Cholesterol-dependent cytolysins (CDCs) are pore-forming proteins that serve as major virulence factors for pathogenic bacteria. They target eukaryotic cells using different mechanisms, but all require the presence of cholesterol to pierce lipid bilayers. How CDCs use cholesterol to selectively lyse cells is essential for understanding virulence strategies of several pathogenic bacteria, and for repurposing CDCs to kill new cellular targets. Here we address that question by trapping an early state of pore formation for the CDC intermedilysin, bound to the human immune receptor CD59 in a nanodisc model membrane. Our cryo electron microscopy map reveals structural transitions required for oligomerization, which include the lateral movement of a key amphipathic helix. We demonstrate that the charge of this helix is crucial for tuning lytic activity of CDCs. Furthermore, we discover modifications that overcome the requirement of cholesterol for membrane rupture, which may facilitate engineering the target-cell specificity of pore-forming proteins.
ore-forming proteins rupture lipid bilayers to kill target cells. They comprise the largest class of virulence factors for pathogenic bacteria and are prevalent in all kingdoms of life. Cholesterol-dependent cytolysins (CDCs) are pore-forming proteins secreted by more than five genera of Gram-positive bacteria, including human pathogens from Streptococcus, Clostridium, and Listeria. Pore-forming proteins also play a crucial role in immune defence, killing Gram-negative bacteria, cancer cells, and phagocytosed microbes. Understanding how these proteins discriminate between self-cells and target membranes will provide insight into fundamental virulence strategies, as well as facilitate the application of engineered CDCs that lyse new cellular targets.

Many pore-forming proteins depend on lipid specificity to select their targets. For example, the immune protein perforin preferentially targets post-synaptic membranes through lipid disorder phases and neutral charge headgroups. Another immune protein, gadermin, specifically binds cardiolipin and phosphoinositide lipids to direct activity against mitochondria and the outer leaflets of eukaryotic plasma membranes, triggering cell death. For CDCs, the requirement of cholesterol directs lytic activity towards plasma membranes of eukaryotic hosts. Since cholesterol is not present in bacterial cells that secrete CDCs, this is one way the pathogen protects itself from damage during infection and toxin production; it also limits the repurposing of CDCs to attack bacterial pathogens. Many CDCs, such as pneumolysin (PLY), bind cholesterol directly through a conserved cholesterol-recognition motif located on a membrane-binding loop. Although cholesterol is a well-known receptor for PLY, it may not be the only one. PLY also binds to a mannose receptor in dendritic cells, downregulating inflammation and promoting bacterial survival. Other CDCs, such as intermedilysin (ILY) secreted by Streptococcus intermedius, achieve species-specificity for their hosts by hijacking the cell surface receptor CD59 to initiate membrane binding. For this subgroup of CDCs, the interaction with CD59 is sufficient for attachment and the role of cholesterol is restricted to pore formation. While differences in membrane targeting may reflect diversity in CDC virulence strategies, lipid dependency for membrane penetration remains highly conserved.

CDCs bind target membranes through interactions located in their domain 4 (D4). Both the cholesterol-recognition loops of PLY and CD59-binding site of ILY lie within this domain. A long extended domain 2 (D2) flexibly links D4 to the membrane attack complex perforin (MACPF)/CDC domain (comprised of D1 and D3 together), which is responsible for pore formation. Membrane binding of CDC monomers triggers a series of conformational rearrangements that are required for oligomerization into an assembly referred to as a prepore. A dramatic vertical collapse of the oligomeric prepore brings two helical bundles (HB1 and HB2) within the MACPF/CDC domain close to the target membrane. HB1 and HB2 residues undergo rearrangement in secondary structure to form adjacent transmembrane β-strands in the pore. These rearrangements are facilitated by stabilizing interactions between a number of amino acids at the MACPF/CDC domain interface, but structural details of intermediate conformations remain unresolved.

Here we report a structure of an early prepore-trapped conformation of the CDC ILY solved by cryo electron microscopy. ILY binds the human immune receptor CD59 as a monomer to initiate attachment to the membrane. CD59 binding then promotes ILY oligomerization and triggers movement of a highly conserved amphipathic helix that encodes activity and lipid specificity of CDC pore formation. This helix can be modified to create cholesterol-independent CDCs or “super-CDCs” with significantly enhanced activity. Hence, our results provide a blueprint for engineering broad-purpose pore-forming proteins with tunable lytic activity.

