Inhibiting Type VI Secretion System Activity with a Biomimetic Peptide Designed To Target the Baseplate Wedge Complex

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ABSTRACT  Human health is threatened by bacterial infections that are increasingly resistant to multiple drugs. A recently emerged strategy consists of disarming pathogenic bacteria by targeting and blocking their virulence factors. The type VI secretion system (T6SS) is a widespread secretion nanomachine encoded and employed by pathogenic strains to establish their virulence process during host invasion. Given the conservation of T6SS in several human bacterial pathogens, the discovery of an effective broad-spectrum T6SS virulence blocker represents an attractive target for development of antivirulence therapies. Here, we identified and validated a protein-protein interaction interface, TssK-TssG, as a key factor in the assembly of the T6SS baseplate (BP) complex in the pathogen enteroaggregative Escherichia coli (EAEC). In silico and biochemical studies revealed that the determinants of the interface are broadly conserved among pathogenic species, suggesting a role for this interface as a target for T6SS inhibition. Based on the high-resolution structure of the TssKFGE wedge complex, we rationally designed a biomimetic cyclic peptide (BCP) that blocks the assembly of the EAEC BP complex and inhibits the function of T6SS in bacterial cultures. Our BCP is the first compound completely designed from prior structural knowledge with anti-T6SS activity that can be used as a model to target human pathogens.

IMPORTANCE  New therapeutic options are urgently needed to fight drug-resistant and life-threatening infections. In contrast to antibiotics that inhibit the growth pathways of bacteria, the antivirulence strategy is a promising approach to disarm pathogens by interfering with bacterial virulence factors without exerting evolutionary pressure. The type VI secretion system (T6SS) is used by many pathogens, including members of the antibiotic-resistant ESKAPE bacteria (Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa, and Enterobacter spp.), to establish their virulence during the invasion of the human host. Although the T6SS is undoubtedly involved in pathogenesis, strategies targeting this virulence factor are crucially lacking. Here, we used a combination of genetics, microbiology, biochemical, biophysics, and bioinformatics approaches to rationally design a biomimetic peptide that interferes with T6SS assembly and functioning. This study represents a novel proof of concept for an antivirulence strategy which aims to interfere with the assembly of the T6SS.

KEYWORDS  bacterial secretion system, type VI secretion system, T6SS, bioinformatic, biomimetic peptide, protein-protein interface, virulence inhibitor
During the past half-century, seven new antibiotic classes have been approved by the Food and Drug Administration (FDA) (1). None of them is efficient against the Gram-negative WHO priority list ESKAPE pathogens, namely, Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa and Enterobacter species, which are responsible for approximately 75% of infections and deaths by antibiotic-resistant bacteria (2, 3). In this postantibiotic era, new therapeutic options are required to fight against drug-resistant and life-threatening infections. The antivirulence strategy is a promising approach that seeks to disarm and neutralize pathogenic bacteria by interfering with bacterial virulence factors instead of growth pathways. Virulence factors are bacterial products that promote disease by either damaging the host or circumventing and evading the immune system (4). As they are not essential for growth, blocking virulence factors does not impose a strong evolutionary pressure on bacteria and hence could serve as an alternative or complement to traditional antibiotics (5–7). Virulence factors are numerous and include secretion apparatus involved in toxin translocation across the bacterial envelope (8). To date, nine secretion systems have been discovered, and the type VI secretion system (T6SS) is one of the most recently recognized examples (9–11). Genes encoding T6SS have been identified in more than 25% of sequenced Gram-negative bacteria, including pathogenic ESKAPE strains (12, 13). The T6SSs of *P. aeruginosa*, *Aeromonas hydrophila*, and *Vibrio cholerae* act on the host cytoskeleton to promote internalization (14) or to impair phagocytic functions (15, 16) and contribute to virulence in mouse models (17, 18). The T6SS promotes the intracellular spread of *Francisella tularensis*, *Yersinia pseudotuberculosis*, *Burkholderia mallei*, and *Edwardsiella tarda*, and T6SS mutants consistently exhibit a virulence defect *in vivo* (19–22). Similarly, the T6SS of *Acinetobacter baumannii* causes a host viability decrease in *Galleria mellonella* (23), while *Salmonella enterica* serovar Typhimurium, *Shigella sonnei*, and *Vibrio cholerae* employ the T6SS to disrupt the intestinal microbiota and colonize the host gastrointestinal tract (24–27). Although the T6SS is undoubtedly involved in pathogenesis initiation, strategies targeting this virulence factor are crucially lacking (28).

The T6SS belongs to the broad family of contractile injection systems, including bacteriophages, R-pyocins, and the metamorphosis-associated contractile structure (MAC) (29–32). The T6SS machinery includes a needle-like structure loaded with effectors and wrapped into a sheath built in an extended metastable conformation from an assembly platform, the baseplate (33). The T6SS contractile sheath is made of the TssB/C subunits surrounding an inner tube composed of HCP proteins and tipped by the VgrG puncturing spike (34–37). The cytoplasmic contractile tail is anchored to the bacterial cell wall through the TssJ-TssL-TssM transenvelope complex (38–40). Contraction of the tail leads to the perforation of the target cell and the delivery of a broad repository of effectors into both eukaryotic and prokaryotic cells (41–44).

The T6SS baseplate is a central piece of the T6SS machinery as it connects the tail to the membrane complex and initiates needle polymerization (33, 45–47). This complex includes the proteins TssF, TssG, and TssE, respectively, homologous to the T4 bacteriophage baseplate proteins gp6, gp7, and gp25, as well as TssK, which shares structural properties with siphophage receptor-binding proteins (36, 45, 46). TssK interacts with TssG, which is stabilized by TssF, resulting in the formation of the TssKFG wedge complex, representing an early T6SS baseplate building block homologous to the T4 baseplate wedge complex (45, 48–50). Recently, the high-resolution structure of the TssKFG complex from the pathogen enteroaggregative *Escherichia coli* (EAEC) has been solved and provided incisive insights into the structural organization of the protein complex (50, 51). This 500-kDa complex is composed of a monomeric TssG serving as a central backbone that interacts with a TssF dimer and two TssK trimers. While TssG contacts TssF proteins throughout its structure, the interaction with the two TssK trimers is localized and mediated by two small loops (foot 1 and foot 2) at the C terminus of TssG. Foot 1 (residues 216 to 252) and foot 2 (residues 300 to 330) form two triangular loops following the C3 symmetry of the TssK trimer and interacts with its N-terminal region (residues 1 to 18). Recently, we showed that overproduction of a
truncated version of TssG lacking foot 1 and foot 2 interferes with baseplate assembly and leads to T6SS function impairment (50).

In this study, we deciphered the TssK-TssG interface of EAEC and assessed its relevance as a target to hinder T6SS-associated virulence. After validating the role of TssG feet in T6SS baseplate assembly and functioning, we investigated the sequence determinants of the TssK-TssG interacting region. Based on the available structures of the TssK-TssG interface and multiple-sequence alignments (MSA) of homologous proteins, we identified interacting motifs that we validated through site-directed mutagenesis coupled to functional assays. Guided by this structural and biochemical knowledge, we rationally designed a cyclic peptide able to interfere with T6SS baseplate biogenesis in EAEC. Furthermore, we carried out a comparative analysis of the predicted protein-protein interaction interface of TssK-TssG on 17 pathogens harboring at least one T6SS gene cluster. This analysis revealed a high conservation level of the TssK motifs targeted by our peptide as well as a preserved TssK-TssG interacting region that we further confirmed through cross-species protein pulldown experiments. Altogether, this study highlighted a conserved protein-protein interface that plays a key role in T6SS baseplate biogenesis and provided, to our knowledge, the first rationally designed T6SS inhibitor that could serve as a model to reinforce our arsenal against clinical pathogens and pave the way for new antivirulence inhibitors.

