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Cryo-EM structure of lysenin pore elucidates membrane insertion by an aerolysin family protein

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Lysenin from the coelomic fluid of the earthworm Eisenia fetida belongs to the aerolysin family of small β-pore-forming toxins (β-PFTs), some members of which are pathogenic to humans and animals. Despite efforts, a high-resolution structure of a channel for this family of proteins has been elusive and therefore the mechanism of activation and membrane insertion remains unclear. Here we determine the pore structure of lysenin by single particle cryo-EM, to 3.1 Å resolution. The nonameric assembly reveals a long β-barrel channel spanning the length of the complex that, unexpectedly, includes the two pre-insertion strands flanking the hypothetical membrane-insertion loop. Examination of other members of the aerolysin family reveals high structural preservation in this region, indicating that the membrane-insertion pathway in this family is conserved. For some toxins, proteolytic activation and pro-peptide removal will facilitate unfolding of the pre-insertion strands, allowing them to form the β-barrel of the channel.

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Lyseenin, the defence protein of the earthworm *Eisenia fetida*, is produced by coelomocytes in the body fluid that form part of the earthworm’s immune system. It is a member of the aerolysin family of small β-pore-forming toxins (β-PFTs) that include key virulence factors of a large number of bacterial pathogens, such as aerolysin produced by *Aeromonas spp*., *Clostridium perfringens* epsilon toxin and *Clostridium septicum* α-toxin, while *Bacillus thuringiensis* parasporin-2 has cytoidal activity against human cancer cells. Despite their low sequence homology, the structures of their water-soluble monomeric forms reveal that they all share a structurally conserved domain, termed the pore-forming module (PFM), which is known to contribute to the β-barrel pore.

Like most PFTs, lyseenin is secreted as an inactive, water-soluble monomer. On binding to sphingomyelin (SM) in the eukaryotic cell membrane, the toxin undergoes significant changes to its secondary structure and forms oligomeric pre-pores on the membrane surface, as identified by electron crystallographic studies and high-speed atomic force microscopy studies, before inserting into the membrane, which leads to pore formation and consequent cell lysis.

The crystal structure of monomeric lyseenin revealed that the toxin is organized into two main structural domains: (1) the elongated PFM at the N terminus that contains a SM-binding site, as well as the membrane-insertion loop, long thought to be the region that reorganizes into a β-hairpin during membrane insertion, and (2) the C-terminal beta-trefoil domain that has been crystallized bound to phosphocholine (POC). Investigation of the structure of monomeric lyseenin bound to SM revealed that both the POC headgroup and one acyl chain tail of SM are required for toxin binding.

Although several structures have been determined in the past for the monomeric forms of toxins from the aerolysin family, there is no reported atomic structure for their active, pore forms. Here we report the structure of lyseenin in its nonameric membrane-inserted form determined by single particle cryo-EM, to 3.1 Å resolution. The structure provides the first atomic resolution view into the membrane-inserted state of a member of the aerolysin family and gives insights into how other members of this family may be activated and assemble into their respective pore forms.

**Results**

**Structure determination of the lyseenin pore.** Lyseenin monomers oligomerize and form SDS-resistant channels on interaction with lipid bilayers containing SM. We reasoned that we could assemble lyseenin oligomers that are biologically active (Supplementary Fig. 1a) in liposomes and then solubilize these complexes in detergent and treat them as single particles for cryo-EM analysis. Solubility screening using ultracentrifugation identified dodecylmaltoside as a suitable detergent (Supplementary Fig. 1b). Because of the tendency of the sample to aggregate at high concentrations (Supplementary Fig. 2a), we imaged particles enriched against a continuous support film. Due to the relatively small size of the expected lyseenin complex (210 kDa for a hexamer and 315 kDa for a nonamer, based on the molecular weight of 35 kDa for the His-tagged monomer), we deposited samples on TEM holey-carbon grids that were overlaid with graphene oxide (Fig. 1c, Supplementary Figs 2b,c and 3).