Results

CryoEM reconstruction of an ILY early prepore. ILY from Streptococcus intermedius targets human cells by binding the GPI-anchored complement regulator CD59 through an extended β-hairpin of D4. To understand how this interaction initiates oligomerization on a target membrane, we trapped ILY in an early prepore state using a disulfide lock that restricts movement between D2 and the MACPF/CDC domain. This disulfide-locked ILY variant binds CD59 and forms SDS-sensitive loosely associated oligomers, analogous to previously characterized CDC early prepore states. We then used cryo-electron microscopy (cryoEM) to visualize this ILY variant in complex with CD59 anchored to cholesterol-containing lipid nanodiscs. We collected data on a Titan Krios microscope equipped with a direct electron detector and processed images using RELION. 2D classification revealed heterogeneity in the number of subunits within a single nanodisc (Supplementary Fig. 1). Extensive 2D and 3D classification resulted in a final reconstruction comprised of 51,041 particles. The data were further refined with local symmetry, resulting in a map with an average resolution of 4.6 Å. The final local resolution-sharpened map illustrates an arc of five ILY–CD59 complexes, with the central subunit best resolved (Fig. 1a, Supplementary Fig. 1). We therefore built a model into the density of the central subunit, which served as the basis for refinement of a three-subunit oligomer (Fig. 1a and Supplementary Fig. 3).

Our cryoEM map reveals an oligomeric complex whereby a single CD59 can interact with two neighboring ILY monomers (ILY and ILY’). Similar to the soluble monomeric ILY–CD59 crystal structures, a β-hairpin of one ILY D4 extends the central β-sheet of CD59. The same β-hairpin of an adjacent monomer (ILY’), together with the tip of D2, sandwiches an α-helix of CD59 to form the second interface (Fig. 1b). This interface is dominated by two CD59 residues decorated with O-linked glycans in the native protein. ILY binds to an O-glycan prevalent on human CD59, suggesting that sugar recognition may play a role in oligomerization and could be engineered to impact target-cell specificity.

CD59 binding initiates the secondary structure rearrangement of the MACPF/CDC outer β-strand (β-5). In our structure, residues within the β-5 strand form a new helix that comprises part of a helix-turn-helix motif (Fig. 1c, Supplementary Fig. 4, and Supplementary Fig. 5). As a result, the β-4 strand is exposed to propagate oligomerization by binding the β-1 strand of a neighboring monomer. In the early prepore, β-4 rotates from its position in the monomeric initiation complex to reveal an intermediate transition along the pore formation pathway (Supplementary Fig. 5). The ILY prepore assembly is most likely further stabilized by complementary charges on neighboring helical faces of the helix-turn-helix motif (Fig. 1c). Similarly, alternating charges of this motif in the PLY pore are thought to stabilize an inner barrel of helices above the membrane.

In our complex, ILY extends to a vertical height of 110 Å, revealing an oligomeric prepare that has not yet collapsed towards the target membrane. Specifically, residues of the two MACPF/CDC helical bundles (HB1 and HB2) that eventually form transmembrane β-hairpins in the pore assembly do not contact the bilayer. Helical bundle 2 (HB2) is comprised of a vertical helix and a horizontal helix (h-helix) (Fig. 1d and Supplementary Fig. 4), which is directly N-terminal to the newly positioned MACPF β-4 strand (Supplementary Fig. 5). Compared with the monomeric ILY–CD59, we observe a lateral shift in these
helices upon oligomerization (Fig. 1d and Supplementary Fig. 4). This movement prevents an otherwise steric clash with a neighboring monomer and allows the oligomer to propagate. Given the moderate resolution of our map, we sought to verify the movement of these helices using a fluorescence-based assay. To investigate how oligomerization influences the local chemical environment of the h-helix, we covalently linked monobromo-bimane (mBBr) to residues on either the bottom (membrane adjacent) or top face of the h-helix in early prepore-locked ILY (Fig. 1e). Fluorescence was then measured in solution (monomer) or in the presence of CD59-containing DOPC:cholesterol liposomes (early prepore). Upon oligomerization, we observed an increase in mBBr fluorescence when the fluorophore was tethered to the top face of the h-helix, consistent with movement of L340 as it packs more closely against the surrounding protein. By contrast, the fluorescence intensity remained the same for labels attached to the bottom face of the h-helix, in agreement with our structural data showing that the packing of membrane-adjacent residues is not affected by this conformational change (Fig. 1e, f). Together, these biochemical data corroborate the structurally observed lateral movement of the HB2 helices.

