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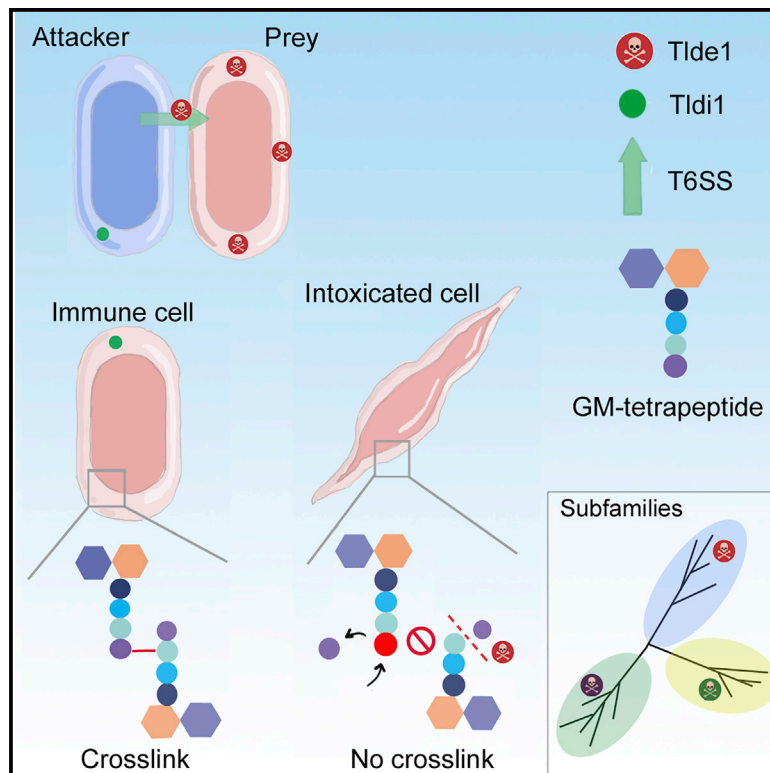
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# A Family of T6SS Antibacterial Effectors Related to L,D-Transpeptidases Targets the Peptidoglycan

## Graphical Abstract



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## In Brief

Antibacterial toxic effectors are an armory used by bacteria to compete against rivals. Sabinelli-Sousa et al. reveal the mechanism by which a family of T6SS effectors interferes with target bacteria peptidoglycan. Tlde1 cleaves and modifies peptide stem precursors and reduces their availability, preventing peptidoglycan synthesis.

## Highlights

- Tlde1 is an antibacterial T6SS effector related to L,D-transpeptidases
- Tlde1 does not form crosslinks between peptide stems
- Tlde1 has L,D-carboxypeptidase and L,D-transpeptidase exchange activity
- Tlde1 reduces the availability of peptidoglycan precursors and impairs synthesis



## Article

# A Family of T6SS Antibacterial Effectors Related to L,D-Transpeptidases Targets the Peptidoglycan

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## SUMMARY

Type VI secretion systems (T6SSs) are nanomachines used by bacteria to inject toxic effectors into competitors. The identity and mechanism of many effectors remain unknown. We characterized a *Salmonella* T6SS antibacterial effector called Tlde1 that is toxic in target-cell periplasm and is neutralized by its cognate immunity protein (Tldi1). Microscopy analysis reveals that cells expressing Tlde1 stop dividing and lose cell envelope integrity. Bioinformatic analysis uncovers similarities between Tlde1 and the catalytic domain of L,D-transpeptidases. Point mutations on conserved catalytic residues abrogate toxicity. Biochemical assays reveal that Tlde1 displays both L,D-carboxypeptidase activity by cleaving peptidoglycan tetrapeptides between meso-diaminopimelic acid<sup>3</sup> and D-alanine<sup>4</sup> and L,D-transpeptidase exchange activity by replacing D-alanine<sup>4</sup> by a non-canonical D-amino acid. Phylogenetic analysis shows that Tlde1 homologs constitute a family of T6SS-associated effectors broadly distributed among Proteobacteria. This work expands our current knowledge about bacterial effectors used in interbacterial competition and reveals a different mechanism of bacterial antagonism.

## INTRODUCTION

Bacteria commonly live in densely populated polymicrobial communities and compete over scarce resources. Several types of contact-dependent antagonistic interactions between bacteria have been described (García-Bayona and Comstock, 2018). The type VI secretion system (T6SS) is a dynamic contractile structure evolutionarily related to bacteriophage tails that delivers protein effectors in a contact-dependent manner into diverse cellular types, including eukaryotic host cells and rival bacteria and fungi (Hachani et al., 2016; Coulthurst, 2019; Trunk et al., 2019). T6SSs were also reported to display contact-independent functions in which secreted effectors facilitate the scavenging of scarce metal ions (Wang et al., 2015; Si et al., 2017a, 2017b; DeShazer, 2019).

The T6SS is anchored in the bacterial envelope and is composed of 13 core structural components that assemble into 3 major complexes: the trans-membrane complex, the baseplate, and the tail (Nguyen et al., 2018). A conformational change in the T6SS baseplate is thought to trigger the contraction of a cytoplasmic sheath, expelling a spear-like structure to puncture target cell membranes (Wang et al., 2017; Salih et al., 2018). The spear is composed of Hcp (hemolysin co-regulated protein) hexamers capped with a trimer of VgrG (valine-glycine repeat protein G) proteins and a PAAR (proline-alanine-alanine-

arginine) protein tip (Mougous et al., 2006; Renault et al., 2018; Shneider et al., 2013). Cargo effector proteins associate through non-covalent interactions with these structural components, while specialized effectors are presented as additional C-terminal domains fused to Hcp, VgrG, or PAAR proteins (Cianfanelli et al., 2016; Jana and Salomon, 2019). Consequently, along with the Hcp-VgrG-PAAR puncturing device, a cocktail of effectors is delivered into the target cell after each contraction event.

Antibacterial effectors delivered by the T6SS induce toxicity by targeting important structural components or affecting target-cell metabolism. Several families of effectors have been described, including peptidoglycan amidases and hydrolases, phospholipases, nucleases, nicotinamide adenine dinucleotide (phosphate) (NAD(P)+)-glycohydrolases, pore-forming proteins, and enzymes that synthesize (p)ppApp (Russell et al., 2011, 2012; Koskiniemi et al., 2013; Whitney et al., 2013; Ma et al., 2014; Altindis et al., 2015; Tang et al., 2018; Ahmad et al., 2019; Jana et al., 2019; Mariano et al., 2019; Wood et al., 2019). An example of an effector inducing toxicity by posttranslational modification (ADP-ribosylation) of the cytoskeleton component FtsZ has also been reported (Ting et al., 2018). To prevent self-intoxication, bacteria have a specific immunity protein for each antibacterial T6SS effector. Immunity proteins are encoded adjacent to their cognate effector, reside in the same cellular compartment where the effector exerts its toxic effect,



and typically work by binding directly to the effector (Hood et al., 2010; Russell et al., 2012).

The peptidoglycan sacculus maintains cell shape and provides mechanical strength to resist osmotic pressure (Vollmer et al., 2008). The mesh-like structure surrounds the cytoplasm and inner membrane and is composed of glycan chains of alternating *N*-acetylglucosamine (NAG) and *N*-acetylmuramic acid (NAM) residues crosslinked by short peptides containing both *L*- and *D*-amino acids. In Gram-negative bacteria, the peptide component is usually made of the following amino acids: *L*-alanine<sup>1</sup>, *D*-isoglutamic acid<sup>2</sup> (*D*-iGlu), *meso*-diaminopimelic acid<sup>3</sup> (*mDAP*), *D*-alanine<sup>4</sup>, and *D*-alanine<sup>5</sup> (Typas et al., 2011). Peptide stems are crosslinked to one another by transpeptidases (TPases) that could be either *D*,*D*-TPases, forming crosslinks between the *D*-Ala<sup>4</sup> of a pentapeptide donor stem and the *mDAP*<sup>3</sup> of a tetrapeptide acceptor stem (4 → 3 crosslink), or *L*,*D*-TPases, forming crosslinks between the *mDAP*<sup>3</sup> of one tetrapeptide and the *mDAP*<sup>3</sup> of another tetrapeptide (3 → 3 crosslink) (Vollmer and Bertsche, 2008). *D*,*D*-TPases (also called penicillin-binding proteins [PBPs]) are the primary enzymes performing crosslinks in the peptidoglycan and are inhibited by  $\beta$ -lactam antibiotics, which mimic the terminal *D*-Ala<sup>4</sup>-*D*-Ala<sup>5</sup> moiety of the donor pentapeptide (King et al., 2016).

T6SS effectors targeting the peptidoglycan distribute into 2 groups: (1) those that act as amidases and cleave within the peptide stems or crosslinks (Russell et al., 2012) and (2) those that act as glycoside hydrolases and cleave the glycan backbone (Whitney et al., 2013). T6SS amidase effectors form 4 phylogenetically distinct families called Tae1–Tae4 (type VI amidase effectors), and the preferred cleavage site within the peptidoglycan varies between each family (Russell et al., 2012). Tae1 and Tae4 cleave between *D*-iGlu<sup>2</sup> and *mDAP*<sup>3</sup> within the same peptide stem, while Tae2 and Tae3 cleave the crosslink bridge between *D*-Ala<sup>4</sup> and *mDAP*<sup>3</sup> of different peptide stems (Russell et al., 2012). An additional effector called TaeX cleaves between NAM glycan and the first *L*-Ala<sup>1</sup> of the peptide stem (Ma et al., 2018). The superfamily of T6SS glycoside hydrolases was divided into 3 families called Tge1–Tge3 (type VI glycoside hydrolase effector) (Whitney et al., 2013). Tge members have a lysozyme-like fold and were shown to cleave the glycoside bond between NAM and NAG (Whitney et al., 2013).

