The *Pseudomonas aeruginosa* T6SS Delivers a Periplasmic Toxin that Disrupts Bacterial Cell Morphology

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**In Brief**
The bacterial type VI secretion system (T6SS) delivers effector proteins into prokaryotic and eukaryotic cells to enhance the survival of the donor cell. Wood et al. describe an antibacterial T6SS toxin family eliciting a profound cell division defect and lysis. The structure of this periplasmic-acting toxin reveals a metallopeptidase fold.

**Highlights**
- The structure of the VgrG2b C-terminal domain presents a metallopeptidase fold
- VgrG2b exerts antibacterial activity in the periplasmic space
- Toxicity of VgrG2b is counteracted by a cognate periplasmic immunity protein
- VgrG2bC-ter-intoxicated prey cells bleb at the midcell and lyse

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SUMMARY

The type VI secretion system (T6SS) is crucial in interbacterial competition and is a virulence determinant of many Gram-negative bacteria. Several T6SS effectors are covalently fused to secreted T6SS structural components such as the VgrG spike for delivery into target cells. In *Pseudomonas aeruginosa*, the VgrG2b effector was previously proposed to mediate bacterial internalization into eukaryotic cells. In this work, we find that the VgrG2b C-terminal domain (VgrG2bC-ter) elicits toxicity in the bacterial periplasm, counteracted by a cognate immunity protein. We resolve the structure of VgrG2bC-ter and confirm it is a member of the zinc-metallopeptidase family of enzymes. We show that this effector causes membrane blebbing at midcell, which suggests a distinct type of T6SS-mediated growth inhibition through interference with cell division, mimicking the impact of ß-lactam antibiotics. Our study introduces a further effector family to the T6SS arsenal and demonstrates that VgrG2b can target both prokaryotic and eukaryotic cells.

INTRODUCTION

Protein secretion systems are used by bacteria to interact with other organisms and exploit the local environment (Filloux, 2011). Proteins transported by these systems shape the behavior of polymicrobial communities and interfere with eukaryotic cells. The type VI secretion system (T6SS) delivers effector proteins into both bacterial and eukaryotic targets in a contact-dependent manner (Ho et al., 2014; Jiang et al., 2014) and has been proposed to be involved in the acquisition of public goods such as metal ions (Lin et al., 2017; Si et al., 2017a, 2017b). Although the T6SS can manipulate signaling pathways and the cytoskeleton of eukaryotes (Aubert et al., 2016; Chen et al., 2017; Hachani et al., 2016), it appears to be predominantly involved in bacterial antagonism, in which toxin delivery into neighboring cells elicits growth inhibition and cell death (Allsopp et al., 2017; Hachani et al., 2014; MacIntyre et al., 2010; Russell et al., 2014).

The T6SS is a contractile injection system anchored in the bacterial envelope, evolutionarily related to bacteriophage tails (Leiman et al., 2009; Pukatzki et al., 2007). A conformational change of the T6SS baseplate platform is thought to trigger the contraction of a cytoplasmic sheath, expelling a spear-like structure to puncture a target cell membrane (Cianfanelli et al., 2016; Salih et al., 2018; Wang et al., 2017). The spear is composed of a stack of Hcp rings capped with a spike complex of a trimer of VgrG proteins and a PAAR protein tip (Renault et al., 2018; Shneider et al., 2013). The trimeric VgrG complex shares extensive structural homology with the bacteriophage gp27-gp5 tail spike, which also serves as a membrane puncturing device (Hachani et al., 2011; Leiman et al., 2009; Pukatzki et al., 2007). The spear is decorated with cargo effector proteins, often associating through non-covalent interactions with the structural components (Cianfanelli et al., 2016; Jiang
et al., 2019; Shneider et al., 2013). Besides these cargo effectors, effector domains can be fused to a Hcp, VgrG, or PAAR protein. Examples of these specialized effectors, also called evolved spear components, include the NAD(P)⁺-hydrolyzing toxin Tse6 from *Pseudomonas aeruginosa* or the peptidoglycan hydrolase extension of VgrG3 VC of *Vibrio cholerae* (Brooks et al., 2013; Hachani et al., 2014; Whitney et al., 2015).

*P. aeruginosa* possesses three T6SSs, designated H1-, H2-, and H3-T6SS (Mougous et al., 2006). Although the H1-T6SS is well characterized as an antibacterial weapon, our understanding of the effectors secreted by the H2-T6SS is still in its infancy. Two phospholipases, PldA and Tle4, are secreted by this system into both bacterial and eukaryotic cells and have consequently been designated trans-kingdom effectors (Jiang et al., 2014, 2016). In addition, the antibacterial nuclease effector TseT and the evolved VgrG2b have been linked to this system (Burkinshaw et al., 2018; Sana et al., 2015). VgrG2b is involved in the internalization of *P. aeruginosa* PAO1 into epithelial cells, because uptake of a vgrG2b mutant is decreased versus the wild-type strain (Sana et al., 2015). Infection of epithelial cells expressing vgrG2b displays enhanced internalization not only of wild-type *P. aeruginosa* but also of an H2-T6SS-deficient strain. Elements of the host cytoskeleton are co-opted for invasion, because chemical inhibition of actin and microtubule polymerization blocks bacterial uptake. Furthermore, interactome analysis of VgrG2b ectopically expressed in host cells identified components of the γ-tubulin ring complex (γTuRC) as binding partners of this evolved spike protein. Despite the microtubule network being implicated in *P. aeruginosa* invasion, the mechanism of this process remains to be understood.

In this study, we report that VgrG2b represents an evolved trans-kingdom T6SS effector. We provide evidence that VgrG2b is directly secreted by the H2-T6SS and that the VgrG2b C-terminal domain (VgrG2bC-ter) possesses antibacterial activity. We show that this domain, the structure of which we present in this work, is a member of a widespread family of metallopeptidases eliciting toxicity in the bacterial periplasm and that it can be neutralized by a cognate immunity protein. The toxicity of VgrG2bC-ter results in profound morphological anomalies characterized by blebbing of the bacterial membrane at the site of septation, a phenotype reminiscent of inhibition of the cell division machinery by β-lactam antibiotics.

**RESULTS**

**VgrG2b Is an Evolved VgrG Protein Secreted by the H2-T6SS**

VgrG2b is encoded in the *hcpC* locus with the hypothetical protein PA0261 and the effector-immunity module Tle3-Tli3 (type VI lipase effector 3-type VI lipase immunity 3) (Figure 1A), where Tle3 is a predicted phospholipase (Barret et al., 2011; Wood et al., 2019). As such, it is not genetically linked to any T6SS core gene clusters. Beyond its canonical trimeric spike-forming...
region containing gp27- and gp5-like domains, VgrG2b harbors a C-terminal extension, rendering it an evolved VgrG protein (Figure 1B). VgrG2b also contains a DUF2345 domain, predicted to adopt a β-helical fold similar to the gp5-like domain (Figure 1C), which can be considered an extension of the spike region (Sana et al., 2015). The C-terminal portion of the VgrG2b spike protein is predicted to function as a metallopeptidase because of the presence of a signature zinc-binding HEXXH motif (Figure 1B). Bioinformatic analysis of the linker region between the DUF2345 and the metallopeptidase domains reveals a transthyretin (TTR)-like fold, implicated in protein-protein interactions. Indeed, C-terminal TTR folds have been described in PAAR and VgrG proteins, with that of VgrG1 in enteroaggregative *Escherichia coli* empirically shown to be important in the delivery of the cargo effector Tle1 (Flaugnatti et al., 2016; Shneider et al., 2013).

A previous study determined that VgrG2b and the H2-T6SS are required in invasion of epithelial cells during infection, but no H2-T6SS-dependent secretion of VgrG2b was effectively demonstrated (Sana et al., 2015). We have recently described in vitro conditions in which the H2-T6SS is active and established that the Hcp2 tail tube protein, encoded by at least one of the identical paralogs hcpA, hcpB, hcpC, and hcp2 in *P. aeruginosa* PA14, is secreted by this system (Allsopp et al., 2017). Here, we employed antibodies raised against peptides within the metallopeptidase domain of VgrG2b to probe for this spike protein in the supernatant of *P. aeruginosa* cultures grown in H2-T6SS-conducive conditions. VgrG2b and Hcp2 are detected in the extracellular milieu of the parental strain (PAO1ΔrsmA); however, deletion of the tssE2 gene, encoding an H2-T6SS baseplate component, abolishes their secretion (Figure 1D). Complementation with tssE2 in trans partially restores secretion of VgrG2b and Hcp2.

Altogether, this confirms that VgrG2b is secreted in an H2-T6SS-dependent manner. Although the H2-T6SS locus of *P. aeruginosa* PAO1 lacks genes encoding the spike and tube proteins of the T6SS spear assembly, this result highlights that the H2-T6SS supports the secretion of several of distally encoded effectors (Barret et al., 2011; Hachani et al., 2011).

**Crystal Structure of VgrG2b C-Terminal Effector Domain**

Next, we purified VgrG2bC-ter (residues 770–1,019) to initiate structural analysis. We solved the structure of VgrG2bC-ter, encompassing residues 833–1,019, using crystals of both the native and the selenomethionine-substituted forms with resolutions of 3.2 and 3.0 Å, respectively (Figure 2A). Two copies of VgrG2bC-ter were found in the asymmetric unit of the crystal, which could be superimposed with a root-mean-square deviation (RMSD) of 0.1 Å (Tables S1 and S2). In solution, however, the purified metallopeptidase domain exists predominantly as a monomer, as determined by size-exclusion chromatography-multi-angle laser light scattering (SEC-MALLS), with a dimeric species also present (Figure S1A). It is likely that the physiologically relevant oligomerization state of the metallopeptidase domain is achieved only in the full-length VgrG2b protein.

The VgrG2bC-ter protomer possesses a single domain composed of six α helices and three β strands (Figure 2A), forming a shallow bowl with multiple small loops. A PDB search using the DALI server identified several thermolysin-like metallopeptidases as structural homologs of the VgrG2bC-ter effector domain (Holm and Laakso, 2016). One such metallopeptidase is human angiotensin-converting enzyme 2 (Z score: 5.0, Cα RMSD of 4.2 Å across 105 residues), with the active sites of these proteins displaying similar architectures (Figure 2B). Although metallopeptidases target diverse substrates, the overall core fold of three α helices (α2, α3, and α5) and two β strands (β2 and β3) is highly conserved and bears the classical HEXXH catalytic motif typical of these zinc-dependent enzymes, where X is any amino acid (Figure 2B).