The h-helix encodes the ability to tune lytic activity. Based on our structure, we hypothesized that the h-helices of a collapsed oligomer could play a role in destabilizing the lipid bilayer. To test this, we first needed to stall pore formation of a collapsed oligomer by trapping ILY in a late prepore assembly intermediate. We generated disulfi de-locked ILY mutants to restrict movement of either HB1 (HB1-lock) or HB2 (HB2-lock) and prevent the formation of transmembrane β-hairpins (Fig. 2a and Supplementary Fig. 6). SDS gel shift assays reveal a portion of HB1 and HB2. This movement prevents steric clashes between adjacent monomers and allows the oligomer to propagate. Given the moderate resolution of our map, we sought to verify the movement of these helices using a fluorescence-based assay. To investigate how oligomerization influences the local chemical environment of the h-helix, we covalently linked monobromo-bimane (mBBr) to residues on either the bottom (membrane adjacent) or top face of the h-helix in early prepore-locked ILY (Fig. 1e). Fluorescence was then measured in solution (monomer) or in the presence of CD59-containing DOPC:cholesterol liposomes (early prepore). Upon oligomerization, we observed an increase in mBBr fluorescence when the fluorophore was tethered to the top face of the h-helix, consistent with movement of L340 as it packs more closely against the surrounding protein. By contrast, the fluorescence intensity remained the same for labels attached to the bottom face of the h-helix, in agreement with our structural data showing that the packing of membrane-adjacent residues is not affected by this conformational change (Fig. 1e, f). Together, these biochemical data corroborate the structurally observed lateral movement of the HB2 helices.

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HB2 mutants remain unlocked. Despite the variable efficiency of disulfide bond formation both HB1-lock and HB2-lock mutants show a significant reduction in lytic activity, which is rescued under reducing conditions (Fig. 2b). Together these data suggest that variants could have a dominant negative effect on the lytic activity of an oligomer containing a mixture of locked and unlocked monomers. To investigate if these ILY variants formed vertically collapsed oligomers, we imaged complexes using atomic force microscopy on supported lipid bilayers containing both cholesterol and CD59. In contrast to monomeric ILY10 and the ILY early prepore (Fig. 2a), which extend 10 nm from the membrane surface, both HB1-lock and HB2-lock mutants are approximately 8 nm from the surface (Fig. 2c). The vertical heights of the HB1-lock and HB2-lock oligomers are analogous to the 8 nm high wild-type pores (Fig. 2c), consistent with collapsed oligomers. Despite undergoing a vertical collapse, both mutants show significantly reduced lytic activity (Fig. 2b). We therefore conclude that HB1-lock and HB2-lock mutants trap collapsed prepore assemblies whereby the helical bundles are brought closer to the target membrane but do not breach the bilayer.

The h-helix is amphipathic, with charged or polar residues along the membrane-adjacent face and hydrophobic residues on the opposite face (Fig. 3a). To test how lytic activity and hence ILY function depend on the distribution of charges along the ILY h-helix, we generated ILY variants where charged or polar residues on the membrane-proximal face were substituted with alanine (ILYno charge) or where hydrophobic residues on the top face were swapped with lysine (ILYcharge+). Removal of charged residues impaired lytic activity of wild-type ILY. By contrast, incorporating positively charged amino acids on the top face of the h-helix increases lysis. Lytic activity of wild-type ILY could also be improved by substituting h-helix residue N342 for tryptophan (Fig. 3b). Taken together, these results demonstrate that the lytic function of ILY can be modulated by changing the physicochemical properties of the h-helix.

Given that similar modifications did not impact lytic activity of the uncollapsed early prepore (Supplementary Fig. 6), we next analyzed the effect of h-helix variations on otherwise dysfunctional (non-lytic) collapsed prepores by introducing mutations in the HB1-lock and HB2-lock backgrounds. Remarkably, we found...
that without altering the proportion of disulfide-locked mono-
mers (Supplementary Fig. 6), the lytic function was fully
recovered by the addition of charged residues (charge +) or the
N342W substitution in the h-helix, as shown on cholesterol-
containing liposomes (Fig. 3b). By contrast, tryptophan sub-
stitutions elsewhere along the helix only partially recovered lytic
activity for the HB1-lock mutant and did not enhance lysis at all
for the HB2-lock mutant (Fig. 3b). We deduce that tryptophan
substitutions along the bottom face of the h-helix augment lysis
by intercalating into the outer leaflet of the bilayer and anchoring
the h-helix to the membrane. The position of the tryptophan
likely affects the ef
f+ci
ency of this mechanism, since substitution
at N342 shows a stronger phenotypic change than at N338.