In *Salmonella enterica*, T6SS gene clusters are encoded within different pathogenicity islands (SPIs), depending on the subspecies and serovar (Blondel et al., 2009; Bao et al., 2019). *S. enterica* subsp. *enterica* serovar Typhimurium (*S. Typhimurium*) encodes only one T6SS that is located within SPI-6. The expression of SPI-6 T6SS genes is not detected under laboratory culture conditions, but it is activated in later stages of macrophage infection (Parsons and Heffron, 2005; Mulder et al., 2012) and in the mammalian gut, where it works as an antibacterial weapon to kill the resident species of the microbiota, contributing to *Salmonella* colonization (Sana et al., 2016). The histone-like nucleoid structuring protein (H-NS) (Brunet et al., 2015) and the ferric uptake regulator (Fur) (Wang et al., 2019) were reported to repress SPI-6 T6SS genes *in vitro*. Only one effector (Tae4) has been described to date as a substrate for SPI-6 T6SS. Tae4 is an antibacterial effector and works as an *L*,*D*-endopepti-

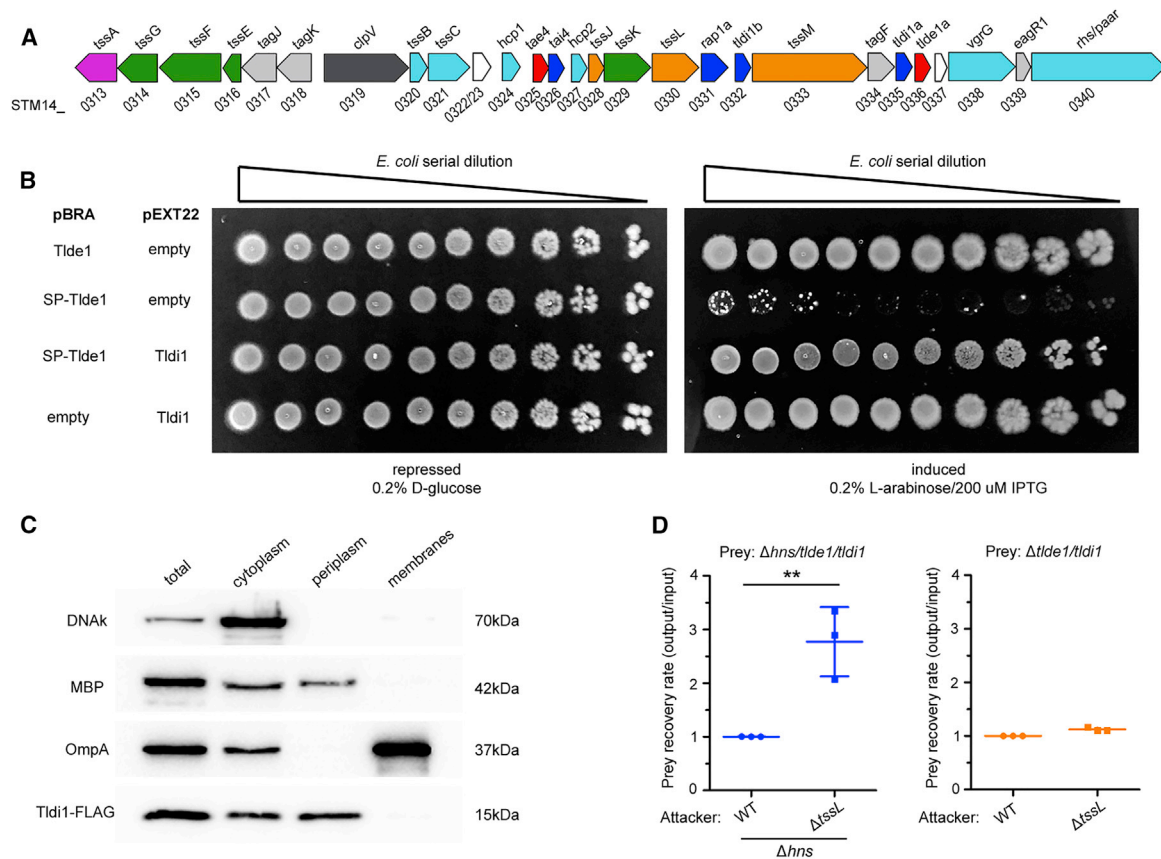
dase cleaving between *D*-iGlu<sup>2</sup> and *mDAP*<sup>3</sup> (Russell et al., 2012; Benz et al., 2013; Zhang et al., 2013).

Here, we set out to identify new SPI-6 T6SS effectors and report the characterization of a family of antibacterial effectors containing the domain of unknown function DUF2778, which displays both *L*,*D*-carboxypeptidase (CPase) and *L*,*D*-TPase exchange activities. This family is evolutionarily related to enzymes that have an *L*,*D*-TPase fold and is broadly distributed among  $\alpha$ -,  $\beta$ -, and  $\gamma$ -Proteobacteria. We also describe a protein containing DUF2195 as the immunity protein (Tldi1). Expression of Tldi1 in the periplasm of target *Escherichia coli* prevents cell division and induces cell elongation, swelling, and lysis. Our study reveals a different mechanism for effector-mediated bacterial antagonism and indicates that Tldi1 targets the peptidoglycan synthesis in two ways: (1) the *L*,*D*-CPase activity reduces the amount of acceptor tetrapeptide stems, thus reducing the formation of new crosslinks by *D*,*D*-TPases; and (2) the *L*,*D*-TPase exchange activity promotes the incorporation of non-canonical *D*-amino acid (NCDAAs) into tetrapeptides affecting their recycling and reducing the availability of cell wall precursors and substrates for *D*,*D*-transpeptidation. These activities together result in an altered formation of the division septum and an overall weakened peptidoglycan structure.

## RESULTS

### Tldi1-Tldi1 Are an Antibacterial Effector-Immunity Pair

To search for new T6SS effectors secreted by *S. Typhimurium*, we inspected the SPI-6 T6SS gene cluster of the 14028s strain looking for bicistrons that could resemble an antibacterial effector-immunity pair (Figure 1A). The gene annotated as *STM14\_0336* (Tldi1) encodes a small protein of 173 amino acids and contains a DUF2778 (PF10908). Upstream to this gene, there is another small protein (138 amino acids) containing a DUF2195 annotated as *STM14\_0335* (Tldi1), which encodes a predicted Sec signal peptide sequence for periplasmic localization (Figure S1A) (Almagro Armenteros et al., 2019). Bastion6 software prediction (Wang et al., 2018) indicates that *STM14\_0336* could be a T6SS effector (score 0.758). To test whether *STM14\_0336* and *STM14\_0335* are a bona fide effector-immunity pair, we cloned these genes into compatible vectors under the control of different promoters. To evaluate the toxicity of *STM14\_0336* upon expression in *E. coli* and to establish in which cellular compartment the toxin exerts its effect, *STM14\_0336* was cloned into the pBRA vector under the control of the  $P_{BAD}$  promoter (inducible by *L*-arabinose and repressed by *D*-glucose), both with and without an N-terminal PelB periplasmic localization sequence (pBRA SP-Tldi1 or pBRA Tldi1). The putative *STM14\_0335* immunity protein was cloned with its endogenous signal peptide into the pEXT22 vector under the control of the  $P_{TAC}$  promoter, which can be induced by isopropyl  $\beta$ -*D*-1-thiogalactopyranoside (IPTG) (pEXT22 Tldi1). *E. coli* strains carrying different combinations of pBRA and pEXT22 plasmids were serially diluted and incubated on LB agar containing either 0.2% *D*-glucose or 0.2% *L*-arabinose plus 200  $\mu$ M IPTG (Figure 1B). Results showed that *STM14\_0336* is toxic when directed to the periplasm of *E. coli* (pBRA SP-Tldi1) but not to the cytoplasm (pBRA Tldi1), and that *STM14\_0335* (pEXT22 Tldi1) could neutralize its toxicity (Figure 1B). To confirm



**Figure 1. Tlde1/Tldi1 Are an Antibacterial Effector-Immunity Pair**

(A) Schematic representation of SPI-6 T6SS gene cluster of *S. Typhimurium* 14028s: membrane complex (orange), baseplate (green), tail (light blue), toxins (red), immunity proteins (dark blue), chaperone and stabilizing protein (pink), ATPase for disassembly (dark gray), and accessory proteins (light gray). Accession numbers are indicated below.

(B) The 4-fold dilutions of *E. coli* containing pBRA and pEXT22 constructs, as indicated, spotted onto LB agar plates. Growth inhibition is observed upon expression of the SP-Tlde1 construct and can be neutralized by co-expression of Tldi1.

(C) *E. coli* cells expressing pEXT22 Tldi1-FLAG were fractionated and analyzed by western blot using anti-FLAG, anti-DNAK (cytoplasmic), anti-OmpA (membranes), and anti-MBP (periplasm) antibodies.

(D) Bacterial competition assay using *S. Typhimurium* LT2 strain (WT,  $\Delta tssL$ , and  $\Delta tld1/tldi1$ ) either in the  $\Delta hns$  (activated T6SS) or the WT background (repressed T6SS). The prey recovery rate was calculated by dividing the colony-forming unit (CFU) counts of the output by the input. Data represent the mean  $\pm$  SD of 3 independent experiments performed in triplicate and were analyzed through comparison with WT that were normalized to 1. \*\* $p < 0.01$  (Student's *t* test). See also Figure S1.

