Article



SOURCE

DATA

RANSPAREN

PROCESS

 $\widehat{\Box}$

OPEN

ACCESS

Marie-Louise R Francis^{1,†,‡}, Melissa N Webby^{1,†}, Nicholas G Housden¹, Renata Kaminska¹, Emma Elliston^{1,¶}, Boonyaporn Chinthammit^{1,§}, Natalya Lukoyanova², Colin Kleanthous^{1,*}

Abstract

Bacteria deploy weapons to kill their neighbours during competition for resources and to aid survival within microbiomes. Colicins were the first such antibacterial system identified, yet how these bacteriocins cross the outer membrane (OM) of Escherichia coli is unknown. Here, by solving the structures of translocation intermediates via cryo-EM and by imaging toxin import, we uncover the mechanism by which the Tol-dependent nuclease colicin E9 (ColE9) crosses the bacterial OM. We show that threading of ColE9's disordered N-terminal domain through two pores of the trimeric porin OmpF causes the colicin to disengage from its primary receptor, BtuB, and reorganises the translocon either side of the membrane. Subsequent import of ColE9 through the lumen of a single OmpF subunit is driven by the proton-motive force, which is delivered by the TolQ-TolR-TolA-TolB assembly. Our study answers longstanding questions, such as why OmpF is a better translocator than OmpC, and reconciles the mechanisms by which both Toland Ton-dependent bacteriocins cross the bacterial outer membrane.

Keywords bacteriocins; cryo-electron microscopy; fluorescent microscopy; Gram-negative bacteria; outer membrane

Subject Categories Membranes & Trafficking; Microbiology, Virology & Host Pathogen Interaction

DOI 10.15252/embj.2021108610 | Received 29 April 2021 | Revised 9 August 2021 | Accepted 17 August 2021 | Published online 13 September 2021 The EMBO Journal (2021) 40: e108610

Introduction

The asymmetric outer membrane (OM) of Gram-negative bacteria, composed of lipopolysaccharides (LPS) in the outer leaflet and phospholipids in the inner leaflet, shields the organism against environmental insults, the immune systems of plants and animals, bile salts in the human gut and several classes of antibiotics (Nikaido, 2003; Whitfield & Trent, 2014; Ranf, 2016; Vergalli et al, 2020). Bacteria have nevertheless evolved potent antibacterial systems to breach the OM that are integral to the internecine warfare between bacterial populations as they compete for space in which to grow and nutrients on which to feed (Granato et al, 2019). Attack strategies are typically of two types, those that depend on physical or close contact between bacterial cells and those mediated by diffusible molecules (Ruhe et al, 2020). The former includes contact-dependent inhibitors, where a toxin projected from an attacking cell binds a surface receptor of a recipient cell prior to import (Aoki et al, 2010), and type VI secretion, where toxins are delivered by a needle that punctures the OM (Basler et al, 2013). Diffusible toxins are generally referred to as bacteriocins and can be peptides or proteins. Here, we focus on a family of protein bacteriocins from the antimicrobial armoury of the Enterobacteriaceae and reveal their convoluted use of porins for OM transport.

THE

IOURNAI

EMR

Protein bacteriocins from the Enterobacteriaceae and Pseudomonadaceae (hereafter referred to as bacteriocins) are plasmid or chromosomally encoded multidomain toxins that kill cells through the action of a C-terminal cytotoxic domain orchestrated by N-terminal regions of the toxin (Kleanthous, 2010). Cytotoxic activity is expressed either in the periplasm, through depolarisation of the cytoplasmic membrane or cleavage of the lipid II peptidoglycan precursor, or in the cytoplasm through cleavage of specific tRNAs, ribosomal RNA or chromosomal DNA. Passage across the OM is catalysed by one of two proton-motive force (PMF)-linked assemblies in the inner membrane. Ton-dependent (group B) bacteriocins contact TonB via TonB-dependent transporters (TBDTs) in the OM, whereas Tol-dependent (group A) bacteriocins contact one or more components of the Tol-Pal system (Cascales et al, 2007). Nearly all Tol-dependent bacteriocins identified to-date enter bacteria using a porin in combination with an outer membrane protein receptor, which can be a neighbouring TBDT or the porin itself (Kleanthous,

¹ Department of Biochemistry, University of Oxford, Oxford, UK

² Department of Biological Sciences, ISMB, Birkbeck College, London, UK

^{*}Corresponding author. Tel: +44 1865 613370; E-mail: colin.kleanthous@bioch.ox.ac.uk

[†]These authors contributed equally to this work

[‡]Present address: NovaBiotics, Aberdeen, UK

[¶]Present address: Arnold & Porter Kaye Scholer LLP, London, UK

[§]Present address: University College London, London, UK

2010). Escherichia coli porins OmpF and OmpC are abundant trimeric β -barrel outer membrane proteins with narrow channels running through them that taper to an eyelet, 7–8 Å at the narrowest point (Nikaido, 2003). The dimensions and electrostatic nature of the eyelet allow the diffusion of nutrients and metabolites (< 600 Da) but excludes large antibiotics such as vancomycin and folded proteins.

Recent work has shown that Ton-dependent bacteriocins specific for Pseudomonas aeruginosa are transported unfolded through TBDTs (White et al, 2017). Import is energised by the PMF via TonB and its associated ExbB-ExbD stator complex. By contrast, controversy surrounds how Tol-dependent bacteriocins exploit porins to cross the OM and whether this transport is energised. The present work focuses on the group A endonuclease (DNase) colicin, ColE9, a member of the E group of colicins (ColE2-E9), all of which use the same import machinery but target different nucleic acids in the cytoplasm (DNA, rRNA or tRNAs) (Kleanthous, 2010). We demonstrate using cryo-EM and live-cell imaging that ColE9 exploits the trimeric porin OmpF to cross the OM of E. coli by a combination of nonenergised and energised transport. We show that following binding to its TBDT receptor, BtuB, ColE9 threads its disordered N-terminus through two subunits of OmpF both as a means of capturing a component of the PMF-coupled Tol-Pal system in the periplasm but also to disengage the toxin from its receptor. ColE9 then translocates unfolded through the narrow lumen of a single OmpF subunit by a process that is entirely driven by the PMF. We also demonstrate that large fluorophores normally excluded by the pores of OmpF can be forcibly brought into the cell by attaching them to the colicin.

Results

Cryo-EM structure of the ColE9 outer membrane translocon

ColE9 is a plasmid-encoded heterodimer comprising a 60-kDa toxin bound to a small immunity protein, Im9. The two proteins are released together from E. coli cells in response to environmental stress (Cooper & James, 1984). Immunity proteins are high-affinity inhibitors that are co-expressed with the colicin and protect the producing cell from its cytotoxic action (Kleanthous et al, 1999), but are jettisoned during import of the bacteriocin (Vankemmelbeke et al, 2013). ColE9 has four functionally annotated domains (Fig 1A) (Klein et al, 2016). At its N-terminus is an 83 amino acid intrinsically unstructured translocation domain (IUTD) that houses three protein-protein interaction epitopes (Housden et al, 2010); a 16-residue TolB-binding epitope (TBE) associates with the periplasmic protein TolB which is flanked by two OmpF-binding sites (OBS1 and OBS2). The IUTD is followed by a folded α/β domain which is thought to be associated with inner membrane translocation and generally referred to as the translocation or T-domain (Sharp et al, 2017). Following the T-domain is a long coiled-coil receptor-binding (R-) domain, the apex of which binds the vitamin B₁₂ transporter BtuB on the cell surface (Fig 1B). Finally, at the C-terminus of the colicin is a cytotoxic DNase to which the immunity protein Im9 is bound. Following import and ejection of Im9, the internalised DNase elicits cell death through random cleavages of the bacterial genome. The ColE9 DNase is a member of the H-N-H/ββα-Me class of metal-dependent endonucleases (Pommer et al, 2001), which includes apoptotic nucleases and the core nuclease of CRISPR/Cas9. Other cytotoxic domains are also delivered by the same toxin chassis, including the RNase of ColE3 that inhibits protein synthesis by site-specific cleavage of the ribosomal A-site (Ng *et al*, 2010). The structures of ColE3 (Soelaiman *et al*, 2001) and ColE9 (Klein *et al*, 2016) have been reported previously as have the binary complexes of ColE3 R-domain-BtuB (Kurisu *et al*, 2003), ColE9 OBS1-OmpF (Housden *et al*, 2010) and ColE9 TBE-TolB (Loftus *et al*, 2006).

Tol-dependent colicins assemble a translocon at the cell surface that involves its periplasmic target along with the OM receptor and translocator which together establish the import pathway of the toxin. No structures have yet been reported for any colicin translocon and so there is little understanding of how these assemblies underpin OM transport. We have reported previously the isolation and purification of the entire OM translocon for ColE9 in which the toxin, bound to TolB, BtuB and OmpF, is solubilised in 1% βoctylglucoside (β -OG) (Housden *et al*, 2013). In addition, the translocon contains an engineered disulphide bond at the periphery of the ColE9 TBE-TolB interface, designed to stabilise the assembled complex without impacting the protein-protein interface (Housden et al, 2013). We followed the same procedure, which combines in vivo and in vitro approaches, but, for ease of assembly (see Materials and Methods), using ColE9 truncated at the end of the Rdomain and hence missing both the DNase and Im9. The assembled translocon was transferred into amphipols and vitrified on graphene oxide coated grids for visualisation of individual particles by cryo-EM (see Materials and Methods for details).

Single particle analysis revealed the presence of two populations of ColE9 translocon complexes, one where the translocon was intact and a second class where BtuB was absent, which we refer to as the partial translocon (Appendix Figs S1 and S2). The final postprocessed map of the full translocon (Fig 1C and D) is at an overall resolution of 4.7 Å, according to the gold-standard FSC method, with local resolutions ranging from 4.7 to 16 Å. The absence of BtuB in the partial translocon, most likely due to receptor dissociation during its deposition on grids, resulted in an improvement in map resolution (Fig 1E and F), which varies from 3.7 to 6 Å (FSC). Masked subtraction of BtuB from the full translocon map and refinement of these subtracted particles did not improve map quality, suggesting that it is the loss of BtuB from the complex which resulted in the increase in map resolution. Density for ColE9 was better resolved in the full compared to the partial translocon where only the T-domain was resolved. This is most likely due to the ColE9 R-domain providing some conformational rigidity in the fully assembled translocon through its interaction with BtuB. This contention is supported by the loss of density for the R-domain in the partial translocon map. By contrast, density for TolB was better resolved in the partial translocon map. The partial translocon map presented here is similar to a negative stain map of a proteolysed version of the translocon (comprising of OmpF, TolB and ColE9 residues 1-122), reported previously at a resolution ~20 Å (Housden et al, 2013). However, the lower resolution of this earlier structure meant little information was forthcoming about ColE9 interactions with OmpF.

Isolated crystallographic structures for the majority of ColE9 translocon components provided a starting point for model building into the full and partial translocon maps. Although the resolution of the full translocon map did not allow for local refinement of docked



Figure 1. Cryo-EM structures of the ColE9 outer membrane translocon.