Since ILY only requires cholesterol to rupture the membrane
and not to bind it 10, the ILY–CD59 system offers an opportunity
to explicitly investigate and potentially modulate the cholesterol-
dependency of CDC membrane lysis. We therefore measured the
lytic activity of our ILY variants on liposomes without cholesterol
(Fig. 3c). Notably, the requirement of cholesterol was abrogated
by the addition of positively charged residues on the top face of
the h-helix (charge +); this modification resulted in high levels of
lysis in all ILY backgrounds (wild-type, HB1-lock, and HB2-lock,
respectively). Individual data points from three independent experimental assays are shown as circles. Error bars represent standard deviation with the center of the bar indicating the mean value. P-value significance determined by one-way ANOVA with a Bonferroni post-test: ns, not significant; *p < 0.05; **p < 0.01; ***p < 0.001.

Fig. 3 Targeted mutations in the horizontal helix (h-helix) tune lytic activity of ILY. a H-helix mutations are mapped onto the ILY structure. Left panel (ILY no charge) highlights residues (gray) exchanged for alanine (E337A, N338A, K341A, N342A) on the membrane proximal face of the h-helix. Middle panel (ILY charge+) indicates residues (cyan) mutated to lysine (Y336K, L340K) on the top face of the helix. Right panel (ILY Trp mutants) shows the location of residues (orange) mutated to tryptophan (Y336W, N338W, L340W, or N342W). Sidechains have been added in COOT 44 for visualization purposes. b, c Activity of ILY h-helix mutants in wild type (WT), HB1-lock or HB2-lock backgrounds were tested after 45 min using a calcein-release liposome lysis assay. Lysis of CD59-decorated liposomes comprised of DOPC:cholesterol lipids is shown in (b). Lysis of CD59-decorated DOPC liposomes is in (c). Statistical significance displayed above each bar is for comparison with the base ILY disulfide-locked variant (WT, HB1-lock, or HB2-lock, respectively). Individual data points from three independent experimental assays are shown as circles. Error bars represent standard deviation with the center of the bar indicating the mean value. P-value significance determined by one-way ANOVA with a Bonferroni post-test: ns, not significant; *p < 0.05; **p < 0.01; ***p < 0.001.
**Discussion**

Our combined structural and biochemical data support a model whereby CDC oligomerization triggers movement of an amphipathic helix that modulates lytic activity (Fig. 5). Soluble CDC monomers associate with membranes through interactions within D4. Binding is mediated either by loops at the tip of D4 with cholesterol, or by engaging cell surface receptors, such as CD59. Membrane binding initiates oligomerization and conformational changes within the MACPF/CDC domain that create new interaction interfaces between monomers. Specifically, the formation of a helix-turn-helix motif exposes the MACPF/CDC β-4 strand to propagate the oligomer; it may also contribute to stability of the assembly. In our structure of an early prepore, the h-helices of ILY are arranged parallel to the membrane with their charged surfaces facing the outer leaflet of the lipid bilayer. Following a dramatic vertical collapse of the oligomeric prepore, the h-helix is brought in close proximity to the target membrane. Lying along the surface, the amphipathic h-helices of the oligomer would then cause local membrane disruption, analogous to the ‘carpet’ model of membrane disruption by cationic antimicrobial peptide (CAMP) transition of CDC transmembrane residues is likely to occur via several intermediate structural states with different local energy minima, including collapsed coil-like intermediates. The transient states of the h-helix may contribute to disruptive forces that prime the membrane for hairpin insertion and to the displacement of lipids by water molecules at the inner hydrophilic surface of the pore, as observed by molecular dynamics simulations. We demonstrate that modifications of the h-helix similarly modulate the lytic activity of a non-CD59-binding CDC, PLY. Furthermore, amphipathic helices are present in and influence the lytic activity of other oligomeric pore-forming proteins such as gasdermin and MPEG-1. Taken together, our model may reveal a fundamental mechanism underpinning membrane rupture by oligomeric pore-forming proteins.

Our results open an exciting perspective for protein engineering of h-helices for the creation of “super-CDCs” with...
significantly enhanced lytic activity or of cholesterol-independent CDCs that can be used to target bacterial pathogens. Specifically, our results show that CDC-mediated lysis is enhanced by the addition of positive charges to the top face of the h-helix, and that this removes the requirement of cholesterol for membrane rupture. We note that the tunable lysis of the h-helix bears similarities to CAMP-mediated cell lysis, where the membrane specificity is encoded by the charge properties of the helix and where the introduction of tryptophan residues can modulate antimicrobial activity of synthetic CAMPs. By combining h-helix modifications with mutations in membrane-binding regions of CDCs, CDCs can be designed to lyse new cellular targets such as plant-pathogenic bacteria. The devastating effects of these pathogens on crop yield and productivity could be eliminated by engineered pore-forming proteins that lyse bacterial membranes but are unable to penetrate a plant cell wall.

