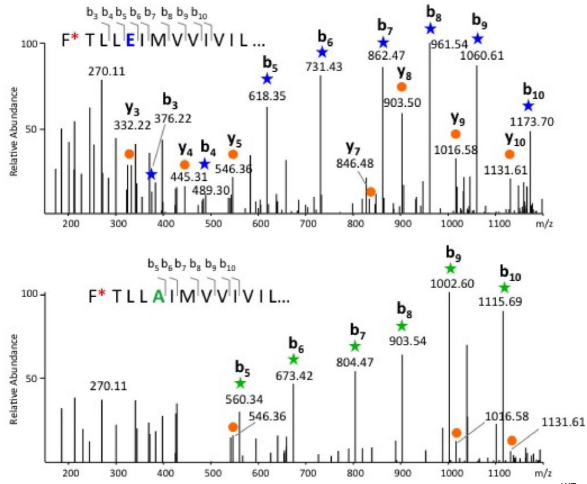
Name	Origin/resistance	Relevant markers	Source/reference
pUT18c	ColE1/Ap ^R	p <i>lacZ</i> -T18	(Karimova <i>et al</i> ., 1998)
pKT25	p15A/Km ^R	p <i>lacZ</i> -T25	(Karimova <i>et al.</i> , 1998)
pUT18c-Zip	ColE1/Ap ^R	GCN4 (Leu zipper region) fused to T18	(Karimova <i>et al</i> ., 1998)
pKT25-Zip	p15A/Km ^R	GCN4 (Leu zipper region) fused to T25	(Karimova <i>et al</i> ., 1998)
pSU18	p15A/Cm ^R	placZ, lacZ'	(Bartolomé <i>et al.</i> , 1991)
pCHAP1205	p15A/Cm ^R	pSU18 <i>pulG</i>	(Possot <i>et al</i> ., 2000)
pCHAP1329	p15A/Cm ^R	pSU18 <i>pulJ</i>	(Possot <i>et al.</i> , 2000)
pCHAP1331	p15A/Cm ^R	pSU18 <i>pulH</i>	(Possot <i>et al.</i> , 2000)
pCHAP1351	p15A/Cm ^R	pSU18 <i>pull</i>	(Cisneros <i>et al</i> ., 2012)
pCHAP1362	p15A/Cm ^R	pSU18 <i>pulG-His</i> ₆	(Kohler <i>et al</i> ., 2004)
pCHAP2393	ColE1/Ap ^R	pUC18 <i>pulM</i>	(Possot <i>et al.</i> , 2000)
pCHAP6117	p15A/Cm ^R	pSU18 <i>pulJ^{E5A}</i>	This study
pCHAP7330	ColE1/Ap ^R	pUT18c <i>pulG</i>	(Nivaskumar et al., 2014)
pCHAP7332	p15A/Km ^R	pKT25 <i>pulG</i>	(Nivaskumar et al., 2014)
pCHAP7785	p15A/Cm ^R	pSU18 <i>pulG^{E5A}-His</i> 6	This study
pCHAP7790	p15A/Cm ^R	pSU18 <i>pulG^{E5A}</i>	This study
pCHAP8113	ColE1/Ap ^R	pUT18c <i>pulC</i>	This study
pCHAP8119	p15A/Km ^R	pKT25 <i>pulC</i>	This study
pCHAP8154	ColE1/Ap ^R	pUT18c <i>pulM</i>	This study
pCHAP8155	p15A/Km ^R	pKT25 <i>pulM</i>	This study
pCHAP8184	ColE1/Ap ^R	pCHAP8185 <i>∆pulG</i>	(Campos et al., 2010)
pCHAP8185	ColE1/Ap ^R	pulS, pulA _{NA} pulB pulCDEFGHIJKLMNO	(Cisneros <i>et al</i> ., 2012)
pCHAP8201	ColE1/Ap ^R	pCHAP8185 pulH::kan	(Cisneros <i>et al</i> ., 2012)
pCHAP8209	ColE1/Ap ^R	pCHAP8185 pulJ::kan	(Cisneros <i>et al</i> ., 2012)
pCHAP8212	ColE1/Ap ^R	pCHAP8185 pulK::kan	(Cisneros <i>et al</i> ., 2012)
pCHAP8218	ColE1/Ap ^R	pCHAP8185 pull::kan	(Cisneros <i>et al</i> ., 2012)
pCHAP8245	ColE1/Ap ^R	pUT18c- <i>pull</i>	(Cisneros <i>et al</i> ., 2012)