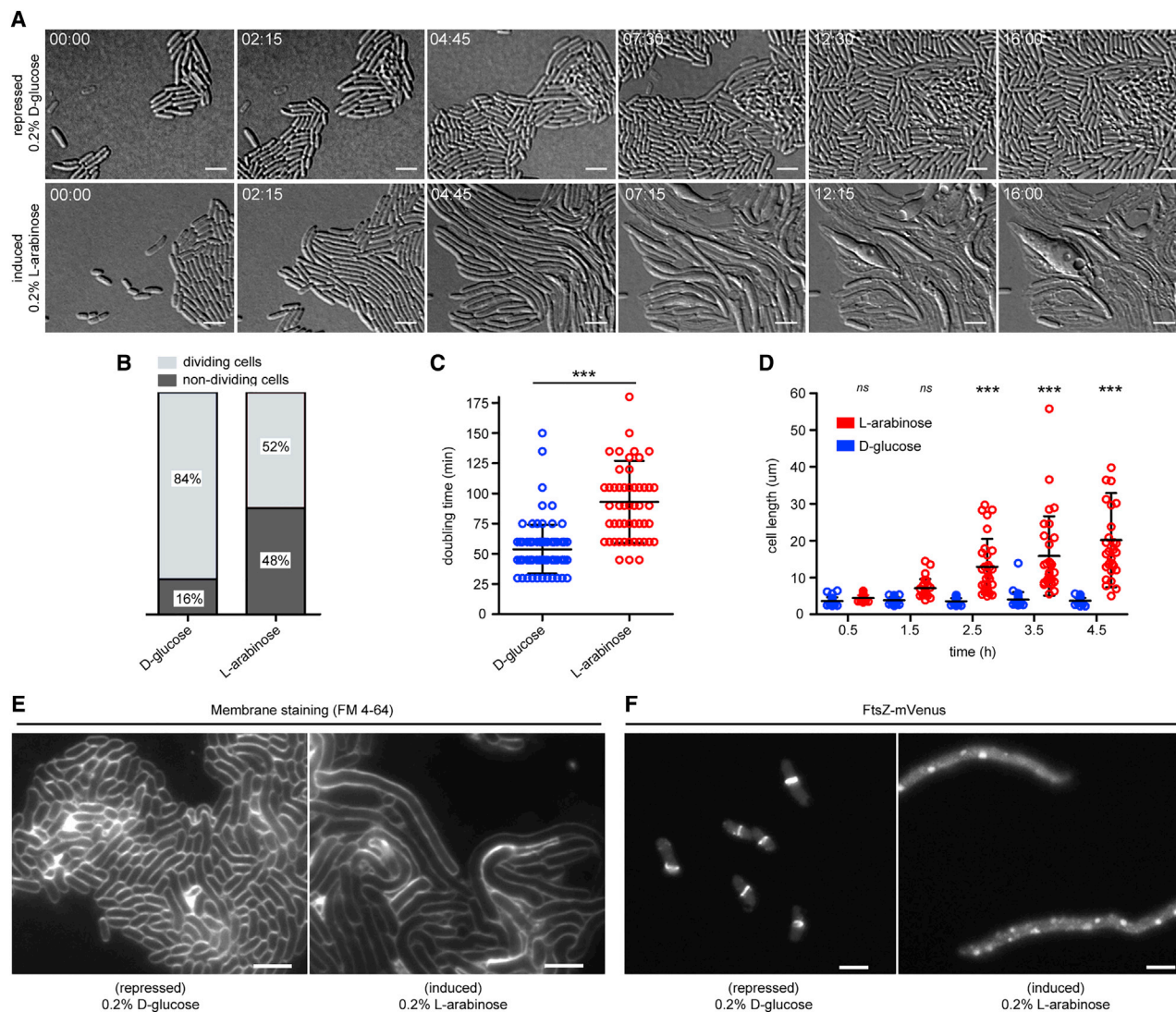
the subcellular localization of the immunity protein, *E. coli* cells expressing a C-terminal FLAG-tagged version of Tldi1 were subjected to subcellular fractionation, and results confirmed its periplasmic localization (Figure 1C). Western blot signals detected with anti-FLAG and anti-maltose-binding protein (MBP) in the cytoplasmic fraction represent a portion of soluble periplasmic proteins that were not completely extracted during the osmotic shock (Figure 1C).

The SPI-6 T6SS of *S. Typhimurium* is repressed in LB media by the silencer protein H-NS, and its deletion activates the T6SS (Brunet et al., 2015). After analyzing published RNA sequencing (RNA-seq) data comparing wild-type (WT) and  $\Delta hns$  strains (LT2 strain background) (Navarre et al., 2006), we confirmed that *VgrG*/STM0289 and *Tae4*/STM0277 mRNA levels were upregulated by 7.2-fold and 4.6-fold, respectively, in  $\Delta hns$  compared to WT. The same pattern was observed for *Tlde1*/STM0288, which

was upregulated by 3.5-fold in  $\Delta hns$  compared to WT. To analyze whether Tlde1 is a T6SS substrate, we performed inter-bacterial competition assays using WT and  $\Delta tssL$  (attacker) versus a mutant strain lacking the effector-immunity pair  $\Delta tld1/tldi1$  (prey), either in the  $\Delta hns$  (activated T6SS) or the WT background (repressed T6SS). Results showed that the prey recovery rate was  $\sim$ 3-fold higher in  $\Delta tssL/\Delta hns$  compared to  $\Delta hns$ , while no difference was observed when the T6SS was repressed (Figure 1D). These results confirm that Tlde1 is an antibacterial effector secreted via the SPI-6 T6SS.

### Tlde1 Intoxication Causes Altered Cell Division, Swelling, and Lysis

To gather further insight on the mechanism by which Tlde1 induces toxicity, we performed time-lapse microscopy to evaluate the growth and morphology of individual *E. coli* cells carrying



### Figure 2. Tlde1 Is a Periplasmic-Acting Toxin That Alters Cell Division and Weakens the Peptidoglycan

(A) Time-lapse microscopy of *E. coli* cells expressing SP-Tlde1 grown on LB agar pads containing either 0.2% D-glucose (repressed) or 0.2% L-arabinose (induced). Scale bar, 5 μm. Timestamps in hours:minutes.

(B) Percentage of dividing and non-dividing cells observed in (A) between 0 and 4.5 h.

(C) Doubling time in minutes of cells that divided quantified in (B); error bars represent the SDs of the means of ~100 cells in each condition. \*\*\**p* < 0.0001 (Student's *t* test).

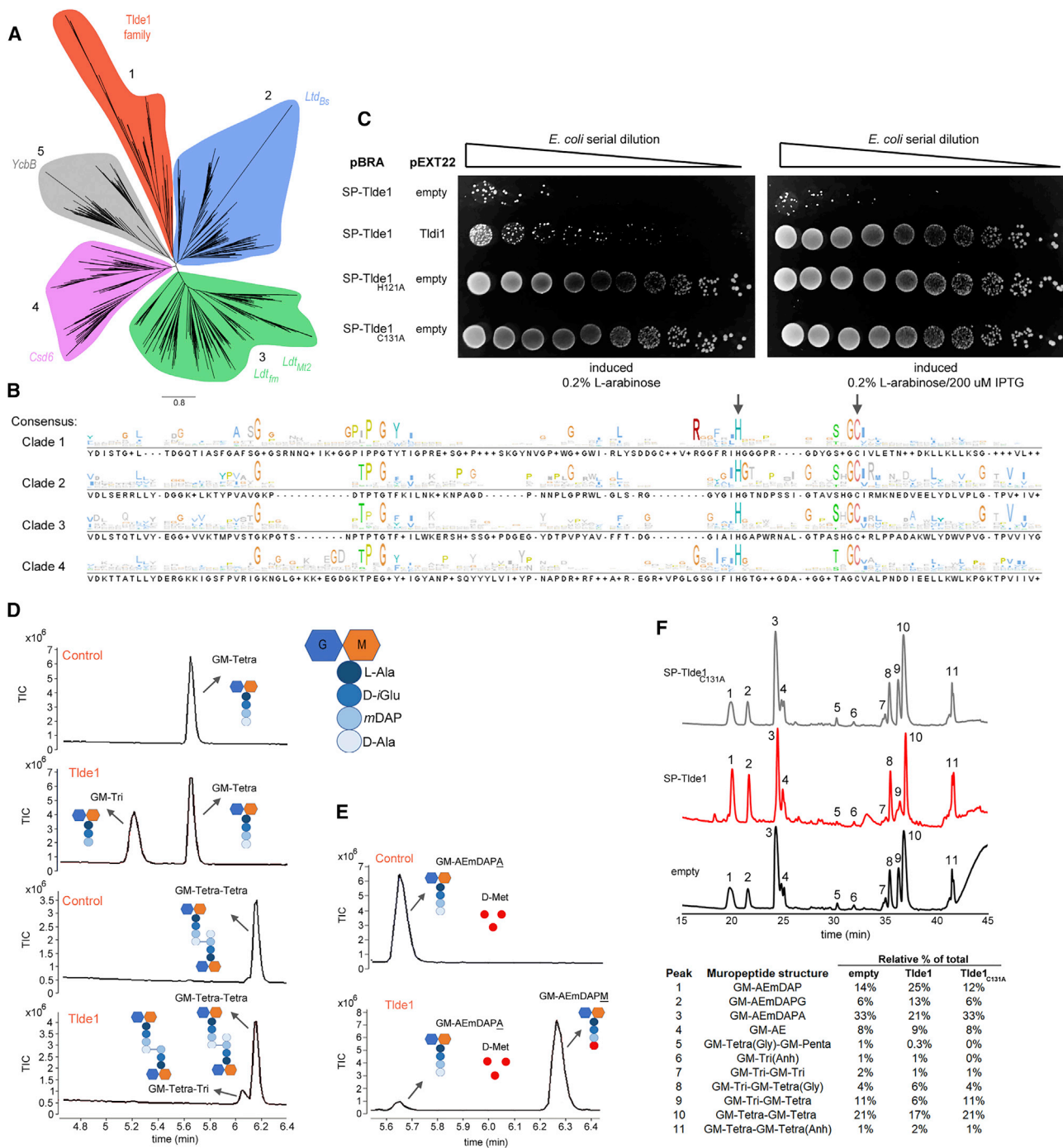
(D) Cell length of cells observed in (A) between 0 and 4.5 h; error bars represent the SDs of the means of ~30 cells measured at each time point, and were analyzed through comparison with D-glucose at 0.5 h by 1-way ANOVA followed by Dunnett's multiple comparison test. \*\*\**p* < 0.0001 and ns, not significant.

(E) Fluorescence microscopy images of *E. coli* cells harboring pBRA SP-Tlde1 labeled with the membrane dye FM 4-64 and incubated with 0.2% D-glucose (repressed) or 0.2% L-arabinose (induced). Scale bar, 5 μm.