**Methods**

**Bacterial strains and plasmids.** Escherichia coli strain DH5α was used to maintain plasmids and for cloning purposes. Strains BL21 DE3 and BL21 DE3 pLYS S were used for expression of ILY and PLY variants, with the latter strain grown in the presence of 25 µg/ml chloramphenicol to maintain the pLYS S plasmid. His-tagged ILY and PLY constructs were cloned and expressed using pTrcHisA and maintained with 100 µg/ml ampicillin. His-tagged MSP2N2 was expressed from pET28-MSP2N2 (Addgene) in the presence of 50 µg/ml kanamycin.

**Generation of ILY and PLY variant constructs.** Wild-type ILY and PLY, for this study, have been codon optimized for *E. coli* and all native Cys in ILY have been mutated to Ser. All ILY and PLY variants were created by site-directed mutagenesis, in which a pTrcHisA plasmid with the template gene was amplified with pairs of primers that contain the desired mutations (Supplementary Table 1 and Q5 DNA polymerase (New England Biolabs). This was followed by DpnI (New England Biolabs) digestion, to remove the template plasmid, heat-shock transformation into *E. coli* DH5α, and verification by sequencing. CDC variants used in this study are summarized in Supplementary Table 2.

**Purification of ILY and PLY variants.** *E. coli* BL21 DE3 containing pTrcHisA with an *ily* or *ply* variant gene were grown to an OD₆₀₀ of ~0.6–0.9 in Luria-Bertani (LB) broth at 37 °C, and induced with a final concentration of 50 µg/ml of isopropyl β-d-thiogalactopyranoside (IPTG) at 18 °C for 18 h. In the case of the over-active variants ILY-WTcharge and ILY-HB2lockcharge, this method resulted in very low levels of protein expression likely due to toxicity. Therefore, these two ILY variants were expressed in *E. coli* BL21 pLYS S and induced with 50 µg/ml IPTG at 18 °C for 1 h. Regardless of the expression protocol, all variants were subjected to the same purification methods. The cells were collected by centrifugation and lysed by sonication in Buffer A. The purity of all ILY and PLY variants was assessed by SDS-PAGE and their functionality was determined by the ability to form oligomers on lipid monolayers (Supplementary Fig. 6).

**Forming the ILY early prepore on CDC-decorated lipid nanodiscs.** MSP2N2 protein was first produced by growing BL21 DE3 cells containing pET28-MSP2N2 (Addgene) to an OD₆₀₀ of ~0.8 in LB and inducing with 0.5 mM IPTG at 37 °C for 2–3 h. Cells were pelleted and reconstituted in 40 mM Tris-HCl pH 7.4 with Complete Protease Inhibitors (Roche), to which Triton X-100 was added to a final concentration of 1% and DNase I to a final concentration of 5 µg/ml. Cells were stored on ice for 10 min, then lysed by sonication and supplemented with 0.3 M NaCl. His-tagged protein was bound to HisPur Cobalt resin (Thermo Fisher Scientific), washed with 40 mM Tris-HCl pH 8.0, 0.3 M NaCl, 0.5% Triton X-100, then washed with 40 mM Tris-HCl pH 8.0, 0.3 M NaCl, 50 mM sodium cholate. Protein was then eluted into 40 mM Tris-HCl pH 8.0, 0.1 M NaCl, 300 mM imidazole and subjected to size exclusion chromatography over a Superdex 200 10/300 column (GE Healthcare) with 40 mM Tris-HCl pH 8.0, 0.1 M NaCl, 0.5 mM EDTA. Nanodiscs were created by mixing 3.44 mM MSP2N2 with 34.4 mM lipids (DOPC:cholesterol, 1:1 molar ratio) solubilized in 40 mM Tris-HCl pH 8.0, 100 mM NaCl, 0.5 mM EDTA, 64 mM sodium cholate, so that the final sodium cholate concentration was 32.5 mM. This mixture was then incubated on ice for 30 min followed by an overnight incubation with activated Bio-beads SM2 (BioRad) at 4 °C with agitation, to remove the detergent and promote nanodisc formation. The nanodiscs were then purified by size exclusion chromatography with 40 mM Tris-HCl pH 8.0, 0.1 M NaCl, 0.5 mM EDTA over a Superose 6 10/300 column (GE Healthcare) and stored at 4 °C.