Indeed, the conformations of the HEXXH residues in our structure are suggestive of metal ion coordination, and the geometry...
suggests that the electron density in the putative metal-binding site corresponds to a divalent cation such as zinc. To confirm that VgrG2b C-ter binds zinc, we employed differential scanning fluorimetry to assess the thermodynamic stabilization of the metallopeptidase in the presence of zinc ions. The melting temperature of the domain increases from 46.3°C in the absence of zinc to a maximum of 55.1°C in the presence of an eight-fold excess of the metal ion (Figure S1B), providing a strong indication of the binding of zinc to the metallopeptidase. The assigned zinc ion is coordinated by the two histidine residues of the catalytic triad and has a third glutamate ligand (E983) at the top of helix a5 (Figure 2B). These features permit classification of VgrG2b C-ter as a thermolysin-like gluzincin within the MA(E) metallopeptidase subclan (Rawlings et al., 2018).

**VgrG2b C-ter Is Part of a Wider Family of Metallopeptidase Effectors**

We assessed the prevalence of the VgrG2b C-ter metallopeptidase domain by exploring its phylogenetic distribution. Homologous metallopeptidases were found throughout the proteobacterial phylum, especially in bacteria possessing a T6SS (Li et al., 2015) (Figure 3A). Intriguingly, the presence of VgrG2b C-ter-like proteins in Sphingomonadaceae, which rarely encode a T6SS, implies that this protein family may have a role beyond constituting a T6SS effector. Nevertheless, many orthologs are encoded within T6SS gene clusters and are classified as DUF4157 domain-containing putative metallopeptidases, found as both putative cargo and specialized effectors. For example, *Burkholderia pseudomallei* encodes an evolved VgrG with this...
metallopeptidase domain, similar to VgrG2b. Yet in Collimonas and Alcanivorax species, it is grafted as an extension of a PAAR protein. Finally, many members of the Enterobacteriaceae family code for this metallopeptidase as a putative T6SS cargo effector (Figures 3B and 3C), underscoring the modularity of T6SS effector proteins.

Further analysis of the distribution of vgrG2b genes uncovered evidence for a remarkable recombination event in the P. aeruginosa BL21 strain, in which the tie3-tli3 module is duplicated within the 3′ end of the vgrG2b gene (Figure 3D). This has given rise to a canonical vgrG gene encoding VgrG2b up to its TTR fold, with the metallopeptidase domain encoded by a separate open reading frame (ORF) downstream of the phospholipase catalytic domain. This genetic reorganization also implies that vgrG genes can acquire or lose their 3′ end through recombination, allowing the exchange of effector domains at the VgrG tip. This is reminiscent of observations concerning C-terminal toxin modules in systems such as contact-dependent growth inhibition (CDI) and Rhs proteins (Ruhe et al., 2017; Wang et al., 1998). Overall, these in silico analyses suggest that VgrG2bC-ter is frequently associated with proteobacterial T6SS loci and that its secretion can be achieved via various effector configurations.

VgrG2b-PA0261 Constitutes an Antibacterial Effector-Immunity Pair

Our bioinformatic analysis of the genetic context of T6SS-associated VgrG2bC-ter orthologs revealed intriguing synteny of this effector gene with a small ORF, often unannotated and encoded downstream, such as PA0261 in P. aeruginosa (Figure 1A), BTH_I2696 in Burkholderia thailandensis, ACIAD0054 in Acinetobacter baylyi, SARI_02726 in Salmonella enterica subs. arizonae, or BST_02183 in Alcanivorax dieselolei (Figure 3C). The juxtaposition of these genes is invariant, prompting the hypothesis that P. aeruginosa VgrG2bC-ter may constitute an antibacterial effector-immunity pair with the PA0261 gene product. Indeed, T6SS effectors whose genes are systematically encoded in tandem with a small ORF are frequently found to exert antibacterial activity, with the adjacent gene encoding its cognate immunity protein (English et al., 2012; Hood et al., 2010; Russell et al., 2011).

We engineered a strain lacking the vgrG2b-PA0261 genes and placed it in competition with the parental strain under conditions in which the H2-T6SS is active. Whereas competition of the parent with itself results in competitive parity, the ΔvgrG2bPA0261 strain exhibits a significant growth disadvantage (Figure 4A). VgrG2b is responsible for the advantage of the parental strain, because a ΔvgrG2b attacker no longer outcompetes the ΔvgrG2bPA0261 prey. Importantly, the attacker and prey strains are isogenic; thus, the vgrG2b-PA0261 module represents the sole difference between these strains. As such, the prey strain is immune to the action of all other T6SS toxins such as Tle3, encoded downstream of PA0261 (Figure 1A). Deletion of tssE2 in the attacker strain, rendering the H2-T6SS non-functional, restores competitive parity of the prey strain.
confirming the H2-T6SS-dependent secretion of VgrG2b (Figure 4A), whereas complementation of tssE2 on a plasmid partially restores the growth advantage of the attacker strain. Furthermore, a P. aeruginosa strain lacking the immunity gene exhibits a growth defect on solid media, but not in liquid culture, consistent with contact-dependent delivery by the H2-T6SS (Figures S2A and S2B).

To assess the role of PA0261 as the immunity protein neutralizing VgrG2b-mediated toxicity, we introduced the PA0261 gene into the ΔvgrG2bPA0261 strain in trans. Constitutive expression of PA0261 in the prey strain abolishes the competitive growth advantage of the attacker, whereas the empty vector provides no such immunity (Figure 4B). Bioinformatic analysis of the PA0261 sequence revealed the presence of a putative N-terminal signal peptide, likely directing the protein into the periplasm (Figure S2C). Provision of PA0261 without the sequence encoding this N-terminal portion (PA0261(no SP)) fails to prevent elimination by the parent (Figure 4B), suggesting that the predicted extracytoplasmic localization of this protein is needed to neutralize VgrG2b, similar to the immunity proteins of other P. aeruginosa periplasmic-acting toxins (Jiang et al., 2016; Russell et al., 2011).

If PA0261 requires its putative N-terminal signal peptide to act as an immunity protein, VgrG2b must elicit its toxicity beyond the cytoplasmic membrane. We ectopically produced VgrG2bC-ter in E. coli and targeted the domain to the periplasm by fusing it to an N-terminal signal peptide from the Tat-dependent substrate TorA (Palmer and Berks, 2012; Santini et al., 2001). Production of VgrG2bC-ter is not detrimental to E. coli when the protein remains in the cytoplasm, because this strain grows similarly to vector control strain (Figure 4C). However, this domain exhibits acute toxicity when targeted to the periplasmic compartment by the Tat-dependent signal peptide. Crucially, substitution of the putative catalytic glutamic acid residue to an alanine (E936A from HEXXH) renders VgrG2bC-ter non-toxic in the periplasm, restoring growth. No differences in the levels of VgrG2bC-ter production are observed between the periplasmic-targeted toxin and its inactive isoform (Figure S2D), indicating that the putative enzymatic activity of the VgrG2bC-ter metallopeptidase is responsible for toxicity. Mutation of the two histidine residues (H935A and H939A) coordinating the zinc ion in our structure also abolish periplasmic toxicity, suggesting our designation of the HEXXH catalytic triad (Figure S2E). Interestingly, although raising the toxicity of media has been reported to rescue growth of bacteria producing the periplasmic-acting T6SS effectors Tse1 and Tse3, the addition of 0.5 M sucrose does not suppress the growth defect of E. coli producing VgrG2bC-ter in the periplasm (Russell et al., 2011) (Figure S2F). This suggests that VgrG2bC-ter may elicit an activity distinct from the D,L-endopeptidase and muramidase activities of Tse1 and Tse3, respectively (described later).

We sought to determine whether toxicity was displayed by orthologs of VgrG2bC-ter and expressed the putative T6SS cargo effector ACIAD0053 from A. baylyi similarly. Surprisingly, induction of periplasmic-targeted ACIAD0053 expression does not hamper E. coli growth under identical conditions to that of intoxication by VgrG2bC-ter (Figure S2G). Nevertheless, reduction of the osmolarity of the medium (low-salt LB [LB-LS]) to increase bacterial susceptibility to lysis results in a drastic reduction in growth of the strain expressing pTat-ACIAD0053. Once more, alanine substitution of the residues of the catalytic triad (ACIAD0053) produces a non-toxic isoform (Figures S2G and S2H). These results indicate that members of the VgrG2bC-ter metallopeptidase effector family act as potent periplasmic antibacterial toxins.

**Functional Characterization of the PA0261 Immunity Protein Family**

To validate the toxin-immunity relationship of the VgrG2bC-ter~ PA0261 pair, we investigated whether PA0261 could protect E. coli from VgrG2bC-ter-mediated toxicity. Coexpression of the immunity gene relieves the growth defect caused by the metallopeptidase (Figure 5A). As a specificity control, we expressed tli4, encoding the cognate immunity protein of the antibacterial H2-T6SS-dependent effector Tle4, which was unable to restore bacterial viability (Jiang et al., 2016). Immunoblot analysis determined that both immunity proteins are produced (Figure S3A). This demonstrates that PA0261 is the cognate immunity determinant of the VgrG2bC-ter toxin. Furthermore, the additional presence of a higher molecular weight band for PA0261-HA is indicative of the precursor form of a periplasmic protein, supporting the predicted localization of this immunity protein, which we later confirm (Figures S3A and S3A). We then used far-western dot blotting (Figure 5B) to investigate whether complex formation with the metallopeptidase domain was the neutralization mechanism of PA0261. We spotted recombinant VgrG2bC-ter (Figure S3B) or the C-terminal TTR domain of VgrG4b as bait proteins on nitrocellulose membrane and incubated them with a bacterial lysate overproducing hemagglutinin (HA)-tagged immunity proteins PA0261 or Tli3. Immunoblotting shows that despite production of both immunity proteins, only PA0261-HA binds to VgrG2bC-ter, and not to VgrG4bC-ter, suggesting a specific interaction with the metallopeptidase domain (Figure 5B). Furthermore, PA0261 binds the inactive metallopeptidase variant VgrG2bC-ter (E936A), suggesting that the inactive variant is fully folded (Figures 5B and S3C). This implies that PA0261 neutralizes VgrG2bC-ter through complex formation, as has been shown for other biochemically characterized T6SS toxin-immunity pairs (Russell et al., 2011; Shang et al., 2012).

In general, immunity proteins of T6SS toxins display little sequence identity with their orthologs, likely because of selective pressure solely on the conservation of toxin binding (Russell et al., 2012). This phenomenon is also apparent in the PA0261 family; however, a multiple sequence alignment of homologous immunity proteins revealed the presence of cysteine residues in partially conserved positions (Figure S4). Because PA0261 is a predicted periplasmic protein, the conservation of cysteine residues engenders the hypothesis that disulfide bond formation may be required for the function of the protein in the oxidizing environment of this compartment. To begin answering this question, we demonstrated through subcellular fractionation that PA0261 is found in the periplasm when expressed in E. coli (Figure S5A). We assessed the ability of PA0261 to protect against the toxicity of periplasmic VgrG2bC-ter in the absence of dsbA, encoding the major oxidoreductase of the disulfide bond formation (Dsb) system in E. coli (Figure 5C). The levels of PA0261 in the ΔdsbA mutant are greatly diminished, hampering the growth
of this strain in the presence of VgrG2bc-ter (Figures 5C and 5D).