(F) Fluorescence microscopy images of *E. coli* FtsZ-mVenus carrying pBRA SP-Tlde1 incubated with 0.2% D-glucose (repressed) or 0.2% L-arabinose (induced). Scale bar, 5 μm.

pBRA SP-Tlde1. *E. coli* cells grew normally when incubated on LB agar pads containing 0.2% D-glucose (repressed) over a time frame of 16 h (Figure 2A; Video S1). However, incubation in the presence of 0.2% L-arabinose induced a series of alterations in cell division and morphology (Figures 2A–2F; Video S2). At early time points (up to 4.5 h) after induction with L-arabinose, intoxicated cells tend to stop dividing or divide with

increased doubling times (Figures 2B and 2C). Between 0 and 4.5 h of incubation in L-arabinose, only 52% of intoxicated cells were observed to undergo at least 1 round of cell division, with a doubling time of  $93 \pm 34$  min, while 84% of cells grown with D-glucose divided in the same time frame, with a doubling time of  $54 \pm 20$  min (Figures 2B and 2C). Although intoxicated cells had clearly impaired cell division, they continued to increase in



**Figure 3. Tide1 Is Evolutionarily Related to L<sub>D</sub>-Transpeptidases**

(A) Maximum likelihood phylogenetic tree of Tide1 (DUF2778) homologs identified using JackHMMER and HMMsearch. Tide1 is a distant homolog of L<sub>D</sub>-TPases with the YkuD domain (PF03734). DUF2778-containing proteins grouped separately (clade 1, red) from known L<sub>D</sub>-TPases.

(B) Partial amino acid sequence alignment of consensus sequences from clades 1–4. Arrows indicate conserved catalytic histidine and cysteine residues of L<sub>D</sub>-TPases.

(C) The 4-fold dilutions of *E. coli* containing pBRA and pEXT22 constructs, as indicated, spotted onto LB agar plates. Growth inhibition is observed upon the expression of SP-Tide1, but toxicity is abolished by H<sub>121</sub>A and C<sub>131</sub>A point mutations.

(D) Reversed phase-HPLC (RP-HPLC) coupled to MS showing purified monomeric GM-tetrapeptides or dimeric GM-tetrapeptide-GM-tetrapeptide incubated with recombinant Tide1. The schematic structure of a muropeptide is shown. L-Ala, L-alanine; D-iGlu, D-isoglutamic acid; mDAP, meso-diaminopimelic acid; D-Ala, D-alanine; G, N-acetylglucosamine; M, N-acetylmuramic acid.

(legend continued on next page)

size (Figure 2D; Video S2). After 4.5 h, cells grown in L-arabinose had an average cell length of  $20 \pm 12 \mu\text{m}$ , while cells grown in D-glucose displayed an average length of  $3.7 \pm 0.7 \mu\text{m}$  (Figure 2D). To analyze whether intoxicated cells were not dividing or dividing but not segregating into daughter cells, we incubated *E. coli* with the membrane dye FM 4-64 (Figure 2E). Results showed that elongated cells lack membrane invaginations indicative of a division septum (Figure 2E). Transformation of an *E. coli* strain in which mVenus was fused to the endogenous *ftsZ* gene (Moore et al., 2016) with our pBRA SP-Tlde1 plasmid showed that the FtsZ-mVenus localization pattern was altered in intoxicated cells (Figure 2F). It is unclear at this point whether the Z-ring assembly was affected by the Tlde1 activity or is an indirect result of a pleiotropic effect such as the loss of membrane potential.

After 4.5 h in L-arabinose, *E. coli* pBRA SP-Tlde1 cells tend to swell and burst, indicating that the integrity of their peptidoglycan structure was compromised (Figure 2A; Video S2). No obvious alterations were observed in *E. coli* pBRA SP-Tlde1 cells grown in the presence of D-glucose during the same time frame (Figure 2A; Video S1).

### Tlde1 Is Evolutionarily Related to L,D-Transpeptidases

To gain insight into the molecular function of Tlde1, we used its amino acid sequence as a query in JackHMMER searches (Potter et al., 2018) to fetch a total of 143,242 sequences with significant similarity (inclusion threshold  $\leq 10^{-3}$  and e-value  $\leq 10^{-6}$ ) from the NCBI nr database (June 7, 2019). Additional JackHMMER searches using selected hits of the first iterative search as queries and the Pfam models DUF2778 (PF10908), YkuD (PF03734), and YkuD\_2 (PF13645) resulted in a total of 153,327 sequences. To reduce data complexity, we clustered amino acid sequences requiring 80% coverage for all pairwise alignments generated by MMseqs (Steinegger and Söding, 2017), and a maximum e-value of  $10^{-3}$ , resulting in 4,113 groups. A single amino acid sequence was chosen as representative from each group with at least 9 members, resulting in 943 sequences representing a sample of 145,969 homologs. These representative sequences were used to build a phylogenetic tree using the maximum likelihood method (Figure 3A). The homologs of Tlde1 clustered into 5 main clades: clade 1 (red, 6,332 sequences), to which Tlde1 belongs, is composed of proteins containing the uncharacterized DUF2778 domain (Tlde1 family); clade 2 (blue, 44,540 sequences) contains the L,D-TPases from *Bacillus subtilis* (Ldt<sub>Bs</sub>, PDB: 1Y7M) (Bielnicki et al., 2006); clade 3 (green, 44,090 sequences) contains L,D-TPases from *Enterococcus faecium* (Ldt<sub>fm</sub>, PDB: 1ZAT) (Biarrotte-Sorin et al., 2006), *Mycobacterium abscessus* (Ldt<sub>Mab</sub>, PDB: 5UWV) (Kumar et al., 2017), and *Mycobacterium tuberculosis* (Ldt<sub>Mt1-3</sub>, PDB: 3TUR, 5DCC) (Erdemli et al., 2012; Bianchet et al., 2017); clade 4 (purple, 25,242 sequences) contains an enzyme from *Helicobacter pylori* (Csd6, PDB: 4XZZ) that has a

catalytic domain resembling L,D-TPases but with L,D-CPase activity (Kim et al., 2015); and clade 5 (gray, 25,765 sequences) contains an L,D-TPase from *E. coli* (YcbB, PDB: 6NTW) (Caveney et al., 2019) and proteins recognized by the Pfam model YkuD\_2.

Multiple amino acid sequence alignments of proteins from each clade revealed conserved residues similar to the conserved motif described for L,D-TPases: HXX<sub>14-17</sub>[S/T]HGCh (underlined letters are conserved catalytic residues and “h” hydrophobic residues) (Erdemli et al., 2012) (Figure 3B). According to the Pfam hidden Markov model (HMM) logo, the number of residues between the conserved catalytic His and Cys is smaller in DUF2778 (PF10908) compared to YkuD (PF3734). To evaluate whether the conserved His in position 121 and Cys in position 131 of Tlde1 are required to induce toxicity, we produced point mutations by substituting these residues for alanine. Plasmids containing point mutations (pBRA SP-Tlde1<sub>H121A</sub> and pBRA SP-Tlde1<sub>C131A</sub>) were transformed into *E. coli* cells and grown in the presence of 0.2% L-arabinose, revealing the complete loss of toxicity (Figure 3C).

To determine the enzymatic activity of Tlde1, we incubated purified recombinant protein (Figure S2A) with purified peptidoglycan muropeptides and analyzed the reaction product by reverse-phase high-performance liquid chromatography (HPLC) coupled to mass spectrometry (MS) (Figures 3D and S2B). Results showed that Tlde1 has L,D-CPase activity and cleaves NAG-NAM-tetrapeptides (GM-tetrapeptide) between *mDAP*<sup>3</sup> and D-Ala<sup>4</sup>, producing GM-tripeptides (45% of substrate was converted) (Figures 3D). The formation of crosslinked GM-tripeptide-GM-tetrapeptide could not be detected, suggesting that Tlde1 does not have standard L,D-TPase activity (Figure 3D). Crosslinked dimeric GM-tetrapeptide-GM-tetrapeptide forms were also provided as substrates and a small proportion of cleavage was detected between the *mDAP*<sup>3</sup>-D-Ala<sup>4</sup> of the acceptor peptide stem during the same incubation period (11% of substrate was cleaved), thus confirming the L,D-CPase activity and suggesting that Tlde1 preferentially uses monomeric GM-tetrapeptide as a substrate (Figure 3D). Also, we could not detect any product indicating that Tlde1 cleaves the D-Ala<sup>4</sup>-*mDAP*<sup>3</sup> crosslink bridge between peptide stems (Figure 3D). In addition, co-incubation of GM-tetrapeptides (GM-AEmDAPA) with 1 mM D-methionine and recombinant Tlde1 produced GM-tetrapeptides with D-Met at position 4 (GM-AEmDAPM) (92% of substrate was converted), showing that Tlde1 is able to exchange the last amino acid of GM-tetrapeptides for a NCDAA, as was reported for other L,D-TPases (Figure 3E) (Mainardi et al., 2005).

The composition of peptidoglycan extracted from *E. coli* cells growing at exponential phase is enriched in monomeric GM-tetrapeptides and dimeric GM-tetrapeptide-GM-tetrapeptides (Glauner et al., 1988). During the synthesis of new peptidoglycan, the GM-pentapeptide precursor works as donor peptide stem

(E) RP-HPLC coupled to MS showing purified monomeric GM-tetrapeptides incubated with recombinant Tlde1 and 1 mM D-methionine. Tlde1 shows L,D-TPase exchange activity and incorporation of non-canonical D-amino acid (NCDAA) into peptidoglycan.

(F) RP-HPLC coupled to MS showing total ion chromatograms of muropeptides obtained after mutanolysin digestion of peptidoglycan extracted from *E. coli* harboring empty pBRA (black), pBRA SP-Tlde1 (red), and pBRA SP-Tlde1<sub>C131A</sub> (gray). Inferred muropeptide structures and relative abundance (relative percentage of total) of each peak was quantified by MassHunter software (bottom).

