Structure and assembly of a bacterial gasdermin pore

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Abstract

In response to pathogen infection, gasdermin (GSDM) proteins form membrane pores that induce a host cell death process called pyroptosis^{1–3}. Studies of human and mouse GSDM pores reveal the functions and architectures of 24-33 protomers assemblies⁴⁻⁹, but the mechanism and evolutionary origin of membrane targeting and GSDM pore formation remain unknown. Here we determine a structure of a bacterial GSDM (bGSDM) pore and define a conserved mechanism of pore assembly. Engineering a panel of bGSDMs for site-specific proteolytic activation, we demonstrate that diverse bGSDMs form distinct pore sizes that range from smaller mammalianlike assemblies to exceptionally large pores containing >50 protomers. We determine a 3.3 Å cryo-EM structure of a Vitiosangium bGSDM in an active slinky-like oligomeric conformation and analyze bGSDM pores in a native lipid environment to create an atomic-level model of a full 52mer bGSDM pore. Combining our structural analysis with molecular dynamics simulations and cellular assays, we define a stepwise model of GSDM pore assembly and demonstrate that pore formation is driven by local unfolding of membrane-spanning β -strand regions and pre-insertion of a covalently bound palmitoyl into the target membrane. These results yield insights into the diversity of GSDM pores found in nature and the function of an ancient post-translational modification in enabling a programmed host cell death process.

Main

In mammalian pyroptosis, active GSDMs bind the inner leaflet of the plasma membrane and assemble into pores that secrete cytokines and induce lytic cell death^{1–5}. Human cells encode six GSDM proteins (GSDMA–E and PJVK), with individual GSDMs capable of targeting not only the plasma membrane but also membranes of mitochondria¹⁰ and intracellular bacteria¹¹. Structures of human and mouse GSDM pores demonstrate that mammalian GSDM proteins form oligomeric assemblies of differing architectures with inner diameters ranging from 150–215 Å^{6–9}. Recent identification of GSDM homologs encoded in invertebrates¹², fungi^{13,14}, and bacterial anti-phage defense systems¹⁵ further suggests that GSDMs may form a large diversity of pore structures to support distinct cellular and antiviral functions.

bGSDM pores are large and diverse To define the mechanism and architectural diversity of GSDM pore formation beyond closely related mammalian proteins, we reconstituted pore formation in vitro using a panel of bacterial GSDM (bGSDM) homologs from diverse bacterial phyla (Fig. 1 and Extended Data Fig. 1a). Human and bacterial GSDM proteins use a shared mechanism of activation where proteolytic cleavage within an interdomain linker releases a pore-forming N-terminal domain (NTD) from a partnering inhibitory Cterminal domain (CTD). Most mammalian GSDM inhibitory CTDs are ~200 amino acids (aa) in length, but bacterial and fungal GSDMs achieve autoinhibition via a short ~20-40 aa CTD14,15. GSDM activation in bacterial cells is controlled by a series of upstream proteins that sense unknown phage infection. aspects of limiting biochemical reconstitution of proteolytic activation and pore formation. We therefore sought to build on previous mammalian GSDM experiments^{4,6,7,11} to engineer bGSDMs for rational activation and enable analysis of pore formation across phylogenetically diverse systems. We first predicted bGSDM CTD cleavage sites based on our previous crystal structures of bGSDMs in inactive states¹⁵, and screened pairs of bGSDM full-length and Δ CTD constructs in vivo for toxicity in a cell growth assay (Extended Data Fig. 1a). For each bGSDM \triangle CTD construct resulting in cellular toxicity we next purified recombinant protein variants with a series of inserted sequencespecific protease motifs in the CTD linker and used a terbium-based liposome rupture assay to identify proteins capable of full