This indicates that the Dsb system contributes to the protection of bacterial cells from T6SS-mediated antagonism, in juxtaposition to a recent study demonstrating that a lack of DsbA in prey cells can indirectly confer protection against T6SS attack because of the improper folding of delivered toxins (Mariano et al., 2013; English et al., 2018). Crystal structures of immunity proteins such as those of the Tai4 family have also revealed the presence of disulfide bonds, extending the relevance of our observation to other families of T6SS immunity proteins (Benz et al., 2013; English et al., 2018). Crystal structures of immunity proteins such as those of the Tai4 family have also revealed the presence of disulfide bonds, extending the relevance of our observation to other families of T6SS immunity proteins (Benz et al., 2013; English et al., 2018).

In silico examination of putative immunity proteins to members of the VgrG2bc-ter family suggests that like PA0261, they are targeted to the cell envelope. However, the mechanisms of attaining this localization seem to be diverse. In Enterobacteriaceae, PA0261 homologs are predicted to be lipoproteins (Figure S5B), whereas ACIA00054, the putative immunity protein to ACIA00053 in A. baylyi, appears to bear hallmarks of an N-terminal transmembrane helix (Figure S4, top line). Irrespective of these differences, in both cases the bulk of the immunity protein would be exposed in the periplasm. The second residue following the lipobox cysteine residue of the predicted immunity lipoproteins in Enterobacteriaceae is often a glycine, likely directing the protein to the outer membrane rather than constituting an avoidance signal for the lipoprotein outer membrane localization (Lol) pathway as described by the +2 rule (Figure S5B) (Narita and Tokuda, 2017; Yamaguchi et al., 1988).

Indeed, membrane fractionation of the E. coli cell envelope using selective detergent treatment showed that the PA0261 ortholog from S. arizonae, SARI_02726, is enriched in the outer membrane fraction (Figure S5C). Likewise, the A. baylyi ortholog ACIA00054 was enriched in the inner membrane fraction as predicted (Figure S5D). Determination of the localization of members of this immunity protein family supports the designation of VgrG2bc-ter-PA0261 as a periplasmic effector-immunity pair while highlighting the diversity of strategies employed by the immunity determinants to gain access to the periplasm, as summarized in Figure S5E.

VgrG2bc-ter Is Bacteriolytic but Does Not Display Peptidoglycan Hydrolase Activity

Next, we investigated the activity of the VgrG2bc-ter effector domain. To determine whether VgrG2bc-ter induces growth arrest or lysis of target bacteria, we employed an outside-in approach, permeabilizing E. coli cells with a sublethal concentration of polymyxin B and measuring whether the toxin reduces culture turbidity. Although VgrG2bc-ter (E936A) does not decrease the turbidity of the suspension, exogenous addition of lysozyme or VgrG2bc-ter elicits suspension clarification, thereby showing that the metallopeptidase effector challenges cell integrity, having gained access to the periplasm (Figure 6A).
Because this bacteriolytic effector is a periplasmic-acting metallopeptidase, we reasoned that it might target either the peptide bonds in the cell wall or a protein within the cell envelope. We probed the action of the toxin toward the bacterial cell wall with various approaches. In a turbidimetric assay, recombinant proteins were incubated with lyophilized Micrococcus lysodeikticus as a substrate, but while lysozyme reduces turbidity, VgrG2b C-ter does not (Figure S6 A). Similarly, zymography demonstrated that lysozyme produces a zone of clearing in a stained gel impregnated with purified E. coli sacculi, yet no such peptidoglycan degradation was apparent for VgrG2b C-ter (Figure S6 B). An analogous assay for crude protease activity was also used, with a similar negative outcome (Figure S6 C). Finally, quantification of muropeptides released through sacculus hydrolysis using high-performance liquid chromatography (HPLC) revealed that treatment with VgrG2b C-ter does not alter the composition of either tetrapeptide-rich or pentapeptide-rich E. coli cell wall preparations (Figures S6D–S6F). Thus, we conclude that VgrG2b C-ter does not exhibit peptidoglycan hydrolase activity.

VgrG2b Induces Lysis by Apparent Perturbation of Cell Division

Next, we turned to single-cell analysis of E. coli producing the periplasmic-targeted metallopeptidase or its inactive variant from the pTat vector. Bacterial morphology was highlighted by staining the membranes with the fluorescent dye FM1-43. Over the course of 2.5 h, no alterations in rod-like E. coli cell shape were observed in bacteria harboring the empty vector or pTat-vgrG2b C-ter (E936A). However, production of the active periplasmic metallopeptidase causes aberrant cellular morphology characterized by membrane blebbing, leading to cell rounding and lysis (Figure 6B). Periplasmic VgrG2b C-ter is still toxic in high-osmolarity conditions (Figure S2F), suggesting that the effector does not cause lysis through global weakening of the cell wall as with Tse1 and Tse3 (Russell et al., 2011). Figure 6C
shows that blebbing predominately originates at midcell, leading
to the cracking of the rod shape before rounding up, suggesting
a defect in cell division similar to observations during β-lactam
antibiotic administration (Chung et al., 2009; Yao et al., 2012).
This observation of site-dependent perturbation in cell shape is
also in opposition to the notion of VgrG2bC-ter exhibiting generic
peptidoglycan hydrolase activity, unless the metallopeptidase
requires a factor or structure specifically localized at the cell
septum for its activation.

To discount the possibility that this phenotype is an
artifact because of toxin overexpression, we monitored the
morphology of P. aeruginosa cells subject to VgrG2b-mediated
killing in competition assays. Upon attack of the susceptible
ΔvgrG2bPA0261 strain by its parent, we observed similar mem-
brane blebbing, whereas no such morphological aberrations
occurred when the attacker strain lacked a functional H2-T6SS
(ΔtsS2) (Figure 6D). In addition, when the strains were incubated
alone, no blebbing occurred. The blebs produced during elimi-
nation of VgrG2b-sensitive prey again localized to the septum,
indicating that the mechanism of killing by the metallopeptidase
domain is similar whether it is delivered by the T6SS or th
effector is artificially targeted to the periplasm (Figure 6E). These
results demonstrate that VgrG2bC-ter-induced blebbing is a
physiologically relevant phenotype and that the target of this
toxin is conserved between E. coli and P. aeruginosa.

Identification of Putative Interacting Partners of
VgrG2bC-ter

Because characterization of the biochemical activity of
VgrG2bC-ter had been unsuccessful, we turned to affinity purifi-
cation-coupled mass spectrometry to investigate the identities
of putative target proteins that interact with the toxin in the peri-
plasm. A tagged inactive VgrG2bC-ter variant was directed to the
periplasm after which cellular proteins were crosslinked in vivo
using dimethyl 3,3′-dithiobispropionimidate (DTBP). Affinity puri-
fication pulled down the tagged inactive metallopeptidase and
any crosslinked interaction partners. Three independent repli-
cates of the affinity-purified protein samples were subject to
mass spectrometry analysis to identify candidate interactors of
the metallopeptidase. This preliminary analysis found 89 pro-
teins enriched in the tagged sample dataset compared with the
untagged VgrG2bC-ter control, which were present in at least
two of the three replicates. Of these, just 19 proteins (21.3%)
were found with MASCOT scores (representing the probability
of true identification of the protein) higher than the threshold
set to exclude background noise. Twelve of these 19 proteins
(63.2%) are localized in the cell envelope and were subsequently
considered candidate interactors of VgrG2bC-ter (Table S3).
Three of the four highest-scoring candidates (MitC, PBP5,
and PBP6a) are involved in peptidoglycan biogenesis. This suggests
that the metallopeptidase may perturb the complex machinery,
which maintains the cell wall, inhibiting cell division and produc-
ing the observed blebbing at midcell. If VgrG2bC-ter acts through
interference with peptidoglycan remodeling, it may be through
direct cleavage of these identified target proteins; however, we
cannot exclude that the interaction with periplasmic binding
partners stimulates a putative peptidoglycan hydrolase activity
of the toxin. The periplasmic action of VgrG2bC-ter prevents
growth of E. coli strains with single deletions in mitC, dacA
(encoding PBP5), or dacC (encoding PBP6a), indicating that
cleavage of one of these enzymes individually is not sufficient
for toxicity (Figure S7).

The dataset is also enriched with lipoproteins (8/12, or 67%),
raising the possibility that VgrG2bC-ter might interact with the
linker region between the acylated cysteine residue of a lipopro-
tein and its main fold, resulting in disruption of its intrinsic
localization. We monitored the abundance of two lipoproteins found
in the dataset, MitC and LolB, in bacterial membranes after incub-
aturation of purified VgrG2bC-ter or VgrG2bC-ter (E936A) with bacte-
rial lysates. The presence of VgrG2bC-ter elicits a reduction in
membrane-associated MitC or LolB in a manner dependent on
its catalytic activity (Figures 7A and 7B). We examined the distri-
bution of RcsF, an outer membrane lipoprotein that was identi-
fied below the significance cutoff in our pull-down experiments (Table S3), and found that its levels were also diminished in the
membrane fraction in the presence of the toxin (Figure 7C). The abundance of the periplasmic protein TEM-1 was unaltered
in this assay (Figure 7D), suggesting that VgrG2bC-ter may prefer-
entially target lipoproteins; specifically, the toxin may cleave lipid
anchors to release lipoproteins from membranes. Because this assay was conducted with non-physiological levels of both toxin
and tagged lipoproteins, it is not possible to currently state
whether MitC, LolB, or RcsF is a physiological target of VgrG2b
delivered by the T6SS. In all, our data indicate that the
VgrG2bC-ter metallopeptidase represents a hitherto-undescribed
family of antibacterial T6SS effectors that appear to dysregulate
the cell division process.

DISCUSSION

Previous work has proposed that VgrG2b is an anti-host effector
that subverts the cytoskeleton (Sara et al., 2015). VgrG2b inter-
acts with members of the γTuRC, whereas VgrG2a, sharing
99.5% identity with VgrG2b across its spike and DUF2345 do-
mains, does not, implying that the interaction occurs through
the C-terminal extension of VgrG2b, encompassing the predicted
metallopeptidase domain. The precise function of this domain
was not explored, so it is unclear whether its catalytic activity is
required for invasion. In addition, although ectopic expression
of vgrG2b within host cells enhances P. aeruginosa uptake, it is
possible that other bacterial factors are involved in the mecha-
nism, because the internalization of neither inert particles nor
other bacteria was shown. For example, other studies have found
that P. aeruginosa can be internalized by non-phagocytic cells in
a process that requires the H2-T6SS phospholipase effectors PldA
and PldB (Jiang et al., 2014). Interestingly, PldA and PldB also
display antibacterial activity, which necessitates a set of immunity
proteins to avoid self-intoxication, and were coined trans-
kingdom effectors (Jiang et al., 2014; Russell et al., 2013). In
this study, we performed an in-depth characterization of VgrG2b
and propose that it is also a trans-kingdom effector.