See also Figure S2.

and is crosslinked to acceptor GM-tetrapeptide stems by the action of  $D,D$ -TPases, forming 4  $\rightarrow$  3 crosslink bridges (Typas et al., 2011). As Tlde1 turns GM-tetrapeptides into GM-tripeptides *in vitro*, we hypothesized that this activity would promote toxicity *in vivo* by depleting the pool of acceptor GM-tetrapeptides, thus reducing the formation of new crosslinks by endogenous  $D,D$ -TPases. To test this hypothesis, we extracted peptidoglycan from *E. coli* cells carrying (1) empty pBRA plasmid, (2) pBRA SP-Tlde1, or (3) pBRA SP-Tlde1<sub>C131A</sub> incubated with 0.2% L-arabinose for 3 h and analyzed the mucopeptides profile by HPLC-MS (Figure 3F). The relative levels of mucopeptides in *E. coli* cells expressing SP-Tlde1 were reduced in GM-tetrapeptide (GM-AEmDAPA) (21%) compared to cells with empty plasmid (33%) or SP-Tlde1<sub>C131A</sub> (33%), comprising a 36% reduction in GM-AEmDAPA relative levels (Figure 3F). Likewise, the proportion of GM-tripeptides (GM-AEmDAP) was enriched in cells expressing SP-Tlde1 (25%) compared to empty and SP-Tlde1<sub>C131A</sub> (13%), which constitutes a 78% increase in relative levels of GM-tripeptides (Figure 3F). Moreover, the proportion of most crosslinked forms were reduced in *E. coli* cells expressing SP-Tlde1 compared to cells carrying the empty plasmid or expressing SP-Tlde1<sub>C131A</sub> (Figure 3F). *In vitro* peptidoglycan synthesis assays showed that GM-tripeptides could work as acceptors for  $D,D$ -TPases (Born et al., 2006; Bertsche et al., 2005; Catherwood et al., 2020). However, a study of PBP1B TPase showed that the  $K_m$  of GM-tripeptides is higher ( $35 \pm 4 \mu\text{M}$ ) than the  $K_m$  of GM-tetrapeptides ( $8.1 \pm 0.3 \mu\text{M}$ ), indicating that GM-tripeptides may not be preferential substrates for PBP1B *in vivo* (Catherwood et al., 2020).

Incorporation of NCDAA into peptidoglycan was reported to negatively regulate the amount of peptidoglycan of a given cell and its strength (Lam et al., 2009; Cava et al., 2011). These changes occur mainly because the incorporation of NCDAA into GM-tetrapeptide impairs its recycling, thus leading to reduced amounts of acceptor GM-tetrapeptides and donor GM-pentapeptides for  $D,D$ -transpeptidation during peptidoglycan synthesis (Templin et al., 1999; Hernandez et al., 2020). In agreement with the  $D$ -amino acid exchange activity detected *in vitro* (Figure 3E), we noticed an increase in GM-AEmDAPG (13%) in *E. coli* cells expressing SP-Tlde1 compared to cells with empty plasmid (6%) or catalytic inactive SP-Tlde1<sub>C131A</sub> (6%), which represents a 116% increase in relative levels (Figure 3F). This result supports the hypothesis that the exchange activity and the incorporation of NCDAA are also relevant *in vivo*. Overall, the combined action of both Tlde1 enzymatic activities impair the peptidoglycan synthesis and its structural strength.

### Tlde1 Family Is Widespread among $\alpha$ -, $\beta$ -, and $\gamma$ -Proteobacteria

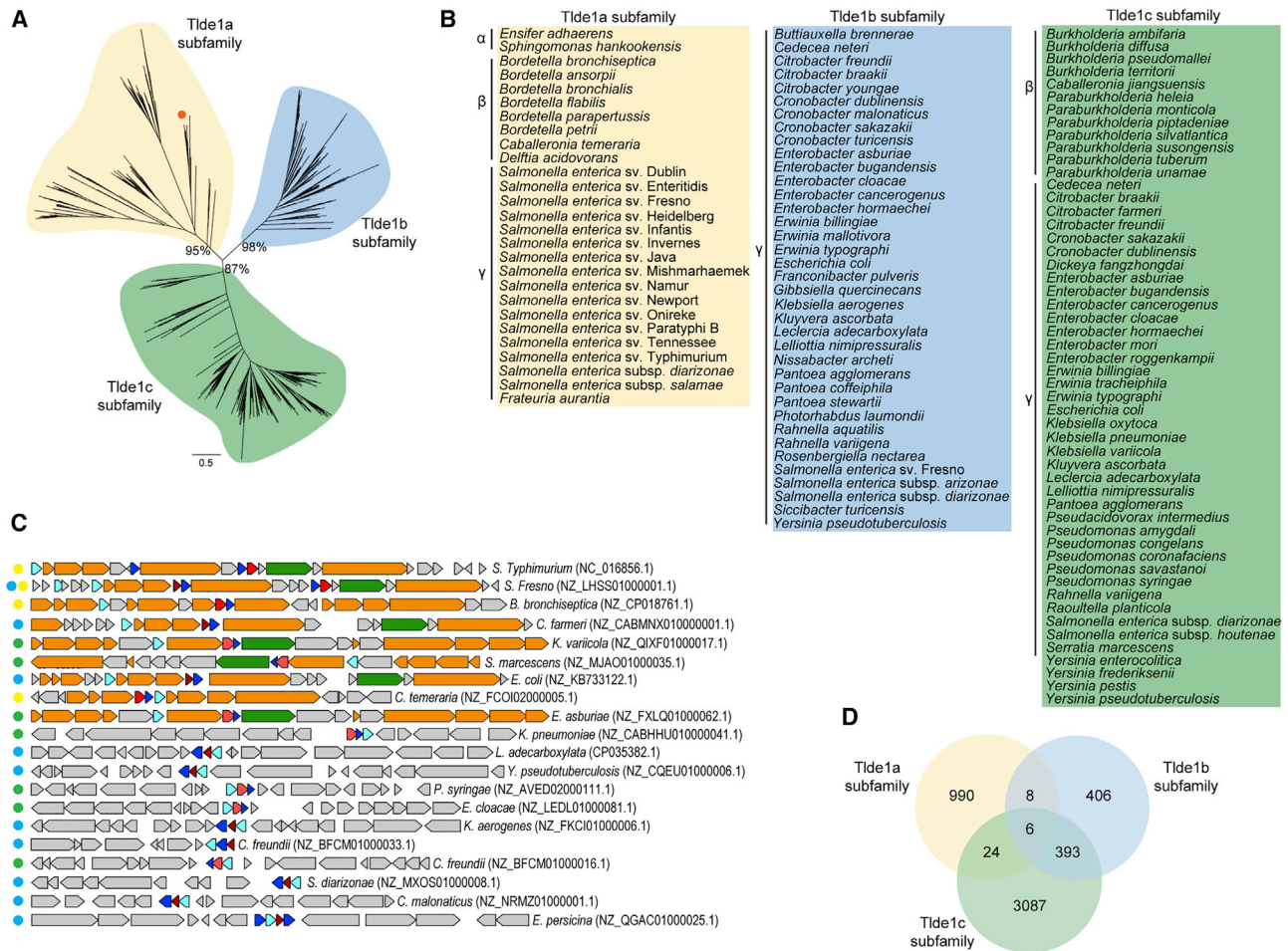
Comprehensive phylogenetic analysis of sequences from clade 1 comprising the Tlde1 family (Figure 3A) showed that proteins segregate into 3 subfamilies (Figure 4A). Given that all subfamilies contain the domain DUF2778, we propose calling them Tlde1a, Tlde1b, and Tlde1c (Figure 4A). The Tlde1a subfamily contains the *S. Typhimurium* Tlde1 effector (Tlde1a<sup>STM</sup>) and is composed mainly of *S. enterica* subsp. *enterica* serovars encoding SPI-6 T6SS, species of *Bordetella*, and some  $\alpha$ -Proteobacteria from Rhizobiaceae and Sphingomonadaceae (Figure 4B; Table S1A).

A few examples of *S. enterica* subsp. *diarizonae* (SPI-20 and SPI-21 T6SSs) and *S. enterica* subsp. *salamae* (SPI-19 T6SS) were detected in the Tlde1a subfamily (Figure 4B; Table S1A). In addition, the Tlde1a subfamily also contains examples of Actinobacteria (Streptomycetaceae) and Cyanobacteria (Synechococaceae, Microcystaceae, Aphanothecaceae); however, it is not clear at this point whether these proteins are secreted by alternative secretion mechanisms, such as the extracellular contractile injection systems (eCIS) (Chen et al., 2019), or simply work in the remodeling of peptidoglycan. The Tlde1b subfamily is abundant in a large variety of species from Enterobacteriaceae, Erwiniaceae, and Yersiniaceae. Examples of *S. enterica* subsp. *arizonae* (SPI-20 and SPI-21 T6SSs) and *S. enterica* subsp. *diarizonae* (SPI-20 and SPI-21 T6SSs) and a few *S. enterica* subsp. *enterica* serovars such as Fresno (SPI-6 T6SS) were detected in the Tlde1b subfamily (Figure 4B; Table S1B). The Tlde1c subfamily is the most widespread and can be found in Enterobacteriaceae, Burkholderiaceae, Pseudomonadaceae, Erwiniaceae, and Yersiniaceae (Figure 4B; Table S1C). Examples of *S. enterica* subsp. *diarizonae* (SPI-20 and SPI-21 T6SSs) and *S. enterica* subsp. *houteanae* (SPI-19 T6SS) were detected in the Tlde1c subfamily.

Examining the genomic context of Tlde1a–Tlde1c members, we observed that at least one member of each subfamily is encoded within a T6SS gene cluster, thus supporting their role as T6SS substrates (Figure 4C; Table S2). In addition, Tlde1b and Tlde1c were found to be associated with orphan Hcp proteins (Figure 4C; Tables S2B and S2C), suggesting that these proteins are secreted as cargo effectors associated with Hcp proteins. Tlde1a members were found outside the T6SS gene cluster together with their putative immunity proteins, although none of these examples were associated with an orphan Hcp (Table S2A). From the total of 1,028 genomes containing a Tlde1a member, 615 (59.8%) are encoded in the vicinity of a T6SS structural gene (Figure 4C). The same high frequency is observed for Tlde1b and Tlde1c subfamilies, with 76.6% (623 of 813) and 42.6% (1,495 of 3,510) of the genomes containing a member in the vicinity of T6SS structural genes, respectively.

Genomic analysis revealed that some organisms encode more than one member of the Tlde1 family (Figures 4C and 4D; Table S1D). *S. Fresno* encodes 2 members within its SPI-6 T6SS, Tlde1a<sup>SF</sup> (WP\_000968384.1) and Tlde1b<sup>SF</sup> (WP\_058115706.1). Tlde1a<sup>SF</sup> is closely related to *S. Typhimurium* Tlde1a<sup>STM</sup> (STM14\_0336), but the Tlde1b<sup>SF</sup> homolog is absent in *S. Typhimurium* 14028s (Figure 4C). A close inspection of the *S. Typhimurium* 14028s SPI-6 T6SS locus revealed that it contains an orphan Tldi1b<sup>STM</sup> immunity protein (STM14\_0332) orthologous to *S. Fresno* Tldi1b<sup>SF</sup> (WP\_077921582.1) in the same genomic location (Figures 1A and 4C). This gene is located next to a Tai4 orphan immunity protein (STM14\_0331) homologous to Rap1a, which confers immunity to the Tae4 family member Ssp1 from *Serratia marcescens* (English et al., 2012). Growth inhibition assays showed that the Tldi1b<sup>STM</sup> immunity protein was not able to neutralize the toxicity of Tlde1a<sup>STM</sup>, thus confirming the specificity of each immunity protein to its effector (Figure S3) and suggesting that Tldi1b<sup>STM</sup> and Rap1a may play a role in immunity against effectors secreted by other bacteria.

