Bacteria have evolved toxins to outcompete other bacteria or to hijack host cell pathways. One broad family of bacterial polymorphic toxins gathers multidomain proteins with a modular organization, comprising a C-terminal toxin domain fused to a N-terminal domain that adapts to the delivery apparatus. Polymorphic toxins include bacteriocins, contact-dependent growth inhibition systems, and specialized Hcp, VgrG, PAAR or Rhs Type VI secretion (T6SS) components. We recently described and characterized Tre23, a toxin domain fused to a T6SS-associated Rhs protein in *Photorhabdus laumondii*, Rhs1. Here, we show that Rhs1 forms a complex with the T6SS spike protein VgrG and the EagR chaperone. Using truncation derivatives and cross-linking mass spectrometry, we demonstrate that VgrG-EagR-Rhs1 complex formation requires the VgrG C-terminal β-helix and the Rhs1 N-terminal region. We then report the cryo-electron-microscopy structure of the Rhs1-EagR complex, demonstrating that the Rhs1 central region forms a β-barrel cage-like structure that encapsulates the C-terminal toxin domain, and provide evidence for processing of the Rhs1 protein through aspartyl autoproteolysis. We propose a model for Rhs1 loading on the T6SS, transport and delivery into the target cell.
To colonize a niche or a host, bacteria deliver toxins that destroy bacterial rivals or hijack the host defenses. Polymorphic toxins are defined as multidomain proteins with a modular organization, composed of a C-terminal toxin domain fused to a N-terminal trafficking domain\(^1,2\). Polymorphic toxins comprise diverse families including bacteriocins, contact-dependent growth inhibition (CDI) CdiA, Multiple adhesin family (Mal), Rearrangement hot spots (Rhs), and specialized type VI secretion system (T6SS) Hcp, VgrG, and PAAR proteins\(^3,9\).

Rhs proteins have a tripartite architecture: a N-terminal region (Rhs\(^{NT}\)) that specifies the mode of secretion, a conserved central domain predicted to fold as a shell-like structure (Rhs\(^{Shell}\)) consisting of Tyrosine/Aspartate-rich (YD) YDxxxDGRL/IT repeats demarcated from a highly variable C-terminal toxin domain (Rhs\(^{CT}\)) by a PxxxxDPxGL motif\(^4,10\). Rhs\(^{CT}\) domains possess various activities including nuclease, metalloproteases, ADP-riboseylases, or deaminases\(^1,2\). These Rhs\(^{CT}\) domains share GC content and codon bias that are different from sequences of the Rhs\(^{NT}\) that speciﬁs the mode of secretion of the Rhs\(^{CT}\) toxin domain. Indeed, Rhs proteins have been shown to confer advantages against bacterial competitors to Dickeya dadantii, Proteus mirabilis, Pseudomonas aeruginosa, Serratia marcescens, Enterobacter cloacae, Aeromonas dhakensis, P. fluorescens, and several pathogenic E. coli strains\(^6,10,14–22\).

YD-repeat-like elements are not only found in proteobacteria but are widely distributed in Bacteria, Archaea, and Eukarya, suggesting that they correspond to an ancient fold that has been adapted to various needs\(^23–26\). Well-known examples of YD-repeat proteins are the insecticidal TcB/TcC complex found in entomopathogenic strains such as Photorhabdus, Yersinia, and Serratia that associate with a third subunit, TcA, to assemble a tripartite Tc-toxin complex targeting insect cells\(^27,28\), the antibacterial WapA \(\tau\)RNases encoded on the genomes of Gram-positive bacteria, such as Bacillus and Listeria species\(^15\), Ss-Rhs1 that is essential for the virulence of the fungus Sclerotinia sclerotiorum toward host plants\(^29\), or the teneurins found in higher metazoans and involved in the establishment of neuronal cell connections during embryogenesis\(^30–33\). YD-repeat proteins structurally characterized so far, Tc-toxins and teneurins, assemble a large \(\beta\)-barrel shell\(^31,32,34–36\). While the teneurin C-terminal domain is positioned at the external surface of the shell, the toxin domains of Tc-toxins are engaged into the \(\beta\)-barrel\(^31,34,35,37\). It is not well determined how the toxin domain is released from the barrel but Rhs and TcC proteins share conserved motifs with the signature of aspartyl proteases that catalyze internal autolyses\(^10,38,39\).

The N-terminal regions of Rhs proteins, Rhs\(^{NT}\), define their mode of secretion. Rhs proteins can be associated with diverse secretion mechanisms such as type VI (T6SS) or type VII (T7SS) secretion systems\(^1,2\). However, most rhs genes are located within T6SS gene clusters or hcp-vgrG islands\(^4\). Indeed, those Rhs proteins are dependent on the T6SS for their delivery into target cells\(^6,14–22\). The T6SS is a multiprotein machine that uses a contractile mechanism to inject an effector-loaded needle into target cells\(^40\). It comprises a membrane complex that serves as a channel for the passage of the needle, on which is docked the assembly platform that initiates the polymerization of the contractile tail and triggers its contraction\(^41–44\). Effectors are usually fused or associated to components of the needle, such as the Hcp tube protein, the VgrG spike, or PAAR, a small conical subunit that sharpens the tip of the VgrG spike\(^3,9,43–49\). Rhs\(^{NT}\) domains of T6SS-associated Rhs proteins share a PAAR motif (Rhs\(^{PAAR}\))\(^47\), suggesting that Rhs proteins associate with the needle tip through VgrG-Rhs\(^{PAAR}\) interactions. Indeed, Rhs\(^{NT}\) deletion abolishes Rhs T6SS-dependent secretion in A. dhakensis\(^10\) and the P. fluorescens, S. marcescens, and A. dhakensis Rhs N-terminal domains have been shown to interact with VgrG\(^1,2,9,20,21\). T6SS-associated Rhs\(^{NT}\) domains may also bear transmembrane helices that are proposed to insert into the membrane of the target cell and that require a dedicated chaperone of the effector-associated gene (Eag, DUF1795) family to be maintained in a preinsertion state\(^19–21,30\).

Recently, we characterized a new toxin domain associated to a Rhs element in Photorhabdus laumondii\(^31\). This Rhs1 toxin, encoded by the plu0353 gene, is delivered by the T6SS, through interactions with VgrG (Plu0355) and the Eag chaperone (EagR, Plu0354). Production of the Plu0353 Rhs\(^{1,2}\)C domain, called Tre23, is toxic in the cytoplasm of E. coli and is counteracted by the Tr23 (Plu0352) immunity protein. Tre23 inhibits protein synthesis through ADP-ribosylation of helix 44 of the ribosomal 23S rRNA, hence preventing protein elongation by blocking the GTPase-associated center. Here we conduct a biochemical and structural characterization of Rhs1. Using a combination of pull-down and cross-link mass spectrometry analyses, we first provide details on the association of the VgrG, Rhs1, and EagR proteins, demonstrating that Rhs1\(^{NT}\) and VgrG gp5-like needle are responsible for VgrG-Rhs1 complex formation, and that EagR binds to two regions located upstream and downstream the PAAR motif of Rhs1\(^{NT}\), including a predicted hydrophobic transmembrane domain. We then provide the cryo-electron microscopy structure of the Rhs1-EagR complex, demonstrating that Rhs1\(^{Shell}\) assembles a \(\beta\)-barrel cage that encapsulates Rhs1\(^{CT}\). We finally demonstrate that Rhs1 undergoes autolysis at two positions located on each side of the Rhs1\(^{Shell}\) barrel, releasing the toxin Rhs1\(^{CT}\) domain.