Compared with amorphous carbon, the near-electron transparent nature of graphene oxide yielded higher signal to noise ratios and improved alignment accuracies. Image analysis and two-dimensional (2D) classification of 53,779 particles led to a data set of 42,830 particles, which were used for an initial 3D reconstruction at 3.4 Å resolution with C9 symmetry or 4.2 Å with no symmetry imposed. Movie frame processing for beam-induced particle motion tracking and B-factor-weighted radiation-damage correction followed by 3D classification resulted in a subset of ‘shiny’ particles used for further refinement. This produced a final symmetrized map at 3.1 Å resolution (Fig. 1a,b and Supplementary Fig. 4), approaching the Nyquist limit of 2.9 Å for the pixel size used. The local resolution in the map varies only very little from the overall estimate, with minor parts at the tip of the complex resolved at 4 Å (Supplementary Fig. 4c). The map allowed unambiguous placement of the C9 main-chain and almost all side chains. No other oligomeric species were identified during the 2D (Fig. 1d) and 3D classification steps, indicating that lyseenin pores are almost exclusively nonameric. This symmetry is consistent with previous cross-linking studies and is the second identified example of a protein from the aerolysin family, alongside monalysin, to form nonameric assemblies.

**Architecture of the lyseenin assembly.** The lyseenin pore resembles the mushroom-shaped pore complexes of the α-haemolysin family of small β-PFTs (Fig. 2a,b and Supplementary Fig. 5), although structures of their water-soluble monomers are fundamentally different. Monomeric lyseenin has previously been divided into two domains: the N-terminal PFM and the C-terminal beta-trefoil domain. However, because of the large structural rearrangement associated with its membrane insertion, we have changed the domain nomenclature. Thus, each of the nine monomers in the membrane-inserted state can be divided into three domains: the N-terminal cap domain (Glu10–Ser33 and Pro108–Ile156), the β-hairpin domain (Val34–Ile107) and the C-terminal receptor-binding domain (Val157–Gly297) (Fig. 2a,b). Neighbouring interactions between the subunits bury ~3,300 Å² and involve 90 residues, which are distributed among the three domains and include hydrogen bonds and several salt bridges. A random coil from Val157 to Glu167 connects the cap
domain to the receptor-binding domain (Fig. 2a, asterisk). This region is flexible and plays an important role in the transition to the membrane-inserted state. The C-terminal receptor-binding domain has remained mostly unchanged relative to the soluble monomer crystal structure, with an r.m.s.d. (Cα–Cα) of 0.6 Å, and consists of 12 β-sheets and a short 310 helix arranged in a β-trefoil motif.

Structure of the β-barrel pore. In contrast to the α-haemolysin family where the β-barrel begins close to the membrane-spanning region (Supplementary Fig. 5), the long β-barrel in lysenin spans the length of the assembly. The membrane-spanning region, measuring ~35 Å, is clearly visible in filtered maps as a detergent-bound shell that delineates the boundaries of the lipid bilayer (Fig. 2c) and is lined on the outside by hydrophobic residues, such as Phe, Tyr, Val and Ile. In the water-soluble lysenin monomer, these residues reside within the hydrophobic core of the PFM and are thus shielded from solvent. Two aromatic rings, one formed by Phe70 and the other by Tyr79 lie on the intracellular and extracellular sides of the transmembrane region, respectively (Fig. 2a). These rings are common to β-barrel membrane proteins and are typically found snorkelling at the interface between the polar lipid head groups and the hydrophobic acyl chains helping to stabilize the orientation of the β-barrel. In addition to the aromatic rings, lysenin also contains a histidine triad just above the Tyr79 ring (histidine residues 58, 81 and 83) (Fig. 2a,d) that is situated within the edge of the detergent belt, suggesting these residues may interact with the extracellular leaflet of the bilayer. The β-turn of each hairpin ends in Glu71, which protrudes towards the intracellular side of the bilayer. The lumen of the β-barrel is lined with charged and polar residues, including 12 threonine and 9 serine residues. An electrostatic potential map of the lumen surface (Supplementary Fig. 6) reveals four regions of negative charge, including one on either entrance to the β-barrel and two in the middle of the pore. The charge distribution is consistent with planar-bilayer channel measurements that suggest a cation-selective pore.