- A Schematic of the ColE9 sequence showing its constituent domains: an intrinsically unstructured translocation domain (IUTD) at the N-terminus is followed by three structured domains involved in translocation (T), receptor (R) binding and cytotoxicity (C). The IUTD houses three linear protein-protein interaction epitopes, two OmpF-binding sites (OBS1, OBS2) flank a TolB-binding epitope (TBE). Residue numbers denote position in ColE9 sequence.
- B Cartoon of the ColE9 OM translocon. ColE9 (*orange*) exploits the vitamin B₁₂ transporter BtuB (*grey*) as its extracellular receptor and the porin OmpF (*green*) for threading its N-terminal IUTD (*solid black line*) through to the periplasm where it captures TolB (*blue*). *Star* represents the primary site on the ColE9 DNase domain (K469C) where fluorophores were covalently attached throughout this study.
- C Cryo-EM map of the fully assembled CoIE9 translocon, with local resolution range 4.5–16 Å. Component proteins are coloured as in panel B. The structure shows extracellular CoIE9 creating a protein bridge between the two OMPs, BtuB and OmpF. The β-barrel of BtuB is tilted 35° relative to that of OmpF. ToIB is located on the periplasmic side of OmpF, with no associations to BtuB. CoIE9 and ToIB regions of the map have weaker density than those of the β-barrels.
- D Model of the intact CoIE9 translocon generated after docking and rigid-body refinement of individual structures of CoIE9 residues 85–580 (PDB ID 5EW5), OmpF (3K19), BtuB (PDB ID 2YSU) and ToIB (PBD ID 4JML).
- E Cryo-EM map of the partial ColE9 translocon with an average resolution of 3.7 Å, which has density consistent with OmpF, TolB, and ColE9 residues 3–314. Map is coloured based on component parts shown in panel B. TolB is much better resolved here than in the full translocon map in panel C, although ColE9 density is weaker. ColE9 and TolB density align on the extracellular and periplasmic side of OmpF, respectively.
- F The refined structure of the partial translocon, generated by docking and refinement of CoIE9 residues 85–580 (PDB ID 5EW5), OmpF (PDB ID 3K19) and ToIB-CoIE9 TBE (PDB ID 4JML). CoIE9 residues 3–67 were built *de novo*.

structures, rigid-body refinement were carried out to generate a molecular model. For the partial translocon model, OmpF was docked and refined locally along with the ColE9 IUTD (residues 3–67), which were built *de novo* into the map. The remaining ColE9 T-domain residues (85–316) and TolB-ColE9 (residues 32–47) were rigid-body refined into the partial translocon map due to the lower resolution associated with these parts of the map.

In the fully assembled translocon model, the 22-stranded β -barrel of BtuB sits at a 35° angle relative to the trimeric β -barrel structure of OmpF (Fig 1C and D). Unmodelled density, most likely that of a single molecule of LPS, is wedged between the two proteins which likely contributes to displacement of BtuB relative to OmpF. The

region of unmodelled density in the full translocon map aligns with an LPS binding site (site A) observed in the structure of the OmpF orthologue from *Enterobacter cloacae* (PDB ID 5FVN), further supporting this as lipid density. Consistent with this interpretation, native state mass spectrometry has shown previously that a single LPS molecule remains associated with the purified translocon complex (Housden *et al*, 2013). Since there are no examples in the PDB of heterologous bacterial outer membrane proteins residing next to each other it is not clear if the 35° tilt reflects the natural position of BtuB in the OM or is a consequence of how the ColE9 translocon was assembled. We note however that if the β -barrels of OmpF and BtuB were flush in the membrane, the 45° trajectory of the ColE9 R-domain-BtuB complex, first shown for the ColE3 Rdomain-BtuB complex (0.95 Å rmsd) (Kurisu *et al*, 2003), would project the T-domain beyond OmpF. The coiled-coil nature of the Rdomain, with BtuB at its apex, means the cytotoxic DNase domain although absent from the current structure would sit ~60 Å above OmpF. Finally, TolB is held on the periplasmic side of OmpF through its association with the ColE9 TBE but is not well-resolved in the full translocon model (Fig 1C and D).

The partial translocon is comprised of OmpF, the IUTD and Tdomain of ColE9 and TolB (Fig 1E and F). Previous work on the translocation mechanism of group A colicins has demonstrated that these colicins contact their periplasmic binding partners (typically TolB and/or TolA) by a direct epitope delivery mechanism following binding to the cell surface receptor (Housden et al, 2010; Jansen et al, 2020). The colicin threads its IUTD through two of OmpF's three pores thereby presenting a tethered protein-protein interaction epitope to the periplasm. In the case of ColE9, the TBE promotes contact between TolB and TolA in the inner membrane which is coupled to the PMF through the stator proteins, TolQ and TolR (Bonsor et al, 2009). The partial translocon structure of ColE9 shows how the IUTD (residues 3-67) navigates through the lumen of one OmpF subunit in order to thread back up into a neighbouring subunit. The end result is that each OBS is lodged within an individual subunit of OmpF: OBS1 (residues 2-18) is bound to subunit 1 while OBS2 (residues 54-63) is bound to subunit 2, leaving subunit 3 free of colicin. The relative positions of the two OBSs in the partial translocon model implies that subunit 2 is the entry port for the ColE9 IUTD following its initial binding to BtuB in the OM.

The resolution of the full translocon does not allow for detailed comparison of OBS binding with that of the partial translocon (details below); however, an overlay of the two models reveals that the ColE9 T-domain undergoes a large-scale movement, rotating by 15° and moving vertically by 12 Å along the rotation axis (Fig 2A–C, Movie EV1). In the full translocon, the orientation of the ColE9 T-domain is fixed by the R-domain's contact with BtuB. Rotation of the ColE9 T-domain in the partial translocon to achieve the orientation adopted in the full translocon structure results in a poorer fit to map density. Importantly, rotation of the ColE9 T-domain from the full to the partial translocon repositions ColE9 from a central position above the OmpF trimer to become localised above subunit 2 within which OBS2 is bound (Fig 3A and C).

TolB also undergoes large-scale movements when transitioning from the full to partial translocon structures, which are likely linked to the re-arrangement of the ColE9 T-domain on the extracellular side of the porin (Fig 2D and E, Movie EV1). The position of TolB is rotated by ~50° between the two structures and in the partial translocon is located closer to OmpF through a ~8 Å upward movement along the axis of rotation. The positioning of TolB closer to OmpF in the partial translocon structure is stabilised by additional interactions between the periplasmic region of OmpF subunit 1 and the IUTD sequence immediately preceding the TBE (Fig 3G). Irrespective of the observed movements between the two cryo-EM models, TolB remains located beneath subunit 2 within which OBS2 is bound.

In summary, we have solved the first cryo-EM structure of a colicin OM translocon. The ColE9 translocon includes the OMPs BtuB and OmpF, as well as the periplasmic target TolB. We have also solved a partial translocon structure in which the primary

receptor BtuB is lost from the complex, which leads to substantial rearrangements of ColE9 and TolB above and below the membrane, respectively. As we demonstrate below, these structural changes within the translocon are integral to the mechanism by which ColE9 translocates across the OM.

ColE9 IUTD interactions with the subunits of OmpF

The higher resolution structure of the partial ColE9 translocon provides a molecular explanation as to how bacteriocins exploit porins to establish their translocon complexes at the OM. The lumen of an OmpF monomer is not a straight channel but akin to an hourglass (Fig 3A-C). The outer (extracellular side) and inner (periplasmic side) vestibules of OmpF have diameters of 25 and 30 Å, respectively, constricting at the eyelet of the channel to ~7 Å. After the IUTD region passes through OmpF subunit 2, OBS1 threads back up into subunit 1 so that it snakes along the bottom of the lumen at its widest point and finishes up at the eyelet (Fig 3A and B). By contrast, OBS2 spans the eyelet of OmpF subunit 2 via a kink created at Gly61, which is enforced by the hydrogen bonding of neighbouring residues, Ser60 and Arg62 (Fig 3C). After OBS2 traverses the eyelet, the remaining ColE9 sequence does not track the OmpF interior as observed for OBS1, instead remaining more centrally located in the inner vestibule.

The bound conformations of the two OBSs are stabilised by electrostatic and hydrogen bonding interactions (Fig 3D-H). The vestibules and eyelet of OmpF are highly charged environments; the porin exhibits slight cation selectivity in solute diffusion studies (Basle et al, 2006). Charged patches within the eyelet of OmpF are matched by opposing charges of the OBS sequences (Fig 3D and E). In addition to these interactions, networks of hydrogen bonds lock OBS1 and OBS2 of ColE9 into defined conformations within their respective OmpF subunits (Fig 3F and H). Key OBS1 residues that interact with subunit 1 include Asp5, Arg7, His9, Thr11 and Ser15, (Fig 3F and G). Previous studies have shown that mutation of Asp5, Arg7 or Thr11 to alanine substantially weakens OmpF binding by OBS1 while an Asp5Ala/Arg7Ala double mutant abolishes OmpF binding, consistent with their essential role in stabilising the OBS1-OmpF subunit 1 complex (Housden et al, 2005, 2010). We have reported previously the 3 Å crystal structure of an OBS1 peptide bound to OmpF (Housden et al, 2010). At the time, there was uncertainty as to the orientation of the peptide, which was refined with the N-terminus facing the periplasm. We re-processed these data and rebuilt the model based on the orientation observed in the cryo-EM data for the partial translocon, validating this re-processed model (Appendix Fig S2K-M). The new model shows conclusively that OBS1 has its N-terminus pointing towards the extracellular environment (Fig 3A) and in both models the conformation adopted by OBS1 is identical. This orientation agrees with recent electrophysiological data and live-cell imaging experiments all of which show OBS1 binding subunit 1 of OmpF from the periplasm (Housden et al, 2018; Lee et al, 2020). In contrast to OBS1, the hydrogen bond network that stabilises OBS2 within subunit 2 of OmpF primarily involves OBS2 backbone atoms and OmpF side-chains (Fig 3H), likely explained by the high glycine content within the OBS2 sequence (6/11 residues are glycine, compared to 6/16 for OBS1). The high glycine content also explains how OBS2 can bind in an extended conformation to OmpF, traversing the narrow eyelet.



Figure 2. Large-scale structural rearrangements accompany the loss of BtuB from the ColE9 translocon.

- A Superposition of the complete and partial CoIE9 translocon structures (*grey*) aligned on OmpF. ToIB in *red* and *orange* denote the full and partial translocons, respectively, CoIE9 T-domain is presented in *crimson* and *pale orange* for the full and partial translocons, respectively.
- B Side view comparison showing the relative positions of the CoIE9 T-domain (residues 85–316) in the two structures and highlighting the 15° rotation that occurs transitioning from the full (*crimson*) to the partial (*pale orange*) translocon.
- C Extracellular view of the CoIE9 T-domain position, with the OmpF trimer shown in the background. The loss of BtuB from the translocon complex elicits a 12 Å movement along the axis of rotation (*black arrow*) that results in repositioning of the T-domain from a central location (*crimson*) to above subunit two of OmpF (*pale orange*).
- D TolB undergoes both rotation and translation when transitioning from the full (red) to the partial translocon (orange). The C-terminal β-propeller domain of TolB, which binds the ColE9 TBE, moves towards OmpF in the OM by '8 Å along the rotation axis.
- E View along the rotation axis from the periplasmic side of OmpF (grey surface) highlighting the 50° rotation that TolB undergoes upon loss of BtuB from the translocon complex.

In conclusion, the bound conformations of the OBS epitopes within the electrostatic pores of OmpF are stabilised by salt bridges, backbone and sidechain hydrogen bonds and aided by their high glycine content. The specific binding modes of each OBS is explained by their specific set of interactions within identical OmpF pores and, in the case of OBS1, through directional insertion into the binding site from the periplasm.

Threading of CoIE9's IUTD through the pores of OmpF disengages the toxin from its primary receptor, BtuB

The structural differences evident between the full and partial ColE9 translocons centre around the loss of BtuB from the complex. Concomitant with this loss is the restructuring of the ColE9 T-

domain and TolB so that, linked by the OBS2 epitope, the two proteins become positioned immediately above and below, respectively, subunit 2 of OmpF. Given the precise nature of these rearrangements, we hypothesised that the partial translocon may be indicative of an intermediate state subsequent to ColE9 docking onto the cell surface. To test this hypothesis, we exploited the observation that the isolated 76-residue R-domain of ColE9 inhibits the growth of an *E. coli* strain dependent on vitamin B₁₂ (Fig 4A) (Penfold *et al*, 2000). *Escherichia coli* 113/3 is a *metE* mutant that requires vitamin B₁₂ to synthesise methionine. Since the binding sites of the ColE9 R-domain and vitamin B₁₂ overlap substantially, and their low nM affinities are similar (Gudmundsdottir *et al*, 1988; Kurisu *et al*, 2003; Housden *et al*, 2005), addition of excess Rdomain to *E. coli* 113/3 cells grown in minimal media supplemented with vitamin B_{12} results in the suppression of cell growth. We repeated this experiment using an identical concentration of intact ColE9 harbouring a single mutation, W39A, in the disordered IUTD. Substitution for alanine inactivates the toxin by abolishing binding to TolB in the periplasm, which in turn uncouples the colicin from the remainder of the PMF-linked TolQ-TolR-TolA complex in the IM (Hands *et al*, 2005). Hence, while all other OM interactions of the colicin are those of the wild-type toxin, including interactions of its coiled-coil R-domain with BtuB, ColE9 W39A cannot translocate and remains on the cell surface. Addition of ColE9 W39A to *E. coli* 113/3 largely restored cell growth under conditions where the isolated R-domain inhibited growth (Fig 4A–C). These data imply, along with the cryo-EM data, that subsequent to binding BtuB in the OM, threading of the ColE9 IUTD through two subunits of OmpF to form the translocon results in the toxin disengaging from BtuB. This translocon-induced dissociation is not an energised process since ColE9 W39A is unable to engage with the PMF-linked Tol system. We next investigated if and when the PMF is engaged in OM transport of the colicin.