The identification of a small ORF downstream of genes encoding
homologs of the VgrG2bC-ter metallopeptidase family is a key
finding in this work. We demonstrated the impact of the VgrG2b
growth at midcell. If VgrG2bC-ter acts through
direct cleavage of these identified target proteins; however, we
cannot exclude that the interaction with periplasmic binding
partners stimulates a putative peptidoglycan hydrolase activity
of the toxin. The periplasmic action of VgrG2bC-ter prevents
metallopeptidase domain on bacterial survival when reaching
the periplasm of prey cells in an H2-T6SS-dependent manner.
The HEXXH motif in its C-terminal domain had led to its designation as a putative zinc-dependent metallopeptidase, and here we present the crystal structure of this domain, which confirms this classification. Our phylogenetic analyses of VgrG2b_C-ter-like proteins also find the corresponding gene within T6SS loci of diverse proteobacteria. It is frequently annotated as DUF4157, which corresponds to a small domain of approximately 80 residues encompassing the HEXXH motif. Our data show that mutations in the catalytic site abrogate the antibacterial activity, although we were unable to identify direct cleavage of a substrate by VgrG2b.

Our characterization of VgrG2b_C-ter as an antibacterial toxin is supported by the recent description of the ACIAD0053 protein from A. baylyi (Ringel et al., 2017). This study showed that an A. baylyi strain lacking ACIAD0053-ACIAD0054 can be eliminated by its parent, leading to the designation of the locus as encoding a type VI metallopeptidase effector-immunity pair, Tpe1-Tpi1. We have additionally demonstrated that similar to VgrG2b_C-ter, Tpe1 exerts its activity in the periplasm, whereas the cognate immunity protein Tpi1 resides in the inner membrane. The ortholog of this immunity determinant in S. arizonae is an outer membrane lipoprotein, but in both cases, the bulk of the immunity protein is exposed to the periplasm; accordingly, we also observe PA0261 of P. aeruginosa localizing to this compartment. It is therefore fitting to propose that the VgrG2b_C-ter-PA0261 family of metallopeptidase effector-immunity pairs constitutes members of the Tpe-Tpi family.

In our study, we revealed the differing ability of VgrG2b_C-ter and the A. baylyi toxin Tpe1 to intoxicate E. coli. Work by Ringel and colleagues demonstrated Tpe1 toxicity in A. baylyi under standard laboratory conditions but failed to see an impact on E. coli (Ringel et al., 2017). We corroborate this observation under standard laboratory conditions but find that a defect in E. coli growth is elicited by Tpe1 in conditions of low salinity, implying that osmolarity, or salt per se, may regulate its action. Recently, a study of effector synergy within a delivered payload indicates that conditional efficacy is a common phenomenon among T6SS toxins; thus, toxin efficiency should be considered in the context of the whole T6SS-delivered payload (LaCourse et al., 2018;...
Division defect mediated by VgrG2b C-ter that we observe appears to reflect a more environmentally relevant condition and imply that VgrG2b C-ter is a more robust enzyme. Salinity is known to regulate periplasmic proteins such as peptidoglycan biosynthetic enzymes, presenting the possibility that these toxins may interact with different members of a family of peptidoglycan remodeling enzymes such as penicillin-binding proteins (PBPs) or lytic transglycosylases (LTases) (Möll et al., 2015; Palomino et al., 2009). Indeed, P. aeruginosa, A. baylyi, and E. coli possess differing quotas of these peptidoglycan hydrolases, rendering species-dependent nuances in substrate targeting an attractive hypothesis (Dik et al., 2017; Pazos et al., 2017).

We observed bulge formation at sites of septation resulting from the activity of VgrG2b C-ter. Recently, a T6SS effector inhibiting bacterial cell division was described in Serratia proteamaculans, for which ADP ribosylation of the essential cell division protein FtsZ prevents Z-ring formation, causing bacterial elongation because of inhibition of septation (Ting et al., 2018). The cell division defect mediated by VgrG2b C-ter that we observe appears to be distinct, characterized by midcell blebbing. This is reminiscent of the inhibition of the cell wall biosynthesis machinery by β-lactam antibiotics, in which the balance of peptidoglycan growth and degradation is disrupted, causing severe morphological aberrations and cell lysis (Cho et al., 2014; Dik et al., 2019; Templin et al., 1992). Class A PBPs possess transpeptidase (TPase) and glycosyltransferase activities to incorporate new monomers into the sacculus, and these biosynthetic processes are coordinated with the degradative action of enzymes such as LTases, amidases, and carboxypeptidases (CPases), which modify the peptidoglycan to regulate its growth and permit septation of daughter cells (Egan et al., 2017). These enzymes form large complexes, whose subcellular localization often regulates their activities, and dysregulation of this process usually results in lysis. For example, the recruitment to midcell and subsequent activity of the TPase LdtD requires the concerted actions of PBPs, PBP6a, and PBP1b to facilitate septation under certain stress conditions, the latter of which is also associated with LTases MltA and MltG and is recruited to the septum by the lipoprotein LpoB (Hugonnet et al., 2016; Montón Silva et al., 2018; Moré et al., 2019; Paradis-Blieu et al., 2010; Typas et al., 2010; Vollmer et al., 1999; Yunck et al., 2016). Similarly, perturbation of the localization of EnvC and NlpD to the division site, which controls the activity of amidases required for separation of daughter cells, results in septation failure (Tsang et al., 2017; Uehara et al., 2009, 2010).

Consequently, it is thought that functional redundancy may help to insulate this crucial process from environmental insults, and correspondingly, individual deletions in LTases and many PBP genes often do not produce phenotypic changes, because the enzymes display partially overlapping functions for robust control of cell division in changing conditions (Denome et al., 1999; Heidrich et al., 2002; Pazos et al., 2017; Peters et al., 2016). Our pull-down assays identified the membrane-associated LTase MltC and the D,D-CPases PBP5 and PBP6a as possible interaction partners of VgrG2b C-ter, yet individually they are dispensable for effector-mediated toxicity, potentially as a consequence of built-in redundancy. The precise biochemical activities and functions of these peptidoglycan biosynthetic enzymes have only started to be unraveled (Artola-Reclons et al., 2014; Dijkstra and Keek, 1996; Meirsonne et al., 2017; Peters et al., 2016), so further work is required to establish their relevance during intoxication by the T6SS.

We found that VgrG2b C-ter depletes several lipoproteins from membranes, implying broad target specificity by associating with the lipidated linker region of these proteins. We speculate that the activity VgrG2b C-ter may disrupt the localization of lipoproteins associated with cell envelope maintenance such as membrane-associated LTases, the cell wall anchoring protein Lpp, or components of the Lom and Bam machineries, thereby perturbing bacterial cell division. Targeting of lipidated substrates has been described for bacterial effectors of the type III secretion system, namely, YopT of Yersinia enterocolitica and IpaJ of Shigella flexneri. These proteases are delivered into host cells, where they cleave the isoprenyl and N-myristoyl anchors of small guanosine triphosphatases (GTPases), respectively, to disrupt host cell signaling and facilitate bacterial survival (Bunaevskiy et al., 2013; Shao et al., 2002). Indeed, these effectors target specific families of GTPases during infection, rather than a single protein; thus, the notion of target promiscuity for VgrG2b C-ter has an intriguing precedent (Bunaevskiy et al., 2015; Fueller and Schmidt, 2008).

In summary, we find that besides its described role in invasion of eukaryotic cells, the evolved VgrG2b spike protein of P. aeruginosa has a direct role in bacterial antagonism. Given the vast dissimilarity between host microtubule manipulation and bacteriolytic activity, reconciliation of VgrG2b metallopeptidase dual role warrants further work to shed light on the activity of this effector within eukaryotic systems. In prokaryotes, the C-terminal metallopeptidase domain elicits acute toxicity when targeted to the periplasmic space, resulting in dysregulation of bacterial morphology and cell lysis. The prevalence of this family of T6SS effector proteins suggests conservation of its target, so elucidation of its antibacterial mechanism may help progress both our knowledge of bacterial cell division and how to target this process with therapeutic compounds.

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.2019.08.094.