RESULTS

Validation of the TssK-TssG interface as a drug target. We analyzed the effect of TssG foot mutation on T6SS biogenesis and functioning in the enteropathogenic organism EAEC. Using a chromosomally encoded and functional fusion protein of TssK and the superfolder green fluorescent protein (sfGFP), we monitored T6SS baseplate biogenesis by fluorescence microscopy in EAEC (45, 50). In this context, the deletion of both TssG feet abrogates the assembly of the baseplate (Fig. 1D and Fig. S1D). We proceeded with an in-frame chromosomal deletion of TssG foot 1 or foot 2, which resulted in the total loss of T6SS activity, highlighting the crucial role played by these two TssG structural elements (Fig. 1A and B). In order to diagnose at which stage the T6SS biogenesis was affected by TssG foot deletion, we biochemically analyzed the behavior of the TssKFGE unit using native gel experiments and observed that deletion of either foot 1 or foot 2 destabilizes the wedge complex (Fig. 1C and Fig. S1A and C). The TssG variants have no impact on the production and stability of TssF and TssK, which ruled out any indirect effect of TssG mutation on the stability of the wedge proteins (Fig. 1C and Fig. S1A and C). Based on the structure of the wedge complex, which displays a very localized interaction between TssG and TssK mediated by the feet, one could hypothesize that the deletion of the feet would uniquely impair the interaction between these two proteins. However, we found that the amount of TssG copurified using TssF-based pulldown was significantly lower (4.6% and 1.1% compared to 74.8%) when either of the 2 feet was deleted (Fig. S1B). We thus concluded that deletion of TssG foot 1 and foot 2 impairs the interaction of TssG with TssK and reduces the association with TssF. This was further confirmed using native polyacrylamide gel electrophoresis (PAGE) analysis, which showed that no complex intermediates were observable when foot 1 and/or 2 was deleted. Previous work suggested the importance of foot 1 and foot 2 in mediating the interaction with TssK through overproduction of TssG-derived interfering domains (50) or heterologous bacterial two-hybrid (BACTH) experiments (51). All this accumulated evidence indicates that any manipulations of TssG feet (i.e., deletion, mutations, and binding interference) has an important effect on the T6SS assembly and function and that the foot region of TssG represents a crucial target for therapeutic intervention that aims to block the T6SS assembly and functioning.

Based on the role of TssG feet in T6SS biogenesis, we hypothesized that targeting the interaction between TssK and TssG through a suitably designed molecule would essentially reproduce the results of TssG manipulations and hence would be detrimental to T6SS function. For this purpose, we decided to perform an in-depth analysis of
TssG foot domains are essential for T6SS functioning. (A) Architecture of the T6SS nanomachine. (Left) The different subcomplexes of the T6SS are presented: membrane complex (MC), baseplate complex (BP), and tail-tube complex (TTC). OM, outer membrane; PG, peptidoglycan; IM, inner membrane. (Center) Assembly of the BP, highlighting the three main components: TssK (red), TssG (yellow), and TssF (blue). The encircled inset shows the TssK-TssG interface. (Right) Structure of the fundamental unit, the wedge complex.

(B) Antibacterial assay. The number of recovered *Escherichia coli* recipient cells (in log_{10} CFU), indicating their survival after 4 h of incubation against the indicated attacker cells. The assays were performed at least three independent times, with technical triplicates, and representative technical triplicate measurements (circles) are shown, with the corresponding average value calculated on all assays (white bar). (B) Antibacterial assay. The number of recovered *Escherichia coli* recipient cells (in log_{10} CFU), indicating their survival after 4 h of incubation against the indicated attacker cells. The assays were performed at least three independent times, with technical triplicates, and representative technical triplicate measurements (circles) are shown, with the corresponding average value calculated on all assays (white bar). (C) Native 4 to 16% gel analyzed by immunoblotting using anti-His antibodies on BL21(DE3) lysate expressing TssK-6×His, TssK-6×His plus TssFGE, TssK-6×His plus FG^{Foot1}E (tssG containing a deletion of the foot 1 domain), TssK-6×His plus FG^{Foot2}E (tssG containing a deletion of the foot 2 domain). The TssK-6×His plus TssFGE produced from BL21(DE3) cells shows the positions of two high-molecular-weight complexes (HMWCs) (*, TssK trimer; **, TssKFGE wedge complex). Formation of the HMWCs was monitored in a variant of TssG with either the foot 1 or foot 2 domain deleted. In each indicated construct, the TssK production level was analyzed by SDS-PAGE followed by immunoblotting and used as loading control (bottom). The native gel experiment was independently performed three times, and results of a representative experiment are (Continued on next page)
the TssK-TssG interface and identify critical motifs that can guide toward the rational design of a biomimetic peptide (52). To better decipher the EAEC TssK-TssG binding region and to highlight molecular properties required for its interference, we conducted a conservation study using MSAs of EAEC close homologs. TssK is a trimeric protein composed of three domains: an N-terminal $\beta$ sandwich shoulder domain (NTD), a 4-helix bundle middle domain, and a C-terminal $\alpha/\beta$ head domain (CTD) (46). Mapping the amino acid conservation onto the TssK and TssG structures revealed a clustering of highly conserved motifs located on the TssK NTD (Fig. 2A and B). Remarkably, the TssK NTD has a conservation level that is higher than that of the full-length TssK and TssG proteins, as well as TssG foot 1 and TssG foot 2, and even exceeds the conservation level of TssB protein, which is one of the most conserved proteins in T6SS (13) (Fig. 2C; also, see Materials and Methods). The high conservation level of the TssK NTD, which directly interacts with the TssG feet, strengthens the hypothesis that this region might serve as a target for therapeutical purposes.

TssG foot 1 has a conservation level lower than that obtained for the TssK NTD, although similar to that of the full-length TssK, while foot 2 is very variable. Nevertheless, both feet possess a conserved repeated pattern consisting of a hydrophobic amino acid (e.g., leucine or methionine) followed by a small or a basic residue (e.g., glycine, serine, arginine, or lysine) alternating with a variable region, i.e., LGXXXX'1'LGXXXX'2'LG (referred to here as the LG repeat). The first variable region (XXXX'1') is heterogeneous in length and composition, while the second (XXXX'2') has a conserved length of about four residues, and it is less variable (Fig. 2D). Substitution of the three hydrophobic residues in the TssG LG repeat motif by an arginine in EAEC has been shown to impair TssK binding in vitro (51). This observation indicates that the interaction between TssK and TssG is stabilized by hydrophobic interplays between the conserved LG repeats of both TssG feet and the TssK NTD. In agreement with these observations, the TssK NTD harbors three highly conserved amino acids (W8, L14, and F19) forming hydrophobic cavities hosting the three LG repeat motifs in each TssG loop (Fig. 2D and E). To analyze the role of these three interfacial motifs, we performed alanine-scanning mutagenesis and created TssK variants (i.e., W8A, L14A, and F19A). In EAEC, a TssK transcomplementation experiment with a plasmid expressing each TssK mutant did not restore T6SS-dependent killing of the prey (Fig. 3A). Biochemically, these three mutations had no impact on the stability or the oligomeric state of TssK (Fig. S1E) but abolished the direct interaction with TssG, as monitored by affinity copurification experiments where streptavidin (Strep)-tagged TssK variants were used to pull down TssG (Fig. 3B). We thus concluded that the TssG-TssK hydrophobic interaction is an absolute determinant for the entire wedge complex stability and is also highly specific, since a mutation of each of the three side chains belonging to the hydrophobic pocket of TssK leads to the complete disruption of the assembly.