Results
Organization of the P. laumondii VgrG-Rhs1-EagR complex
We recently showed that the P. laumondii Plu0353 Rhs1 protein, encoded within a Type VI secretion system (T6SS) gene cluster (Fig. 1a), is delivered by the T6SS and participates in antibacterial competition by inhibiting protein synthesis\(^31\). The full-length Rhs1 protein was shown to form a tripartite complex with the VgrG spike and the EagR chaperone\(^31\). To get further insights on the architecture of this complex, we conducted pairwise co-purification experiments using C-terminally FLAG-tagged Rhs1 (‘R’), N-terminally 6xHis-tagged VgrG (‘V’) or C-terminally Strept-tagged EagR (‘E’). Figures 1b and 1c show that the full-length Rhs1 protein is unstable in the absence of the EagR chaperone (Fig. 1b, c, lanes ‘R’ and ‘V + R’), suggesting that these two proteins interact. Indeed, FLAG-tagged Rhs1 copurifies with strept-tagged EagR (Fig. 1b, c, lane ‘E + R’). By contrast, the EagR chaperone does not pull down VgrG (Fig. 1b, c, lane ‘E + V’). The tripartite complex comprising VgrG, EagR and Rhs1 (Fig. 1b, c, lane ‘E + V + R’) could be readily purified, suggesting that the EagR-stabilized Rhs protein contacts the VgrG spike. We propose a model of the architecture of the tripartite complex in which EagR-bound Rhs1 interacts with VgrG (Fig. 1d).

To provide detailed information onto the architecture of the VgrG-Rhs1-EagR complex, we engineered truncations of the VgrG and Rhs1 proteins, and tested the ability of these truncated variants to form complexes. The P. laumondii Plu0355 VgrG protein shares the prototypical VgrG organization: from N- to C-terminus it comprises a gp27-like domain followed by OB-fold and gp5-like needle domains (Fig. 2a; Supplementary Fig. 1). Figure 2b shows that variants deleted of the gp27 domain only, or
A similar truncation approach was then performed for the Rhs1 protein. The Rhs1 protein comprises, from N- to C-terminus, a short 30-amino-acid sequence, called ‘prePAAR’ domain, followed by a predicted transmembrane domain (TMD1), a PAAR domain, a postPAAR domain comprising a conserved α-helix, the Rhs YD-shell and the Tre23 toxin domain (Fig. 2a; Supplementary Fig. 2). Pull-down of the tripartite complex with Rhs1 N-terminal truncations deleted of prePAAR, prePAAR-PAAR, or prePAAR-PAAR-postPAAR, shows that deletion of prePAAR is sufficient to promote attachment of the EagR-Rhs1 complex to VgrG, while EagR-Rhs1 interaction is only lost when Rhs1 is missing the entire N-terminus composed of prePAAR, PAAR and postPAAR (Fig. 2c). Experiments performed with Rhs1 C-terminal truncations showed that prePAAR is sufficient to interact with EagR, while prePAAR-PAAR is sufficient to recruit VgrG to the EagR-Rhs1 complex, although the VgrG-EagR-Rhs1 complex is fully stabilized when the complete N-terminus of Rhs1 comprising prePAAR-PAAR-postPAAR is produced (Fig. 2d). Finally, the isolated postPAAR region is also co-precipitated by EagR (Supplementary Fig. 3). Taken together, the results of Rhs1 N- and C-terminal truncations suggest that prePAAR-PAAR is sufficient to interact with VgrG, while Rhs1 presents two binding sites for EagR located at TMD1 and postPAAR, respectively. In agreement with these results, and as shown for the P. protegens RhsA protein, stability assays showed that the N-terminal region of Rhs1, comprising prePAAR-TMD1-PAAR-postPAAR, is an unstable domain that requires EagR for its stabilization (Supplementary Fig. 4). To test this model, the purified VgrG-Rhs1-EagR complex was subjected to in vitro cross-linking with NNp9, a chemical cross-linker that carries two NHS carbamate reactive groups and an azido function allowing further enrichment of cross-linked peptides through click-chemistry. Mass spectrometry analysis of cross-linked peptides revealed that the three proteins establish extensive contacts (Fig. 2e; Supplementary Table 1; Supplementary Fig. 5). Notably, interactions between EagR and Rhs1 TMD1 and postPAAR were confirmed. This approach also demonstrated numerous contacts between the VgrG C-terminus and Rhs1 prePAAR, although few contacts between Rhs1 prePAAR and the gp27 base of VgrG were also identified. In addition, intramolecular contacts were revealed between the N-terminal region of the prePAAR domain and PAAR, and between the N-terminus of the Rhs1 shell and the prePAAR and postPAAR domains, as well as with the toxin domain (Fig. 2e). Taken together, these results establish a structural organization of the VgrG-Rhs1-EagR complex in which the prePAAR-PAAR N-terminus of Rhs1 sits on the tip of the VgrG gp5-like β-helix while two EagR molecules embed the Rhs1 TMD1 and postPAAR domains (Fig. 2f). This model agrees with recent data demonstrating that the P. aeruginosa EagR-like EagT6 chaperone interacts with the Tse6 effector TMDs, and that prePAAR and PAAR domains mediate effector-VgrG interactions. In addition, the contacts detected between the Tre23 C-terminal toxin domain and the Rhs1 N-terminus suggest that the toxin domain is encapsulated into the Rhs1Shell barrel (Fig. 2f).
we could build de novo an atomic model of Rhs1Shell from amino acids (aa) 272 to 1357 (Fig. 3a). Rhs1Shell is composed of a continuous strip of anti-parallel sheets (residues 420-1262) defining a hollow barrel, which is closed on each side by two plugs (residues 272–420 and 1263–1357 respectively) (Fig. 3a). The C-terminal plug is deeply embedded in the Rhs1β-barrel and tightly closes one side of the barrel (Fig. 3a, b). This domain corresponds to an aspartyl protease domain that is conserved in homologous Rhs proteins and in YD-repeat proteins such as in the Y. entomophaga ABC toxin34. The N-terminal plug caps the other side of the Rhs1β-barrel but remains at the exterior (Fig. 3a, b). It is composed of an Immunoglobulin-like (Ig-like) domain (336–421) preceded by a small extended domain sealing it to the Rhs1β-barrel (Fig. 3a, b). The β-barrel and the two plugs define a central cavity of ~28,000 Å³ in volume (Fig. 3a, b). In this cavity, densities are visible but not well resolved (Fig. 3b), confirming that the C-terminal toxin domain is encapsulated into the Rhs1β-barrel but suggesting that it is highly flexible and/or partially unfolded.

The N-terminal plug is decorated by an external highly flexible density that corresponds to the N-terminal portion of the Rhs1 protein (Rhs1NT) in complex with an EagR dimer (Fig. 3c).
homology model of the chaperone dimer (Supplementary Fig. 8) could be fitted in the low-resolution density obtained (Fig. 3c).