Figure 3. Distinct modes of CoIE9 OBS1 and OBS2 recognition within the pores of OmpF enable threading and the tethered presentation of the TBE to the periplasm.

- A ColE9 residues 3–67 of the IUTD in the partial translocon structure pass through OmpF subunit 2 and then back up into subunit 1. As a result, OBS1 (gold) docks within subunit 1 and OBS2 (orange) within subunit 2. The ColE9 TBE motif (pink) interacts with TolB and is positioned below subunit 2, above which the ColE9 T-domain is located.
- B Surface representation of OmpF subunit 1 (grey) and ColE9 residues 3–24 (gold) containing OBS1 sequence. ColE9 OBS1 binds the inner vestibule of OmpF subunit 1 such that it runs along the edge of the vestibule, stopping at the eyelet.
- C CoIE9 residues 45–67 (orange) encompassing OBS2 traverse the pore of OmpF subunit 2 (grey). After passing through the eyelet, OBS2 tracks more centrally in the inner vestibule unlike in panel C where residues 3–24 trail the side of the pore.
- D OmpF subunit 1 displayed as an electrostatic surface (same cut-through as in B), revealing a patch of negative charge located on the extracellular side of the eyelet that interacts with OBS1, also shown as an electrostatic surface adjacent to the subunit.
- E OmpF subunit 2 displayed as an electrostatic surface (same cut-through as in C), with OBS2 also shown as an electrostatic surface adjacent to the subunit. Projecting out of the page for OBS2 is a positively charged region flanked by negative charges.
- F A zoom-in of grey box in panel A highlighting hydrogen bonding network between residues 3–7 of OBS1 (gold sticks) and nearby residues within OmpF subunit 1 (grey sticks).
- G OBS1 residues 11–17 (gold sticks) also interact with the base of OmpF subunit 1 (grey sticks) in the periplasm. Region is a zoom-in of blue box in panel A.
- H OmpF subunit 2 residues (*grey sticks*) form a hydrogen bond network with the backbone of residues of CoIE9 OBS2. Region is a zoom-in of brown box in panel A. All electrostatic surfaces shown in panels D and E were calculated using the APBS plugin within Pymol.

The electrical potential of the PMF drives ColE9 import across the OM

Assembly of ColE9's OM translocon occurs without the input of cellular energy, including dissociation from the BtuB receptor. At this point in the import process, ColE9 is composed of multiple folded domains so this raised the question of whether transport across the OM involved the PMF in order that domain unfolding could occur. Earlier studies suggested some form of unfolding to be involved, inferred from the activating effects of urea on colicin toxicity and the inhibitory effects of intramolecular disulphide bonds (Benedetti *et al*, 1992; Housden *et al*, 2005). However, the involvement of the PMF in Tol-dependent nuclease colicin entry is controversial. A recent model, for example, suggests that instead of energised transport, passive, electrostatically driven ratcheting of the colicin nuclease occurs through the one remaining unoccupied subunit of OmpF (subunit 3 in the present work) once the translocon is formed (Cramer *et al*, 2018).

A major issue the bacteriocin field has faced thus far in establishing if transport across the OM is energised is the lack of appropriate measures of transport that are not dependent on cell death as a readout. To address this problem, we developed a live-cell imaging platform using fluorescently labelled ColE9. A single cysteine mutation was engineered at Lys469 in the ColE9 DNase domain (Fig 1B), which would be the last domain of the colicin to enter if the toxin was pulled into the cell by the Tol-Pal system. Lysine 469 is adjacent to the DNase active site (Appendix Fig S3A). The mutant was labelled initially with Alexa Fluor (AF) 647 and excess dye removed by gel-filtration chromatography. Labelling efficiency was typically 90-100% for this and all other fluorophores used in this study (see Materials and Methods for details). The impact of labelling ColE9 K469C on biological activity was assessed by a plate-killing assay, where serial dilutions of the colicin were pipetted onto a lawn of susceptible E. coli JM83 cells and grown overnight (Appendix Fig S4). Wild-type ColE9 typically shows killing activity against E. coli JM83 (a K-12 strain), visible down to picomolar concentrations. Labelling ColE9 K469C with AF647 reduced colicin activity by "three orders of magnitude relative to wild-type ColE9, which nevertheless represents significant toxicity. Interestingly, zones indicative of bacterial killing for ColE9 K469CAF647 took longer to appear than those of wild-type toxin, which is a point we return to below. The decrease in activity could be the result of the fluorophore affecting cytotoxicity and/or import. Alternatively, if labelling completely abolished colicin activity the residual toxicity might simply reflect the presence of unlabelled molecules.

In order to distinguish between these possibilities, we investigated the effect of labelling the same position with fluorophores of differing size and charge (AF488, AF568, AF647). We rationalised that if decreased colicin activity reflected unlabelled molecules then altering the fluorophore would have a negligible effect given that the maleimide chemistry used to attach the fluorophores was the same in each case. None of the fluorophores attached at this position affected significantly the ability of the isolated nuclease domain of ColE9 to degrade dsDNA (Appendix Fig S3B-E). We next repeated the E. coli JM83 cell-killing assay for the ColE9 K469C mutant labelled with each of the fluorophores. These experiments showed that the conjugated fluorophores affected colicin toxicity to varying degrees, with the least-to-most toxic being AF568> AF647>AF488 (Appendix Fig S4). Indeed, ColE9 K469CAF568 displayed very little colicin activity. Finally, we conducted further in vivo tests by assessing the impact of fluorophore labelling ColE9 K469C on the activation of the DNA damage-induced SOS response in E. coli cells (Appendix Fig S5A-D). Transported ColE9 readily activates SOS through cleavage of chromosomal DNA, which can be monitored by lux bioluminescence in an E. coli reporter strain (DPD1718) (see Materials and Methods). As reported previously (Vankemmelbeke et al, 2005), ColE9 above 20 nM completely arrested growth of DPD1718 in liquid culture at 37°C after 90 min and this corresponded to a peak in *lux* bioluminescence, which then subsided to zero as all cells were killed-off (Appendix Fig S5B). Lower concentrations of ColE9 resulted in the same initial increase in lux bioluminescence but did not subside because cell growth begins to outcompete colicin toxicity. While none of the fluorophore labelled ColE9 constructs (at 20 nM) significantly impacted growth of E. coli DPD1718, all resulted in lux bioluminescence albeit with altered kinetics (Appendix Fig S5C and D). The suppression of signal observed in these experiments mirrored their impact in the cell-killing plate assays (Appendix Fig S4). Combined, these experiments imply that fluorophore labelling of ColE9 K469C affects the kinetics of colicin-mediated killing (which is consistent with the



Figure 4. ColE9 threading through the subunits of OmpF disengages the toxin from its outer membrane receptor, BtuB.

- A Cartoon depicting the basis for the control where excess ColE9 R-domain (residues 348–418, *orange*) blocks binding of vitamin B_{12} (*red*) to BtuB (*grey*), impairing growth of *E. coli* 113/3, a B_{12} -dependent strain (Penfold *et al*, 2000).
- B Cartoon schematic of the two possible translocon outcomes for binding of CoIE9 W39A to BtuB and their implications on vitamin B₁₂ import in *E. coli* 113/3. The W39A mutation within the TBE of CoIE9 (*orange*) impairs ToIB (*blue*) binding thereby uncoupling the OM components of the translocon from the energised ToIQ-ToIR-ToIA complex in the inner membrane. If threading of the CoIE9 IUTD through OmpF (*green*) has no impact on CoIE9 binding to BtuB (*grey*) then vitamin B₁₂ (*red*) cannot enter cells to support growth. In contrast, if OmpF threading disengages CoIE9 from BtuB then vitamin B₁₂ can enter cells to support growth.
- C Growth curves for *E. coli* 113/3 in defined media and in the presence or absence of B₁₂ (\blacksquare and O, respectively). When challenged with 40 nM isolated R-domain (\blacktriangledown), cell growth was largely impaired whereas growth in the presence of 40 nM CoIE9 W39A (\square) approached that of the no-colicin control, consistent with OmpF threading causing the disengagement of CoIE9 from BtuB. Error bars represent the standard deviation across three biological replicates.
- Source data are available online for this figure.

slow appearance of killing zones for ColE9 K469C^{AF647}) and that the identity of the fluorophore is a significant factor. The molecular basis of this effect is explored further below. We conclude that the residual toxicity of fluorescently labelled ColE9 reflects a reduced capacity to transport into *E. coli* rather than the activity of unlabelled molecules.

Since fluorescently labelled ColE9 can be imported into *E. coli*, we exploited this approach to probe the entry mechanism and its PMF dependence using fluorescence microscopy. Our experiments focussed primarily on ColE9 K469C modified with AF647 because it circumvented cell auto-fluorescence issues associated with using

AF488 and was more active than AF568 in colicin-mediated killing assays. An active site mutation (H551A hereafter denoted as *: Appendix Fig S3A) was also introduced in the ColE9 K469C background to avoid killing cells during long time courses. Following the labelling of cellular DNA with Hoechst stain, ColE9* K469CAF647 was added to E. coli JM83 cultures, the excess fluorophore removed by repeated wash steps and cells imaged on agar pads by widefield fluorescence microscopy (see Materials and Methods for details). In contrast to whole cell labelling by Hoechst, ColE9* K469CAF647 predominantly labelled the cell periphery (Fig 5A and B). To ensure labelling was specific for the ColE9 receptor BtuB, the experiment was repeated using a *btuB* deletion mutant, which abolished labelling by ColE9* K469CAF647 (Fig 5C). To determine if ColE9* K469CAF647 was surface exposed, labelled cells were treated with trypsin prior to imaging. Following this treatment, significant fluorescence remained associated with the cells suggesting these were imported molecules protected from proteolysis (Fig 5A-C). Finally, we established that AF647 attached to other sites, within the R- and T-domains of ColE9 (K298C and S420C, respectively), were also protected against proteolysis following addition of singly labelled ColE9 constructs to E. coli JM83 cells (Appendix Fig S6). Based on these data, we conclude that the entire ColE9 molecule must be transported across the OM.

The energetics of ColE9 transport across the OM were investigated using ColE9* K469C^{AF647}. When cells were pre-treated with the protonophore CCCP, which disrupts both the proton gradient (ΔpH) and electrical potential ($\Delta\Psi$) components of the PMF, all ColE9* K469C^{AF647} fluorescence was lost from cells following the addition of trypsin (Fig 5A-C), with similar results observed for cells labelled with ColE9* K469C carrying AF488 or AF568 fluorophores (Appendix Fig S7A and B). Consistent with a role for PMF in ColE9* K469CAF647 import, pre-treatment of cells with ColB, a Tondependent pore-forming colicin that depolarises the inner membrane, also blocked uptake of ColE9* K469CAF647 across the OM similar to the effects of CCCP (Appendix Fig S7C and D). As a final test for the involvement of the PMF in ColE9* K469CAF647 import, nigericin and valinomycin, which inhibit the proton gradient (ΔpH) and electrical potential ($\Delta\Psi$), respectively, were added to cells prior to labelling (Appendix Fig S8). Of these, only valinomycin blocked the uptake of the colicin from which it can be inferred that it is the electrical potential of the PMF that drives ColE9 translocation across the OM. We conclude that following formation of its OM translocon at the cell surface, import of ColE9 across the membrane is an energised process that requires the electrical potential of the PMF.