**Design of a biomimetic cyclic peptide.** Inspired by the specific TssK-TssG hydrophobic interaction deciphered above and its influence on the T6SS function, we aimed to design a peptide inhibitor mimicking the arrangement of the TssG foot domains bound to TssK. Structural analysis of the TssKFGE unit revealed that the TssG feet region harbors an intrinsic triangular fold that fits the C3-symmetric shape of the TssK NTD (Fig. 4). Following this observation, we hypothesized that a peptide with a cyclic structure and LG repeats organized as in TssG feet might be sufficient to bind to the TssK NTD and hence compete with TssG association (Fig. 4). We decided to design such a peptide using the TssG foot 1 as a template for two main reasons: (i) foot 1 has a higher conservation level than foot 2 (Fig. 2C) and (ii) foot 1 is less structurally complex than foot 2. In fact, foot 2 has a partially globular fold and more intramolecular
FIG 2  EAEC TssK-TssG interface conservation. (A and B) TssK and TssG residue conservation obtained with aligned EAEC variants mapped on the EAEC wedge complex structure (PDB 6N38). A dense group of highly conserved residues can be observed on the top of TssK trimers. This is particularly visible in panel B, in the top view of the trimer. TssG exhibits another group of conserved residues on its antenna (A), which is the domain involved in the interaction with the two TssF copies of the wedge complex. (C) Conservation level for TssK, TssK NTD, TssG, TssG foot 1, TssG foot 2, and TssB on EAEC. The numbers at the top of the bars indicate the amount of aligned EAEC homolog sequences used to compute the conservation level. (D) Sequence logos (Continued on next page)
Based on the residue conservation analysis (Fig. 4B), we selected the sequence between the second and the third LG repeats of TssG foot 1, SRPVMG (positions 238 to 243), as a repeat unit. The SRPVMG unit was repeated twice to fill the three hydrophobic cavities located within the TssK NTD (Fig. 4C), resulting in the sequence SRPVMG-SRPVMG-SRPVMG. We applied the head-to-tail cyclization to create a triangular shape that could best fit the TssK trimeric fold (Fig. 4C). In parallel and to further substantiate the inhibitory potential of the biomimetic cyclic peptide (BCP), we rationally designed two additional cyclic peptides for control experiments (Fig. S8A): (i) a randomized version (SVRMPG-SPMRVG-SMVRPG) (see Materials and Methods) aiming to evaluate the importance of the amino acid order in the binding to TssK NTD and (ii) a mutated version generated by mutating each methionine to alanine (SRPVAG-SRPVAG-SRPVAG) to explore the importance of the conserved methionine residue in binding to the TssK hydrophobic pocket.

**Molecular properties of the BCP and specificity of its binding to TssK.** In order to characterize the biophysical and binding properties of the peptide, a series of nuclear magnetic resonance (NMR) experiments were performed. These experiments included one-dimensional (1D) $^1$H-$^1$H total correlation spectroscopy ($^1$H-$^1$H-TOCSY), $^1$H nuclear Overhauser effect spectroscopy ($^1$H-NOESY), and $^1$H-$^15$N heteronuclear single quantum coherence (HSQC) spectroscopy using its $^{15}$N natural abundance. $^1$H-$^15$N HSQC spectroscopy is of particular interest in characterizing the peptide, as each H/N correlation peak is associated with the NH group of an amino acid in a particular chemical and magnetic environment. Here, we observed only 5 correlations for the 5 observable amino acids out of 6 (Fig. S7A), the last being a proline. This confirmed that the amino acids of the three repeated (circular) sequences are equivalent (i.e., 3-fold axis symmetry). In addition to

**FIG 2** Legend (Continued)

Illustrating the residue conservation of the TssK-TssG interacting regions for EAEC close homologs. The height of each letter represents the information content of the corresponding amino acid at that position in bits. Black boxes highlight the three hydrophobic residues forming the TssK NTD hydrophobic cavity on top of TssK trimers and the TssG foot LG repeats. (E) TssK-TssG binding sites. The triangular TssG loops, foot 1 and foot 2, bind TssK trimers on its N-terminal region. These contacts are mediated by hydrophobic interactions between the TssK NTD hydrophobic cavity and the conserved LG repeats of both feet.
HSQC spectroscopy, the $^{1}\text{H}-\text{NOESY}$ experiment did not exhibit any indications of a particular 3D folding (Fig. S7B). After addition of a 1:10 final concentration of TssK (1 part TssK to 10 parts BCP), the $^{1}\text{H}-^{15}\text{N}$ HSQC correlations (same conditions as the free-peptide experiment) totally disappear, while 90% of the peptide is supposed to be in the unbound form, in solution. The $^{1}\text{H}-^{15}\text{N}$ HSQC spectrum could not be recovered by a longer data accumulation. This deleterious effect was also confirmed on the 1D $^{1}\text{H}$ proton NMR spectrum (Fig. S7C), exhibiting a strong broadening of the peptides peaks that clearly indicates information transfer between the peptide and the protein either by chemical exchange relaxation (millisecond range for $k_{\text{on}}/k_{\text{off}}$) ($k_{\text{on}}$ [association rate constant] is the rate at which an interaction happens per second in a unimolar mixture; $k_{\text{off}}$ [dissociation rate constant] is the fraction of complex which dissociates per second) or a peptide/protein size cross-relaxation effect. Overall, these experiments confirmed the absence of an internal structure of the free peptide and confirmed our in silico prediction.

To evaluate the direct interaction between the BCP and its target TssK, we performed isothermal titration calorimetry (ITC) experiments. The wild-type BCP and TssK interact with an observable affinity ($K_{D}$) of 60 μM (Fig. 5A). The ITC binding signature of TssK-BCP suggests that it is driven by a favorable enthalpy, presumably based on hydrogen bonds and van der Waals interactions. The positive entropy suggests conformational changes which could be responsible for the low affinity observed. To confirm the specificity of this interaction, we performed a series of controls consisting of (i) mixing the BCP with the TssK-L14A variant, which we previously demonstrated has lost its capacity to interact with TssG, and (ii) incubating TssK with each of the control peptides. None of these conditions led to a detectable ITC binding signal (Fig. 5A and Fig. S8B), demonstrating the direct liaison of BCP to the TssK target, precisely at the TssG foot binding site.