Interaction with lipids. Monomeric lysenin has previously been crystallized bound to SM and POC ligands, although these structures were clearly of the water-soluble state. POC was crystallized bound to the C-terminal β-trefoil domain of lysenin in a pocket lined by Lys185, Ser227, Gln229, Tyr233 and Tyr282 (ref. 9). In the pore form of lysenin this pocket is ideally situated for interaction with the phosphatidylcholine or SM head groups as it lies on the tip of the receptor-binding domain directly above the target cell membrane (Fig. 2a, ‘POC’). SM was identified bound to the N-terminal PFM, and the residues that interacted with SM in the monomer structure (Lys21, Tyr24, Tyr26, Gln117 and Glu128) are buried within the cap domain of the lysenin pore (Supplementary Fig. 4d). Moreover, we did not find additional densities corresponding to lipids in the C9 symmetrized map or
lysenin on planar bilayers, which found a height reduction of up to 3 nm on membrane insertion. This movement induces a bend in the middle of the relatively flat cap domain of the soluble monomer (Fig. 3b, dashed lines) and results in each cap domain being positioned on the top of the adjacent subunit’s receptor-binding domain (Supplementary Movie 1). Extension of the long 74 residue β-hairpin requires the complete rearrangement of 5 β-strands and a 3₁₀ helix from the PFM of monomeric lysenin into 2 new β-strands, as highlighted in Fig. 3a,b. The β-hairpin begins near the extracellular side of the pore lumen and curves towards the intracellular side by almost 270° (Fig. 3d). Thus, the β-barrel pore of lysenin is made up of 3 times as much polypeptide as was previously thought (24 residues).”

**Discussion**

The cryo-EM structure of the 315 kDa lysenin pore at 3.1 Å resolution represents the first atomic resolution pore structure for a member of the aerolysin family of small β-PFTs, unveiling the transition of the water-soluble monomer into its membrane-inserted, oligomeric state. The structure sets the framework for future studies on how other members of this family may achieve pore formation and opens the pathway for the design of new therapeutics aimed at the functional disruption of pore formation for this family.

Members of the aerolysin family share an overall basic domain arrangement as evident by their crystal structures. They typically contain one or more receptor-binding domains and a PFM. The latter domain contains a flexible insertion loop, long thought to be the region that reorganizes into a β-hairpin during membrane insertion, due to its amphipathic character of alternating hydrophilic/hydrophobic residues (Fig. 4, coloured in green). The pore structure of lysenin reveals that the β-barrel recruits polypeptide from both the amphipathic loop and the two β-strands that feed into it (Fig. 3b), from here onwards referred to as the pre-insertion strands. Closer examination of all solved monomer structures from the aerolysin family reveals that the hypothetical insertion loops in these proteins are also flanked by predominantly β-strand polypeptides that lead up to the top of the PFMs of each protein (some examples are shown in Fig. 4, yellow strands). Interestingly, in all cases, these pre-insertion strands are positioned on the edge of the toxins, isolated from the main body of the protein, making limited contact in the form of short β-sheets. It is also notable that the combined lengths of the pre-insertion strands and insertion loops in all these proteins do not vary significantly (mean of 75 residues and a s.d. of 8 residues) and are rich in serine and threonine residues (Thr + Ser = 23–44%) , most of which are found lining the lumen of the lysenin β-barrel pore. Due to the structural homology shared within the PFMs of the aerolysin family, we expect that other members may follow similar strategies for monomer reorganization, which will involve the unfolding of the pre-insertion strands for β-barrel formation and the remainder of the PFM forming the caps of their respective mushroom-shaped oligomers. In agreement with this hypothesis, the predicted membrane-spanning region of the recently crystallized monalysin also begins at the top of the PFM and its deletion abolishes pore formation.