ColE9 translocates through the pore of a single OmpF subunit

The structural changes evident within the ColE9 translocon in the absence of BtuB, whereby the T-domain becomes repositioned immediately above the same OmpF subunit (subunit 2) as TolB in the periplasm, are consistent with OmpF being the port of entry. Indeed, OmpF eyelet mutations are known to confer resistance to Tol-dependent colicins (Jeanteur *et al*, 1994). The absolute requirement for the electrical potential of the PMF to import ColE9 across the OM further supports this contention since this could provide the cellular energy needed to unfold the toxin for passage through the porin. We hypothesised that if ColE9 passes directly through the eyelet then the physicochemical properties of both the colicin and



Figure 5. ColE9* K469C^{AF647} translocation across the outer membrane is PMF-dependent.

ColE9* denotes ColE9 H551A, an active site mutation that inactivates the toxin.

- A Widefield fluorescence microscopy images of *E. coli* JM83 cells labelled with ColE9* K469C^{AF647} (1.5 μM) and Hoechst DNA stain (20 μM) for 30 min at 37°C with or without trypsin treatment with or without prior treatment with CCCP. Each panel shows the same cell in DIC (*grey*), Hoechst DNA stain (*blue*) and ColE9* K469C^{AF647} fluorescence (*red*). Overlays of Hoechst and ColE9* K469C^{AF647} fluorescence are also shown. ColE9* K469C^{AF647} remains bound to the OM in the presence of CCCP but this signal is lost on treatment with trypsin. Trypsin treatment in the absence of CCCP yields some cell-associated ColE9* K469C^{AF647} fluorescence that likely represents internalised molecules (see text). Scale bar, 1 μm.
- B Normalised fluorescence intensity profiles across *E. coli* JM83 cell widths for CCCP and/or trypsin treated cells. Fluorescence was measured in ImageJ for 15 cells per condition and normalised to more clearly show distribution profiles for each condition. CCCP-treated cells showed peripheral ColE9* K469C^{AF647} fluorescence consistent with the colicin being bound at the OM. Trypsin treatment of cells in the absence of CCCP results in loss of this peripheral fluorescence but the presence of mid-cell signal consistent with internalisation. Treatment with CCCP and trypsin removed all ColE9* K469C^{AF647} fluorescence from cells, as in A. Error bars represent % SEM.
- C Box and whisker plots showing mean pixel intensities for CoIE9* K469C^{AF647} fluorescence per cell measured for the indicated cells and condition used, whiskers represent minimum and maximum mean pixel intensity, box shows 1st and 3rd quartile with the median shown as a line. From left to right: *E. coli* JM83 cells; *E. coli* JM83 cells; *E. coli* JM83 cells in the presence of CCCP; *E. coli* JM83 cells following trypsin treatment showing significant CoIE9* K469C^{AF647} fluorescence remains associated with cells indicative of import; *E. coli* JM83 cells treated with CCCP and trypsin showing the complete loss of internalised CoIE9* K469C^{AF647} fluorescence. *n*, number of cells used, typically from 2 to 4 biological replicates. **** indicates a t value < 0.05 in a Student's t-test as a statistically significant result, ns indicates no significant difference as determined by Student's t-test.

the eyelet would be important factors in transport. A clue that this might be the case came from the experiments showing that three fluorophores (AF488, AF568, AF647) differentially affected the ability of ColE9 to induce the SOS response in E. coli DPD1718 cells (Appendix Fig S5); the fluorophores are negatively charged and have molecular weights (720-1,250 Da) significantly greater than are permissive in porin solute diffusion assays (Vergalli *et al*, 2020) yet did not impact colicin activity similarly. A complicating factor in interpreting these experiments however is the presence of both OmpF and OmpC in the OM of typical E. coli K-12 strains, the ratios of which change depending on the osmolarity, pH and temperature of the growth medium (Pratt et al, 1996). The two porins share 60% sequence identity, have similar trimer structures and eyelet dimensions, and can even form mixed trimers in the OM (Gehring & Nikaido, 1989). Where they differ however is the degree of negative charge at the eyelet; OmpC is significantly more electronegative than

OmpF (Fig 6A). We therefore assessed the impact of fluorophore labelling on the ability of ColE9 K469C to kill E. coli strains with either OmpF or OmpC in the OM. Plate-killing assays with wildtype, unlabelled ColE9 showed that cells with only OmpC in the outer membrane are less susceptible than cells with only OmpF (Fig 6B), which is consistent with early colicin literature (Mock & Pugsley, 1982). Labelling ColE9 K469C with AF488, AF568 or AF647 impacted colicin activity on cells with OmpF by up to 10³-fold. By contrast, cytotoxicity was essentially abolished in cells with OmpC in the outer membrane, equating to type="InSpacing_Modifier_Letters">~10⁹-fold loss of activity (Fig 6B). Lastly, we investigated the ability of AF647-labelled ColE9 to translocate into ompF- or ompCexpressing cells using widefield fluorescence microscopy with or without trypsin treatment. Here again, the presence of the fluorophore had a much greater impact on the ability of the colicin to enter cells with OmpC in the outer membrane relative to OmpF

(Fig 6C and D). We conclude that the slightly narrower pore and greater negative charge of the OmpC eyelet act as a significant impediment to ColE9 import when the colicin has a large (> 600 Da), negatively charged fluorophore covalently attached to its C-terminal DNase domain; an attachment that has little or no impact on nuclease activity (Appendix Fig S3). Since ColE9 import is dependent on the PMF and delivery of the PMF occurs via the N-terminal regions of the colicin, these results can only be explained if the entire colicin passes through the eyelet of the porin carrying the fluorophore with it.

The differential effects of fluorophores on ColE9 toxicity and import in cells harbouring OmpC or OmpF raised the question of why, when the dimensions of porin eyelets are similar, bulky fluorophores can still pass through OmpF? We hypothesised that the long time course (overnight) of the cell-killing assays might mask the import effects of bulky fluorophores on *E. coli* cells expressing *ompF*. To determine if more subtle effects were evident on shorter timescales, the kinetics of killing *E. coli* JW2203 cells, which only have OmpF in the outer membrane, were determined after addition of ColE9 K469C labelled with AF488, AF568 or AF647 (Fig 6E). The



Figure 6.

Figure 6. Labelling ColE9 with fluorophores impedes its translocation through the pores of porins.

- A Cut-through molecular surface images showing the distribution of charged residues (acidic, *red*; basic, *blue*) at the eyelets of trimeric OmpF (PDB ID 3K19) and OmpC (PDB ID 2J1N). The eyelet of OmpC is significantly more electronegative than that of OmpF. Electrostatic surfaces were calculated using the APBS plugin within Pymol.
- B Overnight plate assays comparing the cytotoxic activity of wild-type ColE9 with ColE9 K469C labelled with AF488, AF568 or AF647 against a lawn of *E. coli* with either OmpF (JW2203) or OmpC (JW0912) in the outer membrane (shown as +OmpF and +OmpC, respectively). Each plate was spotted with a serial dilution of the colicin (10 μM–2.4 pM, 4-fold dilution per spot). Wild-type ColE9 was more active against *E. coli* cells with OmpF in the outer membrane than OmpC. Labelling ColE9 K469C with fluorophores reduced colicin activity in cells with OmpF in the outer membrane but largely abolished colicin activity in cells with OmpF.
- C Widefield fluorescence microscopy images of *E. coli* JW2203 (+OmpF) and JW0912 (+OmpC) cells labelled with ColE9* K469C^{AF647} (1.5 μM) and Hoechst stain (20 μM) for 30 min at 37°C with or without trypsin treatment. Each panel shows the same cell in DIC (*grey*), Hoechst DNA stain (*blue*) and fluorescence of ColE9* K469C^{AF647} (*red*). Overlays of Hoechst and ColE9* K469C^{AF647} fluorescence are also shown. Data show that following trypsin treatment, significant ColE9* K469C^{AF647} fluorescence remains associated with *ompF*-expressing *E. coli*, whereas little or no fluorescence remains associated with *ompC*-expressing *E. coli*. Scale bar, 1 μm.
- D Box and whisker plots of *E. coli* JW2203 and *E. coli* JW0912 cells labelled with ColE9* K469C^{AF647} with and without trypsin treatment: whiskers represent minimum and maximum mean pixel intensity, box shows 1st and 3rd quartile with the median shown as a line. Microscopy data were collected as in C. The mean pixel intensity of ColE9* K469C^{AF647} per cell was measured for each cell condition. *n*, number of cells, typically from 3 or 4 biological replicates. Data show that ColE9* K469C^{AF647} translocates across the OM through OmpF and that this is impeded when *E. coli* has OmpC in the outer membrane.
- E First-order cell death kinetics for wild-type ColE9 or ColE9 K469C labelled with different AF dyes (AF488, AF568, AF647) against *E. coli* JW2203 with OmpF in the outer membrane. Cultures were incubated with 80 nM toxin at pH 7.5 and 37°C and at various time points the reaction stopped using trypsin at 37°C for 30 min, cells plated out and CFUs recorded. All time courses were conducted in the presence of chloramphenicol (20 µg/ml) to reversibly block cell division, which would otherwise compete with cell killing. Control data shown are for cells with no ColE9 added but where growth was inhibited by the presence of chloramphenicol. Cell-killing half-lives, obtained from the fitted first-order plots shown (error bars represent the standard deviation across two biological replicates), were as follows: wild-type ColE9, 2.5 min; ColE9 K469C^{AF647}, 29.6 min; ColE9 K469C^{AF568}, 78.8 min. The data show that the chemical nature of the fluorophore in the C-terminal DNase domain of ColE9 has a dramatic effect on the cell-killing kinetics of the colicin. The schematic shown alongside the kinetic data is a surface representation of AF568 (model Generated in coot), the fluorophore with the biggest impact on OmpF-mediated killing, manually grafted onto the structure of ColE9 OBS2 bound within subunit 2 of OmpF. The slow cell death kinetics of ColE9 K469C^{AF568} likely reflects the time taken to pass this bulky molecule through the narrow eyelet of the porin.

Source data are available online for this figure.

kinetics of cell death were monitored by measuring colony-forming units (CFUs) as a function of time, following the addition of trypsin to degrade extracellular colicin. Low chloramphenicol concentrations were used throughout these experiments to stop cells dividing but otherwise maintain cell viability thus simplifying interpretation of cell death kinetics. The data in Fig 6E demonstrate that the ColE9-mediated killing of ompF-expressing cells showed a first-order kinetic profile regardless of which fluorophore is attached. The fluorophores nevertheless had a profound impact on the half-life of this process; the half-life for cell death by unlabelled ColE9 was ~2.5 min, at 37°C, which increased almost two-fold for ColE9 K469CAF488, ~10-fold for ColE9 K469C^{AF647} and ~30-fold for ColE9 K469C^{AF568}. The relative order of fluorophore impact on the killing half-life (AF568>AF647>AF488) was the same as the order observed in the plate-killing assay with E. coli JM83 cells (Appendix Fig S4) and the induction of the SOS response in E. coli DPD1718 cells (Appendix Fig S5), where both strains express *ompF* and *ompC*. In conclusion, the kinetic profiles shown in Fig 6E reflect the rate-limited, energised transport of ColE9 molecules through the lumen of a single OmpF subunit following assembly of the OM translocon, which is a concept that is further explored in the Discussion.

Discussion

Model for the force-dependent import of ColE9 through OmpF

Combining cryo-EM structure determination of ColE9 translocon complexes with fluorescence microscopy data, and in conjunction with past biochemical, biophysical and structural data on this and related colicins, it is finally possible to unravel the complex interplay of non-energised processes and those powered by the PMF in the entry mechanism of Tol-dependent colicins (Fig 7).