**Inhibition of the EAEC TssK-TssG interface by the BCP.** We conducted in vitro and in vivo experiments to assess the potential T6SS-blocking activity of our BCP. First, the assembly of the TssKFGE wedge complex was reconstituted in vitro on the surface of a bilayer interferometry biosensor (Fig. 5). Briefly, biotinylated TssK trimers were
FIG 5 Biomimetic peptide interfere with TssK-TssG interface. (A) Isothermal calorimetry was used to demonstrate the specific interaction between TssK and wild-type BCP. (Left) Interaction between 25 μM TssK-WT, TssK-L14A and 1 mM wild type BCP. (Top) Heat exchange upon TssK ligand titration with BCP for TssK WT as ligand and BCP as analyte (blue curve), and for TssK-L14A as ligand and BCP as analyte (red curve). (Bottom) Integrated data with binding isotherms (solid line) fitted to a single-site binding model.

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immobilized onto a biosensor. Normalized bacterial cell lysate without TssFGE plasmid was added to the TssK-bound biosensor to block free Strep sites. To demonstrate the specific binding of TssFGE with TssK, normalized bacterial cell lysate expressing the TssFGE protein partners was added to the bound TssK to form the TssKFGE wedge complex (Fig. 5B and Fig. S2). TssK was shown to interact specifically with its natural partners, with a maximum binding average response from three independent assays of 0.189 ± 0.036 nm at 150 s association. When TssK was preincubated with the different BCP constructs, i.e., the mutated BCP, the randomized BCP, and the wild-type BCP, the maximum binding average responses of three independent assays were 0.169 ± 0.016 nm, 0.160 ± 0.028 nm, and 0.121 ± 0.0102 nm, respectively (Fig. 5B, inset), demonstrating that the wild-type BCP significantly interferes with the TssKFGE formation and the control peptides have a negligible activity.

Modulation of bacterial competition by the BCP. Based on the BCP ability to bind to TssK and to limit the in vitro binding of the TssFGE wedge components onto TssK, we evaluated its T6SS interference potential on bacteria. We set up a competition experiment between EAEC (predator, T6SS+) and E. coli (prey, T6SS-) strains (see Materials and Methods) in the presence or absence of the BCP. Incubation of cells with 1 mM BCP is accompanied by a drastic decrease of prey cells number, similar to what is observed in the absence of BCP. We concluded that the BCP does not affect the killing ability of EAEC when added directly in the competition mixture (Fig. S3C), which can be explained by its physicochemical properties, which prevent its diffusion across biological membranes. To test this hypothesis, we developed a protocol favoring BCP entry into bacterial cells without any electroporation or chemical steps likely to impair T6SS activity (see Materials and Methods). This procedure, relying on an osmotic change, has been applied to the TssK-sfGFP strain, which allowed us to enumerate the number of fluorescent foci and hence assembled baseplates (45, 50) in presence or absence of the BCP. Osmoporation of the TssK-sfGFP strain with 1 mM BCP increased significantly (~30%) the proportion of bacteria with no observable foci compared to peptide-free treated cells (Fig. 5C). In contrast, incubation with a 1 mM concentration of the control peptides did not impact the number of fluorescent foci, which was comparable to that in the free-peptide condition. Importantly, the peptide was not toxic for bacteria up to 1 mM (Fig. S3A), and the osmotic stress did not affect fluorescent-focus assembly (Fig. S3B).

Evolution of the TssK-TssG interaction interface. Our functional analysis and peptide design revealed the determinants of the TssK-TssG interaction, which relies on hydrophobic interplays conserved within EAEC. To assess whether BCP can target other T6SS-harboring bacteria, we systematically compared the sequences of full-length TssK, TssG, TssK-TssG interacting domains (i.e., the TssK NTD and TssG foot 1 and foot 2), and TssB, which was used as a reference. For this purpose, we (i) collected the sequences from bacteria encoding a T6SS directly linked to pathogenicity or with host-associated

FIG 5 Legend (Continued)
activity (Table S1B and C), (ii) clustered the sequences based on the sequence similarity and also their variants, and (iii) computed their conservation levels.

Unexpectedly, TssK and TssG have divergent sequence clustering patterns, while being structural neighbors (Fig. 6). Indeed, clustering of TssB and TssG sequences (Fig. 6D) followed the standard T6SS subtype classification (53) (Fig. S6B), whereas TssK (Fig. 6B) did not. i2 is the only T6SS subtype that is consistently clustered in both TssK and TssG as well as their interacting domains. Importantly, our EAEC T6SS model (referred to here as EAEC3 to differentiate the three EAEC T6SS gene clusters under study) belongs to this specific T6SS subtype. This pattern indicates the existence of a conserved and specific TssK-TssG interface signature and suggests that the interfering activity spectrum of the BCP applies not only to EAEC3 but also to other subtype i2 T6SSs, such as those of Klebsiella pneumoniae (K.p 1 and K.p 2), Yersinia pseudotuberculosis (Y.ps 3 and Y.ps 5), Yersinia pestis (Y.p 3), and Burkholderia cenocepacia (B.c 1) (Table S1A; Fig. S6C).

The overall sequence conservation level was computed for the full-length TssK, TssG, and TssB proteins as well as the interacting domains TssK NTD, TssG foot 1, and TssG foot 2 of each pathogen (Fig. S4 and Fig. 2C). In general terms, the TssK NTD has a higher conservation level than full-length proteins and TssG feet. TssG foot 2 is less conserved than foot 1. Multiple-sequence alignments of TssK NTD for all pathogens and their variants (2,482 sequences) show that the TssK NTD hydrophobic amino acids W8, L14, and F19 are highly conserved, as they are for EAEC3 (Fig. S6A). Likewise, considering the MSA for all pathogens (3,712 sequences), the LG repeat motif is consistently conserved in both feet (Fig. S6A). All these observations indicate that all the TssK-TssG interfaces of pathogenic T6SS follow the same physicochemical rules as in EAEC3, which consists of TssK hydrophobic cavities accommodating the TssG hydrophobic residues. Based on this analysis, we therefore expect that the BCP can bind to a larger repertoire of TssK targets and hence have a greater spectrum of activity.

Proof-of-concept evaluation of the broad-spectrum activity of the anti-T6SS BCP. Based on the high and low conservation levels, respectively, of the TssK NTD and TssG feet, we hypothesized that the TssK-TssG hydrophobic interplay in EAEC3 is a general binding mechanism. To challenge this hypothesis, we investigated the ability of the EAEC TssK target to accommodate several TssG protein partners encoded in strains clustered outside the T6SS i2 subtype. We selected three TssG homologs belonging to different subgroups (Fig. 6A), including the pathogenic species Acinetobacter baumannii (A.b subgroup i4b) and Pseudomonas aeruginosa H2 (P.a2 subgroup i1) as well as a member of the human microbiota, Bacteroides fragilis (B.f subgroup iii). We performed protein copurification experiments with TssK-TssG pairs and evaluated cross-species interactions. No interaction was observed between TssKEAEC and TssG, while TssKEAEC interacts with both TssG and TssG (Fig. 6A and Fig. S5B), and reciprocally, TssGEAEC interacts with both TssK and TssK (Fig. S5D). To validate these interactions, we performed the same experiments using the EAEC TssK variant mutated at the W8 residue. Substitution of the TssK hydrophobic residue tryptophan by an alanine induces the loss of interaction with TssG and TssG, suggesting that these proteins interact through a mechanism similar to that in EAEC (Fig. SSC). Based on the interaction between TssKEAEC and TssG homologs from non-co-occurring species, we proposed that the TssK-TssG binding properties expands beyond the i2 subtype. Overall, the high conservation of hydrophobic characteristics found in both protein sides combined with biochemical experiments revealed that few determinants are required to drive the interaction specificity between TssK and TssG. We thus propose that BCP provides a framework for the design of broad-spectrum T6SS inhibitors targeting multiple bacterial pathogens.