The domain architecture of most Tlde1 members is usually quite simple and composed of a single DUF2778 domain, but



**Figure 4. Tlde1 Family Is Widespread in Proteobacteria**

(A) Maximum likelihood phylogenetic tree of Tlde1 family (clade 1; Figure 3A), showing its division into 3 subfamilies (Tlde1a, Tlde1b, and Tlde1c). Non-parametric bootstrap values at the branches defining each subfamily are indicated. The position of *S. Typhimurium* Tlde1a is highlighted by an orange dot.

(B) List of common bacterial species encoding members of Tlde1a, Tlde1b, and Tlde1c. The toxic effector is widespread and detected among  $\alpha$ -,  $\beta$ -, and  $\gamma$ -Proteobacteria.

(C) Schematic representation of the genomic context of Tlde1a, Tlde1b, and Tlde1c members. T6SS structural genes (orange), VgrG genes (green), Hcp genes (light blue), Tlde1a-c members (red), and their cognate immunity proteins Tldi1a-c (dark blue). Bacterial species and genome accession number are described on the right side of each cluster. Colored dots on the left side denote the subfamily to which the effector belongs: yellow (Tlde1a), blue (Tlde1b), and green (Tlde1c).

(D) Venn diagram representing the total number of bacterial genomes that encode one or more Tlde1a–Tlde1c member.

See also Figure S3 and Tables S1 and S2.

a few examples such as *Shigella sonnei* WP\_052983304.1 present DUF2778 at the C-terminal region of rearrangement hotspot (Rhs) proteins, which are encoded close to PAAR-containing proteins that are secreted via T6SS (Koskiniemi et al., 2013). These Rhs-DUF2778 proteins are mainly represented in the Tlde1c subfamily, in which 11% (387 of 3,510) of the genomes display a member fused to Rhs protein (Table S1C). Moreover, DUF2778-containing proteins could be found located in the vicinity of genes encoding other secretion systems such as T1SS (*abc*, *omf*, *mfp*), T2SS (*gspD-N*), T4SS (*virb1-10*), and T5SS (PF03797, PF03895, translocator) (Table S2) (Abby et al., 2016), suggesting that these DUF2778-containing proteins may have been recruited to be part of the polymorphic toxin cassettes that are known to be associated with various protein

secretion systems (Zhang et al., 2012). These results are in line with bioinformatic analyses that predicted polymorphic toxins with L,D-peptidase domains fused to Rhs repeats or domains related to secretory systems and were linked to the neighboring immunity proteins DUF2750/Imm16 and DUF2750/Imm57 (Zhang et al., 2012).

## DISCUSSION

In this study, we identified and characterized a family of T6SS antibacterial effectors, which is evolutionarily related to L,D-TPases containing the YkuD domain (formerly called ErfK/YbiS/YcfS/YnhG). The founding member of this family, Tlde1 from *S. Typhimurium*, displays both L,D-CPase activity, cleaving

peptidoglycan tetrapeptides between *mDAP*<sup>3</sup> and *D-Ala*<sup>4</sup>, and *L,D*-TPase exchange activity, replacing *D-Ala*<sup>4</sup> by a NCDAA. The specificity and toxic mechanism of Tlde1 are different from previously characterized T6SS effectors targeting the peptidoglycan. T6SS amidase effectors Tae1 and Tae4 cleave the bond between *D-iGlu*<sup>2</sup> and *mDAP*<sup>3</sup> within the same peptide stem, while Tae2 and Tae3 cleave the crosslink bridge between *D-Ala*<sup>4</sup> and *mDAP*<sup>3</sup> of different peptide stems (Russell et al., 2012). T6SS glycoside hydrolases (Tge) cleave the glycoside bond between NAM and NAG (Whitney et al., 2013). In addition, the evolutionary origin of Tlde1 is different from T6SS amidases: while the Tae1–Tae4 catalytic domains are related to CHAP (PF05257) and NlpC/P60 (PF00877) amidases (Anantharaman and Aravind, 2003; Bateman and Rawlings, 2003), the Tlde1a–Tlde1c catalytic domains are related to YkuD of *L,D*-TPases (PF03734) (Biarrotte-Sorin et al., 2006).

Due to the importance of peptidoglycan in providing shape and resistance to osmotic pressure, it is not surprising that a large number of bacterial toxins target the synthesis and/or integrity of the sacculus. The peptidoglycan is made of glycan strands crosslinked by short peptides, whose composition is highly diverse among bacterial species (Vollmer and Born, 2009). The greatest variation occurs at the third position of the peptide stem, which is *mDAP* in almost all Gram-negative bacteria (including some species of *Bacillus*, *Lactobacilli*, and *Mycobacterium*) and *L-Lys* in most Gram-positive bacteria, but other amino acids can occur at this position such as *L-ornithine*, *D-Lys*, *meso-lanthionine*, *L-homoserine*, *L-Ala* and *L-Glu*. Moreover, *mDAP* and *L-Lys* can be hydroxylated or amidated in a few species (Vollmer and Born, 2009). Such variability in the peptidoglycan may be the reason why T6SS-containing bacteria use such a diverse array of effectors to target this important structure. A detailed study of 20 species representing 5 orders of the class Alphaproteobacteria revealed a great diversity in peptide stem composition and crosslink (Espaillat et al., 2016). Modifications in the peptidoglycan of Acetobacteria could provide protection against T6SS amidases Tae1 from *Pseudomonas aeruginosa* and Tae2 from *Acidovorax citrulli*; these species naturally coexist in the soil (Espaillat et al., 2016).

Besides the variability in target-cell peptidoglycan structure, it is known that T6SS effectors display conditional toxicity depending on pH, salinity, temperature, and oxygen availability (LaCourse et al., 2018). Furthermore, some effectors exhibit a synergistic effect and are more proficient in killing target cells when combined with other T6SS effectors (LaCourse et al., 2018). This evidence helps explain why bacteria harbor so many effectors for the same cellular target. In the case of *Salmonella enterica*, most serovars live in a variety of environments such as soil, water, plants, and vertebrate hosts (e.g., chicken, cattle, humans) (Bäumler et al., 1998; Winfield and Groisman, 2003). In light of this vast range of environmental conditions, one can only imagine the great diversity of rival bacterial species encountered, so it makes sense for *Salmonella* to deliver a cocktail of effectors into target cells, which may have different levels of effectiveness depending on the pH, salinity, temperature, and oxygen availability at that specific encounter. This could be the case for *Salmonella* Tae4 effector and the newly identified Tlde1.

*L,D*-TPases form *mDAP*<sup>3</sup>-*mDAP*<sup>3</sup> crosslinks between peptide stems by transferring the peptide bond between the third residue of a tetrapeptide donor stem to the side-chain amide group of a third residue of an adjacent acceptor stem. The catalytic mechanism was proposed to occur in a two-step enzymatic reaction requiring a cysteine residue; it involves the acylation of the enzyme by the penultimate peptide of the donor stem with the release of the C-terminal amino acid residue *D-Ala*<sup>4</sup>, followed by deacylation of this acyl-enzyme intermediate by an acceptor stem (Biarrotte-Sorin et al., 2006; Erdemli et al., 2012). Enzymes displaying *L,D*-TPase activity (*Ldt*<sub>BS</sub>, *Ldt*<sub>fm</sub>, *Ldt*<sub>Mt</sub>) (clades 2 and 3; Figure 3A) have an elongated active site, with the catalytic residues accessible via two paths. However, *Csd6* from *H. pylori*, which displays an *L,D*-TPase domain but only *L,D*-CPase activity (clade 4; Figure 3A), has the active site in a deep pocket, with the catalytic triad positioned at the bottom and accessible via a single narrow path (Kim et al., 2015). The absence of detailed structural information for Tlde1 does not allow us to determine its precise catalytic mechanism at this point; however, we speculate that structural features that restrict access to its active site may explain its activity. The lack of crosslinked GM-tetrapeptide dimer products during *in vitro* assays (Figure 3D) and their *L,D*-CPase and *D*-amino acid exchange activities support the hypothesis that the active site of Tlde1 is not accessible to larger acceptor substrates such as GM-tetrapeptides; instead, Tlde1 preferentially uses small molecules such as water and free amino acids as acceptors.

Peptidoglycan synthesis is differentially controlled during cell division and elongation, with the septal synthesis machinery being favored during cell division (Woldringh et al., 1987; Wientjes and Nanninga, 1989). It has been shown that cell division is mediated by the filaments of FtsZ and FtsA that treadmill circumferentially around the division ring and drive the motion of peptidoglycan-synthesizing enzymes (Bisson-Filho et al., 2017; Yang et al., 2017). The rate of division septum closure is mainly determined by the *D,D*-TPase activity of FtsI (Coltharp et al., 2016). The toxic phenotype induced by periplasmic expression of Tlde1 seems to be the result of a series of events. First, up to 4.5 h of intoxication, cells stop dividing but continue growing in length. Our hypothesis that Tlde1 promotes toxicity by interfering with peptidoglycan synthesis and transpeptidation fits with the finding that division septum closure is limited by peptidoglycan synthesis. Second, the phenotype observed after 4.5 h in which cells form blebs and lyse (Video S2) is also aligned with the idea of diminished peptidoglycan synthesis and crosslink rates; when the peptidoglycan growth rate falls behind that of overall cell growth, the peptidoglycan net stretches, resulting in changes in pore size that alter the integrity of the cell wall (Typas et al., 2011).