**Rhs1 autocleavage.** The cryo-EM structure of the Rhs1-EagR complex shows that the Rhs1 Tre23 toxin domain is encaged inside the Rhs1 β-barrel (Fig. 3b), suggesting that drastic conformational changes or cleavage(s) should occur to release the toxin. Interestingly, the Rhs1 protein from *A. dhakensis* has recently been demonstrated to undergo two autocleavages, located at each side of the β-barrel10. The profile of the purified *P. laumondii* Rhs1 protein reveals four major bands corresponding to full-length Rhs1, and three cleaved products (Fig. 4a). Mass spectrometry analyses and N-terminal sequencing of these products defined that they correspond to N-terminal (Rhs1NT), central (Rhs1Shell) and C-terminal (Rhs1CT) fragments. The 29-kDa Rhs1NT N-terminal fragment comprises prePAAR, TMD1, PAAR, postPAAR, and stops after residue Pro-289 (Fig. 4b).
~120-kDa Rhs1\textsubscript{Shell} fragment starts at Ile-290 and ends at Leu-1342 and corresponds to the \(\beta\)-barrel with N- and C-terminal plugs that are folded inside the first and last rings of the barrel (Fig. 4b–f). Finally, the 16-kDa Rhs1\textsubscript{CT} C-terminal fragment comprises the Tre23 toxin domain that starts at residues Ala-1343 (Fig. 4b). Notably, while EagR interacts only with Rhs1\textsubscript{NT} TMD1 and postPAAR, all Rhs1 fragments are pulled-down by EagR, suggesting that the Rhs1 NT, Rhs1\textsubscript{Shell} and Rhs1\textsubscript{CT} fragments remain associated together after cleavage, as observed with the \(P.\) \textit{dhakensis} Rhs protein\textsuperscript{10}. The C-terminal cleavage site is less clear as no sequence corresponds to the signature of aspartyl proteases. However, a sequence alignment of the 250 best BLAST hits of Plu0353 revealed a \textsuperscript{289}PVYVAGE signature (Fig. 4b) that shares conserved residues with the PVSMVTGE motif defined by Pei et al\textsuperscript{10}. Interestingly, Pro-289 is located immediately downstream a highly conserved aspartate residue, such as in the DPLGL motif of aspartyl proteases (Fig. 4b). In agreement with these observations, the D288N mutation abolished N-terminal cleavage (Fig. 4a). Finally, the double D288N-D1338N Rhs1 variant remained intact (Fig. 4a). To test whether these cleavages are physiologically relevant to liberate the Tre23 toxin domain, the toxicity of the full-length \(P.\) \textit{laumondii} Rhs1 protein, and of its D288N, D1338N, and D288N-D1338N variants, was assayed in \textit{E. coli} cells. Figure 4g shows that while the full-length Rhs1 protein is highly toxic, all the cleavage variants are inactive, suggesting that the Tre23 domain needs to be released from the Rhs1 encapsulation cage to exert its lethal activity. Interestingly, in the cryo-EM map, densities are unresolved before D288 and after D1338, indicating that the Rhs1\textsuperscript{NT} and C-terminal toxin domains are flexible. However, even after autolyses, it is likely that the two lid domains will remain...
inside the barrel, on each side of the Rhs1\textsuperscript{CT} toxin, such as in the \textit{Y. entomophaga} ABC toxin\textsuperscript{34}. Further events will therefore be needed to liberate the toxin domain.

**Discussion**

In this work, we report the architecture and structure of a Rhs polymorphic toxin associated with the type VI secretion system in \textit{P. laumondii}. We show that Rhs1 is composed of three main regions: a Rhs1\textsuperscript{NT} N-terminal region that comprises a PAAR domain followed by a transmembrane segment, a PAAR domain, and a postPAAR domain comprising a conserved a-helix, a Rhs1\textsuperscript{Shell} central domain that folds as a \(\beta\)-barrel closed by plugs on each side, and a Rhs1\textsuperscript{CT} C-terminal domain that comprises the Tre23 ADP-ribosyltransferase toxin activity. The EagR chaperone binds Rhs1\textsuperscript{NT}, specifically at the level of TMD1 and postPAAR. Rhs1\textsuperscript{NT}. The prePAAR and PAAR domains are involved in the interaction with the \(\beta\)-needle tip of the VgrG transporter protein. The Tre23 toxin domain is encapsulated inside the \(\beta\)-barrel. We also demonstrate that Rhs1 undergoes two aspartate-catalyzed self-cleavages located at both sides of the Rhs1\textsuperscript{Shell}.

**A model for T6SS-dependent delivery of Rhs toxins.** While the mode of translocation of Rhs toxins into recipient cells remains enigmatic, we propose a model (Fig. 5) in which the PAAR-containing Rhs protein is stabilized in the cytoplasm of the attacker cell through interactions with its cognate EagR chaperone (Fig. 5a). Upon assembly of the T6SS, Rhs is loaded onto the VgrG spike through contacts between the Rhs prePAAR-PAAR region and the tip of the VgrG gp5-like needle (Fig. 5b). In this conformation, the attacker cell is protected from the action of the Tre23 toxin by both its cognate Tri23 cytoplasmic immunity\textsuperscript{51} and by the \(\beta\)-barrel cage that encapsulates Tre23. Such a double-lock mechanism has been already demonstrated for the EAEC Tle1 phospholipase active site, which is blocked by the Tli1 immunity protein\textsuperscript{52}, and by steric hindrance when loaded onto the VgrG spike\textsuperscript{53}. Whether the EagR chaperone remains associated on the tip complex or dissociates after Rhs loading on VgrG is yet to be experimentally determined. Contraction of the T6SS sheath fires the complex into the recipient target cell, where it is delivered into the cytoplasm or periplasm (Fig. 5c). If the Rhs complex is delivered into the periplasm, we propose that it dissociates from VgrG and that the Rhs\textsuperscript{NT} TMD inserts into the inner membrane to translocate the Rhs\textsuperscript{Shell} \(\beta\)-barrel across the inner membrane (Fig. 5d). Autocleavages at the N- and C-termini of the \(\beta\)-barrel liberates the toxin into the cytoplasm (Fig. 5e) where it can target 23S rRNAs to inhibit translation (Fig. 5f). Alternatively, the Rhs cage can be opened into the periplasm, and the released Tre23 C-terminal domain can be translocated through a channel formed by the Rhs TMD, or by an inner membrane protein/complex of the host. Many of these steps are hypothetical and should be experimentally addressed, e.g., does the EagR chaperone interact with Rhs prior to its loading onto the T6SS? Is the EagR chaperone dissociated prior secretion or in the target cell? Is the Rhs protein delivered into the periplasm or cytoplasm of the target cell? Does the Rhs \(\beta\)-barrel translocate across the inner membrane, insert into it or does it interact with an inner membrane protein/complex? How does the Rhs-EagR complex dissociate from VgrG? How is the toxin released or extracted from the Rhs\textsuperscript{Shell} cage?

**Fig. 5 Model for Rhs loading and delivery by the type VI secretion system.** a The EagR chaperone (blue) binds to the Rhs\textsuperscript{NT} hydrophobic domains to stabilize the Rhs protein (red) prior loading on the T6SS. b The Rhs protein is loaded on the C-terminus of the VgrG spike (orange) via its N-terminal prePAAR-PAAR domain. c Sheath contraction propels the inner tube (black) to inject the VgrG loaded with Rhs into the target cell. It is not known whether the EagR chaperone remains in the attacker cell or whether it is translocated together with the VgrG-Rhs complex. d The Rhs1 protein is directly delivered in the cytoplasm, or if delivered in the periplasm, the Rhs\textsuperscript{NT} hydrophobic transmembrane domains insert into the inner membrane of the target cell and are used to translocate the Rhs \(\beta\)-barrel and/or the toxic domain into the cytoplasm of the target cell. Alternatively, the Rhs \(\beta\)-barrel itself inserts into the membrane. e N- and C-terminal autoproteolysis opens the Rhs shell and releases the Rhs\textsuperscript{CT} toxin into the target cell cytoplasm. f The \textit{P. laumondii} Rhs1 C-terminal toxin domain, Tre23, arrests translation by ADP-ribosylation of the 23S rRNA.