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AUTHOR CONTRIBUTIONS

T.E.W. and A. Filoux conceived the study and wrote the manuscript; T.E.W., S.A.H., A. Förster, L.M.N., E.M., N.P.B., and H.C.L.Y. performed experiments; and A.H., R.D.H., J.C.W., W.V., and P.S.F. helped with experiment design.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Mouse monoclonal anti-RpoB | Biolegend | Cat#663006; RRID: AB_2565555 |
| Rabbit polyclonal anti-Hcp2 | Jones et al., 2014 | N/A |
| Rabbit polyclonal anti-VgrG2b | Sana et al., 2015 | N/A |
| Rabbit polyclonal anti-LasB | Gift from Romé Vouloux | N/A |
| Rabbit polyclonal anti-VgrG4b | Allsopp et al., 2017 | N/A |
| Mouse monoclonal anti-Myc | Millipore | Cat#05-724; RRID: AB_309938 |
| Mouse monoclonal anti-HA | Biolegend | Cat#MMS-101P; RRID: AB_2314672 |
| Mouse monoclonal anti-TEM-1 | QED Bioscience | Cat#15720; RRID: AB_129940 |
| Rabbit polyclonal anti-DesbA | Gift from Despoina Mavridou | N/A |
| Rabbit polyclonal anti-SecA | Gift from Jan Tommassen | N/A |
| Rabbit polyclonal anti-PhoE | Gift from Jan Tommassen | N/A |
| **Bacterial and Virus Strains** |        |            |
| Pseudomonas aeruginosa PAO1 | Dieter Haas Laboratory | N/A |
| P. aeruginosa PAO1ΔrsmA | This paper | N/A |
| P. aeruginosa PAO1ΔrsmAΔtssE2 | Wettstadt et al., 2019 | N/A |
| P. aeruginosa PAO1ΔrsmAΔvgrG2b | Wood et al., 2019 | N/A |
| P. aeruginosa PAO1ΔrsmAΔvgrG2bPA0261 | This paper | N/A |
| P. aeruginosa PA14 | Lee et al., 2006 (Whitney lab wild type) | N/A |
| P. aeruginosa PA14ΔrsmAΔamrZ | This paper | N/A |
| P. aeruginosa PA14ΔrsmAΔamrZΔPA14_03210 (PA0261PA14) | This paper | N/A |
| P. aeruginosa PA14ΔrsmAΔamrZΔPA14_03210-20 (vgrG2bPA0261PA14) | This paper | N/A |
| P. aeruginosa PA14ΔrsmAΔamrZΔclpV2ΔPA14_03210 | This paper | N/A |
| Escherichia coli DH5α | Lab Stock | N/A |
| E. coli BL21 (DE3) | Lab Stock | N/A |
| E. coli B834 (DE3) | Lab Stock | N/A |
| E. coli CC118(pir) | Herrera et al., 1990 | N/A |
| E. coli Sm10(pir) | Miller and Mekalanos, 1988 | N/A |
| E. coli 1047 pRK2013 | Hachani et al., 2014 | N/A |
| E. coli XL1-Blue | Agilent | Cat#200249 |
| E. coli MC4100 | Casadaban, 1976 | N/A |
| E. coli MC1061 | Casadaban and Cohen, 1980 | N/A |
| E. coli CS703/1 | Meberg et al., 2001 | N/A |
| E. coli MC1000 | Mavridou et al., 2012 | N/A |
| E. coli MC1000 dsbA::Km | Mavridou et al., 2012 | N/A |
| E. coli BW25113 | Keio collection | N/A |
| E. coli BW25113ΔmltC | Keio collection | N/A |
| E. coli BW25113ΔdacA | Keio collection | N/A |
| E. coli BW25113ΔdacC | Keio collection | N/A |
| Acinetobacter baylyi ADP1 | Suzana Salcedo | N/A |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| Arabinose | Sigma Aldrich | Cat#A3256 |
| Cellosyl | (Gift from Hoechst, Germany) | N/A |
| FM1-43 | ThermoFisher | Cat#T3163 |
| HA Peptide | Sigma Aldrich | Cat#11666975001 |
| IPTG | Melford | Cat#MB1008 |
| Lysozyme | Roche | Cat#10837059001 |
| Proteinase K | QIAGEN | Cat#19131 |
| Thrombin Protease | Sigma Aldrich | Cat#GE27-0846-01 |
| VgrG4bC-ter | Allsopp et al., 2017 | N/A |
| X-gal | Melford | Cat#M1001 |
| **Critical Commercial Assays** | | |
| SuperSignal West Pico PLUS Chemiluminescent Substrate | ThermoFisher | Cat#34580 |
| **Deposited Data** | | |
| VgrG2bC-ter crystal structure | This paper | PDB: 6H56 |
| VgrG2bC-ter Crystal diffraction data | This paper | Zenodo: 10.5281/zenodo.3246345 |
| **Oligonucleotides** | | |
| See Table S4 for full list of primers | N/A | |
| **Recombinant DNA** | | |
| pBBR1-MCS-4 | Kovach et al., 1995 | N/A |
| pBBR1-MCS-4-tssE2-his6 | This paper | N/A |
| pET28a | Novagen | Cat#69864 |
| pET28a-vgrG2bC-ter | This paper | N/A |
| pET28a-vgrG2bC-ter (E936A) | This paper | N/A |
| pET28a-mltC | This paper | N/A |
| pET28a-loIB | This paper | N/A |
| pET28a-rcsF | This paper | N/A |
| pTat | This paper | N/A |
| pTat-vgrG2bC-ter-myc | This paper | N/A |
| pTat-vgrG2bC-ter (E936A)-myc | This paper | N/A |
| pET22b | Novagen | Cat#69744 |
| pET22b-PA0261-HA | This paper | N/A |
| pET22b-tfl4-HA | This paper | N/A |
| pBBR1-MCS-5 | Kovach et al., 1995 | N/A |
| pBBR1-MCS-5-PA0261-HA | This paper | N/A |
| pBBR1-MCS-5-PA0261(no SP)-HA | This paper | N/A |
| pBBR1-MCS-5-tfl3-HA | Wood et al., 2019 | N/A |
| pTat-ACIAD0053-FLAG | This paper | N/A |
| pTat-ACIAD0053*-FLAG | This paper | N/A |
| pET22b-SARI_02726-HA | This paper | N/A |
| pET22b-ACIAD0054-HA | This paper | N/A |
| pET22b-vgrG2ac-ter-streplll | This paper | N/A |
| pET22b-vgrG2bc-ter (E936A) | This paper | N/A |
| pET22b-vgrG2bc-ter (E936A)-streplll | This paper | N/A |
| pSChraB2-CV | Cardona and Valzano, 2005 | N/A |
| pSChraB2-CV-peri-vgrG2b(847-1019)P14 | This paper | N/A |
| pSChraB2-CV-peri-vgrG2b(847-1019)P14 (H935A) | This paper | N/A |
| pSChraB2-CV-peri-vgrG2b(847-1019)P14 (E936A) | This paper | N/A |

(Continued on next page)
LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Alain Filloux (a.filloux@imperial.ac.uk). All reagents generated in this work shall be shared upon request without restrictions.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Bacterial strains were grown at 37°C in lysogeny broth (LB) with agitation unless stated otherwise. All cultures were supplemented with antibiotics and other supplements where necessary. The following antibiotic concentrations were employed for *E. coli*: 50 μg/ml kanamycin, 50 μg/ml ampicillin, 34 μg/ml chloramphenicol, 15 μg/ml tetracycline, 200 μg/ml trimethoprim and 15 μg/ml gentamicin. For *P. aeruginosa*, 50 μg/ml gentamicin, 2 mg/ml streptomycin, 50 μg/ml tetracycline and 100 μg/ml carbenicillin were used. Deletion mutants were constructed as previously described (Vasseur et al., 2005). Briefly, splicing by overlap extension polymerase chain reaction (PCR) was used to generate DNA fragments of regions of the *P. aeruginosa* genome with in-frame gene deletions introduced, which were cloned into the suicide vector pKNG101. After mobilisation into *P. aeruginosa* by three-partner conjugation from *E. coli* CC118 pir and with the 1047 pRK2013 helper strain, selection of conjugants was achieved on Vogel-Bonner medium (20 mM magnesium sulfate heptahydrate, 200 mg anhydrous citric acid, 1 g potassium phosphate dibasic and 350 mg ammonium sodium phosphate dibasic tetrahydrate) supplemented with 1.5% (w/v) agar and 2 mg/ml streptomycin. Counter-selection on solid LB medium containing 20% (w/v) sucrose at ambient temperature for 72 h lead to plasmid excision and generation double recombinants. All mutants were confirmed by PCR and sequencing.

METHOD DETAILS

Secretion assays and immunoblot analysis

*P. aeruginosa* secretion assays were conducted as previously described (Allsopp et al., 2017) with the following modifications. Cultures were inoculated into 25 mL tryptic soy broth (TSB) at OD₆₀₀ 0.1 and grown for 8 h at 25°C with agitation. Supernatants were cleared of cells by four rounds of centrifugation at 4000 g at 4°C, successively taking the uppermost supernatant. Proteins were precipitated with 10% trichloroacetic acid supplemented with 0.03% sodium deoxycholate overnight at 4°C. Separation of protein samples by SDS-PAGE and subsequent immunoblot analysis was conducted as previously described (Hachani et al., 2011) with the exception of culture supernatants being loaded at 20x concentrated in comparison to the whole cell extracts.
Here, bacterial culture samples were normalized to an OD$_{600}$ 1.0. Gels containing 6%, 8%, 10%, 12% or 15% polyacrylamide were used to separate protein samples by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) in Tris-glycine-SDS buffer (0.3% (w/v) Tris, 1.44% (w/v) glycine, 0.01% (w/v) SDS) at 180 V, depending on the size resolution desired. Gels were prepared using MiniPROTEAN Tetra handcast systems (BioRad) and the Precision Plus Protein Kaleidoscope Prestained Protein Standards marker (Bio-Rad) was loaded to show the migration of molecular weight standards. Samples were stained with Coomassic Brilliant Blue R250 (Sigma) or subjected to immunoblotting. For immunoblot analysis, protein samples were transferred to 0.2 μm nitrocellulose membrane (Amersham Biosciences) after incubation in transfer buffer (10% (v/v) Tris-glycine-SDS buffer, 20% (v/v) ethanol) using a TransBlot SD semi-dry transfer cell (BioRad) at 24 V and 0.18 A for 44 min. Membranes were blocked using a solution of 5% (w/v) skimmed milk powder (Sigma), 50 mM Tris-HCl pH 8.0, 150 mM NaCl and 0.1% (v/v) Tween-20 for 1 h before the addition of primary antibodies. Primary separation from non-cleaved protein on a Ni$_2^+$-NTA column in the flow-through fractions. Proteins tag was cleaved by thrombin protease (Sigma Aldrich) and dialysed into low imidazole buffer (50 mM Tris-HCl pH 8.0, 500 mM sodium chloride, 20 mM imidazole) before separation from non-cleaved protein on a Ni$_2^+$-NTA column in the flow-through fractions. Proteins were concentrated in Amicon Ultra Centrifugal Filter Units (Millipore) before further purification by SEC using a Superdex S200 10/300 GL column (GE Healthcare). Purity of elution fractions was determined by Coomassie staining of proteins separated by SDS-PAGE.

**Protein purification**

Purification of proteins was performed as described previously (Allsopp et al., 2017). Briefly, E. coli BL21 (DE3) pET28a-vgrG2bC-ter or pET28a-vgrG2dC-ter (E936A) were grown in terrific broth at 37°C for 2 h prior to induction of gene expression by addition of 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) and growth overnight at 18°C. B834 (DE3) was the host for selenomethionine incorporation. Cell harvesting and protein purification was done as described, but without protease inhibitors. The C-terminal hexahistidine tag was cleaved by thrombin protease (Sigma Aldrich) and dialysed into low imidazole buffer (50 mM Tris-HCl pH 8.0, 500 mM sodium chloride, 20 mM imidazole) before separation from non-cleaved protein on a Ni$_2^+$-NTA column in the flow-through fractions. Proteins were concentrated in Amicon Ultra Centrifugal Filter Units (Millipore) before further purification by SEC using a Superdex S200 10/300 GL column (GE Healthcare). Purity of elution fractions was determined by Coomassie staining of proteins separated by SDS-PAGE.