The PFMs in aerolysin, epsilon toxin and parasporin-2 also contain N- or C-terminal peptides (CTPs), which have to be proteolytically removed for activation. These pro-peptides have been implicated in chaperoning the secreted protoxins and controlling oligomerization, and their removal is most likely to occur subsequent to receptor binding and on monomer–monomer interactions. The most striking feature of the pre-insertion strands is that they interact directly with the CTPs.
of aerolysin and epsilon toxin. In both cases, the CTPs create a bridge between the pre-insertion strands and the main body of the toxin by participating in β-sheets with both domains (Fig. 4, CTPs coloured in red). Thus, it is evident that removal of the CTP in aerolysin and epsilon toxin will reduce the energy required for the pre-insertion strands to disengage from the main body of the toxin and rearrange into the long β-hairpin. In the case of aerolysin, one of the pre-insertion strands is predicted to partially unfold in the absence of the CTP, possibly acting as a switch to initiate membrane insertion32, providing further evidence for the involvement of the pre-insertion strands in pore formation.

Finally, removal of the CTP is likely to expose surfaces in the cap domains of these proteins required for complete interaction between the subunits and to promote oligomerization prior to pore formation31.

The mode of lysenin membrane insertion identified in this study is similar to that of the α-haemolysin family of small-βPFTs. The staphylococcal haemolysins and clostridial NetB toxin undergo a simple extension of a pre-stem region, which is tucked against the β-sandwich domain, to form the β-barrel pores without any significant changes to the remainder of the structure23,25,26,35. This mechanism is remarkably well-conserved within this family even in cases where the symmetry, subunit stoichiometry and receptor-binding regions vary27. In a similar way, the lysenin pre-insertion strands unfold away from the PFM and rearrange into a long β-barrel. However, in contrast to α-haemolysin the PFM of lysenin undergoes a conformational change resulting in a collapse of ~20 Å and an inwards compaction (Supplementary Movie 1). The pore model for aerolysin proposed by Degiacomi and co-workers31 derived from an 18 Å cryo-EM map combined with molecular dynamics of mutants locked in a pre-pore and quasi-pore conformations also bears similarities to the lysenin pore structure in that the overall domain arrangement of the receptor-binding domain 2 and the PFM (domains 3 and 4) are positioned similar to lysenin. The extension of the aerolysin β-hairpins, however, may begin, as in the case of lysenin, at the top of the PFM and include the pre-insertion strands. As more pore structures of the aerolysin family are solved to high resolution, it will become clearer how these toxins vary and to what extent the insertion mechanisms are conserved.

Based on our findings and previous work in the field, a model for pore formation by lysenin is presented in Fig. 5. The initial binding of soluble monomers to the membrane will be mediated by interactions between the receptor-binding domain and POC head groups. On membrane binding, the local concentration increase of monomers will promote oligomerization, leading to a pre-pore assembly. Depending on the orientation of the monomers, the height of the pre-pore may be as high as 100 Å (the length of a lysenin monomer). The membrane-inserted form of lysenin solved in this study will extend ~65 Å above the membrane surface. The hypothetical pre-pore shown here was assembled by imposing ninefold symmetry to the monomeric crystal structure aligned in reference to the receptor-binding domain of the membrane-inserted form.
through the receptor-binding domain to POC head groups (Fig. 5a) (ref. 9 and this study). This will additionally help to orient the toxin prior to pre-pore formation. Recognition of SM may also occur at this point. This will be followed by oligomerization to a pre-pore state (Fig. 5b), as identified by AFM10,36 and electron crystallography 9, through monomer–oligomerization to a pre-pore state (Fig. 5b), as identified by 9 and satisfies the β-strand hydrogen-bonding potential in a hydrophobic environment such as the lipid bilayer. Finally, β-barrel formation will create a pore in the bilayer (Fig. 5c), leading to the disruption of membrane homeostasis and ultimately to cell lysis.