A recombinant extracellular domain of CD59 modified with a lipid-anchoring peptide was gifted by R.A.G. Smith (King’s College London). Briefly, the cytoplasmic domain of human CD59 modified with an additional C-terminal cysteine was expressed in *E. coli* and purified from inclusion bodies. The cytoplastic modification reagent bis-myristyl lysyl SSSKSPSKDDKPGD (S-2-thiopyridyl)-cysteine acid (APT3146, Cambridge Research Biochemicals) was covalently linked to the C-terminal cysteine and the modified protein was purified by hydrophobic interaction chromatography and ammonium sulfate precipitation.

Based on the peak absorbance reading for the nanodisc purification, 50 mM MSP2N2 nanodiscs were incubated with 330 mM CD59 at room temperature for 20 min in Buffer A, followed by another 20-min incubation after the addition of 200 µg/ml ILY-prepore. Assuming 100% incorporation, each nanodisc likely contains 6 molecules of CD59; however, the size of the ILY oligomer may also be limited by the nanodisc diameter. The mixture was immediately washed twice with Buffer A on an Amicon Ultra 0.5 ml 100 kDa concentrator column (Merck) to remove unbound protein and concentrated by a factor of 5. To avoid particle aggregation, the final sample was immediately used to prepare EM grids.

**Negative-stain EM.** All EM samples were first screened by negative-stain EM to assess particle integrity and distribution. 2.5 µl of ILY-prepore-CD59 on nanodiscs was applied to glow-discharged, carbon-coated copper grids (Agar Scientific) and stained with 0.1% uranyl acetate. Samples were imaged on a 120 keV Tecnai T12 microscope (Thermo Fisher Scientific) with a 2 k Eagle camera (FEI) at a nominal magnification of ×30,000 for evaluation.

**CryoEM grid preparation and data collection.** ILY–CD59 nanodisc complexes were imaged using holey carbon grids coated with graphene oxide. To coat R1/2/1.3 Quantifoil grids with graphene oxide, grids were first glow-discharged for 1 min, then, 0.2 mg/ml of a graphene oxide solution (Sigma) in water was applied to the glow-discharged, to the top face of the grid, followed by blotting by filter paper on the bottom face of the grid. This process was repeated twice, followed by two washes of the top face of the grid with 20 µl of water. Grids were then fixed, and used within one hour of graphene oxide coating. Immediately following condensation, 2.5 µl of the early prepore-locked ILY–CD59 oligomers on nanodiscs was adsorbed on graphene oxide-coated grids and blotted for 2.5 s at “blot force” 3 and plunged...
frozen in liquid ethane cooled to liquid nitrogen temperatures with a Vitrobot mark IV (Thermo Fisher Scientific). Electron micrograph movies were collected on a 300 kV Titan Krios (Thermo Fisher Scientific) with a cryo-electron detector (Thermo Fisher Scientific) in linear mode with image acquisition software EPU (Thermo Fisher scientific). Specific collection details for all three datasets are summarized in Supplementary Table 3.

Initial model generation. An initial model was generated in RELION19 from a dataset collected on a 300 kV Titan Krios (Thermo Fisher Scientific) with a Quantum K2 Summit direct electron detector (Gatan) in counting mode with a magnified pixel size of 1.08 Å. Manually picked particles from motion corrected and ctf-estimated micrographs were subjected to rounds of 2D classification and class curation, resulting in 4241 particles which were used to generate an initial model in RELION19. 3D refinement with this initial model produced a reconstruction with an average resolution of 8.6 Å. This density was then used as the initial model for the first individual 3D refinement of datasets 1, 2, and 3. Particles used to generate the initial model were not included in the refinement of subsequently collected datasets 1, 2, and 3.

CryoEM data analysis and density reconstruction. The overall data analysis and reconstruction strategy are summarized in Supplementary Fig. 2. All data analysis and reconstruction were completed via RELION19 unless otherwise stated. Datasets 1, 2, and 3 were treated separately for the initial stages of data processing. For each dataset, 100 low-dose micrographs were aligned using MotionCor238 as CTF parameters were estimated with CTFFind49. Any flattened movies containing low figure of merit scores, crystalline ice, low contrast, or substantial drift were removed from further analysis. Particles were picked with a combination of manual picking, autopicking in RELION, and CRyOLo40, and duplicate particles were removed. All CTF-corrected particle sets were density reconstruction, resulting in an average resolution of 4.8 Å. To improve homogeneity in the particles, they were classified into 4 groups with local angular filters were applied to the top region resulted in a reconstruction with the best local resolution and density features of the central monomer. The optimized transformation for only the top region resulted in a reconstruction with the best local resolution—of—220 Å. Local resolution estimates were calculated using RELION and the local resolution-filtered map was used for building and refining the structural model. Maps were rendered using UCSF Chimera42.