Past work has shown that E group colicins assemble their OM translocons by first binding with nM affinity to BtuB at a 45° angle relative to the plane of the membrane from where the N-terminal IUTD captures a neighbouring OmpF (Fig 7A) (Kurisu et al, 2003; Housden et al, 2005). The cryo-EM structure of the intact ColE9 translocon reveals that the colicin also has to overcome a 35° tilt of BtuB relative to OmpF, which it does by having an R-domain with long enough reach. From this overhanging position, the IUTD associates with the pore of an OmpF subunit (designated subunit 2 in our structure), most likely due to its high local concentration and electrostatic attraction between the pore which is mildly cationic selective and the N-terminus of the colicin. The disordered polypeptide chain then reptates through subunit 2 of OmpF to deliver the TBE to the periplasm, capturing TolB at the expense of its endogenous binding partner, the OM lipoprotein Pal (Fig 7B) (Bonsor et al, 2007, 2009). ColE9 mimics the interactions of Pal; unlike Pal however, the colicin induces TolB to expose its Nterminus which houses the TolA binding site.

Concomitant with (or following) binding to TolB, the ColE9 OBS1 sequence inserts into subunit 1 of OmpF and OBS2 becomes stably bound within subunit 2. Threading through two subunits of an OmpF trimer is accompanied by three structural changes within the translocon. First, the R-domain of ColE9 dissociates from BtuB, which is consistent with the return of vitamin B₁₂-dependent growth for E. coli metE cells when contact between BtuB-bound ColE9 and the energised Tol system is abolished (Fig 4). We speculate that induced dissociation of the ColE9-BtuB complex may have its basis in the acute angles between the colicin's coiled-coil R-domain and BtuB and the R-domain-T-domain connection. Threading of the IUTD through the subunits of OmpF might apply a pulling force via the T-domain on the coiled-coil, the apex of which contacts BtuB, thereby disrupting the ColE9-BtuB complex. Second, BtuB dissociation removes an architectural constraint on repositioning of the colicin's T-domain, from a central position above OmpF to one that is



Figure 7. Model of CoIE9 translocon assembly and energised OM transport (see text for details).

- A ColE9 R-domain binds BtuB with nM affinity, positioning the T-domain and IUTD above a neighbouring OmpF trimer.
- B The CoIE9 IUTD translocates through subunit 2 of OmpF to deposit the TBE in the periplasm and capture ToIB at the expense of the OM lipoprotein Pal. The TBE allosterically promotes displacement of ToIB's N-terminus which constitutes the ToIA binding site. We propose this complex equates to the full translocon cryo-EM structure.
- C ColE9 OBS1 binds subunit 1 of OmpF from the periplasm. Threading through two of OmpFs three subunits triggers dissociation of the ColE9 R-domain-BtuB complex and reorients the T-domain above subunit 2 below which TolB is positioned in the periplasm. The docking of OBS1 also results in TolB moving closer to the opening of subunit 2. This complex, which we propose equates to the partial translocon cryo-EM structure, is now primed for contact with TolA in the inner membrane. TolA extension through the periplasm is coupled to the PMF via its stator proteins, TolQ and TolR.
- D Retraction of TolA (the molecular mechanism of which remains to be established) provides the driving force for pulling ColE9 bound to TolB into the periplasm through subunit 2 of OmpF, accompanied by unfolding of its constituent domains. It is at this point the immunity protein Im9 (not shown) would be displaced at the cell surface.
- E The ToIA-ToIB complex is thought to retract through the cell wall, which would bring CoIE9 close to the cytoplasmic membrane. The toxin likely refolds prior to transport across the cytoplasmic membrane, which involves the AAA⁺ATPase/protease FtsH (not shown) (Walker *et al*, 2007).

now directly above subunit 2 (Fig 7C). Third, OmpF threading is accompanied by rotation of the ColE9 TBE-TolB complex in the periplasm and formation of additional stabilising interactions between the colicin and the porin, all of which position the ColE9 TBE-TolB complex close to the periplasmic entrance of OmpF subunit 2.

It is at this point that optimal engagement between PMF-linked TolQ-TolR-TolA in the inner membrane and the ColE9 TBE-TolB complex likely occurs (Fig 7C). The physiological role of the TolQ-TolR-TolA assembly is to drive dissociation of TolB-Pal complexes at the OM, releasing Pal to bind the cell wall (Szczepaniak *et al*, 2020). As a consequence of this PMF-driven cycle, TolB is thought to be pulled through the cell wall by TolA. Hence, by virtue of its association with TolB, ColE9 is dragged unfolded into the periplasm through subunit 2 of OmpF, its C-terminal DNase domain the last to enter the cell (Fig 7D and E). Previous colicin translocation models have suggested Tol-dependent bacteriocins remain associated with their OM components while killing the cell (Benedetti *et al*, 1992; Duche, 2007). Our data suggest this not the case. Protease-

protection assays using ColE9 fluorescently labelled in all its constituent domains demonstrate that the entire ColE9 molecule enters the periplasm, driven by interactions at its N-terminus with the PMF-driven Tol-Pal system. Subsequent transport of the nuclease domain across the inner membrane requires the AAA⁺ATPase/protease FtsH. FtsH also proteolytically processes the domain as it enters the cytoplasm, although the underlying mechanism is unknown (Walker *et al*, 2007; Chauleau *et al*, 2011).

Forced unfolding of ColE9 during OM transport is consistent with several lines of evidence. First, our cryo-EM data, showing the structural changes that accompany ColE9 translocon formation, in which the T-domain and ColE9 TBE-TolB complex become aligned over the same OmpF subunit, suggesting they are the means by which the bacteriocin maximises its access to PMF-mediated force transduction through OmpF. Second, internal disulphide bonds within the coiled-coil domain block ColE9 entry, which can be explained by the cross-links blocking the unfolding necessary to allow entry through OmpF (Penfold *et al*, 2004). Third, *in vivo* experiments have shown that

release of Im9 (which forms a fM complex with the E9 DNase) at the cell surface is dependent on the electrical potential of the PMF (Vankemmelbeke *et al*, 2013), which is also the case for the entry of fluorescently labelled ColE9 in our experiments. These observations are consistent with the release of Im9 accompanying the force-dependent unfolding and import of the colicin DNase through OmpF. Fourth, *in vitro* atomic force microscopy measurements have demonstrated that when forces equivalent to those attainable by the PMF are applied to the isolated ColE9 DNase-Im9 complex the DNase domain melts, releasing Im9 but only when the force is applied at the N-terminus of the domain (Farrance *et al*, 2013). Viewed in the context of a translocating ColE9 molecule at the OM, the force exerted by the PMF would similarly be applied at the N-terminus of the toxin, with the surface of OmpF acting as the barrier against which the toxin is unfolded and Im9 displaced.

We conclude that the transport mechanism uncovered for ColE9 is likely to be the basis for the import of most Tol-dependent bacteriocins, the majority of which require porins. This will certainly be the case for nuclease colicins (E2-E9) that use BtuB as their receptor, and share high sequence identity with ColE9, but deliver rRNases or tRNases as well as DNases into the cell. This is also likely to be the case for pore-forming bacteriocins such as ColA, ColK and ColN, which use porins as translocators but different OM receptors (Klean-thous, 2010).

Ton- and Tol-dependent bacteriocins exploit analogous PMF-dependent mechanisms to cross the outer membrane through protein pores

The basic import principles we have uncovered for ColE9 highlights similarities with those recently described for the Ton-dependent bacteriocins pyocins S2 and S5, which target Pseudomonas aeruginosa. Both S2 and S5 associate with the common polysaccharide antigen on the cell surface and then exploit different TBDTs, FpvAI and FptA, respectively, as translocators to the periplasm (White et al, 2017; Behrens et al, 2020). Translocation of these bacteriocins involves two energised events. The first displaces the plug domain that occludes the central pore of the TBDT by a TonB-dependent mechanism, akin to that accompanying ligand uptake through the transporter. The second involves transport of the unfolded toxin through the now-open TBDT pore following delivery of the toxin's N-terminal TonB box to the periplasm. The mechanism we describe for Tol-dependent bacteriocins is analogous to the second of these steps. The similarity is underscored by the way in which force is exerted by TonB and TolA on their endogenous OM protein targets. The C-terminal domains of both proteins form structurally related, parallel β -sheets with their partners involving β -strand augmentation; TonB with the N-termini of TBDTs, TolA with the N-terminus of TolB (Pawelek et al, 2006; Szczepaniak et al, 2020). Such complexes are resistant to mechanical deformation (Chen et al, 2015). Hence, when the PMF-coupled stators ExbB-ExbD and TolQ-TolR, respectively, exert force on them instead of dissociating they remain bound, descending towards the inner membrane as TonB and TolA retract (Hickman et al, 2017). By hijacking these systems, both Ton- and Tol-dependent bacteriocins use the PMF to import themselves across the OM in unfolded form through either preexisting pores (porins) or pores opened transiently by the PMF (TBDTs).

Rate-limited transport of ColE9 through porins can be used to import large chemical entities into *E. coli*

The single-hit kinetic profiles observed for colicin-mediated killing of E. coli have been known since the 1960s (Reeves, 1965). However, the basis of this characteristic phenomenon has remained elusive. Our data reveal that this kinetic behaviour reflects the ratelimited transport of the colicin through the OM and not, as widely assumed, the killing activity of a single cytotoxic colicin molecule within the cell. The cell-killing half-life for ColE9 targeting E. coli cells with OmpF as the only porin in the OM (~2.5 min at 37°C and pH 7.5) changes significantly when Alexa Fluor dyes are attached to the DNase domain, by up to 30-fold in the case of AF568. We suggest that much of this additional time reflects repeated attempts by the Tol-Pal system, in conjunction with the PMF, to bring this single, bulky, negatively charged adduct through the narrow eyelet of OmpF subunit 2 (Fig 6E). The task of bringing this adduct through OmpC becomes essentially impossible because of the evelet's greater negative charge.

Glycopeptide antibiotics like vancomycin do not kill Gramnegative bacteria because they cannot partition into the OM and are too large to diffuse through porins. This impermeability problem is a major factor in multidrug resistance amongst Gram-negative pathogens. One approach being developed to circumvent these issues is to conjugate complex antibiotics to siderophores (so-called Trojan horse antibiotics) so that they can be actively transported through specific TBDTs, energised by the PMF-linked Ton-system (Klebba *et al*, 2021). Our data show that large, bulky, organic fluorophores that approach the size and complexity of vancomycin can be actively transported through the porins of *E. coli* by piggy-backing on a colicin. Colicins might therefore represent a new vehicle for the delivery of antibiotics that cannot otherwise penetrate the OM of Gramnegative bacteria.

Materials and Methods

Plasmids, bacterial strains and media

Whole plasmid site-directed mutagenesis was used to introduce a solvent accessible cysteine K469C in the cytotoxic domain of ColE9. Im9His6, cloned into pET21a to give pMLF07. pMLF07 was used to introduce H551A within the cytotoxic domain to give pMLF08, a cytotoxic inactive construct. Plasmid pEE01 was used to express ColB-His6 in pACYCduet-1. Plasmid pBC1 expressing ColE9 DNase K469C. Im9His6, containing ColE9 DNase domain with K469C single cysteine, in pET21d, was generated from pRJ353. Plasmid pREN151 was amplified from a pCS4 derivative and inserted between NdeI and XhoI sites of pET21a to enable expression and purification with C-terminal His6 tag. All E. coli strains and plasmids used in this study are listed in Appendix Tables S2 and S3. Strains were routinely grown at 37°C in either Lysogeny broth (LB) (Miller 1972) or M9 minimal media or plated on LB-agar. Supplemented M9 minimal media contained 47.78 mM Na₂HPO₄, 22 mM KH₂PO₄, 8.56 mM NaCl, 2 mM MgSO₄, 0.1 mM CaCl₂, 18.7 mM NH₄Cl, 0.4% (w/v) D-glucose, 0.05% (w/v) casamino acids and 0.0002% (w/v) thiamine.