Methods

Recombinant protein production and purification. For expression of lysenin, recombinant plasmid pParallel-Lys was transformed into E. Rosetta 2 (DE3) cells (Merck, Darmstadt, Germany) and expression of lysenin was induced using the autinduction system38 as follows. Cells (21) were grown in ZYM-5052 autoinducing medium supplemented with 100 μg ml−1 ampicillin and 34 μg ml−1 chloramphenicol and cultured at 37°C for 3 h at 300 r.p.m., then for a further 24 h at 20°C, 300 r.p.m. Cells were harvested by centrifugation and the cell pellet was lysed by 200 μl BugBuster Protein Extraction Reagent (Merck) containing 200 μl r-lysozyme (1 KU ml−1) (Merck) and 200 μl Benzonase Nuclease (25 U μl−1) (Merck). The cell suspension was incubated on a rotating mixer for 25 min at room temperature and centrifuged at 16,000g for 20 min at 4°C to separate soluble and insoluble fractions. The supernatant was loaded onto HisGravITrap columns (GE Healthcare Life Sciences, Little Chalfont, UK) following the manufacturer’s guidelines. In brief, His-tagged proteins were bound to the affinity column using a buffer composed of 20 mM sodium phosphate, 500 mM NaCl and 20 mM imidazole, pH 7.4. The column was washed with a buffer composed of 20 mM sodium phosphate, 500 mM NaCl and 60 mM imidazole, pH 7.4. A second buffer devoid of imidazole but containing 500 mM sodium phosphate, 500 mM NaCl and 500 mM imidazole, pH 7.4. All purification steps were carried out at 4°C. For buffer exchange and sample clean up, toxin containing eluate was applied to a PD-10 Desalting Column (GE Healthcare Life Sciences) and eluted in DPBS buffer pH 7.0–7.2 (Invitrogen). Protein concentrations were determined using the BCA assay (Fisher Scientific UK Ltd, Loughborough, UK).

Haemolysis assay. Fresh whole blood from healthy individuals was collected by venepuncture from the University of Exeter Medical School, NIHR Exeter Clinical Research Facility, Diabetes and Vascular Medicine Centre into neutral tubes and 1 ml whole blood was immediately transferred into 20 ml DPBS buffer pH 7.0–7.2 supplemented with 500 μg ml−1 bovine serum albumin (BSA). Cells were washed three times with DPBS supplemented with 1 mg ml−1 BSA and resuspended to 3 × 105 cells per ml in DPBS. Washed cells (3 × 105 cells per well) were incubated with a twofold dilution series of purified recombinant lysenin (ranging from 500 g ml−1 to 0.015 mg ml−1) in DPBS in round-bottomed 96-well plates in a final volume of 200 μl DPBS and 0.9% Triton X-100 were used as controls. Following incubation at 37°C for 30 min and for 30 min on ice, intact cells were removed by centrifugation at 1,500g for 3 min at 4°C and the supernatants (100 μl) were transferred to a flat-bottom plate to measure haemoglobin release by absorbance at 415 nm using a Model 680 Microplate Reader (Bio-Rad Laboratories Ltd., Hemel Hempstead, UK). The absorbance values for each sample were normalized by subtracting the absorbance value obtained for untreated cells and haemolytic activity (%) was calculated. The toxin dose required to lyse 50% of the cells (CT50) was determined reverse side). Grids were then used for plunge-freezing without further treatment.

Specimen preparation and data collection. Specimens were plunge-frozen using a custom-fabricated plunging at 4°C. Lysenin oligomers (50 μM in 50 mM Tris pH 7.4) were applied to copper 300 mesh Quantifoil R1.2/1.3 holey-carbon grids (Quantifoil Micro Tools, GmbH) overlaid with graphene oxide (see below) and left to adhere for 30 s. The grids were then blotted from the specimen side for 10 s before being plunge-frozen in liquid ethanol. Specimens were imaged on an FEI Titan Krios transmission electron microscope operating at an accelerating voltage of 300 kV. Micrographs were recorded in super-resolution counting mode using a Gatan K2 Summit direct electron detector at the end of a Gatan Quantum energy filter in zero-loss mode and an energy selecting slit width of 20 eV. The total dose on the specimen was 47 e− per Å2 fractionated over 20 frames with a calibrated pixel size of 0.715 Å for the super-resolution micrographs.