Structural model building and refinement. To initiate refinement of the ILY early prepore transition state, we used a combination of templates comprised of the monomer-locked ILY–CD59 complex (PDBID 4BK17) and the PLY pore model (PDBID 5LY643) (Supplementary Fig. 3). The crystal structure for soluble, monomer-locked ILY (PDBID 4BK17) was placed in the middle monomer of the local resolution-filtered density with COOT44. The model was initially refined as 3 rigid bodies. The MACP/CDC domain was refined with Namdinator45 for only the top region resulted in a reconstruction with the best local resolution and density features of the central monomer. The optimized transformation matrices are expressed as three Euler angles (8 141.2, 4 7.6 and 134.0) and three translations (X – 19.6 Å, Y 14.4 Å, Z 1.8 Å). The reconstruction was further refined and the local solvent density of the macropore was maintained. Density was added to the macropore in COOT44. The final density map is 4.6 Å (Supplementary Fig. 1) with a global B-factor of –220 Å. Local resolution estimates were calculated using RELION and the local resolution-filtered map was used for building and refining the structural model. Maps were rendered using UCSF Chimera42.

Gel shift assay. Disulfide bond formation was assessed by incubating 45 µg/ml of each ILY variant with reducing (62.5 mM Tris pH 6.5, 10% glycerol, 0.05% b-mercaptoethanol, 2%, SDS, 75 mM DTT, final concentrations) or non-reducing (62.5 mM Tris, pH 6.5, 10% glycerol, 0.05% b-mercaptoethanol, 2%, SDS, final concentration) sample buffer at 90 °C for 5 min. 225 ng of each ILY variant was analyzed by running 12–16% BisTris NuPAGE gels (Thermo Fisher Scientific) with Bolt MOPS SDS Running Buffer (Life Technologies). Gels were then stained with Quick coomassie stain (Generon) and destained in water overnight before imaging.

Lipid monolayer preparation and negative staining. Integrity and folding of all ILY variants were assessed by ring and arc formation on lipid monolayers. 10 µg/ml ILY and 1.6 µg/ml CD59, final concentrations, were mixed in Buffer A in a multi-well Tullon plate. The protein mixture was then overlaid with 2 µl of chloroform containing 1 mg/ml DOPC/cholesterol (1:1 molar ratio) and incubated at room temperature for 2 min to allow chloroform evaporation. Carbon-coated copper grids (Agar Scientific) were applied to the top of each well before incubation at 37 °C for 15 min in a sealed petri dish with a Buffer A-soaked paper towel, to prevent evaporation. Grids were then stained with 2% uranyl acetate and imaged, as described earlier (Negative-stain EM).

Liposome lysis assay. Lytic activity of CDC variants was assessed using a calcein-release liposome lysis assay. Liposomes containing calcein were first prepared by rehydrating lipids (DOPC or DOPC/cholesterol, 1:1 molar ratio) in Buffer A with 50 mM calcein followed by extrusion through a 0.1 µm filter. Unilamellar calcein-containing liposomes were purified in this case by exclusion chromatography (Superdex G-50 resin (Sigma) in Buffer A with 500 mM sucrose, as previously described10. The resultant liposomes were filled with self-quenching calcein dye while external, unquenched dye was replaced by Buffer A and sucrose. To determine the activity of ILY variants, liposomes were first incubated with a final concentration of 1.0 µg/ml CD59 for 20 min at room temperature, followed by the addition of ILY at a final concentration of 9.0 µg/ml. The activity of PLY variants was determined by adding a final concentration of 48.4 µg/ml of PLY to DOPC cholesterol liposomes. Fluorescence intensity was read with a CLARIOstar plate reader (BMG labtech) at an excitation wavelength of 490 nm and emission wavelength of 530 nm. Liposome lysis was achieved by preincubation of 10% n-dodecyl-beta-maltoside (DDM) and a freeze/thaw cycle at –80 °C. To calculate percent lysis, the fluorescence value from a buffer well was first subtracted from all raw fluorescence readings at a single time point between 30 and 60 min. Each blanked-corrected experimental reading was divided by the corresponding total liposome fluorescence reading (also blank-corrected). Activity of disulfide-locked CDC variants was assessed by preincubating the protein with or without 2 µM DTT for 30 min at room temperature.