DISCUSSION

The T6SS is a phage-related contractile machinery responsible for toxin secretion into both eukaryotic and prokaryotic cells. This secretion apparatus is found in more than 25% of Gram-negative bacteria, including the ESKAPE pathogens K. pneumoniae,
**FIG 6** Phylogeny and conservation of the TssK-TssG interface among pathogenic bacteria. (A) The TssK-TssG interface shares common features among pathogenic bacteria. Summary of the results of copurification experiments presented in detail in Fig. S5. EAEC3 or P.a2

| TssK  | TssG  | Interact | T6SS group   |
|-------|-------|----------|--------------|
| A.b   | +     | i2(K); i4b(G) |
| EAEC3 | B.f   | +        | i2(K); iii(G) |
| P.a2  | -     | i2(K); i11(G) |
| P.a2  | +     | i11(K); i11(G) |

- **A**: Summary of the results of copurification experiments presented in detail in Fig. S5.
- **B**: Heatmap of TssK interaction with different subtypes.
- **C**: Heatmap of TssK-NTD interaction with different subtypes.
- **D**: Heatmap of TssG interaction with different subtypes.
- **E**: Heatmap of TssG foot1 interaction with different subtypes.
- **F**: Heatmap of TssG foot2 interaction with different subtypes.

(Continued on next page)
A. baumannii, P. aeruginosa, and Enterobacter species, where it plays a significant role in either colonizing and damaging the host or manipulating and evading the immune system (12, 23–27, 54–57). Although the T6SS is structurally well characterized and meets all the criteria as a promising virulence factor candidate, a very limited number of studies have attempted to develop anti-T6SS molecules or peptides. Certain studies have used phenotypic screening (58), and others have employed a targeted approach, such as blocking the interaction between the two sheath proteins (TssB and TssC) or the dissociation of a toxin immunity complex (28, 59). Another study employed a nanobody intracellularly expressed to block the assembly of the membrane complex (MC) (60), which is limited in terms of a possible therapeutic application, since nanobodies cannot cross the bacterial outer membrane. All these studies lack a molecular and structural description of the hit binding site, which precluded hits-to-lead optimization and a broad applicability to a large number of human pathogens, including the ones from the ESKAPE list. In our study, we decided to combine structural, molecular, and bioinformatics analysis with the aim of targeting a conserved molecular interface in the T6SS baseplate that is crucial for the functioning of the whole nanomachine.

The TssK-TssG interface plays a pivotal and primordial role in the early stages of the T6SS baseplate assembly, specifically, the two small unfolded TssG domains interacting with the N-terminal region of TssK (50, 51). Using BACTH experiments, previous studies demonstrated the total loss of interaction between TssK and TssG variants mutated in either the foot 1 or foot 2 domain, suggesting that TssG requires two functional feet to bind to TssK (51). In our study, we demonstrated that deletion of either TssG foot 1 or foot 2 abrogates the formation of the TssKFGE wedge complex and is associated with a loss of T6SS activity (Fig. 1), which revealed early-stage T6SS baseplate assembly as a promising target. Overall, we provide strong evidence that TssG foot domains are essential for T6SS activity and bind to key hydrophobic residues representing a docking surface on the TssK NTD. Interfering with this recognition step can be envisaged to inhibit T6SS functioning through a peptide mimicking one of the two TssG feet with hydrophobic characteristics required for TssK cavity binding.

The role of unstructured loops in macromolecular complex stability was reported previously. For instance, the T4 phage TssG homolog gp7 protein connects the tail fibers (gp9, gp10, and gp11) to the baseplate, allowing the communication of important structural transitions. During contraction, a loop formed by residues 841 to 862 of gp7 performs a “jump-rope”-type motion transferring the signal received from the fibers to the gp2S-(gp6),gp7-like wedge module responsible for the conformational switch (49). In our T6SS model, TssG harbors two unfolded domains. The TssG foot 1 domain is a loop that is structurally independent from the rest of the protein. In contrast, TssG foot 2 is a partially folded region characterized by less structural autonomy. Multiple-sequence alignments of TssG in EAEC and other T6SS-harboring pathogens indicated a higher conservation level of the TssG foot 1 than TssG foot 2. Besides these differences, both TssG feet impact T6SS activity to the same extent, which suggests that they are good candidates for inhibition strategies.

While a number of studies have targeted $\alpha$-helix and $\beta$-strand epitopes to block specific protein-protein interaction interfaces (PP2I) (52, 61), loop-mimicking PP2I inhibitors represent an enormous challenge, and very few bioinformatics tools have been designed to help in this quest (62). The inhibition of a protein-protein interaction is challenging. It requires either a robust and costly methodology to screen thousands of compounds from a library or the rational design of interfering molecules guided by

FIG 6 Legend (Continued)

TssK was tested for the interaction with A.b, B.f, or P.a2 TssG. A positive interaction is reported with a plus sign, whereas a negative interaction is reported with a minus sign. T6SS subtypes are indicated in the right column for each pair. (C to F) Hierarchical clustering of TssK, TssG, and corresponding interacting domains using the closeness metric (see Materials and Methods). Diagonal values represent the conservation level of a protein or a domain for a given pathogen, while out-of-diagonal values represent the similarity of a protein or a domain calculated between two different pathogens. Gene cluster nomenclature is fully reported in Table S1A. For example, EAEC1 represents the first T6SS gene cluster found in the EAEC genome, while EAEC3 is the third. Clusters of pathogens that follow the standard subtype classification for T6SS are indicated by the color of the matching square.
the molecular description of the target (63). Usually, targeting the interaction between TssK and TssG through a suitably designed molecule would have required the systematic screening of ligands able to bind to the target TssK, hence competing with TssG foot docking (64–66). Instead, we decided to base our rational design on the structure of the TssKFG wedge complex that we have obtained recently (50). We decided to design a peptide interfering with the T6SS activity by targeting the crucial TssK-TssG interaction. The TssK NTD is structured, has cavities, and is well conserved, revealing a hot spot-interacting region. Remarkably, the TssK NTD can interact with the two different TssG LG repeat domains, which vary in length and composition. This observation, reflected by the lower conservation level of the TssG foot domains across EAEC variants, suggests that TssK can accommodate different partners as long as they harbor a hydrophobic motif with a triangular fold. Based on these criteria, we designed a circular peptide inhibitor inspired by the EAEC TssG foot 1 domain. The biomimetic cyclic peptide binds to TssK and interferes with wedge protein TssFGE recruitment, which consequently impairs baseplate biogenesis in permeabilized EAEC cells (Fig. 5C).