Fig. 1 | Bacterial gasdermins form pores of different sizes. a, Schematic of bGSDM engineering and screen for functional pore formation. Predicted cleavage site loops were mutagenized to site-specific protease cleavage motifs and tested in liposome leakage assays monitoring release of terbium chloride (TbCl₃). b, Liposome leakage assay of engineered bGSDMs with matched site-specific proteases. The species of bGSDMs (and protease sites) are as follows: Unclassified Bacteroidetes (TEV site), Nostoc sp. Moss4 (WELQ site), Vitiosangium sp. GDMCC 1.1324 (thrombin), and Ideonella sp. 201-F6 (WELQ). Error bars represent the SEM of three technical replicates. n.s. \ge 0.05; ****P < 0.0001. c, Size distributions of bGSDM pore inner diameters determined using negative-stain EM micrographs of pore-liposomes. The number of pores measured (n) for each species is as follows: Bacteroidetes (n = 117), Nostoc (n = 58), Runella (n = 189), Vitiosangium (n = 56), Ideonella (n = 123). The black bar represents the average inner diameter of measured pores. The grey dashed lines represent the average inner diameters mammalian gasdermin pores as previously measured^{6,11,17}. d, Example negative-stain EM micrographs of bGSDM pores from five species. Scale bar = 50 nm.

membrane pore formation *in vitro* (Fig. 1b and Extended Data Fig. 2). Using this approach, we successfully engineered controlled membrane pore formation for four bGSDM proteins from phylogenetically diverse bacteria (Fig. 1b and Extended Data Fig. 2).

We next reconstituted each successful pore-forming bGSDM variant into liposomes and visualized the membrane pores using negative-stain EM. Bacterial GSDM membrane pores are exceptionally large and surprisingly diverse in architecture (**Fig. 1c,d** and **Extended Data Fig. 3**). *Bacteroidetes* bGSDM forms the smallest pores in our panel, with a mammalian-like inner diameter of ~180 Å. In contrast, *Ideonella* bGDSM pores are massive, spanning on average ~300 Å with some pores measuring larger than 400 Å similar to the pneumolysin cholesterol-dependent cytolysin pores produced by *Streptococcus pneumoniae* to kill target eukaryotic cells¹⁶. Notably, each bacterial GSDM protein forms a relatively narrow range of pores with fixed overall diameters, demonstrating that the large variations in GSDM pore size are a programmed trait controlled by protein sequence specificity. Together, these results demonstrate that the size range of bGSDM pores architecture is a diverse feature across GSDM protein evolution.

Cryo-EM structural analysis of bGSDM oligomerization

To further compare the molecular basis of bacterial and human GSDM pore formation we extracted bGSDM assemblies from pore-liposome samples and performed single-particle cryo-EM analysis. We focused on *Bacteroidetes* bGSDM pores that form the smallest assemblies observed in our panel and *Vitiosangium* bGSDM pores that uniformly form some of the largest observed bGSDM assemblies. Cryo-EM and 2D classification analysis of extracted *Bacteroidetes* bGSDM pores reveals a rigid 30–31 protomer assembly that is remarkably similar to the human GSDMD 33-mer pore¹⁷ (**Extended Data Fig. 4a,b**). We confirmed assembly of predominantly 31mer *Bacteroidetes* bGSDM pores directly in the native lipid environment, demonstrating that some bGSDM proteins form pores sharing the overall size of GSDM proteins in human immunity (**Extended Data Fig. 4a,b**). In contrast to the smaller *Bacteroidetes* bGSDM pores, cryo-EM analysis of *Vitiosangium* bGSDM samples demonstrated giant assemblies form through incorporation of ~48–54 protomers (**Fig. 2a** and **Extended Data Fig. 5a**).