pCHAP8246	ColE1/Ap ^R	pUT18c <i>pulJ</i>	This study
pCHAP8247	ColE1/Ap ^R	pUT18c <i>pulK</i>	This study
pCHAP8248	p15A/Km ^R	pKT25 <i>pull</i>	(Cisneros <i>et al.</i> , 2012)
pCHAP8249	p15A/Km ^R	pKT25 <i>pulJ</i>	(Cisneros <i>et al.</i> , 2012)
pCHAP8250	p15A/Km ^R	pKT25 <i>pulK</i>	(Cisneros <i>et al</i> ., 2012)
pCHAP8256	ColE1/Ap ^R	pUT18c <i>pulH</i>	This study
pCHAP8257	p15A/Km ^R	рКТ25 <i>риІН</i>	(Cisneros <i>et al.</i> , 2012)
pCHAP8364	ColE1/Ap ^R	pUT18c <i>pulF</i>	This study
pCHAP8365	p15A/Km ^R	pKT25 <i>pulF</i>	This study
pCHAP8418	p15A/Cm ^R	pSU18 <i>pulH^{E5A}</i>	This study
pCHAP8420	p15A/Km ^R	pKT25 <i>pulG^{E5A}</i>	(Nivaskumar et al., 2014)
pCHAP8434	p15A/Km ^R	рКТ25 <i>риІН^{Е5А}</i>	This study
pCHAP8446	ColE1/Ap ^R	pUT18c <i>pulH^{E5A}</i>	This study
pCHAP8472	ColE1/Ap ^R	pUT18c <i>pulL</i>	This study
pCHAP8484	p15A/Km ^R	pKT25 <i>pulL</i>	This study
pCHAP8568	p15A/Cm ^R	pSU18 <i>pulG</i>	This study
pCHAP8639	p15A/Cm ^R	pSU18 <i>pulK^{M5E}</i>	This study
pCHAP8670	ColE1/Ap ^R	pUT18c <i>pulG^{E5A}</i>	(Nivaskumar et al., 2014)

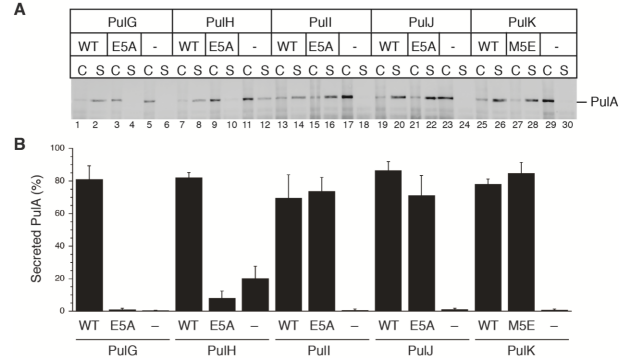




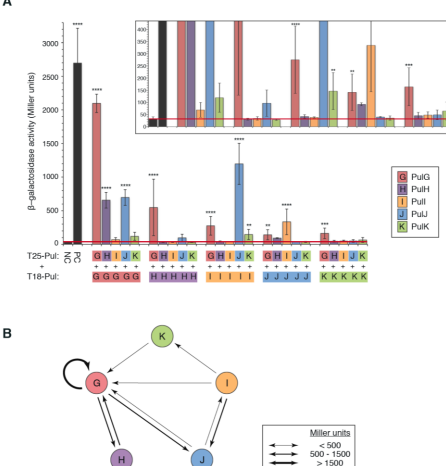
796 Figure 1. PulG^{E5A} is N-methylated and co-assembled into pili with PulG^{WT}. A. SDS-PAGE 797 and Coomassie Blue stained sheared fractions from equivalent amounts of bacteria of strain 798 PAP7460 carrying two compatible plasmids (Table 1), as indicated: lane 1, pCHAP8185 799 (encoding the complete Pul T2SS) and pCHAP7790 (encoding PulG^{E5A}); lane 2, pCHAP8184 (encoding Pul T2SS lacking PulG) and pCHAP7790; lane 3, pCHAP8185 and pCHAP1205 800 (encoding PulG^{WT}); and lane 4, pCHAP8184 and pCHAP1205. Only the relevant portions of 801 802 the gel are shown, with the molecular mass markers indicated on the left. The expected 803 molecular mass of PulG is around 14.6 kDa. The squares and arrowheads indicate samples 804 that were further analysed by mass spectrometry. B. Deconvoluted high-resolution mass spectrum (in Mr) obtained for the sample in lane 4 (PAP7460 carrying $pulG^{WT}$ allele) indicating the presence of a minor peak at 14607.56 Da corresponding to PulG^{WT} and a 805 806 major peak at 14621.54 Da corresponding to methylated PulG^{WT}; C. Deconvoluted high 807 resolution mass spectrum (in Mr) obtained for the sample in lane 1 (PulG^{WT} + PulG^{E5A}) 808 809 indicating the presence of supplementary peaks corresponding to PulG^{E5A} (14549.49 Da) and methylated PulG^{E5A} (14563.53 Da). 810 811