We evaluated the potential spectrum of our peptide through TssK and TssG sequence analysis. The analysis of T6SS bacterial sequences revealed that our model EAEC with other i2 subtypes encoded in Y. pestis, Y. pseudotuberculosis, K. pneumoniae and B. cenocepacia and provided a first hypothetical picture of the spectrum of BCP activity. However, LG repeats and the conserved hydrophobic cavity can be found beyond the i2 subtype, and they are a general property of the TssG-TssK interaction, including that in A. baumannii and B. fragilis, which has been confirmed by biochemical cross-species interaction experiments. The human microbiota member B. fragilis encodes T6SS clusters with a significant role in gut homeostasis and colonization resistance (67–69). The potential BCP interfering activity on B. fragilis T6SS needs to be carefully considered to limit any collateral damage ensuing from antivirulence treatment. However, enterotoxigenic B. fragilis (ETBF) is involved in colorectal cancer (CRC) development (70), and more broadly, B. fragilis is found in a number of antibiotic-resistant bacterial infections (71). Consequently, targeting the B. fragilis T6SS might represent an alternative way to fight this pathogen. The bioinformatics analysis predicting the presence of LG repeats and hydrophobic cavities in the P. aeruginosa H2 TssK-TssG interface contrasted with our biochemical experiments. This unexpected observation indicates that molecular properties different from the LG repeats could additionally drive TssK-TssG specificity and highlights the need to experimentally explore and validate interface conservation across bacteria.

Several improvements can be considered for the BCP, for instance, its limited molecular diffusion in cell membranes. The BCP has a molecular weight exceeding 600 Da, which is the porin-mediated diffusion limit (72). To overcome this obstacle, we could consider fusing our BCP with either siderophores (73) or cell-penetrating peptides (CPPs). Examples of CPP-aided delivery include an inhibitor of bacterial FtsZ altering the growth of S. aureus (74). Implementation of permeation rules increasing drug spectrum activity toward Gram-negative bacteria can be an alternative to rationally modifying BCP diffusion properties (75). In addition to its low diffusion property, the inhibitory effect of the peptide has been tested with a relatively high working concentration (1 mM). This, coupled with our observation that the BCP effect is maximum within the first hour after peptide entry, suggests that both its low affinity and stability are important to consider in the framework of future improvements. In spite of these limitations, we think that the BCP can be used as a scaffold to design a class of clinically relevant inhibitors with increased diffusion properties, better affinity for TssK, and improved activity against T6SS function.

In general, this work provides an in-depth understanding of the EAEC TssK-TssG interface, which has been proven to be a promising target for T6SS inhibition. We highlighted molecular determinants required for the TssK-TssG interference that guided the rational design of a biomimetic peptide for which the activity against EAEC validated the feasibility of our approach. Bioinformatics analysis of the TssK and TssG proteins over 17 pathogens and corresponding sequence variants (2,482 for TssK and 3,712 for TssG) revealed the high conservation
level of the TssK target as well as the preserved triangular hydrophobic core binding motifs, suggesting a broad inhibitory potential. We propose that our peptide could serve as a scaffold to design large-spectrum antivirulence molecules and hence pave the way for the development of anti-T6SS inhibitors.

**MATERIALS AND METHODS**

Data gathering. TssK, TssG, and TssB EAEC sequences were obtained from the genome of EAEC strain S5989 (Table S1C). Importantly, a single species might have more than one T6SS gene cluster. Using KEGG (76) together with SecReT6 (77) annotations, we selected TssK, TssG, and TssB sequences from the 3 different T6SS gene clusters on the EAEC genome (53, 78, 79). The first gene cluster (EAEC1) belongs to subtype i1, the second (EAEC2) to subtype i4b, and the third (EAEC3) to subtype i2. In the present study, we used EAEC3 as our T6SS model thanks to the available structural data of the TssK-TssG interface (wedge complex cryo-electron microscopy [EM] atomic structures; PDB IDs 6GIY and 6N3B). TssK, TssG, and TssB sequences for other pathogenic T6SSs examined were also classified into their corresponding T6SS subtypes following the information retrieved from KEGG and SecReT6 databases (Table S1B and C). Uniref90 (80) was downloaded and converted into a BLAST database using the makeblastdb application (81). We also downloaded all bacterial genomes contained in the SGB collection metagenomic database (82). Using ORFM (83), we identified all possible genes and translated their open reading frames from which putative protein sequences were converted into a BLAST database, using makeblastdb.

Homologous search. The search of homologous sequences for TssK, TssG, and TssB was conducted using BLASTp (81) on the Uniref90 (80) and SGB databases (82). We queried the TssK, TssG, and TssB sequences from T6SS EAEC1, EAEC2, and EAEC3 gene clusters, as well as all the gene clusters from a selected list of pathogenic T6SSs (Table S1C). All these sequences, belonging to different gene clusters and pathogens, are referred as to the query sequences here. Homologous sequences whose length was less than 50% or more than 150% of the query sequence length were discarded. All the sequences resulting from the multiple BLASTp searches were merged and clustered by 0.9 sequence identity using cd-hit (84).

Multiple sequence alignment. TssK, TssG, and TssB multiple-sequence alignments (MSAs) were generated by multiple rounds of (i) homologous sequence alignment performed by MUSCLE (85) and (ii) removal of outliers by EvalMSA (86). After the last iteration, we manually curated the MSAs using JalView (87).

Clusters of variants. Each sequence from the MSA of a given protein was assigned to the most similar query sequence, using blosum62 as a similarity score (88). The assignment allowed us to cluster all the sequences, where each cluster is a set of closely homologous variants of the corresponding query. Clusters of variants were not overlapping, i.e., they did not have any sequence in common. For each cluster, we removed the outlier sequences of the cluster by removing the lowest-scoring 25% of sequences. This metric computes the similarity level between clusters of sequences when aligned sequences. This metric computes the similarity level between clusters of sequences when scored as 

\[ c(X,Y) = \frac{\text{idenscore}(X,Y) - \text{randscore}(X,Y)}{\text{idenscore}(X,Y) - \text{score}(X,Y)} \]

where the score between X and Y is

\[ \text{score}(X,Y) = \frac{\sum_{i=1}^{n} \sum_{j=1}^{m} \text{blosum62}(s_i, s_j)}{n \times m} \]

and \( \text{blosum62} \) is the blosum62 score between the aligned sequences \( s_i \) and \( s_j \). \( \text{randscore} \) is the minimum score that one can get between two clusters of sequences:

\[ \text{randscore}(X,Y) = \frac{\sum_{i=1}^{n} \sum_{j=1}^{m} \text{blosum62}(r(s_i), r(s_j))}{n \times m} \]

where the function \( r(s) \) randomizes the sequence \( s \), keeping the same length and amino acid composition. \( \text{idenscore} \) is the score obtained using identical sequences; thus, it is the highest score one can get:

\[ \text{idenscore}(X,Y) = \frac{\sum_{i=1}^{n} \text{blosum62}(s_i, s_i) + \sum_{i=1}^{m} \text{blosum62}(s_i, s_j)}{n + m} \]

Note that the closeness \( c(X,Y) \) is a normalized quantity, i.e., it ranges between 0 (as far as random clusters) and 1 (as close as identical clusters). Furthermore, the metric is independent of the length of the aligned sequences. This metric computes the similarity level between clusters of sequences when
X is not equal to \( x \), and \( Y \) is not equal to \( y \). The conservation level of a cluster is set to 0.