Protein purification

ColE9 and TolB variants were expressed from ITPG inducible plasmids in *E. coli* BL21 (DE3) cells and purified, as described previously (Garinot-Schneider *et al*, 1996, 1997; Loftus *et al*, 2006; Housden *et al*, 2013). Protein concentrations were determined by UV absorbance at 280 nm using sequence-based extinction coefficient (Gill & von Hippel, 1989).

Preparation of the ColE9 translocon for Cryo-EM

Intact ColE9 translocon complex was prepared as previously described using disulphide trapped translocon complex lacking the C-terminal DNase domain of ColE9 (Housden *et al*, 2013). The purified ColE9 translocon in 20 mM MES pH 6.5 and 1% β -OG (100 μ l), was diluted 10-fold into buffer containing 20 mM potassium phosphate, pH 7.9, 100 mM NaCl and 1% β -OG. Amphipol was added to protein in a ratio of 10 mg amphipol:1mg protein, and the sample was left to incubate at room temperature for 2 h. Detergent was removed through the addition of BioBeads (Bio-Rad) at a ratio of 10 mg beads to 1 mg of detergent. After 2 h incubation at room temperature, the sample was injected onto a superose 6 increase 10/300 column, equilibrated in 20 mM potassium phosphate, pH 7.9, 100 mM NaCl. Fractions were analysed by SDS–PAGE to validate all components of the complex were present, before spin concentrating the sample to 2.0 mg/ml.

The quality and concentration of samples were confirmed by negative stain EM prior to cryo-EM sample preparation. C-flat grids (Protochips, USA; 1.2/1.3 300 mesh) and UltrAuFoil (Quantifoil, Germany; 1.2/1.3 300 mesh) were negatively glow discharged using PELCO Easiglow (Ted Pella, USA) and coated with graphene oxide as described previously (Wang *et al*, 2020). Prior to application to graphene-coated quantifoil grids, translocon preparation was diluted to 0.05 mg/ml with buffer containing 20 mM potassium phosphate, pH 7.9, 100 mM NaCl. 3 μ l of sample was applied to grid and vitrified in liquid ethane using a Vitrobot Mark IV (Thermo Fisher Scientific, USA) at 4°C and 94% humidity.

Cryo-EM image collection and processing

Cryo-EM data were collected in ISMB Birkbeck EM facility using a Titan Krios microscope (Thermo Fisher Scientific, USA) operated at 300 keV and equipped with a BioQuantum energy filter (Gatan, USA). The images were collected with a post-GIF K2 Summit direct electron detector (Gatan, USA) operated in counting mode, at a nominal magnification of 130,000 corresponding to a pixel size of 1.047 Å. The dose rate was set to 4.99 e $Å^2$ per second, and a total dose of 49–50 e/Å² was fractionated over 50 frames. An energy slit with a 20 eV width was used during data collection. Data were collected using EPU software with a nominal defocus range -1.5 to -3.5 µm. Relion 3.1 software package (Scheres, 2012a, 2012b; Zivanov et al, 2018, 2020) was used for data motion correction and dose weighting with MotionCor2 (Zheng et al, 2017). Data were then imported into cryoSPARC (Punjani et al, 2017, 2020) and the contrast transfer function (CTF) estimated using GCTF (Zhang, 2016). After removing micrographs with non-vitreous ice, broken graphene oxide support or poor CTF fit, blob picking followed by particle extraction, and 2D classification were used to generate templates for template-based picking within cryoSPARC. Templatepicked particles were extracted with a 280-pixel box and sorted using 2D classification. From here, the processing workflow for the two datasets diverged.

For data set one (654,865 particles), five ab initio classes were generated following 2D classification, and classes were refined using heterologous refinement within cryoSPARC. In two of the resulting classes, the density was consistent with a translocon complex, these were selected for further processing using non-uniform refinement, prior to import into relion3.1. After 3D refinement, the full translocon map reported an overall resolution of 5.0 Å and the partial translocon map, 3.7 Å. To improve the quality of the partial translocon map, the detergent shell was subtracted and the resulting particles were refined to generate a new map with a reported resolution of 3.9 Å. In parallel to 3D refinement, subtracted particles were further sorted using a 3D classification step without alignment and sorting into four classes. All four classes were refined independently, with only one class showing map density for both ColE9 and TolB. After a 3D refinement of this class, a map was generated with a reported resolution of 3.7 Å which, following post-processing and local resolution estimation, was sharpened using either LocScale (Jakobi et al, 2017) or deepEMhancer (preprint: Sanchez-Garcia et al, 2020). Visual inspection of the sharpened maps confirmed the selection of the deepEMhancer tight map (Appendix Fig S1) as the final map for the partial translocon. Similar sorting and refinement procedures were applied to the full translocon particles; however, no improvements in map resolution and quality were observed.

Data set two (434,671 particles after 2D classification) was processed using cryoSPARC. Three *ab initio* maps were generated and subject to heterologous refinement. Of these classes, only one class showed weak density for both TolB and ColE9, at higher threshold levels, in addition to the density consistent with the OmpF trimer. This map was then further processed using non-uniform refinement. The resulting map reported a resolution of 3.2 Å; however, upon visual inspection of the map it revealed poor density for TolB and ColE9 relative to the partial translocon map from dataset one and also had non-isotropic density within OmpF; therefore, no further processing was carried out.

After independent processing of both datasets, the particles were combined to give 1,089,536 particles which were sorted using a second 2D classification step. Classes were divided into three groups, full translocon, partial translocon and junk. The particles in the full and partial translocon classes were independently processed within cryoSPARC. Two maps of the full translocon were generated from ab initio reconstruction. One map had a more intact structure with density for all components and was therefore refined further using non-uniform refinement, to give a map with a reported resolution of 4.0 Å. Visual inspection of the map showed that only the OmpF portion of the map extended to 4.0 Å with the majority of the remaining map having poorly resolved secondary structure and weak density associated with ColE9 and TolB. To improve the quality of the map, the detergent shell was subtracted and the map locally refined. This generated a map with a reported resolution of 4.6-16 Å; however, the density of ColE9 and TolB had improved. This map was sharpened with either LocScale or deepEMhancer, with the deepEMhancer tight mask map generating the final map of the full translocon. To assess the degree of conformational heterogeneity, 3D variability analysis was implemented using particles

input into the final map. From the analysis, five maps were generated, which showed movement within TolB and ColE9. Comparison of these maps with that of the partial translocon did not show a map whereby the conformation of ColE9 or TolB matched that seen in the partial translocon map, suggesting that the partial translocon and full translocon were discrete states.

After combining datasets, particles assigned to the partial translocon class were processed using a similar method to that applied to the full translocon above. Four classes were generated from *ab initio* reconstruction and heterologous refinement. Of these four classes, only one showed density for both ColE9 and TolB, this class was processed further using non-uniform refinement, local refinement, detergent shell subtraction, and a final local refinement step, to generate a map with a reported resolution of 3.6 Å. This map was input into 3D variability analysis generating five maps to assess the degree of conformational heterogeneity. Once again, density for ColE9 consistent with the full translocon map was absent.

To build a partial translocon model, OmpF (PDB ID 3K19) was docked into the partial translocon map from relion using chimera (Pettersen et al, 2004) and rigid-body refined using Phenix (Afonine et al, 2018a, 2018b; Liebschner et al, 2019). ColE9 residues 2-67 were manually built (stubbing residues where density for side chain was weak) into the remaining map density present within the OmpF pores using coot (Emsley & Cowtan, 2004), and several cycles of rigid-body and local refinement was carried out using Phenix. TolB-ColE9 32-47 (PDB ID 2IVZ) were then docked into map using chimera and rigid body refined using phenix. Due to the weak density in the TolB portion of the map, the ColE9 peptide was not clearly resolved; however, after rigid-body fitting the terminal residues (32 and 47) were proximal to the ColE9 chain built prior to TolB docking. Therefore, residues 32-47 of ColE9 bound in the TolB crystal structure were substituted for ab initio built sequence and the whole complex (OmpF, TolB and ColE9 residues 2-67) refined using phenix. Only rigid-body refinement was applied to the TolB portion of the map due to the lower resolution in this region of the map. The ColE9 T-domain residues 85-314 (PDB ID 5EW5) were docked into the map using chimera, merged with remaining parts of the translocon, and rigid body refined in phenix to give the final partial translocon model. Due to the weak density following on from ColE9 residue 67 the disordered N-terminal domain could not be reliably built past this point to connect residues 67 and 85.

For the full translocon model, OmpF and ColE9 residues 2–66 from the partial translocon structure were docked into the final full translocon map within chimera. Additionally, BtuB bound to the R-domain of ColE3 (PDB ID 2YSU) was also docked into the map, prior to running refinement within Phenix. Only rigid-body refinements were run for this model due to the lower resolution of the map. The crystal structure of ColE9 (PDB ID 5EW5) was docked into the map, through alignment to the R-domain of ColE3 within chimera. This created an anchor point for the R-domain that prevented steric clashes with BtuB. The TolB-ColE9 32–47 crystal structure was once again docked into the map and merged with ColE9 2–66, and the whole complex rigid-body refined in phenix, to generate the final model.

Map-to-model parameters from Phenix, model validation and map processing workflow are summarised in Appendix Figs S1–S3 and Appendix Table S1. Structure representation figures were made using chimeraX (Goddard *et al*, 2018; Pettersen *et al*, 2021) and pymol.

Vitamin B₁₂ competition assay

The assay was carried out as described previously (Penfold *et al*, 2000). In brief, *E. coli* 133/3 cells grown were overnight in 5 ml minimal media (1× M9 salts, 100 μ g/ml L-arginine, 1 mM MgSO₄, 0.2% glucose) and supplemented with vitamin B₁₂ (1 nM). Cells were harvested by centrifugation and re-suspended in minimal media (5 ml), to wash out vitamin B₁₂. This wash step was repeated twice before diluting cells 1:100 in 50 ml of minimal media supplemented with either vitamin B₁₂ (1 nM) and ColE9 R-domain or ColE9 W39A (40 nM), as appropriate. Cells were maintained at 37°C with shaking for a total time of 6 h. Samples (1 ml) were removed every 30 min and optical density at 600 nm (OD₆₀₀) recorded. Average OD₆₀₀ values and standard deviation across triplicate measurements are reported.

Colicin cytotoxicity assay

Colicin cytotoxicity was assayed by plate-based growth inhibition assays that were carried out as described previously (Atanaskovic *et al*, 2020; Jansen *et al*, 2020) Colicin-induced cell killing was indicated as zones of clearance in the bacterial lawn.

Fluorophore labelling of proteins

Single cysteine mutants of ColE9 or the isolated ColE9 DNase domain were desalted on a 5 ml HiTrap column (GE Healthcare) equilibrated with binding buffer (20 mM potassium phosphate pH 7, 500 mM NaCl) to remove DTT. The protein was diluted to 50 μ M and labelled with a 5:1 molar excess of fluorophore:protein with either Alexa Fluor 488 C5-Maleimide (Invitrogen) (AF488), Alexa Fluor 568 C5-Maleimide (Invitrogen) (AF568), or Alexa Fluor 647 C2-Maleimide (Invitrogen) (AF647). Maleimide dyes were resuspended in DMSO and stored as a 10 mM stock. Labelling was allowed to proceed for 1-2 h at room temperature in the dark, and then quenched using 2 mM DTT for 15 min, before overnight dialysis at 4°C (20 mM potassium phosphate pH 7, 500 mM NaCl) to remove excess dye (Spectra Por 10K MWCO). Labelled protein was applied to a Superdex 200 10/300 GL column (GE Healthcare) equilibrated with 20 mM potassium phosphate pH 7, 500 mM NaCl, to remove residual dye. Protein-containing fractions were identified by SDS-PAGE. The protein concentration and labelling efficiency (typically > 90%) was estimated from absorbance readings using sequence-based extinction coefficients, molecular weights, and dye correction factors (Invitrogen).

Plasmid nicking assay

pUC19 DNA was used at 1 μ g/ml in 50 mM triethanolamine (Fisher Scientific) pH 7.5 with 20 mM MgCl₂ and 200 nM ColE9 DNase domain. 5 μ l samples were taken at timepoints, the reaction quenched with EDTA (37.5 mM, pH 8.0) and products separated on an agarose gel (1.2%) following the addition DNA loading dye (Invitrogen).