NATURE COMMUNICATIONS | https://doi.org/10.1038/s41467-021-27388-0
NATURE COMMUNICATIONS | (2021) 12:6998 | https://doi.org/10.1038/s41467-021-27388-0 | www.nature.com/naturecommunications
Rhs structure. We have shown that the Rhs1 central region folds as a β-barrel structure in which the two extremities are obstructed by plug domains. Interestingly, it shares the same modular architecture as Tc-toxins and eukaryotic teneurins that protect their effector domains within a barrel that defines a shell-like compartment (Supplementary Fig. 9). In Tc-toxins and Rhs, this β-barrel maintains the toxin domain confined in a closed environment and most probably maintains it partially unfolded to prevent its action in the producing cell. The volume of the Rhs1 β-barrel, ~28,000 Å³, is compatible with the accommodation of a <25 kDa folded domain, in agreement with the observation that Rhs C-terminal toxin domains have sizes ranging from 9 to 20 kDa (Supplementary Fig. 10). By contrast, the C-terminal domain of teneurins is exposed at the surface of the shell31,35 (Supplementary Fig. 9). The architectures of Rhs1, Tc-toxins, and teneurins diverge at the level of the N-terminal plug (Supplementary Fig. 9). In Tc-toxins, this plug is made of another protein that extends the β-barrel and closes it. It also contains a β-propeller domain that is essential for the opening of the shell to extract the toxin. In teneurins, the N-terminal part of the protein forms a plug made of two Ig-like domains and a β-propeller domain. The T6SS-associated Rhs proteins therefore define another class with a N-terminal plug made of an Ig-like domain and a seal (Supplementary Fig. 9).

Mounting Rhs on the T6SS Needle. As previously evidenced for the S. marcescens, E. cloacae, and P. fluorescens Rhs proteins20–22, we have shown in this work that P. laumondii Rhs1 assembles a tripartite complex with the EagR chaperone and the VgrG spike. Rhs1 is thus likely transported as a VgrG cargo, such as other effectors including the P. aeruginosa Tse6 PAAR-effector or the enteroaggregative E. coli Tle1 phospholipase30,52,53. However, the mode of interaction with the spike is likely to be different. While Tle1 associates to a C-terminal extension of the VgrG spike52,53, our truncation studies demonstrate that Rhs1-VgrG complex formation requires the EagR chaperone for Rhs1NT stabilization, and the Rhs1NT and VgrG gp5-like β-prism. The result agrees with the observation that deletion of the Rhs N-terminal region in A. dhakensis abolishes Rhs secretion10. Further delineation showed that the prePAAR and PAAR regions of Rhs1NT are sufficient to support the interaction with VgrG. This result differs from the A. dhakensis VgrG-Rhs complex that requires both RhsNT and RhsCT for efficient interaction10 but is identical to the formation of the P. protegens VgrG1-RhsA complex that only requires RhsNT21. Negative-stain electron microscopy analyses of the P. protegens VgrG1-RhsA-EagR1 complex showed that the prePAAR-TMD-PAAR region of RhsA mediates contacts with the VgrG tip21. Based on structure modeling, Ahmad et al. suggested that the Tse6 and RhsA PAAR domains are not complete, and that prePAAR complements the PAAR fold, explaining why prePAAR is important for VgrG-RhsA or VgrG-Tse6 complex formation24. The Tse6 prePAAR-PAAR model shows that the His residue of the prePAAR AARxxDxxxH motif is properly positioned to participate to the tetrahedrally coordinated zinc2+ atom21. The P. laumondii Rhs1 prePAAR region also bears a PAAARxxDxxxH motif that comprises the PAAR/PAAR signature and the coordinating His residue (Supplementary Fig. 2c). Interestingly, our XL-MS data fully agree with this model as several contacts between Rhs prePAAR and PAAR were observed (Fig. 2e). Taken together, the current model proposes that the Rhs1NT prePAAR and PAAR regions form a bona fide PAAR conical structure that sits on the VgrG spike, such as canonical PAAR subunits, hence anchoring Rhs1 to the tip of the T6SS needle. While the position of Rhs1 on the needle is now better defined, it remains to be determined whether Rhs1 is loaded onto the VgrG spike during T6SS biogenesis. Recent information on the Enterobacter cloacae T6SS have shown that Rhs is required for assembly of the apparatus22. However, whether Rhs interact with VgrG before its incorporation into the baseplate, or whether it is recruited to the fully assembled baseplate remains to be determined.

The EagR chaperone and the Rhs1 TMD. Previous pull-down and two-hybrid studies showed that the EagR chaperone interacts with Rhs, and more specifically with RhsNT in S. marcescens, E. cloacae, P. fluorescens, and S. enterica Typhimurium20–22. Eag chaperones are required for the secretion and stability of PAAR domain-containing effectors such as P. aeruginosa Tse6, P. fluorescens Tne2 and S. marcescens and P. protegens RhsA19–21,56,57. Similar to our observation, Tse6 copurifies with EagT6 when individual TMDs are deleted, but not when both TMDs are deleted50, suggesting that Eag chaperones bind to both TMDs of PAAR-containing T6SS effectors. Indeed, the low-resolution electron microscopy structure of the VgrG-Tse6-EagT6 complex showed that two EagT6 dimers wrap the TMDs of the Tse6 effector30. The recent crystal structures of Eag chaperones from P. aeruginosa and S. enterica Typhimurium demonstrated that the Eag dimer has an overall horseshoe-like shape that embraces the RhsNT TMD helix bundle in a claw-like manner21. Binding of the Eag dimer to the RhsNT helix bundle is mediated by a combination of knob-hole hydrophobic interaction and hydrophilic residues via bifurcated hydrogen bonds21. A structural model of the P. laumondii EagR dimer shows that EagR has an identical horseshoe-like fold with a central hydrophobic cleft that can accommodate the Rhs TMDs (Supplementary Fig. 8a–c). Indeed, mapping of the Rhs cross-link contacts on the P. laumondii EagR dimer model shows that Rhs TMD1 and the conserved α-helix of postPAAR interact with the concave face of EagR (Supplementary Fig. 8d). Eag chaperones therefore protect the hydrophobic regions of RhsNT and other PAAR-containing effectors to prevent aggregation or premature insertion into membranes. Interestingly, Tse6 spontaneously inserts into lipid vesicles causing EagT6 chaperones to be released suggesting that EagT6 maintains the N-terminal TMDs in a preinsertion state until it reaches the inner membrane of the target cell50. However, this model is challenged by a study that has shown that the EagR protein is not detected in S. marcescens cell supernatant, contrarily to its cognate Rhs protein, suggesting that the EagR-Rhs interaction is not maintained after loading onto the T6SS apparatus20.