**Crystallization experiments and structure determination**

Purified VgrG2bC-ter was concentrated to 12 mg/ml and centrifuged for 20 minutes at 4°C to remove dust and aggregates. The protein was crystallized by vapor diffusion at 20°C, with small crystals growing in 100 Mm bis-tris pH 6.5 and 45% polypropylene glycol P400. Their space group was P3$_1$ and they diffracted to 3.2 Å. The structure was solved by experimental phasing, whereby crystals of selenomethionylated protein were grown in 100 mM sodium citrate pH 4.7, 20 mM magnesium chloride, 50 mM sodium chloride and 29% PEG400. These crystals had space group P42$_2_1_2$ and diffracted to 3.0 Å. Data were collected at the beamlines i04 and i04 of Diamond Light Source (Didcot, UK) from crystals flash-frozen in liquid nitrogen without additional cryoprotection. Diffraction data from both experiments can be downloaded from Zenodo (https://doi.org/10.5281/zenodo.3246345). Data were processed in XDS (Kabsch, 2010). The structure was solved by single anomalous dispersion based on the selenomethionine signal, although due to limited resolution and low anomalous signal, four datasets were combined in Blend (Foadi et al., 2013) to determine the anomalous substructure in SHELX (Sheldrick, 2010). A partial model comprising secondary structure elements was built by phenix.autobuild over several rounds of rebuilding and refinement before being used for molecular replacement with the P3$_1$ dataset in Phaser (McCoy et al., 2007). The new model was rebuilt in Coot and refined in Refmac5 (Murshudov et al., 2011), phenix.refine (Adams et al., 2010) and Buster-TNT (Blanc et al., 2004) until convergence. The crystallographic statistics are summarized in Table S1. Interface analysis was performed on the Proteins, Interfaces, Structures and Assemblies (PIASA) server while molecular graphics figures were prepared with PyMol (Schrödinger).

**Differential scanning fluorimetry**

Analysis of the thermal stability of recombinant VgrG2bC-ter protein in the presence and absence of zinc ions was determined using a Mx3005P qPCR instrument (Agilent). VgrG2bC-ter was used at 1.25 μM, in a buffer of 10 mM HEPES pH 7.4 and 100 mM NaCl supplemented with zinc acetate as indicated. The fluorescent dye SyproOrange (Sigma Aldrich) was used at 1/1000 dilution and the thermal unfolding of VgrG2bC-ter was monitored between 25-98°C at a rate of 1.5K/min. Excitation occurred at 492 nm and emission fluorescence at 610 nm was measured every 40 s. Non-linear least-squares fitting (Kemmer and Keller, 2010) was used to analyze the raw data and the melting point of each sample was determined from the inflection point of the fitted curve.

**SEC-MALLS**

The oligomeric state of recombinant VgrG2bC-ter was determined using SEC-MALLS and refractometry. VgrG2bC-ter was loaded onto a Superdex 75 10/30 column (GE Healthcare) and separated at 20°C with a flow rate of 0.5 ml/min. MALLS detection was performed with a miniDAWN TRESOS detector (Wyatt Technology) using a laser emitting at 657.3 nm. The refractive index of the solution was measured with an Optilab T-REX detector (Wyatt Technology) with the refractive-index increment (dn/dc) set to 0.185 ml/g. Weight-averaged molar masses were calculated using ASTRA software (Wyatt Technology).
Bacterial competition assays

The *P. aeruginosa* strains used in intraspecies competitions were differentiated by integration of Mini-CTX-lacZ at the chromosomal att site of the prey strains, permitting blue/white screening on 5-bromo-4-chloro-3-indolyl-D-galactopyranoside (X-gal)-containing solid media. Bacteria were grown overnight in 5 mL LB with appropriate antibiotics before normalization to an OD600 3.0 in 1 mL sterile PBS. Attacker and prey strains were mixed at a 1:1 ratio and 5 μl competition drops were spotted onto dry 3% low salt LB agar (10 g bactopeptone, 5 g yeast extract, 30 g bacteriological agar per liter). After drying, competitions were incubated at 25°C for 24 h. Both the input and output of competition spots were serially diluted in PBS and plated on LB containing 100 μg/ml X-gal for enumeration of colony forming units (CFU). The competitive index was calculated as the ratio between the input and output attacker/prey ratios. For self-intoxication assays on solid media, overnight cultures of *P. aeruginosa* strains were normalized and spotted onto nitrocellulose membrane containing gentamicin and 300 μM IPTG. Cultures were grown at 25°C with agitation in a 96-well plate, and OD600 readings were taken every 30 minutes for 18 hours using a Synergy 4 Microplate Reader (Biotek Instruments).

Bacterial intoxication assays

*E. coli* DH5α was transformed with pTat plasmids harboring variants of T6SS metallopeptidase genes to target the gene products to the periplasmic space and BL21 (DE3) was the host when pET28a and pET22b plasmids were used. Investigation into the role of cell wall biosynthetic enzymes employed the corresponding deletion strains and the BW25113 parent strain from the Keio collection as hosts (Baba et al., 2006). The host for the peri-vgrG2b (847–1019) plasmids was *E. coli* XL1-Blue harboring the immunity gene under a leaky promoter, since the presence of the plasmid containing the wild-type peri-vgrG2b (847–1019) construct was otherwise not tolerated. Overnight cultures of the strains harboring the vectors of interest were grown in LB, normalized and serially diluted. Dilutions were spotted on LB agar containing inducer (100 μM IPTG, 0.2% arabinose or 0.3% rhamnose) or repressor (0.2% glucose). To modify the osmolality of solid media, low tonicidy was achieved with low salt LB agar, while to raise it LB agar was supplemented with 500 mM sucrose.

Subcellular fractionation of bacterial cells

The localization of proteins within bacterial cells was probed by membrane fractionation and periplasmic extraction procedures. To extract the periplasm, *E. coli* cells producing the proteins of interest were normalized to OD600 20 and resuspended in 200 μl spheroplast buffer (200 mM Tris-HCl pH 8.0, 500 μM EDTA, 500 mM sucrose) to which 50 μg hen egg-white lysozyme (Roche) was added and incubated at 4°C for 15 min. To this, 720 μl half-strength spheroplast buffer was added and incubated for a further 15 min to release the periplasmic contents. Centrifugation at 5000 g at 4°C for 5 min separated the spheroplasts from the periplasmic fraction. To assess the presence of proteins in the inner and outer membrane, *E. coli* cultures were resuspended in sonication buffer (50 mM Tris-HCl pH 8.0, 1 mM EDTA) and lysed using a Vibra-Cell sonicator (Sonics & Materials, Inc.) on ice. Centrifugation at 4000 g cleared the soluble fraction from the cellular debris before ultracentrifugation at 100 000 g for 1 h at 4°C pelleted the total membrane fraction. After washing the membranes with sonication buffer, solubilisation buffer (15 mM Tris-HCl pH 7.4, 2% sodium lauryl sarcosinate) was added to differentially solubilise the inner and outer membrane proteins before further ultracentrifugation to pellet the insoluble outer membrane protein fraction. The membrane fractions were separated, and the pellet was washed with solubilisation buffer. All fractions were normalized prior to analysis by SDS-PAGE.

Far-western dot blot analysis

Protein-protein interactions were probed by far-western dot blotting as described previously (McCarthy et al., 2017). Briefly, 10 μg purified protein of interest or HA peptide (Sigma Aldrich) was spotted onto nitrocellulose membrane and air-dried before blocking. Lysates of *E. coli* with C-terminal HA tags were obtained by sonication and normalized to OD600 10 in protein binding buffer (20 mM Tris-HCl pH 7.6, 100 mM sodium chloride, 10% glycerol, 0.1% Tween-20, 2% skimmed milk powder). Membranes were incubated with the lysates or protein binding buffer overnight at 4°C prior to washing three times and immunoblotting for the bait and prey proteins.

Turbidometric and colorimetric analysis of VgrG2b activity

Generic protease activity of 10 μg recombinant VgrG2b, its inactive mutant and a proteinase K control was assessed using 1% proteinaceous substrates bovine serum albumin and gelatin in 2% bacteriological agar plates. After incubation at 37°C for 24 h, plates were stained with amido black solution (Vermelho et al., 1996) to visualize zones of proteolysis. Lysozyme activity of recombinant proteins utilized the lyophilised substrate *Micrococcus lylodeikticus* (Sigma Aldrich) as previously documented (Losi et al., 2011). The polymyxin B-mediated permeabilisation of the bacterial outer membrane to permit access of exogenous proteins to the periplasmic space was performed as described elsewhere (Brooks et al., 2013), except *E. coli* was employed. *E. coli* DH5α was resuspended in turbidity buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl) at OD600 0.2 and incubated with 4 μg/ml polymyxin B. After addition of 10 μg purified protein, the turbidity at OD600 was monitored at five-minute intervals for 1 h.
Zymography
Peptidoglycan was purified from 500 mL E. coli MC4100 culture for zymographic analysis as described elsewhere (Brooks et al., 2013; Santin and Cascales, 2017). Briefly, 10 µg purified protein was separated by SDS-PAGE in gels impregnated with 0.1% peptidoglycan before the gel was washed with ddH2O and equilibrated in renaturation buffer (10 mM Tris-HCl pH 7.5, 10 µM zinc acetate, 0.1% Triton X-100). Renaturation of the proteins occurred overnight at 37°C and after washing the peptidoglycan was stained with methylene blue (0.1% methylene blue, 0.01% potassium hydroxide) to visualize zones of clearing corresponding to peptidoglycan hydrolase activity.

Peptidoglycan hydrolase assay
Peptidoglycan was isolated from E. coli MC1061 (Casadaban and Cohen, 1980) and CS703/1 (Meberg et al., 2001) strains as described previously (Glauner, 1988). Briefly, E. coli cultures were grown to mid-exponential phase before centrifugation to pellet the cells. Cell membranes were solubilised by adding the cells in ice-cold water drop-wise into a boiling 8% (w/v) SDS solution under agitation. After cooling overnight, cell sacculi were collected by centrifugation at 130 000 g for 1 h at room temperature and washed four times to remove residual SDS. Glycogen and covalently-linked proteins were released by treatment with amylase and pronase, respectively, before boiling in SDS once more. Purified peptidoglycan was washed four more times in distilled water before resuspension in 20 mM sodium phosphate buffer pH 4.8. Reactions were carried out in 50 mM Tris-HCl pH 7.5 containing 150 mM NaCl, 0.05% Triton X-100 and 1 mM ZnCl2. Purified VgrG2bC-term or VgrG2bC-term (E936A) (5 µM) was incubated with 0.1 mg/ml purified peptidoglycan in a final volume of 100 µl for 4 h at 37°C in a thermostaker at 850 rpm. Reactions were terminated by boiling the samples at 100°C for 10 min. Next, 20 µl of 20 mM sodium phosphate buffer pH 4.8 and 1 µM cellulosyl muramidase (kindly provided by Hoechst, Germany) were added and incubated over night at 37°C. Samples were centrifuged at 13 000 rpm for 10 min and the supernatant containing the soluble muropeptides was collected. Samples were reduced using sodium borohydride and adjusted to pH 4-5 before reduced muropeptides were separated by reversed-phase HPLC (Glauner, 1988).