Data analysis, visualization, and storage. Residue conservation was computed with JalView (87) and mapped onto the wedge complex structure (PDB ID 6N38) using Chimera (89) (Fig. 2A and B). Residue frequencies (Fig. 2C and D and Fig. S6E) were mapped on the alignment using the WebLogo (90) server. MSA figures (Fig. S8) were generated with JalView. Hierarchical clustering figures (Fig. 6 and Fig. S7) were generated with seaborn (open-source Python package for statistical data visualization) (91). All molecular visualizations were rendered by UCSF Chimera (89). All Python and shell scripts specifically developed for this work are available together with the initial genomic data (TssK, TssB, and TssG sequences) and the resulting MSA (TssK, TssK NTD, TssB, TssG, TssG foot 1, and TssG foot 2) at https://gitlab.pasteur.fr/ffilella/closeness-metric.

Peptide design. We started by selecting a six-residue sequence based on the EAEC3 foot 1 sequence (Fig. 2D). We selected the sequence residues of the second variable region (SRPV) together with the third hydrophobic repeat (MG): SRPVMG (positions 238 to 243). Then, we used this sequence to create a cyclic peptide, as follows. The structure of the chosen sequence was cloned twice, applying C3 symmetry around the trimeric TssK axis of symmetry. The three resulting peptide structures were linked through the backbone, introducing N-C bonding restraints (bond, angle, and dihedral restraints introduced by the PATCH command of CHARMM). We included an N-to-C-terminus bonding restraint, resulting in a head-to-tail cyclization of the peptide. Finally, the cyclic peptide was relaxed during a short molecular dynamics (MD) simulation to ensure its stability (Fig. 4).

Control peptide design. Two different control peptides were rationally designed: a mutated BCP and a randomized BCP. The mutated BCP was obtained by simply mutating the three Met residues in the 5th position of each BCP repeat to Ala. The randomized peptide was selected from a pool of 13,824 candidates. This pool of candidates was generated by independently permuting the four central residues of each BCP repeat (RPVM), while keeping the Gly and Ser in their original positions to conserve their propensity to form a turn (like the one observed in the foot 1 TssK structure). To design a negative-control peptide, we selected a sequence which we predicted to poorly bind to the trimeric TssK NTD. We ran short MD simulations of all candidates and ranked them according to their total energy after a final minimization. Finally, to avoid any functional redundancy with the original BCP peptide, we selected the peptide that ranked the worst and at the same time did not have any residue at the same position in the BCP sequence (except for the G5 residues). The MD simulations of the peptide bound to the TssK NTD were done with CHARMM (92), using the CHARMM22 force field. The initial backbone structure was built from the original TssG foot 1-TssK cryo-EM model, by symmetrizing on the TssG 238-239-243 backbone of the second TssK repeat. The modeled TssK NTD sequence was starting from Leu7 and truncated at Phe19. To increase the efficiency of the simulations, we used a distance-dependent dielectric that mimicked the presence of the solvent. To stabilize the structure of the TssK NTD, we applied an N-terminal acetylation and a C-terminal amidation and restrained the heavy atoms of their terminal residues (residues 7 and 8 and residues 17 to 19) with a harmonic potential centered on their initial positions. The simulations, which ran for 200 ps for each candidate, generated a total of 100 snapshots per sequence. The snapshots were subsequently minimized, and the median of the 100 final total energies was used to rank the peptides (Fig. S8).

Coevolution analysis. Residue coevolution identified between TssG and the TssK NTD was detected by combining RaptorX (93) complex contact prediction and Gremlin (94) monomer contact prediction. First, we extracted the concatenated MSA generated by RaptorX when TssK and TssG EAEC sequences were input. Second, we trimmed the TssK NTD and TssG foot 1 from the Raptor MSA and used the resulting concatenated MSA as input for a Gremlin monomer contact prediction search. Finally, we selected the TssK-TssG pairs with a probability above a given threshold and a distance between their side chains below 8 Å on the atomic structure of the wedge structure (PDB ID 6N38).

Peptide synthesis. The biomimetic cyclic peptide was synthesized by GenicBio, Ltd.

Plasmid construction. PCRs were performed using Q5 high-fidelity DNA polymerase (New England Biolabs). Restriction enzymes and T4 DNA polymerase were purchased from New England Biolabs and used according to the manufacturer’s instructions. Custom oligonucleotides were synthesized by Sigma-Aldrich and are listed in Table S1A. E. coli DH5α, Acinetobacter baumannii 17978, and P. aeruginosa PA01 chromosomal DNA were used as templates for PCRs. E. coli DH5α was used for cloning procedures. Plasmid construction. Plasmids were engineered by a two-fragments sequence- and ligase-independent cloning (SLIC) strategy (95, 96). Briefly, each DNA fragments were amplified by PCR using two pairs of oligonucleotides (FWD1/REV1 and FWD2/REV2). PCR products were digested (DpnII), cleaned (Macherey-Nagel PCR cleaning kit), and mixed together with the T4 DNA polymerase and its buffer (NEBuffer r2.1) at room temperature. The reaction was stopped after 2 min 45 s, and the mixture was put on ice. The annealing product was transformed into competent E. coli DH5α, and recombinant strains were selected on the appropriate antibiotic. Substitutions in pTRC-99A-TssKH and pRSF-TssKS plasmids were introduced by site-directed mutagenesis using complementary pairs of oligonucleotides. All constructions were verified by DNA sequencing.

Protein production and purification. Plasmids expressing the genes of interest were cotransformed into E. coli BL21(DE3), as described in "Plasmid construction" above. Cells were grown at 37°C in lysogeny broth (LB) to an \( A_{600} \) of \( \sim 0.7 \), and gene expression was induced by the addition of 1 mM IPTG (isopropyl-\( \beta \)-D-thiogalactopyranoside) for 16 h at 16°C. Cell pellets were suspended in 50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1 mM EDTA, 10 mM MgCl₂, supplemented with 10 \( \mu \)g/ml of DNase I, 100 \( \mu \)g/ml of lysozyme, and EDTA-free protease inhibitor (Roche) to an \( A_{600} \) of \( \sim 125 \). Cells were broken using an Emulsiflex-C5 instrument (Avestin) and clarified by centrifugation for 30 min at 20,000 \( \times \) g. The
supernatant was loaded onto a 5-ml StrepTrap HP (GE Healthcare) column on an Äkta Pure system (GE Healthcare) equilibrated in affinity buffer (50 mM Tris-HCl [pH 8.0], 150 mM NaCl). The column was then washed using the affinity buffer, and the proteins were eluted in the same buffer supplemented with 2.5 mM desthiobiotin (IBA Technology). The lysate, flowthrough, wash, and elution fractions were collected, suspended in Laemmli loading buffer supplemented with 1 mM 2-mercaptoethanol, and heated for 10 min at 96°C prior to analyses by SDS-PAGE and immunoblotting. For the study of the interaction between TssK and TssG variants, copurification was carried out by pulling down Strep-tagged TssK and evaluating the presence of copurified TssG by Western blotting.