In T6SS<sup>+</sup> organisms, Tlde1 proteins are encoded in bicistrons with their putative immunity proteins (Figure 4C). Divergence among immunity proteins was considerably higher than that found within the effector families. The immunity protein for Tlde1a subfamily has a DUF2195 domain (Tlde1a), while Tlde1b and Tlde1c immunity proteins are encoded by genes with no annotated domain. Tlde1a and Tlde1b immunity proteins encode a Sec signal peptide for periplasmic localization, while Tlde1c immunity proteins have one or more transmembrane domains

(Figure S1) (Krogh et al., 2001). A similar diversity in localization signals of T6SS immunity proteins was observed for homologs of the toxin encoded at the C-terminal of VgrG2b (DUF4157) of *Pseudomonas aeruginosa* (Wood et al., 2019). The greater sequence divergence in immunity proteins has been reported previously and is likely due to less restrictive selective pressure compared to effectors; while immunity proteins are selected for effector binding, effectors are selected for both immunity protein binding and catalysis (Russell et al., 2012). Due to the small length of these immunity proteins and their diversity, which impeded automated bioinformatic analysis, we did not attempt further assessment of their phylogeny.

T6SSs translocate effectors by decorating the Hcp-VgrG-PAAR puncturing device (Cianfanelli et al., 2016; Jana and Salomon, 2019). Small effector proteins composed of only one toxic domain, such as Tlde1, usually interact with Hcp proteins for secretion as their size favors fitting inside the narrow tube formed by Hcp hexamers (Silverman et al., 2013; Jana and Salomon, 2019). Further supporting this notion, our bioinformatic analysis showed that Tlde1 family members associated with Hcp proteins (Figure 4C). Initial attempts to detect protein-protein interaction between Tlde1a<sup>STM</sup> and *S. Typhimurium* Hcp1–3 using bacterial two-hybrid and co-immunoprecipitation were unsuccessful.

The distribution of Tlde1a, Tlde1b, and Tlde1c effectors between different organisms gives an idea of how these toxins are used by different species to intoxicate competitors. It is interesting to note that most *S. enterica* subsp. *enterica* serovars encode only a Tlde1a member (e.g., *S. Typhimurium*). According to our analysis, Tlde1b members are mainly represented among species of the Enterobacteriaceae, indicating that Tlde1b/Tldi1b effector/immunity pairs must be a commonly used ammunition in the gut environment. Therefore, it does not seem to be a great competitive advantage for *S. Typhimurium* to keep a Tlde1b effector in its repertoire as many species already living in the gut encode a Tldi1b immunity. Similarly, despite that most *S. enterica* subsp. *enterica* serovars encode only Tlde1a, they frequently encode an orphan Tldi1b immunity, suggesting that although not attacking competitors using Tlde1b, *Salmonella* are protected from the attack of members of the microbiota encoding a Tlde1b effector. Overall, from the perspective of an enteric pathogen that needs to kill competitor species in the gut to colonize the environment, *S. enterica* subsp. *enterica* serovars are well equipped with a Tlde1a member, which is not present in most members of Enterobacteriaceae.

## STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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## SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.celrep.2020.107813>.

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## AUTHOR CONTRIBUTIONS

S.S.-S., J.T.H., and E.B.-S. outlined the study. S.S.-S., J.T.H., B.Y.M., and E.B.-S. performed the experiments and analyzed the data. A.P. and S.M. performed the peptidoglycan enzymatic assays and analyzed the data. G.G.N. and R.F.d.S. performed the bioinformatic analyses. S.S.-S., J.T.H., S.M., G.G.N., R.F.d.S., C.R.G., and E.B.-S. contributed to the scientific discussions. S.S.-S. and E.B.-S. wrote the manuscript with input from other authors.

## DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE   | SOURCE                    | IDENTIFIER         |
|---|---------------------------|--------------------|
| <b>Antibodies</b>   |                           |                    |
| Rabbit polyclonal anti-FLAG                                       | Sigma-Aldrich             | Cat#F7425          |
| Mouse monoclonal anti-DNAK  | Abcam                     | Cat#ab69617        |
| Mouse monoclonal anti-maltose binding protein (MBP)               | New England BioLabs       | Cat#E8032L         |
| Rabbit polyclonal anti-outer membrane protein A (OmpA)            | David Holden              | N/A                |
| <b>Bacterial and Virus Strains</b>                                |                           |                    |
| <i>Salmonella</i> Typhimurium str. 14028s wild-type               | Marcelo Brocchi           | ATCC 14028         |
| <i>S. Typhimurium</i> str. LT2 wild-type                          | Eric Cascales             | ATCC 700720        |
| <i>S. Typhimurium</i> str. LT2 $\Delta$ tssL (STM0282)            | This paper                | N/A                |
| <i>S. Typhimurium</i> str. LT2 $\Delta$ tld1/tldi1 (STM0287/0288) | This paper                | N/A                |
| <i>S. Typhimurium</i> str. LT2 $\Delta$ hns (STM1751)             | This paper                | N/A                |
| <i>Escherichia coli</i> DH5 $\alpha$                              | Lab Stock                 | N/A                |
| <i>E. coli</i> SHuffle T7   | New England Biolabs       | Cat#C3026J         |
| <i>E. coli</i> XL10-Gold  | Agilent Technologies      | Cat#200314         |
| <i>E. coli</i> BW27783 strain ftsZ::ftsZ-mVenus                   | Moore et al., 2016        | N/A                |
| <i>E. coli</i> $\Delta$ 6ldt                                      | Meberg et al., 2001       | N/A                |
| P22 phage lysate $\Delta$ hns                                     | Beraud et al., 2010       | N/A                |
| <b>Chemicals, Peptides, and Recombinant Proteins</b>              |                           |                    |
| FM4-64  | Molecular Probes          | Cat#T3166          |
| Pronase   | Roche                     | Cat#10 165 921 001 |
| Mutanolysin   | Sigma-Aldrich             | Cat#M9901          |
| <b>Critical Commercial Assays</b>                                 |                           |                    |
| QuikChange II XL Site-Directed Mutagenesis Kit                    | Agilent Technologies      | Cat#200521         |
| <b>Oligonucleotides</b>   |                           |                    |
| See Table S3 for full list.                                       | N/A                       | N/A                |
| <b>Recombinant DNA</b>  |                           |                    |
| pKD46   | Datsenko and Wanner, 2000 | N/A                |
| pKD4  | Datsenko and Wanner, 2000 | N/A                |
| pCP20   | Datsenko and Wanner, 2000 | N/A                |
| pBRA  | Souza et al., 2015        | N/A                |
| pBRA-SP   | Bayer-Santos et al., 2019 | N/A                |
| pBRA Tld1   | This paper                | N/A                |
| pBRA SP-Tld1  | This paper                | N/A                |
| pBRA SP-Tld1(H121A)   | This paper                | N/A                |
| pBRA SP-Tld1(C131A)   | This paper                | N/A                |
| pEXT22  | Dyxhoorn et al., 1996     | N/A                |
| pEXT22 Tld1a  | This paper                | N/A                |
| pEXT22 Tld1a-FLAG   | This paper                | N/A                |
| pEXT22 Tld1b  | This paper                | N/A                |
| pET28a  | Novagen                   | Cat#69864-3        |
| pET28a-Tld1   | This paper                | N/A                |

(Continued on next page)

**Continued**

| REAGENT or RESOURCE     | SOURCE                      | IDENTIFIER  |
|-------------------------|-----------------------------|---|
| Software and Algorithms |                             |   |
| Neighborhood script     | Robson F. de Souza          | upon request  |
| MassHunter              | Agilent Technologies        | N/A   |
| FIJI                    | Schindelin et al., 2012     | <a href="https://fiji.sc/">https://fiji.sc/</a>   |
| MMseqs                  | Steinegger and Söding, 2017 | <a href="https://search.mmseqs.com/search">https://search.mmseqs.com/search</a>               |
| FastTree 2              | Price et al., 2010          | <a href="http://www.microbesonline.org/fasttree/">http://www.microbesonline.org/fasttree/</a> |

**RESOURCE AVAILABILITY**

**Lead Contact**

Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Ethel Bayer Santos ([ebayersantos@usp.br](mailto:ebayersantos@usp.br)).

**Materials Availability**

Strains and plasmids generated in this study are available upon request to the Lead Contact, Ethel Bayer Santos ([ebayersantos@usp.br](mailto:ebayersantos@usp.br)).

**Data and Code Availability**

The code for data collection and analysis of gene neighborhood used in this study is available upon request to the Lead Contact, Ethel Bayer Santos ([ebayersantos@usp.br](mailto:ebayersantos@usp.br)).

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

A list of bacterial strains used in this work can be found in [Key Resources Table](#). Strains were grown at 37°C in Lysogeny Broth (10 g/L tryptone, 10 g/L NaCl, 5 g/L yeast extract) under agitation. Cultures were supplemented with antibiotics in the following concentration when necessary: 50 µg/mL kanamycin, 100 µg/mL ampicillin, 50 µg/mL streptomycin, 15 µg/mL chloramphenicol.

**METHOD DETAILS**

**Cloning and mutagenesis**

*S. Typhimurium* mutant strains were constructed by λ-Red recombination engineering using a one-step inactivation procedure (Datsenko and Wanner, 2000). All primers are listed in Table S3. *STM14\_0336* was amplified by PCR and cloned into pBRA vector under the control of P<sub>BAD</sub> promoter (Souza et al., 2015) with or without PelB signal peptide sequence from pET22b (Novagen) (Bayer-Santos et al., 2019). Immunity proteins (*STM14\_0332* and *STM14\_0335*) were cloned into pEXT22 under the control of P<sub>TAC</sub> promoter (Dykhorn et al., 1996). For protein expression and purification, *STM14\_0336* residues between 2-174 were cloned into pET28a (Novagen), including a N-terminal His-tag. Point mutations were created using QuikChange II XL Site-Directed Mutagenesis Kit (Agilent Technologies) and pBRA SP-Tide1 plasmid was used as template. All constructs were confirmed by sequencing.

**Growth inhibition assay**

Overnight cultures of *E. coli* DH5α co-expressing effectors for cytoplasmic (pBRA-Tide1) or periplasmic (pBRA SP-Tide1) localization and immunity proteins (pEXT22-Tidi1) were serially diluted in LB (1:4) and 5 µL were spotted onto LB agar (1.5%) containing either 0.2% D-glucose or 0.2% L-arabinose and 200 µM IPTG plus streptomycin and kanamycin and incubated at 37°C. Images were acquired after 24h.