Atomic force microscopy sample preparation. For the preparation of supported lipid bilayers, pure lipids were dissolved in chloroform at 10 mg/ml and mixed in solution to give a lipid mixture of DOPC/cholesterol 2:1 molar ratio. The lipid-in-chloroform solution was then dried in a glass vial under a stream of nitrogen gas to give a lipid film. The lipid film was then incubated in buffer (20 mM Tris, 200 mM NaCl, pH 7.5), vortexed and bath sonicated to give a cloudy lipid suspension. The suspension was then passed through a 50 nm polycarbonate
membrane (GÉ Healthcare Lifesciences) 15 times to yield a clear suspension of small unilamellar vesicles. Supported lipid bilayers were formed by injecting 4.5 µL of the lipid vesicle suspension to a freshly cleaved mica disk (6 mm diameter) under 18 µL of incubation buffer (hydration buffer plus 10 mM CaCl₂ solution) at 37 °C; this induces the rupture of the vesicles onto the mica support over an incubation period of approximately 30 min. Excess vesicles were then removed from the supernatant by rinsing with 500 µL of the hydration buffer, to yield a uniform bilayer free of adsorbed vesicles (as assessed by AFM imaging). The supported lipid bilayers were next incubated with a final concentration of 50 ng/ml CD59 for 5 min, and thereafter with a final concentration of 100 µg/ml ILY for 15 min, all at 37 °C, and next washed with 500 µL of the hydration buffer.

Atomic force microscopy imaging and analysis. AFM imaging was performed in buffer solution using a Multimode 8 AFM (Bruker, Santa Barbara, USA) and MSLN-E and PEAKFORCE-HIRS-F-B cantilevers (Bruker), in off-resonance tapping/fast force-feedback imaging (Bruker’s PeakForce Tapping) mode where force-distance curves were recorded at 2 kHz, with amplitudes of 10–20 nm, with the maximum imaging forces estimated between 50 and 100 pN. The experiments were performed at room temperature. Images were typically recorded at a few minutes per frame on an E scanner (Bruker, Santa Barbara, CA, USA).

We note that (membrane-inserted) pore assemblies are readily imaged by AFM, as they make contact, through the membrane, with the underlying solid support; but that pore preparations are rather hard to image by AFM. Loosely packed assemblies may be broken apart under the mechanical stress of the AFM tip. In addition, the supported lipid bilayers generally too fluid a support to retain the assemblies in place, except when the assemblies form larger clusters. In this case, the mobility of HB1/HB2-lock prepore assemblies may also be constrained by the dense coverage of CD59 on the membrane surface. Images were processed using open-source SPM analysis software, Gwyddion (v2.53) for first-order plane-fit background subtraction and second-order line-by-line flattening. The membrane surface was referenced as zero height. The images in Fig. 2c were cropped from 500 × 500 nm². Assembly heights (mean ± standard deviation) were estimated based on 100 height profiles for 10 different assemblies from each 500 × 500 nm² image, referenced with respect to the membrane surface, as exemplified in Fig. 2c.

Statistical analysis. The Student’s t test (assuming equal variance) was used to assess statistical significance of differences between monomer and early prepore fluorescence quenching of h-helix-labeled ILY (ILY-preporeN342C or ILY-preporeL346C). The average of the ratio of C2-TPG fluorescence quenching of h-helix-labeled ILY (ILY-preporeN342C or ILY-preporeL346C) to that of NMR-labeled lipids was determined by line-scan analysis in RELION-3. Differences in fluorescence quenching were analyzed by one-way ANOVA followed by a Bonferroni post-test for pairwise comparisons. A single-factor ANOVA test was used to determine the significance of variance within datasets describing lytic activity of h-helix mutants (ILY and PLY). A Bonferroni post-test was used for pairwise comparisons within each set of h-helix variants (ILY-WT, ILY-HB1lock, ILY-HB2lock, PLY).

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

Data availability
Supporting the findings of this manuscript are available from the corresponding authors upon reasonable request. All ILY and PLY mutants created in this study are made available at Addgene. A reporting summary for this article is available as a Supplementary Information file. The accession numbers for the EM map and model of the ILY-CD59 early prepore complex reported in this paper are EMD 11172 and PDB 6ZD0. Source data are provided with this paper.

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Author contributions

N.R.S. conducted cryoEM work, generated ILY mutants, and performed ILY fluorescence and lysis experiments. N.R.S. and D.B. conceived the ideas, analyzed the results, and wrote the manuscript. T.B.V. generated PLY mutants and ILY preporecharge, conducted PLY and ILY preporecharge lysis experiments, analyzed these data, and prepared figures. T.B.V., E.S.P., and B.W.H. analyzed AFM data. E.S.P. performed AFM experiments. C.M.B. generated the ILY HB1lock mutant. All authors assisted with manuscript editing.

Competing interests

The authors declare no competing interests.

Additional information

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