Microscopy techniques
E. coli DH5α harboring pTat plasmids were grown to OD600 0.5, resuspended in LB containing chloramphenicol, 0.2% arabinose and 5 µg/ml FM1-43 lipophilic dye, and 1 µl was spotted onto a coverslip under an agarose pad for microscopic analysis. Growth of bacteria was monitored every 30 min in a heated chamber at 37°C using an Axio Observer Z1 epifluorescent microscope (Zeiss). Image processing and cell length measurements were performed with FIJI software. Analysis of P. aeruginosa cell morphology was conducted similarly; however, competitions were initialised as described in the section “Bacterial competition assays.” To investigate the role of DsbA in PA0261-mediated neutralisation of periplasmic VgrG2bC-term, E. coli MC1000 or its dsbA mutant derivative harboring the relevant plasmids were normalized to OD600 1 after overnight growth. Serial dilution followed and spots were plated on LB agar containing the appropriate antibiotics in inducing or repressing conditions. Images of the spots were acquired with a Leica M205FA stereomicroscope with a 1x objective and colony area measurements were also determined within FIJI software.

Mass spectrometry analysis
Mass spectrometry analysis was conducted by the Plateforme Protéomique Structurale et Fonctionnelle at the Institut Jacques Monod, Paris. Briefly, samples underwent on-bead digestion with 12.5 µg/ml sequencing-grade trypsin (Promega), before peptide analysis on an Orbitrap Fusion Trubrid mass spectrometer coupled to an Easy-spray nanoelectrospray ion source and an Easy nano-LC Proxeon 1000 liquid chromatography system (all Thermo Scientific). Chromatographic separation of the peptides was achieved by an Acclaim PepMap 100 C18 pre-column and a PepMap-RSLC Proxeon C18 column at a flow rate of 300 nl/min. The solvent gradient consisted of 95% solvent A (water, 0.1% (v/v) formic acid) to 35% solvent B (100% acetonitrile, 0.1% (v/v) formic acid) over 98 minutes. An Orbitrap mass spectrometer analyzed the peptides in full ion scan mode, with the resolution set at 120 000 with a m/z mass range of 350 - 1550. High energy collision-induced dissociation activation with a collisional energy of 28% permitted fragment acquisition with the quadrupole isolation width of 1.6 Da. The linear ion trap was employed in top-speed mode to acquire the m/z with a mass tolerance of 7 ppm. Precursor ions and 0.5 Da for fragments. Identified modifications included acetylation (N-terminal), oxidation (Met) and phosphorylation (Ser, Thr or Tyr), and two missed trypsin cleavage sites were permitted. The E. coli SwissProt database (August 2017) was searched using the MS/MS data, while searching the P. aeruginosa database confirmed the presence of the bait proteins. A false-discovery rate of 1% was determined with the percolator algorithm, below which the identities of peptides could be assigned.

Analysis of lipoprotein abundance
Overnight cultures of E. coli BL21 (DE3) harboring pET28a-mtIC, -lolB, rcsF or pET22b were back-diluted in LB containing the appropriate antibiotics and construct expression was induced with 100 µM IPTG at OD600 0.4 for 3 h. Cells were harvested by centrifugation at 8 000 g for 3 min and resuspended in lysis buffer (100 mM Tris-HCl pH 7.5, 50 mM NaCl, 10 µM zinc acetate) for cell disruption by sonication. Lysates were clarified of cellular debris by sonication three times at 4 000 g for 15 min at 4°C before incubation with 10 µg VgrG2bC-term or VgrG2bC-term (E936A) at 37°C for 1 h. Ultracentrifugation at 100 000 g for 1 h at 4°C separated the soluble and
membrane fractions, to which Laemmli buffer was added for subsequent SDS-PAGE and immunoblot analysis. Production of MltC, LolB and RcsF was monitored using antibodies against the C-terminal hexahistidine tag, while production of TEM-1 from pET22b was detected using a monoclonal antibody. Densitometry was conducted in FIJI software.

**Bioinformatic and phylogenetic analysis**

Protein sequences were obtained from the non-redundant NCBI database using the BLASTp algorithm, alignments were done in Jalview using MAFFT and sequence logos were generated using the Weblogo server (Crooks et al., 2004; Katoh et al., 2017). The prediction of bacterial lipoprotein signal peptides, type I signal peptides and transmembrane helices was undertaken using LipoP 1.0, SignalP 4.1 and TMHMM server v2.0, respectively (Krogh et al., 2001; Nielsen, 2017). The search algorithm Jackhammer (EMBL-EBI) was used to search for homologs of the VgrG2b metallopeptidase domain. Three iterations querying the UniProtKB database were conducted, retrieving 240 sequences. Sequences were aligned and MEGA7 was employed for phylogenetic analysis, using the maximum-likelihood method of tree generation with 1000 bootstrap replicates (Kumar et al., 2016).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

The statistical analyses described in this work were conducted using Prism 8.0 software (GraphPad) and the statistical parameters, including sample sizes (where $n$ indicates the number of independent biological replicates, technical replicates or bacterial cells), $p$ values and statistical tests performed are indicated in the relevant figure legends. The means of biologically independent replicates were compared using the following statistical tests: Student’s t test, a one-way ANOVA followed by Dunnett’s or Tukey’s multiple comparisons tests. The spread is reported with the standard error of the mean (SEM), or by the median and quartiles for pooled single cell analyses.

**DATA AND CODE AVAILABILITY**

The accession numbers for the crystal structure and diffraction datasets of VgrG2b$_{C\text{-ter}}$ reported in this paper are PDB: 6H56 and Zenodo: 10.5281/zenodo.3246346, respectively.
Supplemental Information

The *Pseudomonas aeruginosa* T6SS Delivers a Periplasmic Toxin that Disrupts Bacterial Cell Morphology

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Figure S1: The VgrG2b metallopeptidase exists as monomeric and dimeric species in solution and is stabilised by zinc. Related to Figure 2.

A SEC-MALLS chromatogram of VgrG2bC-ter purified from E. coli. Solid lines show UV traces of fractions pooled from two peaks obtained after preparatory size-exclusion chromatography. Dashed lines across the peaks show the MALLS-derived apparent masses. The calculated weight-average molecular masses are 56.7 kD (left peak) and 29.1 kD (right peak) corresponding to the dimeric and monomeric species, respectively.

B Thermostability of the purified VgrG2bC-ter domain in the presence of different concentrations of zinc ions as determined by calculation of the melting temperature using differential scanning fluorimetry. Individual points represent the mean of three technical replicates with error bars showing the SEM (n = 3 technical replicates).
Figure S2: Characterisation of the VgrG2b-PA0261 effector-immunity pair family. Related to Figure 4.

A Self-intoxication assay of *P. aeruginosa* on solid media. Viability of the indicated strains assessed by enumeration of colony forming units (CFU). Here, the parental strain is *P. aeruginosa* PA14ΔrsmAΔamrZ. A statistically significant difference in recovered prey *versus* the parental strain was determined using a one-way ANOVA followed by Dunnett’s multiple comparisons test (*n* = 3 biological replicates; *p* < 0.05).

B Self-intoxication assay of *P. aeruginosa* in liquid culture. Viability was measured by monitoring turbidity (OD$_{600}$) in a 96-well format. The parental strain is the same as in panel A. Mean values are plotted from three biologically independent experiments, with error bars representing the standard deviation (*n* = 3).

C The putative N-terminal signal peptide of PA0261. The upper panel shows the prediction output of the SignalP 4.1 server. The lower panel indicates the predicted signal peptide (in pink) and putative signal peptidase cleavage site within the sequence of the PA0261 polypeptide.

D Immunoblot showing that both active and inactive VgrG2b$_{C_{ter}}$ variants are produced in *E. coli* from the pTat plasmids. Both isoforms are detected with anti-myc antibodies to a C-terminal tag. RpoB is a loading control.

E *E. coli* periplasmic toxicity assay showing the requirement of the metallopeptidase catalytic triad. VgrG2b metallopeptidase constructs (residues 847-1019) were targeted to the periplasm by an N-terminal PelB signal peptide. Strains were serially diluted on solid media under inducing conditions. Image is representative of three independent experiments.

F Plating efficiency of *E. coli* harbouring pTat, pTat-vgrG2b$_{C_{ter}}$ or pTat-vgrG2b$_{C_{ter}}$(E936A) in hypertonic conditions (LB agar with 500 mM sucrose). Expression from the plasmid is repressed with 0.2% glucose or induced with 0.2% arabinose.

G Targeting the *A. baylyi* VgrG2b$_{C_{ter}}$ metallopeptidase homologue ACIAD0053 to the *E. coli* periplasm using the TorA Tat-dependent signal peptide of the pTat vector. ACIAD0053* is the catalytic mutant of the effector where the HEMGH motif (residues 102-106) has been mutated to AAMGA. Plating efficiency of ten-fold serially diluted *E. coli* cultures on LB agar or low-salt LB agar (LB-LS) containing 0.2% arabinose inducer are shown. Image is representative of three independent experiments.

H Immunoblot detecting the FLAG-tagged ACIAD0053 toxin and its inactive AAMGA variant (designated by the asterisk). RpoB is a loading control.
Figure S3: Production of immunity proteins and purification of VgrG2b\textsubscript{C-ter}. Related to Figure 5.

A Immunoblot showing the production of the Myc-tagged VgrG2b\textsubscript{C-ter} effector and HA-tagged immunity proteins expressed in E. coli. RpoB acts as a loading control.

B and C SDS-PAGE analysis of recombinant VgrG2b\textsubscript{C-ter} (B) and VgrG2b\textsubscript{C-ter} (E936A) (C) elution after size-exclusion chromatography following Ni\textsuperscript{2+}-NTA affinity chromatography. Proteins were visualised by Coomassie staining. Fraction numbers are shown for each lane.
Figure S4: Alignment of immunity proteins in the PA0261 family. Related to Figure 5.

Multiple sequence alignment of putative immunity proteins for metallopeptidase effectors of the VgrG2bCter family. Due to low sequence homology, the alignment was manually curated after initial alignment by MAFFT. Omitted residues are shown in grey and cysteine residues are highlighted with red boxes. Darker coloration of the residues reflects the increasing conservation of identity, with the scores for each position shown below the alignment.
Figure S5: Members of the PA0261 immunity protein family utilise diverse mechanisms to access the periplasmic compartment. Related to Figure 5.

A Immunoblot analysis of the subcellular localisation of HA-tagged PA0261. RpoB is used as a marker of the soluble fraction (Sol), DsbA for the periplasm (PP), SecA for the inner membrane (IM) and PhoE for the outer membrane (OM). WCE: whole cell extract; TMF: total membrane fraction. Immunoblots are representative of three independent experiments.