Autocleavage and toxin release. PrePAAR-containing effectors usually being cytoplasmic-acting toxins19,21, the TMDs are proposed to insert into the inner membrane to promote entry of the toxin domain in the cytoplasm. In the case of Rhs effectors, the toxin domain is encapsulated into the Rhs β-barrel and hence needs to be released upon translocation. Such as E. cloacae and A. dhakensis Rhs10,22, the P. laumondii Rhs1 protein undergoes two aspartyl autoproteolysis, yielding three fragments corresponding to Rhs1NT, Rhs1Shell and the Rhs1CT toxin. Autocleavage likely occurs when the Rhs protein reaches the target cell, as no autocleavage was detected inside E. cloacae22, suggesting that certain conditions associated with the periplasm and/or inner membrane of the target activate the intramolecular aspartyl protease activity. Interestingly, TcC polymorphic toxins also undergo autocleavage34,36,35, and some teneurins carry C-terminal sequences that resemble neuroendocrine signaling peptides adjacent located to predicted furin cleavage sites, suggesting that the C-terminal peptides are released36. As shown for insecticidal TcC proteins34,36,35, our results demonstrated that substitutions
preventing N- or C-terminal cleavage significantly reduced the toxic activity of Rhs1, and hence that auto-accumulation are required to liberate the Rhs1 CT domain in order to exert its toxic activity. In A. dhakensis, mutations blocking Rhs auto-accumulation do not impact T6SS-dependent secretion but reduces toxicity10. However, we and others have shown that the three fragments remain in complex upon auto-accumulation as they can be all pulled-down by the EagR chaperone that only binds to Rhs1NT10,22. In our conditions, and despite the two auto-accumulations, the Tre23 toxin is not fully liberated from the β-shell. These observations agree with our Rhs1 structure, which shows that the encapsulated Rhs1 CT is blocked into the Rhs1 cage by two plug regions that prevent its release from the β-barrel. While additional experiments are required to better understand how the toxin domain is liberated from the Rhs cage, one can hypothesize that specific conditions or factors in the target cell facilitate the separation of Rhs1 CT from Rhs1NT, or that large conformational changes to eject the lids out of the Rhs shell or to laterally open the β-barrel shell, occur to release the Rhs1 CT toxin.

Methods

**Bacterial strains, growth conditions, media, chemicals, and antibodies**

Bacterial strains, plasmids, and oligonucleotides used in this study are listed in Supplementary Table 3. E. coli DH5α strain was used for cloning and E. coli BL21(DE3) for protein production and purification. P. lachnii subsp. larryonii TTO1 DNA was used as the template for cloning. Bacteria were grown in Lysogeny Broth (LB, with agitation at 37 °C. When required, media were supplemented with ampicillin (100 μg mL−1), streptomycin (100 μg mL−1) or kanamycin (50 μg mL−1). Gene expression from PET vectors was induced by adding the mixture 500 μM isopropyl-β-D-thiogalactopyranoside (IPTG). SDS-PAGE. Western-blots and immunodetection have been performed using standard procedures. Gels were stained using InstantBlue™ (Sigma-Aldrich) or transferred onto nitrocellulose membrane, and immunodetected with His-Tag (clone 1B7G5, Proteintech catalogue #66005-1-Ig), Strep-Tag Classic (clone Strep-Tag II, Bio-Rad catalogue #MCA2489, anti-FLAG (clone M2, Sigma-Aldrich catalogue #F3165) antibodies, and secondary antibodies coupled to the Alkaline Phosphatase, and revealed using 5-bromo-4-chloro-3-indolyl phosphate/nitro blue tetrazolium (BCIP/NBT) in presence of 10 mM MgCl₂ in alkaline buffer (pH 9).

**Plasmid construction**

PET derivatives plasmides producing N-terminally 6×His-TEV-tagged VgrG (PET-His-VgrG), C-terminally Strep-tagged EagR (pRSF-EagR-ST) and C-terminally FLAG-tagged Rhs1 (pCfD-Rhs-FL) have been previously described55. The conserved sequence corresponding to the VgrG and Rhs1 truncated derivatives were cloned by restriction-ulation into PET-Duet™-1, and PET-Duet™-1. The sequence corresponding to full-length Rhs1 was cloned by restriction-igation into pBAD33-rbs, a pBAD33 vector derivative in which a consensus ribosome binding site has been cloned. Briefly, vgrG and rhs1 DNA fragments were amplified from E. coli BL21(DE3) corresponding to the T30 tag polymerase (NEB) and oligonucleotides inserting restriction sites. The PCR fragments were purified using the NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel) before and after digestion with BmtI and HindIII (PET-His-VgrG derivatives) or SacI-Sall (pCfD-Rhs-FL derivatives), pBAD33-rbs derivatives. Digested fragments and plasmids were then ligated with T4 DNA ligase (NEB) and transformed into E. coli DH5α cells. To prevent toxicity, substitutions of Tre23 toxin catalytic residues, Y1351 and Y1375 were introduced by site-directed mutagenesis using complementary pairs of oligonucleotides bearing the desired mutation. PCR-amplified fragments were then purified, phosphorylated by T4PKN (NEB), ligated with T4 DNA ligase and transformed into E. coli DH5α cells. Colonies were performed by colony PCR. Plasmids were isolated using Wizard Plus SV Miniprep kit (Promega) and their sequences were checked by DNA sequencing (Eurofins).

**Protein and complex purification**

VgrG-Rhs1-EagR complex purification was done as previously described55. Briefly, E. coli BL21(DE3) cells freshly transformed with pET-His-VgrG, pRSF-EagR-FL, and pCfD-Rhs-ST or with empty cloning vectors to yield strains with the same combination of antibiotics resistance marker and diluted 1/100 into 50 mL of LB supplemented with ampicillin, kanamycin and streptomycin. Expression of VgrG, eagR and rhs1 was induced at 30 °C with 0.5 μM IPTG for 18 h at 16 °C. Cells were collected by centrifugation at 4,000 × g and resuspended in buffer A (50 mM Tris–HCl pH 8.5, 250 mM NaCl, 1 mM TCEP, Complete™ protease inhibitor cocktail (Sigma)). Cells were broken by sonication on ice, and a cleared cell extract was obtained by centrifugation for 45 min at 20,000 × g. One hundred μL of Strep-Tactin Superflow resin (IBA Technology) equilibrated in buffer A was then added into the cell extract. After incubation for 1 h at 4 °C with gentle mixing, the resin was washed 5 times with 300 μL of buffer A, and proteins were eluted with 100 μL of elution buffer (buffer A supplemented with 2.5 mM desethylbistobutioin). Ten μL of 10× diluted protein extract and 7.5 μL of elution material was separated by SDS-PAGE and proteins were stained with InstantBlue™ (Sigma-Aldrich) or transferred onto nitrocellulose membrane. Western blots were performed by standard procedures, in PBS supplemented with 0.1% Tween-20 and 3% of Bovine Serum Albumin. Coated proteins were then incubated with commercial monoclonal His-Tag (clone 1B7G5, Proteintech catalogue #66005-1-Ig), Strep-Tag Classic (clone Strep-Tag II, Bio-Rad catalogue #MCA2489) or anti-FLAG (clone M2, Sigma-Aldrich catalogue #F3165) antibodies, and secondary antibodies coupled to the Alkaline Phosphatase, and revealed using 5-bromo-4-chloro-3-indolyl phosphate/nitro blue tetrazolium (BCIP/NBT) in presence of 10 mM MgCl₂ in alkaline buffer (pH 9).