Image processing. Micrograph frame stacks were binned by two, subjected to drift-correction using MOTIONCORB39 and contrast transfer function (CTF) parameters were determined using GCTB40. An initial model for refinement was generated using EMAN2 (ref. 41). All subsequent image processing steps were performed using RELION42. Semi-automated particle picking was performed using the documented procedures, resulting in 53,779 particles from 268 micrographs.

Two rounds of 2D classification were used to remove incorrectly selected particles, particularly ones at the graphene layer faces. The remaining 42,830 particles were then used for initial refinement and reconstruction that led to a map with a resolution of 3.4 Å with C9 symmetry or 4.2 Å with no symmetry imposed. Movie particle extraction followed by per particle beam-induced motion correction and radiation-damage weighting resulted in a polished data set that was then subjected to 3D classification. Two of the 3 resulting 3D classes were combined and the resulting 29,329 particles were used for further refinement. The final map from these particles was sharpened with a B-factor of 61 Å2 and the ‘gold standard’ resolution was 3.14 Å at a Fourier shell correlation (FSC) of 0.143 (Supplementary Fig. 4a). The accuracy of rotation and translation during refinement was 0.86 pixels and 0.33 pixels, respectively. Local resolution estimation was performed using RESMAP44 as implemented in RELION.

Model building and refinement. The wild-type lysenin crystal structure29 (PDB ID 32XD) was used as a template for de novo modelling of the N terminus after the C-terminal 150 residues, which form the β-trefoil receptor-binding domain. All model building was performed in Coot45. The model is lacking the N-terminal residues, for which we could not see density. Refinement of the model to improve the data was carried out using automatic refinement carried out using the documented procedures43, resulting in 53,779 particles from 268 micrographs. Three rounds of 2D classification were used to remove incorrectly selected particles, particularly ones at the graphene layer faces. The remaining 42,830 particles were then used for initial refinement and reconstruction that led to a map with a resolution of 3.4 Å with C9 symmetry or 4.2 Å with no symmetry imposed. Movie particle extraction followed by per particle beam-induced motion correction and radiation-damage weighting resulted in a polished data set that was then subjected to 3D classification. Two of the 3 resulting 3D classes were combined and the resulting 29,329 particles were used for further refinement. The final map from these particles was sharpened with a B-factor of 61 Å2 and the ‘gold standard’ resolution was 3.14 Å at a Fourier shell correlation (FSC) of 0.143 (Supplementary Fig. 4a). The accuracy of rotation and translation during refinement was 0.86 pixels and 0.33 pixels, respectively. Local resolution estimation was performed using RESMAP44 as implemented in RELION.

Graphene oxide specimen support preparation. Graphene oxide dispersion in H2O (Sigma) was diluted to 0.2 mg ml−1 in H2O and spun at 300g for 30 s to remove large aggregates. Quantifoil R1.2/1.3 holey grids were glow discharged for 1 min and 3 μl of the graphene suspension was added to the grids for 1 min. Grids were subsequently blotted briefly using Whatman No1 filter paper and washed three times on 20 μl drops of H2O (twice on the graphene side and once on the reverse side). Grids were then used for plunge-freezing without further treatment.

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Author contributions
M.B.-B. purified the protein, conducted the haemolysis assays, analysed the data and wrote the paper. T.G.M. optimized the graphene support methodology and collected the data. C.E.N. analysed the data and contributed to model building. A.K.B. and R.W.T. analysed the data and wrote the paper. C.G.S. isolated the pores, collected the data, performed the image processing and model building, analysed the data and wrote the paper.

Additional information
Accession codes: The cryo-EM maps of the lysenin pore have been deposited to the Electron microscopy Data Bank under accession number EMD-8015 and the refined atomic coordinates have been deposited to the Protein Data Bank under accession number 5GAQ.

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