B Multiple sequence alignment of the N-terminal region of PA0261 orthologues within Enterobacteriaceae. The red box demarcates a putative lipobox sequence and the predicted signal peptidase II (SPII) cleavage site is marked. Light to dark grey shading indicates an increasing level of residue conservation.

C and D Immunoblot of fractionated E. coli cells producing HA-tagged PA0261 orthologues from S. arizonae (SARI_02726; C) and A. baylyi (ACIAD0054; D). See panel A for details of fraction markers.

E Model showing the subcellular localisation of the three metallopeptidase immunity proteins PA0261, SARI_02726 and ACIAD0054. These immunity determinants, depicted in green, are a soluble periplasmic protein, an outer membrane lipoprotein and an inner membrane protein, respectively. The periplasmic VgrG2bc-
ter-family metallopeptidase is represented as an orange circle. Immunoblots are representative of three independent experiments.
Figure S6: VgrG2bC_ter displays no detectable peptidoglycan hydrolase or generic protease activity.
Related to Figure 6.
A Recombinant VgrG2bC_ter does not cause lysis of M. lysodeikticus as measured by turbidometry. Incubation with lysozyme acts as the positive control, while the buffer-only condition and addition of recombinant VgrG4bC_ter act as the negative controls. Points and error bars represent the mean ± SEM (n = 3 technical replicates).
B Recombinant VgrG2bC_ter, VgrG2bC_ter (E936A), VgrG4bC_ter and lysozyme were assessed for peptidoglycan hydrolytic activity by zymography with a purified E. coli peptidoglycan substrate. Samples were run on a 12% SDS-PAGE gel containing 0.1% peptidoglycan and stained with a methylene blue solution to visualise zones of clearing, indicative of peptidoglycan hydrolysis. A Coomassie-stained gel acts as a loading control for the purified proteins. The arrows highlight the location of the band corresponding to VgrG2bC_ter.
C 10 µg VgrG2bC_ter, VgrG2bC_ter (E936A) or proteinase K was spotted onto bacteriological agar plates containing 1% bovine serum albumin or gelatin. After incubation at 37 °C for 24 h, amido black staining was used to visualise zones of clearing indicative of proteolytic activity.
D and E HPLC chromatograms showing the profiles of the major muropeptides (left panels) released from tetrapeptide-rich E. coli MC1061 (D) and pentapeptide-rich CS703/1 (E) sacculi after incubation with VgrG2bC_ter, VgrG2bC_ter (E936A) or a buffer-only control. The composition of muropeptide species in control and assay reactions is also shown (right panels).
F Structures of the muropeptides shown in panels D and E. Abbreviations: GcNAc, N-acetylglucosamine; MurNAc(r), N-acetylmuramitol; L-Ala, L-alanine; D-Glu, D-glutamic acid; mDAP, meso-diaminopimelic acid; D-Ala, D-alanine.
Figure S7: MltC, PBP5 and PBP6a individually dispensable for VgrG2bC-ter-mediated toxicity. Related to Figure 7.

Plating efficiency of *E. coli* strains harbouring pTat vector, pTat-vgrG2bC-ter or pTat-vgrG2bC-ter (E936A). Strains are wild type BW25113, mltC, dacA (encoding PBP5) and dacC (encoding PBP6a) mutants. Serial dilutions assessed toxicity in repressive (0.2% glucose) and inducing (0.2% arabinose) conditions. Images are representative of three biological replicates.
Table S1: Data collection and refinement statistics for the crystal structure of the VgrG2b metallopeptidase domain. Related to Figure 2.

| Data processing | VgrG2b (833-1013) |
|-----------------|------------------|
| Wavelength (Å)  | 0.9795           |
| Resolution Range (Å) | 43.9 - 3.2 (3.42 - 3.2)ᵃ |
| Space Group     | P 3_1            |
| Unit Cell       |                  |
| a, b, c (Å)     | 78.3, 78.3, 115.3 |
| α, β, γ (°)     | 90, 90, 120      |
| Total Reflections | 45339            |
| Unique Reflections | 12782        |
| Multiplicity    | 3.5 (3.6)ᵃ       |
| Completeness (%)| 99.6 (99.9)ᵃ     |
| I/σI            | 12.5 (2.0)ᵃ      |
| d_{min} for I/sig(I) > 1.0 (Å) | 3.1 |
| Wilson B-factor | 123.5            |
| R_{sym} (%)     | 5.7 (46.2)ᵃ      |

Model refinement

| R value       | 0.223 |
| R_{free}      | 0.232 |
| Number of Atoms | 2745 |
| Protein Residues | 360  |
| Water Molecules | 0    |
| Bond lengths (Å) | 0.009 |
| Bond angles (°)  | 1.01  |
| Ramachandran Favoured (%) | 95.5% |
| Ramachandran Outliers (%) | 0    |
| Clash Score    | 4     |

ᵃ Values within parentheses denote the resolution of the highest resolution shell
Table S2: Data collection and processing statistics for crystals of selenomethionine-derivatised VgrG2b metallopeptidase domain. Related to Figure 2.

|                      | Pos 1 | Pos 4 | Pos 5 | Pos 6 | All data |
|----------------------|-------|-------|-------|-------|----------|
| **Data collection**  |       |       |       |       |          |
| Wavelength (Å)       | 0.97949 |       |       |       |          |
| Oscillation width (°) | 0.15  |       |       |       |          |
| Number of images     | 2400  | 9600  |       |       |          |
| **Data processing**  |       |       |       |       |          |
| Space group          | P 4 2 2 |       |       |       |          |
| Unit cell parameters |       |       |       |       |          |
| Resolution range (Å) | 40-2.7 (2.86-2.7) \(^a\) | 40-2.8 (2.97-2.8) \(^a\) | 40-2.9 (3.07-2.9) \(^a\) | 40-2.8 (2.97-2.8) \(^a\) | 40-2.7 (2.85-2.7) \(^a\) |
| Unique reflections   | 60251 (9706) \(^a\) | 54432 (8760) \(^a\) | 48913 (7849) \(^a\) | 53864 (8712) \(^a\) | 60184 (8967) \(^a\) |
| Multiplicity         | 14.0 (13.5) \(^a\) | 13.6 (12.1) \(^a\) | 13.7 (13.6) \(^a\) | 13.5 (12.2) \(^a\) | 49.3 (20.6) \(^a\) |
| R\(_{\text{meas}}\) (lowest resolution) | 0.073 | 0.061 | 0.071 | 0.079 | 0.095 |
| ISa                  | 14.7 | 16.7 | 21.8 | 14.2 | N/A      |
| R\(_{\text{meas}}\)  | 0.323 (6.17) \(^a\) | 0.304 (9.26) \(^a\) | 0.413 (8.15) \(^a\) | 0.345 (6.53) \(^a\) | 0.381 (7.60) \(^a\) |
| Completeness         | 0.999 (0.998) \(^a\) | 0.999 (0.999) \(^a\) | 0.999 (0.997) \(^a\) | 1.000 (1.000) \(^a\) | 0.999 (0.999) \(^a\) |
| (I/σ(I))             | 6.66 (0.13) \(^a\) | 7.80 (0.16) \(^a\) | 7.38 (0.22) \(^a\) | 6.65 (0.23) \(^a\) | 9.91 (0.13) \(^a\) |
| CC\(_{1/2}\)         | 0.997 (0.277) \(^a\) | 0.997 (0.124) \(^a\) | 0.996 (0.201) \(^a\) | 0.997 (0.256) \(^a\) | 0.999 (0.251) \(^a\) |
| d\(_{\min}\) for I/σ(I) > 1.0 (Å) | 3.0 | 3.1 | 3.2 | 3.1 | 3.0 |
| d\(_{\min}\) for SigAno\(^b\) > 1.0 (Å) | 4.65 | 4.82 | 3.88 | 4.19 | 4.27 |

\(\text{a. Values within parentheses denote the resolution of the highest resolution shell}
\)

\(\text{b. SigAno} = |F^+ - F^-|/\sigma(F)\)
| UniProt Accession | Protein* | Replicate 1 | Replicate 2 | Replicate 3 |
|-------------------|----------|-------------|-------------|-------------|
|                   | Coverage (%) | Unique Peptides | MASCOT Score (a) | MASCOT Score (b) | Coverage (%) | Unique Peptides | MASCOT Score (a) | MASCOT Score (b) | Coverage (%) | Unique Peptides | MASCOT Score (a) | MASCOT Score (b) | MASCOT Score (c) |
| Cell envelope     |          |             |             |             |
| Q916M7            | 88       | 26          | 14949       |             |
| P0C066            | 62       | 12          | 489         |             |
| P76537            | 37       | 4           | 155         |             |
| P08506            | 58       | 14          | 46          | 455         |
| P0AEB4            | 58       | 18          | 215         | 417         |
| P18392            | 13       | 4           | 237         |             |
| P0A938            | 76       | 3           | 199         |             |
| P69778            | 33       | 2           | 131         | 53          |
| A7ZKY3            | 54       | 9           | 227         |             |
| P0AC04            | 33       | 6           | 119         | 250         |
| P77774            | 47       | 10          | 77          | 189         |
| P0A903            | 22       | 5           | 14          | 153         |
| P0AAS1            | 33       | 4           | 31          | 195         |
| Cytoplasmic       |          |             |             |             |
| P0ACZ8            | 52       | 11          | 326         | 824         |
| P13035            | 69       | 28          | 56          | 1063        |
| P26616            | 32       | 9           | 20          | 287         |
| P67095            | 36       | 4           | 39          | 154         |
| P0AGK8            | 38       | 3           | 35          | 108         |
| P0AEZ1            | 23       | 5           | 58          | 117         |
| P00561            | 30       | 13          | 49          | 224         |
| Cell envelope (below threshold) |          |             |             |             |
| P69412            | 26       | 3           | 158         |             |

*: BL21 pET22b-vgrG2bC-ter (E936A)
b: BL21 pET22b-vgrG2bC-ter (E936A)-strepII
c: BL21 pET22b-vgrG2aC-ter-strepII

d: Lipoprotein
Table S3: Interactome of VgrG2bC-ter (E936A) in the *E. coli* cell envelope. Related to Figure 7.
The MASCOT scores indicate the enrichment of the proteins pulled down within the three independent replicates, presenting the interactome of periplasmic VgrG2bC-ter (E936A) in *E. coli*. Proteins localised in the cytoplasm were discounted as relevant interaction partners. The UniProt accession number, defining identity of interacting partners, is also shown alongside the coverage and the number of unique peptides found in each replicate. The MASCOT score corresponds to $-10\log(P)$, where $P$ is the calculated probability that the observed match between the experimental data and the database sequence is a random event and serves as a statistical representation of the positive identification of the protein in the sample, with higher scores being more significant. Note that replicates were entirely independent so quantitative comparisons may not be conducted.