**Cross-linking mass spectrometry**

*Sample preparation and cross-linking.* The VgrG-Rhs1-EagR complex was purified (reson 100 fold, culture as detailed above), the 10 most intense precursor ions that share B x 10×M HEPES pH 7.5, 250 mM NaCl, 1 mM TCEP, Complete™ protease inhibitor cocktail (Sigma) was used instead of buffer A to avoid the Tris buffer that interferes with chemical cross-linking. One hundred nmoles of the NNP9 cross-linker56 were added to 200 μg of the complex. After incubation for 30 min at 4 °C, the reaction was quenched by the addition of 50 mM ammonium bicarbonate for 15 min at 4 °C. Samples were flash frozen in liquid nitrogen until further analyzes. Protein digestion, cross-linked peptides enrichment, and mass spectrometry analysis were performed as previously described56.

**Enrichment of labeled peptides.** Peptides were purified by click-chemistry on a photocleavable support. Photo-cleavable alkyne (PCA) agarooses beads (1 μL, 10 nmol) were mixed with the tryptic peptides, in presence of copper(II) sulfate (51, Tris(benzyltriazolylmethyl)amine (THPTA; 251) and sodium ascorbate (501) (ratio) were added to remove unlabeled peptides. PCA agaroose beads were centrifuged for 90 min at 10,000 rpm for 90 min and supernatant was removed. The bead was washed with 200 μL of 70% methanol twice and twice with 0.1% formic acid to remove unlabeled peptides. PCA agaroose beads were washed four times with PCA buffer and to remove unlabeled peptides. PCA agaroose beads were centrifuged for 90 min at 10,000 rpm for 90 min and supernatant was removed. PCA agaroose beads were washed four times with 100 μL of 50 mM ammonium bicarbonate. Peptides were digested overnight on the bead, with a trypsin:protein ratio of 1:100 (wt/wt). Tryptic peptides were recovered from the bead after centrifugation.

**Data analysis.** All raw data were processed using MassSpec Studio V2.40. Data were analyzed using the “E-Value Generator” algorithm with PercentEValue-Threshold set at 0.1. Cross-linked peptide identification were confirmed manually and discrepancies in the cross-linked sites were resolved to ensure that 2 chromatographic peaks did not share the same pair of sites.

**Purification of EagR-Rhs1 complex for structural studies.** E. coli BL21(DE3) cells freshly transformed with pRSF-EagR-ST and pCfD-Rhs-FL carrying the D1338N, Y1351A and Y1375A mutations to avoid autocleavage and toxicity were grown overnight in LB and diluted 1/100 into 4 L of LB supplemented with kanamycin and streptomycin. At A600 = 0.8, expression of eagR and rhs1 was induced by the addition of 0.5 μM IPTG for 18 h at 16 °C. Cells were collected by centrifugation and diluted 1/10 resuspended in buffer A and broken using an EmulsiFlex-C5 (Avestin). The protein extract was cleared by centrifugation for 30 min at 20,000 × g, filtered through a 0.45 μM membrane (ClearLine) and loaded on a 1-mL StrepTrap HP column (Cytiva) equilibrated in buffer A. The column was washed with 20 mL of buffer A and proteins were eluted with 10 mL of buffer B (50 mM Tris–HCl pH 8.5, 250 mM NaCl, 1 mM TCEP, Complete™ protease inhibitor cocktail) and a peptide mixture was loaded and collected as indicated in the Experimental Section. Fractions of highest peptide concentrations were pooled and loaded on a Superose 6 10/300 column (Cytiva) equilibrated with 50 mM Tris–HCl pH 8.5, 250 mM NaCl. The EagR-ST-Rhs1L elution peak was further purified against we.

**Cross-linked peptides MS analysis.** Peptides eluted from PCA agaroose beads were analyzed by nanoLC-MS/MS using an EASY-nLC™ 1200 system (Thermo-Scientific) coupled to the nanoelectrospray ion source of an Orbitrap Q Exactive HF mass spectrometer (Thermo-Scientific). Peptides were loaded on an in-house packed nano-HPLC column (75 μm × 25 cm) containing C18 resin (Aeris PEPTIDE XB-C18, 1.7-μm particle size, Merck). They were separated using a 70-μm inner diameter (resolution 60k at m/z 200), the 10 most intense precursor ions above 4.2 × 105 were selected for MS/MS analysis. Cross-linked peptides enrichment, and mass spectrometry analysis were performed as previously described56.
complex eluted as a single monodisperse peak and was separated from the EagRST excess.

Cryo-EM sample preparation, data collection, and processing
Sample preparation. Quantifoil Cu300 mesh R2/2 grids were glow discharged at 0.3 mbar vacuum and 2 mA current for 35 s. Then, 4 µL of purified protein complex was added to the grid and blotted for 1.5 s in a Vitrobot instrument (FEI), at 100% humidity and 4 °C, at blot force 0, before being plunge-frozen in liquid ethane and transferred/stored in liquid nitrogen.

Data collection and processing. Data collection and processing flowchart are summarized in Supplementary Fig. 11. Data collection was performed in a 200 kV Talos Arctica (FEI) equipped with a K2 camera (Gatan) at 45k magnification, pixel size of 0.93 Å, total dose of 50 e-/Å² and 3.7 s exposition, using SerialEM version 3.8. A total of 8550 movies were collected and processed using the Relion 3.1 software. Two thousand particles were manually picked and submitted to 2D classification in order to generate initial templates, which were in turn used for template-based picking. 2,623,687 particles were extracted with a 280-pixel box size resized to 96 pixel and submitted to several rounds of 2D classification, resulting in 1,036,869 selected particles, which were re-extracted with a box size of 280 pixels with original pixel size. A global 3D refinement resulted in a 3.72 Å resolution map. The refinement was repeated with a mask on the Rhs1 barrel region, which led to a 3.46 Å consensus map. After per-particle CTF refinement and Bayesian polishing, this same dataset retrieved a 3.1 Å map, but in which the C-terminal region of the Rhs1 barrel was suboptimally defined. This map was used for a 3D classification without sample alignment using a mask on the Rhs1 barrel and a separation in eight classes, which retrieved two major classes and six classes containing poorly-aligned particles. The best-defined class was selected, containing 140,577 particles, which led to the final map at 3.1 Å upon masked refinement. The final map was sharpened using the Autosharpen tool from the Phenix package26 using the two half maps as input. De novo model building was performed by tracing the main chain manually in Coot63 and submitting it to the map-to-model tool from the Phenix package, followed by manual building in Coot, optimization in ISODLE, and finished using the Real-Space-Refine tool from the Phenix package. Model validation was analyzed in Coot and then verified using the Comprehensive validation tool from the Phenix package.

Bioinformatic analyses and modeling. Sequence conservation of Rhs1-like proteins was estimated by BlastIP and sequence alignment was made with standard algorithm parameters (NCBI). The alignment was then treated with JalView 2.11.1.4 to generate a consensus sequence logo. Transmembrane segments were predicted using TMHMM server v.2.065 and hydrophobicity was analyzed by ProtScale (Exasy) using Kyte & Doolittle scale.66 VgGR and EagGR 3D models were obtained using Galaxy Homomer66 tool on GalaxyWEB server using the P. aeruginosa VgGR(PDB: 4MTK; unpublished) and EagGR (PDB: 1TU; unpublished) structures as templates, and by imposing trimeric (VgrG) and dimeric (EagR) oligomeric states. Rhs1NT model was made using AphaFold267. Structures were visualized and figures were prepared using Chimera88.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The final cryo-EM density map of the P. laumondii Rhs1-EagR complex has been deposited to the Electron Microscopy Data Bank (EMDB) under the accession code EMD-13587. The final atomic model was deposited into the Protein Data Bank (PDB) under the accession code 6W6M. The proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD028652. The data are provided in the Supplementary Information files or can be obtained from the corresponding authors upon request. Uncropped gels and Western-blot are shown in Supplementary Fig. 12.