**SDS-PAGE, protein transfer, immunostaining, and antibodies.** SDS-PAGE was performed on Bio-Rad Mini-Protean systems using standard protocols with homemade 12.5% polyacrylamide gels. For immunostaining, proteins were transferred onto 0.2-μm nitrocellulose membranes (Amersham Protran) with a Mini-Trans Blot cell (Bio-Rad). The membrane was then saturated in 5% milk and probed with primary antibodies. Mouse secondary antibody coupled to alkaline phosphatase was added and developed in alkaline buffer in the presence of 5-bromo-4-chloro-3-indolylphosphate and nitroblue tetrazolium. The antihemagglutinin (HA) (HA-7 clone; Sigma-Aldrich), anti-Flag (M2 clone; Sigma-Aldrich), anti-Strept (Sigma-Aldrich), anti-5His (Sigma-Aldrich) monoclonal antibodies, and mouse secondary antibodies (Millipore) were purchased as indicated.

**Fluorescence microscopy.** Fluorescence microscopy experiments were performed as described elsewhere (97, 98). Briefly, cells were grown overnight in LB and diluted to an \( A_{600} \) of ~0.04 in SIM (sulfur, indole, and motility) medium. Exponentially growing cells \( A_{600} = 0.8 \) to 1 were harvested, washed in phosphate-buffered saline (PBS) buffer, resuspended in PBS to an \( A_{600} \) of ~50, spotted on a 2% agarose pad, and covered with a coverslip. Fluorescence and phase-contrast micrographs were captured using an AxioImager M2 microscope (Zeiss) equipped with an AxioCam digital camera (Hamamatsu). Fluorescence images were acquired with a minimal exposure time to reduce bleaching and phototoxicity effects, typically 500 ms for TssK-sGFp. Noise and background were reduced using the Subtract Background (20 pixels, rolling ball) and Band plugins of ImageJ (National Institutes of Health). The sGFp foci were automatically detected using the MicrobeJ plugin ([https://www.microbej.com/](https://www.microbej.com/)). Box plots representing the number of detected foci for each strain were made using the Web tool BoxPlotR ([http://shiny.chemgrid.org/boxplotr/](http://shiny.chemgrid.org/boxplotr/)). Microscopy analyses were performed at least at three times, each in technical triplicate, and results of a representative experiment are shown.

**Strain construction.** Deletions of the TssG foot 1 and foot 2 domains were engineered at the native locus on the chromosome by allelic replacement using the pKO3 suicide vector (99) in the enteroaggregator E. coli 17-2 and E. coli 17-2 TssK-sGFp strains. Briefly, the E. coli strains were transformed with a pKO3 plasmid harboring the domain deletion that has been cloned (see below). Insertion of the plasmid into the chromosome was selected on chloramphenicol plates at 42°C. The recombinant plasmid in which a portion of TssG harboring the domain deletion has been cloned (see below).

**Targeting the T6SS Baseplate by a Biomimetic Peptide**
hydrated for 10 min in HEPES buffer before binding of the biotinylated TssK. Biotinylated TssK was loaded at 1.2 μM onto the SAX biosensors for 120 s and then the excess TssK was dissociated in HEPES buffer for 120 s. For the remaining kinetic assays, we used kinetic buffer (PBS [pH 7.3] plus 0.02% Tween 20, 0.1% bovine serum albumin [BSA], 0.05% sodium azide) (Forté Bio number 18-5032) with orbital shaking at 2,200 rpm. Kinetic buffer (KB) was used to dilute the lysates and the peptide to decrease nonspecific signals. The unbound Strep sites were blocked with 2 μM control lysate for 120 s and washed with KB for 120 s. The kinetic assays were performed using three steps with different times: baseline, 30 s; association, 120 s; and dissociation, 120 s. All the following assays were performed in triplicate. Four independent assays were performed to demonstrate the specific binding of TssK with its natural partners TssF-TssG-TssE and the effect of the wild type biomimetic peptide to block or hinder the specific binding of the analytes. In the first assay, we added 47 μM lysate expressing the analytes TssG, TssF, and TssE to the bound TssK. In the second assay, we added 1 mM wild-type (WT) BCP to the bound TssK before adding 47 μM TssF-TssG-TssE. In the third assay, to validate the specificity of the WT BCP, we added 1 mM mutant BCP that lacks the binding residues to the bound TssK before adding 47 μM TssF-TssG-TssE. Last, to redemonstrate the specificity of the BCP-WT, we added 1 mM randomized BCP that contains the binding residues but out of order to the bound TssK before adding 47 μM TssF-TssG-TssE.

**ITC**. Isothermal titration calorimetry (ITC) was performed to demonstrate the specific interaction of TssK with WT BCP and the lack of interaction with TssK L14A. The working buffer for both proteins and peptide was 50 mM HEPES (pH 7.0), 150 mM NaCl to avoid buffer mismatch. The experiments were performed at 25°C using the MicroCal PEAQ-ITC instrument (Malvern Panalyticals, Malvern, UK) in duplicate with 19 injections, first with an initial injection of 0.4 μl followed by 18 injections of 2 μl. Two assays were performed. In the first, 25 μM WT TssK ligand was in the cell and 1 mM WT BCP analyte was in the syringe. In the second, 25 μM TssK L14A ligand was in the cell and 1 mM WT BCP analyte was in the syringe. The reaction was performed with a constant stirring speed of 750 rpm; each injection lasted for 4 s with a 150 s space between injections. A constant heat control (offset) was removed from the raw data to account for heat dilution before integration. The data were fitted using the One Set of Sites model in the PEAQ-ITC analysis software.

**TssK focus in vivo inhibition**. Fluorescence microscopy experiments were performed as previously described (97, 98). An overnight culture in LB of EAEC strain with a chromosomal GFP fusion to TssK was diluted 1/200 in SIM medium and grown at 37°C until the culture reached an OD560 of ~0.8. Ten optical density units (ODU) was pelleted for 10 min at 3,500 × g, resuspended at 1 ODU/ml in the same SIM medium supplemented with 0.8 M NaCl, and equilibrated for 1 h at room temperature. Cells were pelleted for 10 min at 3,500 × g, resuspended in 100 μl of classical SIM without NaCl supplemented with 1 mM peptide of buffer, and incubated for 20 min at room temperature. Cells were diluted in classical SIM medium and 4 ODU/ml and incubated at 37°C. The nuclear focus of buffer, and incubated for 20 min at room temperature. Cells were diluted in classical SIM medium and 1 M TssK L14A ligand was in the cell and 1 mM WT BCP analyte was in the syringe. The reaction was performed with a constant stirring speed of 750 rpm; each injection lasted for 4 s with a 150 s space between injections. A constant heat control (offset) was removed from the raw data to account for heat dilution before integration. The data were fitted using the One Set of Sites model in the PEAQ-ITC analysis software.

**Supplemental Material**

Supplemental material is available online only.

**FIG S1**, TIF file, 1.7 MB.

**FIG S2**, TIF file, 2.1 MB.

**FIG S3**, TIF file, 2.1 MB.

**FIG S4**, TIF file, 1.7 MB.

**FIG S5**, TIF file, 1.7 MB.

**FIG S6**, TIF file, 1.7 MB.

**FIG S7**, TIF file, 2.1 MB.

**FIG S8**, TIF file, 1.7 MB.
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E.D., R.P., and Y.C. conceived the project. E.D., R.P., Y.C., V.S., D.B., I.F.-M., S.B., X.M., and M.N. designed the experiments. Y.C., V. Schmidt, V. Gloglupi, and R.C. performed the in vitro and in vivo studies. I.F.-M. performed the in silico studies. E.D., R.P., Y.C., and M.N. wrote the manuscript with contribution from all of the authors.

We declare no competing interest.

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