Bacterial competition assays were performed using *S. Typhimurium* LT2 strain (WT, Δ*tssL* and Δ*tide1/tidi1*) either in the Δ*hns* (activated T6SS) or WT background (repressed T6SS). LT2 strain has a point mutation in *rpoS* gene which helps support the detrimental effect of *hns* deletion (Navarre et al., 2006; Lucchini et al., 2006). Due to the high frequency of spontaneous compensatory mutations, Δ*hns* strains were freshly prepared at every experiment by P22 phage transduction (Beraud et al., 2010). Overnight cultures of attacker and prey cells were subculture (1:10) until reaching OD<sub>600nm</sub> 1, cultures were mixed 4:1 attacker:prey (OD<sub>600nm</sub> 0.5), 5 µL spotted onto 0.22 µm nitrocellulose membranes (1 × 1 cm), and incubated on LB agar for 16 h at 30°C. The membranes containing the bacterial mixture was placed on 1.5 mL tubes containing 1 mL LB, homogenized by vortex, serially diluted, and plated on selective plates with antibiotics. Prey recovery rate was calculated by dividing the CFU counts of the output by the input. Data represent the mean ± SD of three independent experiments performed in triplicate and were analyzed through comparison with WT that were normalized to 1.

### Subcellular fractionation

Subcellular fractionation was adapted from [Gauthier et al. \(2003\)](#) and [English et al. \(2012\)](#) and is based on osmotic shock and ultracentrifugation. Briefly, *E. coli* cells harboring pEXT22-Tide1-FLAG grown overnight with 200  $\mu$ M of IPTG were harvested (aliquot corresponding to total proteins was collected for analysis), washed twice with phosphate buffer saline (PBS) and resuspended in 1 mL of 50 mM Tris-HCl pH 7.4, 20% sucrose, 10 mM EDTA and protease inhibitor. Cells were incubated for 10 min at 30°C and recovered by centrifugation (10 min, 8000 g, 22°C). Pellets were resuspended in 1 mL of ice-cold water and incubated for 10 min on ice. After centrifugation (10 min, 8000 g, 4°C), 900  $\mu$ L of the supernatant was retained for analysis (enriched in periplasmic proteins). The pellet containing cytoplasmic and membrane proteins was resuspended in 1 mL of sonication buffer (10 mM Tris-HCl pH 7 and protease inhibitor), sonicated 6 rounds of 15 s with amplitude 30% and centrifuged (15 min, 16000 g, 4°C) to remove insoluble proteins. Supernatant was transferred to an ultracentrifuge tube and centrifuged (1 h, 50000 g, 4°C). Supernatant (900  $\mu$ L) corresponding to the cytoplasmic fraction was retained for analysis, and the pellet with total membrane proteins was washed with sonication buffer once, centrifuged (1 h, 50000 g, 4°C) and resuspended in SDS-PAGE buffer. Fractions were precipitated with 4 volumes of acetone and resuspended in equivalent volumes of SDS-PAGE buffer. Protein extracts were separated by SDS-PAGE and analyzed by western blot with anti-FLAG (Sigma-Aldrich #F7425), anti-DNAK (Abcam #ab69617), anti-maltose binding protein (MPB) (New England Biolabs #E8032L) and anti-outer membrane protein (OmpA) antibodies.

### Microscopy

For time-lapse microscopy, LB agar pads were prepared by cutting a rectangular piece out of a double-sided adhesive tape which was taped onto a microscopy slide as described previously ([Bayer-Santos et al., 2019](#)). *E. coli* DH5 $\alpha$  harboring pBRA SP-Tide1 was spotted onto 1.5% LB agar pads supplemented either with 0.2% D-glucose or 0.2% L-arabinose and antibiotics. Phase contrast images were taken every 15 min for 24 h using a Zeiss AxioVert.Z1 microscope fitted with an AxioCam MRm camera and an  $\alpha$  Plan-Apochromat 63x oil objective. Images were analyzed using FIJI software ([Schindelin et al., 2012](#)). To quantify the percentage of dividing or non-dividing cells and the cell doubling time, approximately 100 cells were analyzed in each condition. To determine cell length, approximately 30 cells were measured at each time point (from 30 min to 4.5 h), totaling 150 measurements. To visualize membrane labeling, overnight cultures of *E. coli* carrying pBRA SP-Tide1 were harvested by centrifugation (3 min, 8000 g), washed twice with LB and normalized to OD<sub>600nm</sub> 0.5. Membrane dye FM 4-64 (Molecular Probes) were mixed to cells at 1:1 and 4  $\mu$ L were spotted onto 1.5% LB agarose pads supplemented with 0.2% D-glucose or 0.2% L-arabinose. After 20 h, cells were imaged using a Zeiss AxioVert.A1 microscope fitted with an AxioCam ICm1 camera and a FLUAR 100x oil objective. To analyze FtsZ localization, overnight cultures of *E. coli* FtsZ-mVenus expressing pBRA SP-Tide1 were diluted 1:100 and grown in liquid LB media to an OD<sub>600nm</sub> 0.2. Cells were harvested by centrifugation (3 min, 8000 g), washed twice with LB and added 0.2% D-glucose or 0.2% L-arabinose. At OD<sub>600nm</sub> 0.5 cells were harvested, washed twice with PBS and 4  $\mu$ L were spotted onto 1.5% PBS agarose pads supplemented with 0.2% D-glucose or 0.2% L-arabinose. Cells were imaged using a Zeiss AxioVert.A1 microscope fitted with an AxioCam ICm1 camera.

### Recombinant protein expression and purification

*E. coli* SHuffle cells expressing pET28a-Tide1 were subcultured in LB and grown at 37°C to OD<sub>600nm</sub> 0.7 prior to induction with 0.4 mM IPTG for 16 h at 16°C (150 rpm). Cells were harvested by centrifugation, resuspended with buffer (20 mM Tris-HCl pH 7.35, 200 mM NaCl) and lysed by 10 passages in a French Press system. The lysate soluble fraction was loaded onto a 5 mL HiTrap chelating HP column (GE Healthcare) immobilized with 100 mM cobalt chloride and equilibrated with the lysis buffer. After the removal of unbound proteins, the fusion protein was eluted with lysis buffer supplemented with 400 mM imidazole. Purified proteins were concentrated in Amicon Filter Units (Millipore) before purification by size exclusion chromatography using a HiLoad 26/600 Superdex 75 column (GE Healthcare).

### Peptidoglycan purification and enzymatic assays

Peptidoglycan was purified as described by [Mesnage et al. \(2008\)](#) with small modifications. Briefly, bacterial cells were grown in 1 L of LB to OD<sub>600nm</sub> 0.7 and harvested by centrifugation (15 min, 4000 g, 15°C). Pellets were washed with PBS and resuspended in 20 mL of PBS. Cell suspensions were added dropwise to a glass flask containing 80 mL of boiling 5% SDS and incubated for 30 min. Lysates were washed 4 times with water and treated with pronase 2 mg/mL overnight at 60°C. The following day, 1% SDS was added, incubated for 10 min at 95°C and washed 4 times with water. Muropeptides were obtained by digestion with mutanolysin, followed by reduction with sodium borohydride in borate buffer and separation in reverse-phase high performance liquid chromatography (RP-HPLC) ([Mesnage et al., 2008](#)). Monomeric and dimeric substrates were purified from reduced disaccharide-peptides obtained from the *E. coli*  $\Delta$ 6ldt mutant ([Meberg et al., 2001](#)). Briefly, peptidoglycan was digested by mutanolysin and muropeptides reduced prior to their separation by RP-HPLC using a water-acetonitrile gradient. The fragments corresponding to the monomer GM-tetrapeptide and dimer GM-tetrapeptide-GM-tetrapeptide were collected and freeze-dried, later their concentrations were estimated by nuclear magnetic resonance (NMR) using trimethylsilylpropanoic acid (TSP) as a reference. For *in vitro* assays, 10  $\mu$ M of purified recombinant Tide1 protein was added to 100  $\mu$ M of monomeric or dimeric GM-tetrapeptide in buffer containing 50 mM Tris pH 7.5 and 50 mM NaCl and incubated for 4 h at 37°C. Digestion products were analyzed by RP-HPLC-MS as described previously ([Rodríguez-Rubio et al., 2016](#)). For the amino acid exchange assay, reactions were performed in the same conditions with the addition

of 1 mM D-methionine. For the mucopeptide profile analysis, *E. coli* carrying empty pBRA, pBRA SP-TIde1 and pBRA SP-TIde1<sub>C131A</sub> were used. Cells were grown overnight, subcultured 1:100 in 250 mL of LB and grown until OD<sub>600nm</sub> 0.5 with 50 μg/mL streptomycin and 0.2% D-glucose. Cells were washed twice with 20 mL of LB, added 0.2% L-arabinose and incubated for 3 h until peptidoglycan was extracted as described above. MS data were analyzed with MassHunter software (Agilent Technologies).

### Bioinformatic analysis

Iterative profile searches using JackHMMER (Potter et al., 2018) with a cutoff e-value of  $10^{-6}$  and a maximum of twenty iteration were performed to search for similar sequences in the non-redundant (nr) protein database from the National Center for Biotechnology Information (NCBI). Similarity-based clustering of proteins was carried out using MMseqs software (Steinegger and Söding, 2017). Sequences alignments were produced with MAFFT local-pair algorithm (Kato et al., 2005) and non-informative columns were removed with trimAl software (Capella-Gutiérrez et al., 2009). Approximately-maximum-likelihood phylogenetic tree were built using FastTree 2 (Price et al., 2010). Proteins were annotated using the Pfam database (El-Gebali et al., 2019) and protein secretion systems were identified using models from TXSSdb (Abby et al., 2016) and the HMMER package (Potter et al., 2018). To collect the neighborhood of the genes of interest an in-house python script was used based on information downloaded from the complete genomes and nucleotide sections of the NCBI database.

### QUANTIFICATION AND STATISTICAL ANALYSES

Statistical tests, number of events quantified, standard deviation of the mean, and statistical significance is reported in figure legends. Statistical analysis has been performed using GraphPad Prism5 software and statistical significance is determined by the value of  $p < 0.05$ .