Received: 30 June 2021; Accepted: 15 November 2021; Published online: 01 December 2021

References
1. Zhang, D., de Souza, R. F., Anantharaman, V., Iyer, L. M. & Aravind, L. Polymorphic toxin systems: Comprehensive characterization of trafficking modes, processing, mechanisms of action, immunity and ecology using comparative genomics. Biol. Direct 7, 18 (2012).
2. Jamet, A. & Naffis, X. New players in the toxin field: polymorphic toxin systems in bacteria. mBio 6, e00285–00215 (2015).
3. A total of 8550 movies were collected and processed using the Relion 3.1 software. Two thousand particles were manually picked and submitted to 2D classification in order to generate initial templates, which were in turn used for template-based picking. 2,623,687 particles were extracted with a 280-pixel box size resized to 96 pixel and submitted to several rounds of 2D classification, resulting in 1,036,869 selected particles, which were re-extracted with a box size of 280 pixels with original pixel size. A global 3D refinement resulted in a 3.72 Å resolution map. The refinement was repeated with a mask on the Rhs1 barrel region, which led to a 3.46 Å consensus map. After per-particle CTF refinement and Bayesian polishing, this same dataset retrieved a 3.1 Å map, but in which the C-terminal region of the Rhs1 barrel was suboptimally defined. This map was used for a 3D classification without sample alignment using a mask on the Rhs1 barrel and a separation in eight classes, which retrieved two major classes and six classes containing poorly-aligned particles. The best-defined class was selected, containing 140,577 particles, which led to the final map at 3.1 Å upon masked refinement. The final map was sharpened using the Autosharpen tool from the Phenix package26 using the two half maps as input. De novo model building was performed by tracing the main chain manually in Coot63 and submitting it to the map-to-model tool from the Phenix package, followed by manual building in Coot, optimization in ISODLE, and finished using the Real-Space-Refine tool from the Phenix package. Model validation was analyzed in Coot and then verified using the Comprehensive validation tool from the Phenix package.

Bioinformatic analyses and modeling. Sequence conservation of Rhs1-like proteins was estimated by BlastIP and sequence alignment was made with standard algorithm parameters (NCBI). The alignment was then treated with JalView 2.11.1.4 to generate a consensus sequence logo. Transmembrane segments were predicted using TMHMM server v.2.065 and hydrophobicity was analyzed by ProtScale (Exasy) using Kyte & Doolittle scale.66 VgGR and EagGR 3D models were obtained using Galaxy Homomer66 tool on GalaxyWEB server using the P. aeruginosa VgGR(PDB: 4MTK; unpublished) and EagGR (PDB: 1TU; unpublished) structures as templates, and by imposing trimeric (VgrG) and dimeric (EagR) oligomeric states. Rhs1NT model was made using AphaFold267. Structures were visualized and figures were prepared using Chimera88.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The final cryo-EM density map of the P. laumondii Rhs1-EagR complex has been deposited to the Electron Microscopy Data Bank (EMDB) under the accession code EMD-13587. The final atomic model was deposited into the Protein Data Bank (PDB) under the accession code 6W6M. The proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD028652. The data are provided in the Supplementary Information files or can be obtained from the corresponding authors upon request. Uncropped gels and Western-blot are shown in Supplementary Fig. 12.

Received: 30 June 2021; Accepted: 15 November 2021; Published online: 01 December 2021

References
1. Zhang, D., de Souza, R. F., Anantharaman, V., Iyer, L. M. & Aravind, L. Polymorphic toxin systems: Comprehensive characterization of trafficking modes, processing, mechanisms of action, immunity and ecology using comparative genomics. Biol. Direct 7, 18 (2012).
2. Jamet, A. & Naffis, X. New players in the toxin field: polymorphic toxin systems in bacteria. mBio 6, e00285–00215 (2015).
3. A total of 8550 movies were collected and processed using the Relion 3.1 software. Two thousand particles were manually picked and submitted to 2D classification in order to generate initial templates, which were in turn used for template-based picking. 2,623,687 particles were extracted with a 280-pixel box size resized to 96 pixel and submitted to several rounds of 2D classification, resulting in 1,036,869 selected particles, which were re-extracted with a box size of 280 pixels with original pixel size. A global 3D refinement resulted in a 3.72 Å resolution map. The refinement was repeated with a mask on the Rhs1 barrel region, which led to a 3.46 Å consensus map. After per-particle CTF refinement and Bayesian polishing, this same dataset retrieved a 3.1 Å map, but in which the C-terminal region of the Rhs1 barrel was suboptimally defined. This map was used for a 3D classification without sample alignment using a mask on the Rhs1 barrel and a separation in eight classes, which retrieved two major classes and six classes containing poorly-aligned particles. The best-defined class was selected, containing 140,577 particles, which led to the final map at 3.1 Å upon masked refinement. The final map was sharpened using the Autosharpen tool from the Phenix package26 using the two half maps as input. De novo model building was performed by tracing the main chain manually in Coot63 and submitting it to the map-to-model tool from the Phenix package, followed by manual building in Coot, optimization in ISODLE, and finished using the Real-Space-Refine tool from the Phenix package. Model validation was analyzed in Coot and then verified using the Comprehensive validation tool from the Phenix package.

Bioinformatic analyses and modeling. Sequence conservation of Rhs1-like proteins was estimated by BlastIP and sequence alignment was made with standard algorithm parameters (NCBI). The alignment was then treated with JalView 2.11.1.4 to generate a consensus sequence logo. Transmembrane segments were predicted using TMHMM server v.2.065 and hydrophobicity was analyzed by ProtScale (Exasy) using Kyte & Doolittle scale.66 VgGR and EagGR 3D models were obtained using Galaxy Homomer66 tool on GalaxyWEB server using the P. aeruginosa VgGR(PDB: 4MTK; unpublished) and EagGR (PDB: 1TU; unpublished) structures as templates, and by imposing trimeric (VgrG) and dimeric (EagR) oligomeric states. Rhs1NT model was made using AphaFold267. Structures were visualized and figures were prepared using Chimera88.
Pissaridou, P. et al. The Pseudomonas aeruginosa T6SS-VgrG1b spike is an IHS-repeat-containing protein encapsulation device. *Nature 501*, 547–550 (2013).

Li, J. et al. Structural Basis for Teneurin Function in Circuit-Wiring: A Toxin Motif at the Synapse. *Cell 173*, 735–748.e15 (2018).

Roderer, D. & Raums, S. Tc Toxin Complexes: Assembly, Membrane Permeation, and Protein Translocation. *Annu. Rev. Microbiol. 73*, 247–265 (2019).

Landsberg, M. J. et al. 3D structure of the Yersinia entomophaga toxin complex and implications for insecticidal activity. *Proc. Natl Acad. Sci. USA 108*, 20544–20549 (2011).

Guo, L. et al. Photorhabdus luminescens W-14 insecticidal activity consists of at least two similar but distinct proteins. Purification and characterization of toxin A and toxin B. *J. Biol. Chem. 274*, 9836–9842 (1999).

Liu, D. et al. Insect resistance conferred by 283-kDa Photorhabdus luminescens protein TcdA in Arabidopsis thaliana. *Nat. Biotechnol. 21*, 1222–1228 (2003).