Bioluminescence assay

Escherichia coli DPD1718 cells were inoculated from an overnight culture, into 10 ml LB and grown at 37° C until OD₆₀₀ ~0.4. Cells were

diluted 1:2 in 100 μ l with warmed LB (37°C), into a black Nunc microwell 96-well optical bottom polystyrene plate. ColE9 was used at 0, 0.5, 1, 2 and 20 nM. The plate was loaded into a plate incubator (Freedom Evo 150 robot) at 37°C with gentle shaking. After loading of plates into Tecan Infinite M200 Pro plate reader cell growth (600 nm) and bioluminescence (490 nm) measured every 10 min.

Cell preparation for live-cell imaging by fluorescence microscopy

Escherichia coli cells from an overnight LB culture were used to inoculate 10 ml of supplemented M9 minimal media (with the appropriate antibiotics) and grown at 37°C until they reached OD₆₀₀ of 0.4. Cells (1.6 ml) were pelleted by centrifugation (7,000 *g* for 3 min at 17°C) and re-suspended in 200 μ l M9 containing 1.5 μ M of fluorescently labelled colicin and 20 μ M Hoechst 33342 (Thermo Fisher). Cells were incubated for 30 min at room temperature with rotary inversion in the dark and then pelleted by centrifugation and resuspended in 400 μ l supplemented M9 minimal media. Pelleting followed by resuspension was repeated twice more to remove unbound label. Finally, pelleted cells were re-suspended in a small volume (10–100 μ l) of M9 and 10 μ l loaded onto 1% (w/v) M9 minimal media agarose pad using a 1.5 × 1.6 cm Gene Frame matrix, solidified on a 1.2–1.5 mm thick microscope slide (Thermo Fisher).

For CCCP experiments, pelleted cells were re-suspended in 400 μ l of 100 μ M CCCP (Sigma) and incubated for 5 min at room temperature prior to the addition of ColE9. CCCP remained present in media throughout wash steps and in the agarose pad. For nigericin and valinomycin experiments, cells were permeabilised using 1 mM EDTA pH 8.0 for 5 min before treatment with 5 μ M nigericin (Sigma) or 5 μ M valinomycin (Sigma) in the presence of 150 mM KCl and incubated for 5 min at room temperature, to inhibit the proton concentration gradient (Δ pH) or the electrical potential ($\Delta\Psi$), respectively, before addition of colicin. Nigericin and valinomycin remained present throughout the washes and in the agarose pad.

For trypsin experiments, labelled and washed cells were resuspended in 200 μ l M9 media containing trypsin (Sigma) to achieve a 2:1 colicin:trypsin ratio. Cells were incubated with trypsin for 1 h at room temperature with rotary inversion before washing three times in M9.

Widefield microscopy data acquisition and analysis

Live cells were imaged at room temperature using a $60 \times$ oilimmersion objective (Oil UPlanSApo) (NA 1.42) on an OMX V3 BLAZE Optical microscope with Deltavision OMX software. The 405 nm (DAPI filter), 488 nm (FITC filter), 593 nm (615/24 filter) or 642 nm (Cy5 filter) laser lines were used. Lasers were used at 10–31.5% laser intensities and an exposure time of 30 ms. Differential interference contrast (DIC) was used to image the cell. The fluorescence channels were used to acquire single images, recorded at a frame rate of 30 ms/frame.

Widefield data in the fluorescence channels (405, 488, 593 and 642 nm) were smoothed using ImageJ (Schneider *et al*, 2012). The data were normalised and set at a minimum of ~100 units/pixels to remove background. Fluorescence intensity distribution profiles across the width of cells were determined using ImageJ (Schneider *et al*, 2012) plot profile function. A total of 15 cells for each condition were analysed and data normalised to generate profiles on a

comparable scale and the average plotted with the standard error of mean (SEM). Image intensity quantification was obtained by drawing cell outlines and determining the mean pixel intensity from the 16-bit images within ImageJ. A total of 37–126 cells were analysed per condition. Student's *t*-tests were performed to determine statistical significance of differences in relative intensities between samples.

Escherichia coli cell death kinetics

Escherichia coli JW2203 (*ompF*-expressing) cells (25 ml) were grown in LB at 37°C to OD₆₀₀ ~0.4 (~10⁸ cells) prior to addition of chloramphenicol (20 µg/ml), to inhibit cell division, and either wild-type ColE9 or ColE9 K469C labelled with AF488, AF568 or AF647 (80 nM) added to the culture. A no-colicin control was also included. Colicin import was stopped at specific time points by removing 900 µl of culture and adding 100 µl of trypsin (1 mg/ml) and incubated at 37°C for 30 min. 100 µl of each sample was diluted in LB (10⁰–10⁶ fold), before plating (90 µl) on LB-agar plates (with appropriate antibiotics) and incubating for 16 h at 37°C. Colony-forming units (CFUs) were obtained and averaged from across triplicate plates. The natural log of each CFU value was plotted against time (min), and the rate constant (*k*) of the reaction was calculated assuming a first-order rate equation. Cell-killing rates are quoted as half-lives ($t_{1/2}$).

Data availability

The data supporting the findings of the study are available in the article, available upon request from the corresponding author or available from the protein structure data bank PDB (https://www.rc sb.org/, identifiers 7NSU and 7NST) or electron microscopy data bank EMDB (https://www.ebi.ac.uk/emdb/, identifiers EMD-12577 and EMD-12576).

Expanded View for this article is available online.

Acknowledgements

We are indebted to Patrice Rassam (Strasbourg) and Patrick Inns (Oxford) for help and advice regarding microscopy experiments, David Staunton (Molecular Biophysics Suite, Oxford) for help with biophysical analysis of purified and fluorescently labelled proteins, and Jeremy Keown and Loic Carrique for assistance with computational analysis of cryo-EM data. Cryo-EM data were collected at the ISMB EM facility at Birkbeck College, University of London with financial support from Wellcome Trust (202679/Z/16/Z and 206166/Z/17/Z). The work was funded by the European Research Council (Advanced grant 742555; OMPorg) and the Wellcome Trust (201505/Z/16/Z). MLRF was supported by the BBSRC Oxford Interdisciplinary Bioscience DTP. We are also indebted to Nadia Halidi and the Micron Advanced Bioimaging Unit (Wellcome Strategic Awards 091911/B/10/Z and 107457/Z/15/Z) for their support & assistance in this work. We thank the Oxford Biomedical Research Computing (BMRC) facility, a joint development between the Wellcome Centre for Human Genetics and the Big Data Institute (supported by Health Data Research UK and the NIHR Oxford Biomedical Research Centre).

Author contributions

NGH, MNW, M-LRF, NL and CK designed research; RK, EE and BC provided materials; MLRF, MNW and NL performed research; M-LRF and MNW analysed data; CK and MNW wrote the paper.

Conflict of interest

The authors declare that they have no conflict of interest.

References

- Afonine PV, Klaholz BP, Moriarty NW, Poon BK, Sobolev OV, Terwilliger TC, Adams PD, Urzhumtsev A (2018a) New tools for the analysis and validation of cryo-EM maps and atomic models. *Acta Crystallogr D Struct Biol* 74: 814–840
- Afonine PV, Poon BK, Read RJ, Sobolev OV, Terwilliger TC, Urzhumtsev A, Adams PD (2018b) Real-space refinement in PHENIX for cryo-EM and crystallography. *Acta Crystallogr D Struct Biol* 74: 531–544
- Aoki SK, Diner EJ, de Roodenbeke CT, Burgess BR, Poole SJ, Braaten BA, Jones AM, Webb JS, Hayes CS, Cotter PA *et al* (2010) A widespread family of polymorphic contact-dependent toxin delivery systems in bacteria. *Nature* 468: 439–442
- Atanaskovic I, Mosbahi K, Sharp C, Housden NG, Kaminska R, Walker D, Kleanthous C (2020) Targeted Killing of Pseudomonas aeruginosa by Pyocin G Occurs via the Hemin Transporter Hur. J Mol Biol 432: 3869–3880
- Basle A, Rummel G, Storici P, Rosenbusch JP, Schirmer T (2006) Crystal structure of osmoporin OmpC from *E. coli* at 2.0 A. *J Mol Biol* 362: 933–942
- Basler M, Ho BT, Mekalanos JJ (2013) Tit-for-tat: type VI secretion system counterattack during bacterial cell-cell interactions. *Cell* 152: 884–894
- Behrens HM, Lowe ED, Gault J, Housden NG, Kaminska R, Weber TM, Thompson CMA, Mislin GLA, Schalk IJ, Walker D et al (2020) Pyocin S5 import into Pseudomonas aeruginosa reveals a generic mode of bacteriocin transport. MBio 11: e03230-19
- Benedetti H, Lloubes R, Lazdunski C, Letellier L (1992) Colicin A unfolds during its translocation in *Escherichia coli* cells and spans the whole cell envelope when its pore has formed. *EMBO* / 11: 441–447
- Bonsor DA, Grishkovskaya I, Dodson EJ, Kleanthous C (2007) Molecular mimicry enables competitive recruitment by a natively disordered protein. J Am Chem Soc 15: 4800–4807
- Bonsor DA, Hecht O, Vankemmelbeke M, Sharma A, Krachler AM, Housden NG, Lilly KJ, James R, Moore GR, Kleanthous C (2009) Allosteric betapropeller signalling in TolB and its manipulation by translocating colicins. *EMBO J* 28: 2846–2857
- Cascales E, Buchanan SK, Duché D, Kleanthous C, Lloubès R, Postle K, Riley M, Slatin S, Cavard D (2007) Colicin biology. *Microbiol Mol Biol Rev* 71: 158–229
- Chauleau M, Mora L, Serba J, de Zamaroczy M (2011) FtsH-dependent processing of RNase colicins D and E3 means that only the cytotoxic domains are imported into the cytoplasm. *J Biol Chem* 286: 29397–29407
- Chen Y, Radford SE, Brockwell DJ (2015) Force-induced remodelling of proteins and their complexes. *Curr Opin Struct Biol* 30: 89–99
- Cooper PC, James R (1984) Two new E colicins, E8 and E9, produced by a strain of *Escherichia coli. J Gen Microbiol* 130: 209
- Cramer WA, Sharma O, Zakharov SD (2018) On mechanisms of colicin import: the outer membrane quandary. *Biochem J* 475: 3903–3915
- Duche D (2007) Colicin E2 is still in contact with its receptor and import machinery when its nuclease domain enters the cytoplasm. *J Bacteriol* 189: 4217–4222
- Emsley P, Cowtan K (2004) Coot: model-building tools for molecular graphics. Acta Crystallogr D Biol Crystallogr D60: 2126–2132
- Farrance OE, Hann E, Kaminska R, Housden NG, Derrington SR, Kleanthous C, Radford SE, Brockwell DJ (2013) A force-activated trip switch triggers rapid dissociation of a colicin from its immunity protein. *PLoS Biol* 11: e1001489