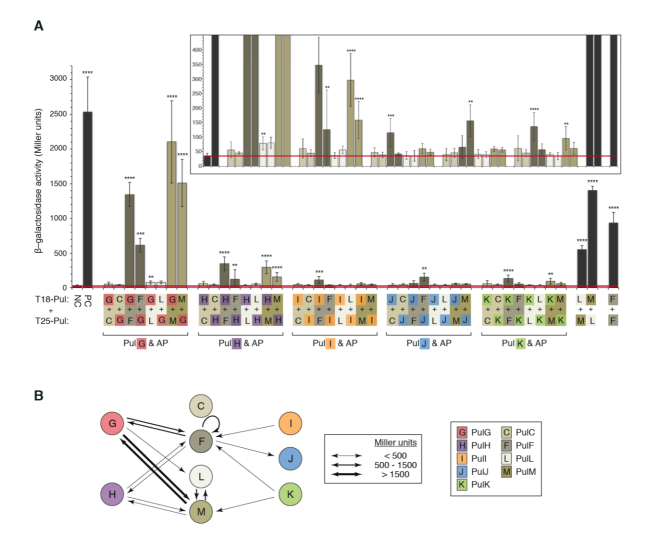
813 200 300 400 500 600 700 800 900 1000 1100 m/z
Figure 2. High-energy Collision Dissociation (HCD) analysis of methylated PulG^{WT} and
PulG^{E5A} variants co-assembled into pili. HCD fragmentation spectra of the (14+) ion of
methylated PulG^{WT} at 1046.05 (top) and methylated PulG^{E5A} at 1041.83 (bottom).
Methylation of the N-terminal residue is indicated with an asterisk. The series of N-terminal btype ions obtained in both cases and marked with a star unambiguously delineates the
presence of an E at the 5th position in PulG^{WT} and an A in PulG^{E5A}. C-terminal y-type ions are
depicted with a circle.



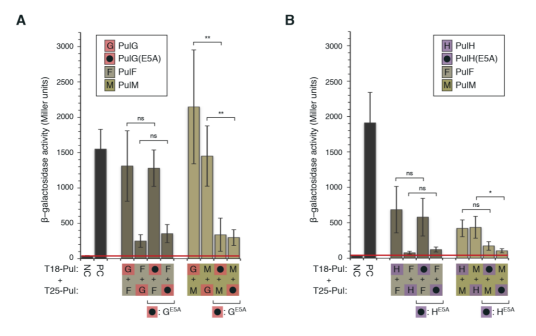
822 823 Figure 3. Effect of 5th residue substitutions on pseudopilin function. Pullulanase (PuIA) 824 secretion was assayed in E. coli strain PAP5207 containing pCHAP8185 derivatives with 825 single deletions of pseudopilin genes was complemented with pSU18 plasmid (-) or its 826 derivatives expressing the wild type (WT) or the mutant (E5A or M5E) allele of the missing 827 pseudopilin gene as indicated above the lanes. A. A representative of three independent experiments is shown. Equivalents of 0.05 OD_{600nm} of cell (C) or supernatant (S) fractions 828 829 were analysed on 9% Tris-Tricin SDS-PAGE and detected using anti-PuIA antibodies. B. 830 Quantification of the fraction of secreted PulA in the three independent experiments shown in 831 panel A and in Figure S1. Bar graphs represent the mean values and the error bars indicate 832 standard deviation.