Brackmann, M., Nazarov, S., Wang, J. & Basler, M. Using Force to Punch Holes: Mechanics of Contractile Nanomachines. *Trends Cell Biol. 27*, 623–632 (2017).

Zoued, A. et al. Architecture and assembly of the Type VI secretion system. *Nat. Rev. Microbiol. 16*, 664–667 (2018).

Coulthrust, S. The Type VI secretion system: a versatile bacterial weapon. *Microbiol. (Read.) 165*, 503–515 (2019).

Cherrak, Y., Flaugnatti, N., Durand, E., Journet, L. & Cascales, E. Structure and Activity of the Type VI Secretion System. *Microbiol. Spectr. 7*, 0031–2019 (2019).

Wang, J., Brodmann, M. & Basler, M. Assembly and Subcellular Localization of Bacterial Type VI Secretion Systems. *Annu. Rev. Microbiol. 73*, 621–638 (2019).

Cascales, E. The type VI secretion toolkit. *EMBO Rep. 9*, 735–741 (2008).

Durand, E., Cambillau, C., Cascales, E. & Journet, L. VgrG, Tae, Tle, and beyond: the versatile arsenal of Type VI secretion effectors. *Trends Microbiol. 22*, 498–507 (2014).

Shneider, M. M. et al. PAAR-repeat proteins sharpen and diversify the type VI secretion system spike. *Nature 500*, 350–353 (2013).

Pissardou, P. et al. The Pseudomonas aeruginosa T6SS-VgrG1b spike is topped by a PAAR protein eliciting DNA damage to bacterial competitors. *Proc. Natl Acad. Sci. USA 115*, 12519–12524 (2018).

Hernandez, R. E., Gallegos-Monterroza, R. & Coulthurst, S. J. Type VI secretion system effector proteins: Effective weapons for bacterial competitiveness. *Cell Microbiol. 22*, e13241 (2020).

Quentin, D. et al. Mechanism of loading and translocation of type VI secretion system effector Tse6. *Nat. Microbiol. 3*, 1142–1152 (2018).

Juez-Alcubierre, D. Photorhabdus antibacterial Rhs polymorphic toxin inhibits translation through ADP-ribosylation of 235 ribosomal RNA. *Nucleic Acids Res 49*, 8348–8395 (2021).

Flaugnatti, N. et al. A phospholipase A1 antibacterial Type VI secretion effector interacts directly with the C-terminal domain of the VgrG spike protein for delivery. *Mol. Microbiol. 99*, 1099–1118 (2016).

Flaugnatti, N. et al. Structural basis for loading and inhibition of a bacterial T6SS phospholipase effector by the VgrG spike. *EMBO J. 39*, e104129 (2020).

Nury, C. et al. A novel bio-orthogonal cross-linker for improved protein/protein interaction analysis. *Anal. Chem. 87*, 1853–1860 (2015).

Meusch, D. et al. Mechanism of Tc toxin action revealed in molecular detail. *Nature 508*, 61–65 (2014).

Whitney, J. C. et al. An interbacterial NAD(P)(+) glycohydrolase toxin requires elongation factor Tu for delivery to target cells. *Cell 163*, 607–619 (2015).

Tang, J. Y., Bullen, N. P., Ahmad, S. & Whitney, J. C. Diverse NADase effector families mediate interbacterial antagonism via the Type VI secretion system. *J. Biol. Chem. 293*, 1504–1514 (2018).

Kenzelmann, D., Chiquet-Ehrismann, R. & Tucker, R. P. Teneurins, a transmembrane protein family involved in cell communication during neuronal development. *Cell Mol. Life Sci. 64*, 1452–1456 (2007).

Rey, M. et al. Advanced In Vivo Cross-Linking Mass Spectrometry Platform to Characterize Proteome-Wide Protein Interactions. *Anal. Chem. 93*, 4166–4174 (2021).

Serpe, V. et al. High Sensitivity Crosslink Detection Coupled with Integrative Structure Modeling in the Mass Spec Studio. *Mol. Cell Proteom. 15*, 3071–3080 (2016).

Zivanov, J. et al. New tools for automated high-resolution cryo-EM structure determination in RELION-3. *Elife 7*, e42166 (2018).

Adams, P. D. et al. PHENIX: building new software for automated crystallographic structure determination. *Acta Crystallogr D. Biol. Crystallogr 75*, 87–124 (2019).

Emseley, P. & Cowtan, K. Coot: model-building tools for molecular graphics. *Acta Crystallogr D. Biol. Crystallogr 60*, 2126–2132 (2004).

Krogh, A., Larsson, B., von Heijne, G. & Sonnhammer, E. L. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. *J. Mol. Biol. 305*, 567–580 (2001).

Kyte, J. & Doolittle, R. F. A simple method for displaying the hydropathic character of a protein. *J. Mol. Biol. 157*, 105–132 (1982).

Baek, M., Park, T., Heo, L., Park, C. & Seok, C. GalaxyHomoser: a web server for protein homo-oligomer structure prediction from a monomer sequence or structure. *Nucleic Acids Res 45*, W320–W324 (2017).

Jumper, J. et al. Highly accurate protein structure prediction with AlphaFold. *Nature 596*, 583–589 (2021).

Petterson, E. F. et al. UCSF Chimera-a visualization system for exploratory research and analysis. *J. Comput. Chem. 25*, 1605–1612 (2004).

Acknowledgements
We thank the members of the Cascales, Fronzes and Chamot-Rooke laboratories for discussions and support, Amaury Payelleville, Julien Brilland and Alain Givaudan (DGIMI, Montpellier, France) for material and discussions, Pascal Mansuelle and Régine Lebrun (Proteomic facility of the Mediterranean Institute of Microbiology, Marseille) for mass spectrometry analyses and N-terminal Edman sequencing, Artemis Kosta (Microscopy facility of the Mediterranean Institute of Microbiology, Marseille) and Eric Durand for technical assistance and advices for preliminary EM analyses, and Moly Ba, Isabelle Bringer, Annick Brun, Audrey Gozzi and Mathilde Valade for technical support. This work was funded by the Centre National de la Recherche Scientifique, the Aix-Marseille Universitè, the Bordeaux Univ., the Institut Pasteur and by grants from the Agence Universitaire de la Recherche (ANR-17-CE11-0039, ANR-20-CE11-0017), the Fondation pour la Recherche Médicale (FRM, DEQ20180339165), and the Fondation Bettencourt Schueller to EC. J.D. was supported by a FRM post-doctoral fellowship (SPF201809007142). This work has benefitted from the facilities and expertise of the Biophysical and Structural Chemistry Platform (BPCS) at IECB.

Author contributions
D.J., L.T.R., M.R., J.C-R., R.F. and E.C. contributed to research. J.C-R., R.F. and E.C. provided tools. D.J., L.T.R., M.R., J.C-R., R.F. and E.C. designed research. D.J., L.T.R. and M.R. performed research. J.C-R., R.F. and E.C. analyzed mass spectrometry analyses and N-terminal Edman sequencing, Artemis Kosta for technical support. J.C-R., R.F. and E.C. wrote the manuscript with contributions from L.T.R., M.R., J.C-R. and R.F. All authors contributed to and approved the final revision of the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-27388-0.

Correspondence and requests for materials should be addressed to Dukas Jurénas, Rémi Fronzes or Eric Cascales.

Peer review information *Nature Communications* thanks the anonymous reviewer(s) for their contribution to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.