- Garinot-Schneider C, Penfold CN, Moore GR, Kleanthous C, James R (1997) Identification of residues in the putative TolA box which are essential for the toxicity of the endonuclease toxin colicin E9. *Microbiology* 143(Pt 9): 2931
- Garinot-Schneider C, Pommer AJ, Moore GR, Kleanthous C, James R (1996) Identification of putative active-site residues in the DNase domain of colicin E9 by random mutagenesis. *J Mol Biol* 260: 731
- Gehring KB, Nikaido H (1989) Existence and purification of porin heterotrimers of *Escherichia coli* K12 OmpC, OmpF, and PhoE proteins. *J Biol Chem* 264: 2810–2815
- Gill SC, von Hippel PH (1989) Calculation of protein extinction coefficients from amino acid sequence data. *Anal Biochem* 182: 319-326
- Goddard TD, Huang CC, Meng EC, Pettersen EF, Couch GS, Morris JH, Ferrin TE (2018) UCSF ChimeraX: meeting modern challenges in visualization and analysis. *Protein Sci* 27: 14–25
- Granato ET, Meiller-Legrand TA, Foster KR (2019) The evolution and ecology of bacterial warfare. *Curr Biol* 29: R521 R537
- Gudmundsdottir A, Bradbeer C, Kadner RJ (1988) Altered binding and transport of vitamin B12 resulting from insertion mutations in the *Escherichia coli* btuB gene. *J Biol Chem* 263: 14224
- Hands SL, Holland LE, Vankemmelbeke M, Fraser L, Macdonald CJ, Moore GR, James R, Penfold CN (2005) Interactions of TolB with the translocation domain of colicin E9 require an extended TolB box. *J Bacteriol* 187: 6733–6741
- Hickman SJ, Cooper REM, Bellucci L, Paci E, Brockwell DJ (2017) Gating of TonB-dependent transporters by substrate-specific forced remodelling. *Nat Commun* 8: 14804
- Housden NG, Hopper JTS, Lukoyanova N, Rodriguez-Larrea D, Wojdyla JA, Klein A, Kaminska R, Bayley H, Saibil HR, Robinson CV *et al* (2013) Intrinsically disordered protein threads through the bacterial outermembrane porin OmpF. *Science* 340: 1570–1574
- Housden NG, Loftus SR, Moore GR, James R, Kleanthous C (2005) Cell entry mechanism of enzymatic bacterial colicins: Porin recruitment and the thermodynamics of receptor binding. *Proc Natl Acad Sci USA* 102: 13849–13854
- Housden NG, Rassam P, Lee S, Samsudin F, Kaminska R, Sharp C, Goult JD, Francis M-L, Khalid S, Bayley H *et al* (2018) Directional porin binding of intrinsically disordered protein sequences promotes colicin epitope display in the bacterial periplasm. *Biochemistry* 57: 4374–4381
- Housden NG, Wojdyla JA, Korczynska J, Grishkovskaya I, Kirkpatrick N, Brzozowski AM, Kleanthous C (2010) Directed epitope delivery across the *Escherichia coli* outer membrane through the porin OmpF. *Proc Natl Acad Sci USA* 107: 21412–21417
- Jakobi AJ, Wilmanns M, Sachse C (2017) Model-based local density sharpening of cryo-EM maps. *Elife* 6: e27131
- Jansen KB, Inns PG, Housden NG, Hopper JTS, Kaminska R, Lee S, Robinson CV, Bayley H, Kleanthous C (2020) Bifurcated binding of the OmpF receptor underpins import of the bacteriocin colicin N into *Escherichia coli*. J Biol Chem 295: 9147–9156
- Jeanteur D, Schirmer T, Fourel D, Simonet V, Rummel G, Widmer C, Rosenbusch JP, Pattus F, Pages JM (1994) Structural and functional alterations of a colicin-resistant mutant of OmpF porin from *Escherichia coli. Proc Natl Acad Sci USA* 91: 10675–10679

Kleanthous C (2010) Swimming against the tide: progress and challenges in our understanding of colicin translocation. *Nat Rev Microbiol* 8: 843–848

Kleanthous C, Kuhlmann UC, Pommer AJ, Ferguson N, Radford SE, Moore GR, James R, Hemmings AM (1999) Structural and mechanistic basis of immunity toward endonuclease colicins. *Nat Struct Biol* 6: 243–252

- Klebba PE, Newton SMC, Six DA, Kumar A, Yang T, Nairn BL, Munger C, Chakravorty S (2021) Iron acquisition systems of gram-negative bacterial pathogens define TonB-dependent pathways to novel antibiotics. *Chem Rev* 121: 5193–5239
- Klein A, Wojdyla JA, Joshi A, Josts I, McCaughey LC, Housden NG, Kaminska R, Byron O, Walker D, Kleanthous C (2016) Structural and biophysical analysis of nuclease protein antibiotics. *Biochem J* 473: 2799–2812
- Kurisu G, Zakharov SD, Zhalnina MV, Bano S, Eroukova VY, Rokitskaya TI, Antonenko YN, Wiener MC, Cramer WA (2003) The structure of BtuB with bound colicin E3 R-domain implies a translocon. *Nat Struct Biol* 10: 948–954
- Lee S, Housden NG, Ionescu SA, Zimmer MH, Kaminska R, Kleanthous C, Bayley H (2020) Transmembrane epitope delivery by passive protein threading through the pores of the OmpF porin trimer. *J Am Chem Soc* 142: 12157–12166
- Liebschner D, Afonine PV, Baker ML, Bunkóczi G, Chen VB, Croll TI, Hintze B, Hung LW, Jain S, McCoy AJ *et al* (2019) Macromolecular structure determination using X-rays, neutrons and electrons: recent developments in Phenix. *Acta Crystallogr D Struct Biol* 75: 861–877
- Loftus SR, Walker D, Mate MJ, Bonsor DA, James R, Moore GR, Kleanthous C (2006) Competitive recruitment of the periplasmic translocation portal TolB by a natively disordered domain of colicin E9. *Proc Natl Acad Sci USA* 103: 12353–12358
- Mock M, Pugsley AP (1982) The BtuB group col plasmids and homology between the colicins they encode. J Bacteriol 150: 1069
- Ng CL, Lang K, Meenan NA, Sharma A, Kelley AC, Kleanthous C, Ramakrishnan V (2010) Structural basis for 16S ribosomal RNA cleavage by the cytotoxic domain of colicin E3. *Nat Struct Mol Biol* 17: 1241–1246
- Nikaido H (2003) Molecular basis of bacterial outer membrane permeability revisited. *Microbiol Mol Biol Rev* 67: 593–656
- Pawelek PD, Croteau N, Ng-Thow-Hing C, Khursigara CM, Moiseeva N, Allaire M, Coulton JW (2006) Structure of TonB in complex with FhuA, *E. coli* outer membrane receptor. *Science* 312: 1399–1402
- Penfold CN, Garinot-Schneider C, Hemmings AM, Moore GR, Kleanthous C, James R (2000) A 76-residue polypeptide of colicin E9 confers receptor specificity and inhibits the growth of vitamin B12-dependent *Escherichia coli* 113/3 cells. *Mol Microbiol* 38: 639
- Penfold CN, Healy B, Housden NG, Boetzel R, Vankemmelbeke M, Moore GR, Kleanthous C, James R (2004) Flexibility in the receptor-binding domain of the enzymatic colicin E9 is required for toxicity against *Escherichia coli* cells. *J Bacteriol* 186: 4520–4527
- Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, Ferrin TE (2004) UCSF Chimera—A visualization system for exploratory research and analysis. *J Comput Chem* 25: 1605–1612
- Pettersen EF, Goddard TD, Huang CC, Meng EC, Couch GS, Croll TI, Morris JH, Ferrin TE (2021) UCSF ChimeraX: structure visualization for researchers, educators, and developers. *Protein Sci* 30: 70–82
- Pommer AJ, Cal S, Keeble AH, Walker D, Evans SJ, Kuhlmann UC, Cooper A, Connolly BA, Hemmings AM, Moore GR *et al* (2001) Mechanism and cleavage specificity of the H-N-H endonuclease colicin E9. *J Mol Biol* 314: 735
- Pratt LA, Hsing W, Gibson KE, Silhavy TJ (1996) From acids to osmZ: multiple factors influence synthesis of the OmpF and OmpC porins in *Escherichia coli. Mol Microbiol* 20: 911–917
- Punjani A, Rubinstein JL, Fleet DJ, Brubaker MA (2017) cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination. *Nat Methods* 14: 290–296

- Punjani A, Zhang H, Fleet DJ (2020) Non-uniform refinement: adaptive regularization improves single-particle cryo-EM reconstruction. *Nat Methods* 17: 1214–1221
- Ranf S (2016) Immune sensing of lipopolysaccharide in plants and animals: same but different. *PLoS Pathog* 12: e1005596
- Reeves P (1965) The Adsorption And Kinetics Of Killing By Colicin Ca42-E2. Aust J Exp Biol Med Sci 43: 191–200
- Ruhe ZC, Low DA, Hayes CS (2020) Polymorphic toxins and their immunity proteins: diversity, evolution, and mechanisms of delivery. *Annu Rev Microbiol* 74: 497–520
- Sanchez-Garcia R, Gomez-Blanco J, Cuervo A, Carazo J, Sorzano C, Vargas J (2020) DeepEMhacer: a deep learning solution for cryo-EM volume postprocessing. *bioRxiv* 2020.2006.2012.148296 [PREPRINT]
- Scheres SHW (2012a) A Bayesian view on Cryo-EM structure determination. J Mol Biol 415: 406-418
- Scheres SHW (2012b) RELION: implementation of a Bayesian approach to cryo-EM structure determination. J Struct Biol 180: 519-530
- Schneider CA, Rasband WS, Eliceiri KW (2012) NIH Image to ImageJ: 25 years of image analysis. *Nat Methods* 9: 671–675
- Sharp C, Bray J, Housden NG, Maiden MCJ, Kleanthous C (2017) Diversity and distribution of nuclease bacteriocins in bacterial genomes revealed using Hidden Markov models. *PLoS Comput Biol* 13: e1005652
- Soelaiman S, Jakes K, Wu N, Li CM, Shoham M (2001) Crystal structure of colicin E3: implications for cell entry and ribosome inactivation. *Mol Cell* 8: 1053–1062
- Szczepaniak J, Holmes P, Rajasekar K, Kaminska R, Samsudin F, Inns PG, Rassam P, Khalid S, Murray SM, Redfield C et al (2020) The lipoprotein Pal stabilises the bacterial outer membrane during constriction by a mobilisation-and-capture mechanism. Nat Commun 11: 1305
- Vankemmelbeke M, Healy B, Moore GR, Kleanthous C, Penfold CN, James R (2005) Rapid detection of colicin E9-induced DNA damage using *Escherichia coli* cells carrying SOS promoter-lux fusions. *J Bacteriol* 187: 4900–4907
- Vankemmelbeke M, Housden NG, James R, Kleanthous C, Penfold CN (2013) Immunity protein release from a cell-bound nuclease colicin complex requires global conformational rearrangement. *MicrobiologyOpen* 2: 853–861
- Vergalli J, Bodrenko IV, Masi M, Moynié L, Acosta-Gutiérrez S, Naismith JH, Davin-Regli A, Ceccarelli M, van den Berg B, Winterhalter M *et al* (2020) Porins and small-molecule translocation across the outer membrane of Gram-negative bacteria. *Nat Rev Microbiol* 18: 164–176
- Walker D, Mosbahi K, Vankemmelbeke M, James R, Kleanthous C (2007) The role of electrostatics in colicin nuclease domain translocation into bacterial cells. *J Biol Chem* 282: 31389–31397
- Wang F, Liu Y, Yu Z, Li S, Feng S, Cheng Y, Agard DA (2020) General and robust covalently linked graphene oxide affinity grids for high-resolution cryo-EM. *Proc Natl Acad Sci USA* 117: 24269–24273
- White P, Joshi A, Rassam P, Housden NG, Kaminska R, Goult JD, Redfield C, McCaughey LC, Walker D, Mohammed S *et al* (2017) Exploitation of an iron transporter for bacterial protein antibiotic import. *Proc Natl Acad Sci USA* 114: 12051–12056
- Whitfield C, Trent MS (2014) Biosynthesis and export of bacterial lipopolysaccharides. Annu Rev Biochem 83: 99–128
- Zhang K (2016) Gctf: real-time CTF determination and correction. J Struct Biol 193: 1-12
- Zheng SQ, Palovcak E, Armache J-P, Verba KA, Cheng Y, Agard DA (2017) MotionCor2: anisotropic correction of beam-induced motion for improved cryo-electron microscopy. *Nat Methods* 14: 331–332

- Zivanov J, Nakane T, Forsberg BO, Kimanius D, Hagen WJH, Lindahl E, Scheres SHW (2018) New tools for automated high-resolution cryo-EM structure determination in RELION-3. *Elife* 7: e42166
- Zivanov J, Nakane T, Scheres SHW (2020) Estimation of high-order aberrations and anisotropic magnification from cryo-EM data sets in RELION-3.1. *IUCR J* 7: 253–267



License: This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.