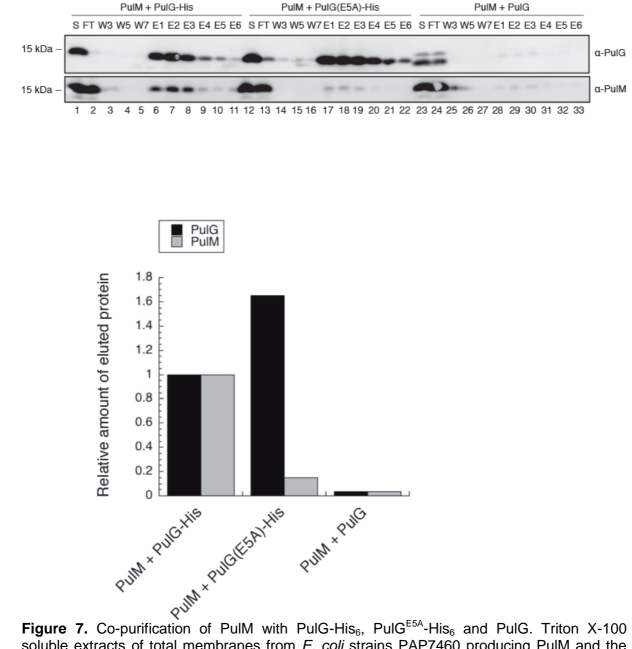
835 836 Figure 4. Pseudopilin interaction network determined by the bacterial two-hybrid analysis. A. 837 Beta-galactosidase activities of DHT1 bacteria co-producing indicated T18 and T25 chimera 838 determined as described in Experimental Procedures and expressed in Miller units. NC, 839 activity of bacteria producing T18 and T25 as negative control; PC, activity of positive control 840 strain producing T18-Zip and T25-Zip chimera. Each bar represents mean betagalactosidase activity value from at least 3 independent colonies obtained by co-841 842 transformation of pKT25 and pUT18c derivatives encoding pseudopilin chimera (T25 or T18 843 fused to the N-terminus of mature pseudopilins) indicated by the colour code in the inset: 844 PulG, red; PulH, purple; Pull, orange; PulJ, blue; and PulK, green. The colours of the bars 845 correspond to the T25-fused proteins. The red horizontal line indicates the background mean 846 beta-galactosidase activity measured in the negative control. Error bars indicate standard 847 deviation. Statistical significance relative to the negative controls is indicated above graphs 848 as follows: **** = p < 0.001; *** = P < 0.01; * = P < 0.1; ns= non-significant. The inset above the 849 main graph shows the part of the same graph with the scale of beta-galactosidase activities 850 expanded in the low range (from 0 to 500 Miller units). B. The results are summarised 851 schematically in a pentagram depicting pseudopilin arrangement in a putative right-handed 852 helical complex. Arrows are oriented from T18- to T25- chimera and their line thickness 853 indicates the strengths of significant interactions, in the range of beta-galactosidase activities 854 (in Miller units) defined as shown in the legend: weak (50>mean>500), strong 855 (500>mean>1500) and very strong (mean>1500).



857 858 Figure 5. Interactions of pseudopilins with assembly platform components identified using 859 bacterial two-hybrid analysis. A. Beta-galactosidase activities of DHT1 bacteria producing 860 indicated T18-Pul or T25-Pul hybrids as bait in the presence of AP component chimera. The Pul components analysed are indicated by the single letter and colour code in the inset. NC, 861 negative control strain co-producing T18 and T25 fragments; PC, positive control co-862 863 producing T18-Zip and T25-Zip chimera. The bars represent mean values from at least 3 864 independent colonies resulting from co-transformation of strain DHT1 with indicated pKT25 865 and pUT18c derivatives. The colours of the bars correspond to the AP proteins tested. The red line indicates the background beta-galactosidase activity measured in the NC. Error bars 866 867 indicate standard deviation. Statistically significant mean values relative to the negative 868 control (Experimental Procedures and Supplementary Dataset 1) are indicated above bars. 869 The inset above the main graph shows the part of the same graph with the expanded scale 870 of beta-galactosidase activities in the low range (from 0 to 500 Miller units). B. Summary of 871 the interaction data, with the arrows oriented from T18- to T25- chimera and with line 872 thickness corresponding to arbitrary range of interaction strengths indicated in the legend, as 873 in Figure 4. 874



876 **Figure 6.** The effect of E5A substitution on PulG and PulH interactions with PulF and PulM. **A.** Bacterial two-hybrid analysis of PulG and PulG^{E5A} interactions with PulF and PulM. **B.** Bacterial two-hybrid analysis of PulH and PulH^{E5A} interactions with PulF and PulM. Each bar represents the mean value from at least 3 independent colonies obtained by transformation of pKT25 and pUT18c derivatives containing indicated inserts. The colours of the bars correspond to the AP proteins. Error bars indicate standard deviation. NC, negative control and PC, positive control as indicated in the legend of Figure 4. The red line indicates the background beta-galactosidase activity of the NC. Statistical significance of multiple pairwise comparisons is indicated by stars, as in Figure 4.



886 887 Figure 7. Co-purification of PulM with PulG-His₆, PulG^{E5A}-His₆ and PulG. Triton X-100 888 soluble extracts of total membranes from E. coli strains PAP7460 producing PulM and the 889 indicated PulG variants were subject to affinity chromatography on Ni-NTA matrix (Experimental Procedures). A. SDS-PAGE and immuno-detection of PulG and PulM in 890 891 fractions of the co-purification. S, solubilised membrane fractions; FT, flow-through; W1, W3 892 and W7, wash fractions; E1-E6, elution fractions. B. Quantification of the relative amount of 893 eluted PulG and PulM relative to wild type levels, according to the densitometric analysis of 894 the bands in (A).

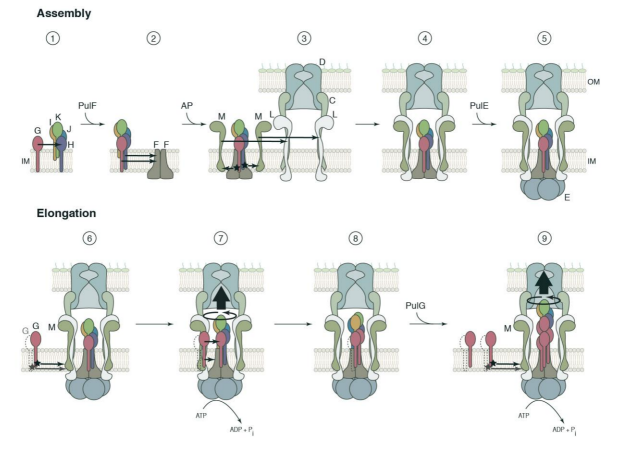


Figure 8. The working model of pseudopilus assembly and elongation. Assembly (top panel): 898 Mature major pseudopilin PulG interacts with the guaternary complex composed of the priming PulJ-Pull-PulK trimer bound to PulH (step 1). Transmembrane regions of PulG and 899 PulH drive the association of the pentameric proto-pseudopilus to a PulF dimer (2). PulM 900 901 binds to this complex via the specific interactions of PulM with E5 residues of PulG and PulH, 902 depicted as stars (3). The PulF-pseudopilin-PulM complex associates with PulL within the 903 pre-assembled secretin PuID in the OM bound to the IM protein PuIC. The initiation complex (4) recruits hexameric ATPase PulE via the cytoplasmic regions of PulL and possibly via 904 905 PulF, thus resulting in a complete, functional T2S machine (5). Elongation (bottom panel): 906 Fibre elongation begins with the recruitment of PulG subunits, likely in the form of dimers, by 907 PulM (6). As in step 3, this recruitment requires a direct contact between the E5 residue of 908 PulG and PulM (6). PulG enters the assembly platform, docks to PulF via its TM segments and associates with PulG^{P+1} protomer of the proto-pseudopilus via their globular domains (7), 909 through electrostatic contacts described previously (Nivaskumar et al., 2014). ATP hydrolysis 910 911 causes conformational changes in PulE that are transmitted to PulF through direct contact, 912 driving the rotation of the proto-fibre (7). The incoming PulG protomer is spooled into the 913 fibre, which results in an overall extension of the pseudopilus (7 and 8). Fibre growth 914 comprises multiple cycles of PulG recruitment, docking and extraction (steps 6 to 9).