Previously, X-ray crystal structural analysis of the Vitiosangium bGSDM revealed the unexpected self-palmitoylation of an N-terminal cysteine and defined how bGSDMs are held in an inhibited conformation prior to proteolytic activation¹⁵. To determine the mechanism of how proteolysis and palmitoylation lead to assembly of giant bGSDM pores, we focused on determining a high-resolution structure of Vitiosangium bGSDM in the active conformation. Due to orientation bias of the exceptionally large Vitiosangium bGSDM pores, single-particle 3D reconstruction was not possible for these samples. While screening a series of detergents and extraction methods we observed in addition to single ring pores that Vitiosangium bGSDM samples adopted double-ring assemblies and other open-ring assemblies including cracked slitshaped oligomers as seen previously for human GSDM by cryo-EM, atomic force microscopy, and molecular dynamics (MD) simulation experiments^{6,18,19}. Unexpectedly, in DDMAB detergent we observed that open-ring assemblies of Vitiosangium bGSDM extend to form slinky-like oligomers with uniform radial dimensions closely mirroring the full pore (Fig. 2b,c and Extended Data Fig. 5b). Mixtures of closed-ring pores and slinkies were observed in HECAMEG-extracted samples after prolonged storage, indicating that Vitiosangium bGSDM slinkies likely represent a stable non-native conformation derived from breaking closed-ring pores (Fig. 2c). Imaging Vitiosangium bGSDM slinkies by cryo-EM yielded 2D classes with excellent orientation distribution (Fig. 2b and Extended Data Fig. 6), and subsequent 3D classification and refinement of an ~25-protomer turn enabled us to determine a complete 3.3 Å structure of Vitiosangium bGSDM in the active oligometric conformation (Fig. 2d,e and Extended Data Figs. 5b,c and 7).



Fig. 2 | Structure of an active bGSDM oligomer from detergent extracted pores. a, Cryo-EM micrograph and representative 2D class averages of HECAMEG detergent-extracted *Vitiosangium* bGSDM pores. The yellow arrowhead points to an example pore. Numbers in the upper left-hand corner of 2D classes represent the number of bGSDM protomers observed in that class. Scale bars = 50 nm. **b**, Cryo-EM micrograph and representative 2D class averages of DDMAB detergent-extracted *Vitiosangium* bGSDM slinkies. The yellow arrowhead points to an example slinky. Scale bars = 50 nm. **c**, Model of structural relationship between bGSDM pores and slinky oligomers, with the pitch of a helical turn indicated. **d**, Cryo-EM map of an ~25 protomer turn of the *Vitiosangium* bGSDM slinky. For the bottom map, the dimension of the inner diameter is shown. **e**, bGSDM active model overlaid on a portion of the slinky map. Left, 5-protomer model fit into the bGSDM slinky map. A single protomer. The dimensions for protomer size are shown.

Evolutionary conservation of GSDM activation

The *Vitiosangium* bGSDM active-state cryo-EM structure reveals dramatic conformational rearrangements and an ancient mechanism of activation shared between bacterial and human GSDM proteins. Using DALI²⁰ to compare the *Vitiosangium* bGSDM model against all structures in the PDB, we observed active-state structures of hGSDMD and mouse GSDMA3 (mGSDMA3) as the top hits confirming that the *Vitiosangium* slinky-derived conformation corresponds to the bGSDM active oligomeric state (**Fig. 3a**). Notable structures more distantly related to the *Vitiosangium* bGSDM active-state include the inactive conformation of mammalian GSDMs and members of the membrane attack complex perforin-like/cholesterol dependent cytolysin (MACPF/CDC) family or pore-forming proteins. Strikingly, despite <15% sequence identity, the *Vitiosangium* bGDSM active-state model reveals strict conservation of the final overall 'and-shaped' architecture of active mammalian GSDM proteins including the α 1 'thumb', the globular 'palm', the β 1– β 2 'wrist' loop, and the membrane-spanning 'fingers' (**Fig. 3b**)^{6,7}. Previous comparisons of the mammalian GSDM active-state structures revealed a key difference in the protein region between the globular domain and fingers region that controls the bend of the beta-sheet fingers and directs the pitch of membrane insertion^{6,7}. In contrast to the gentle bend of the

'knuckles' regions of the hGSDMD and mGDSMA3 active-state structures, the fingers and palm of the bGSDM active-state structure intersect at a nearly perpendicular angle and form a flat plateau from the inner palm to the knuckles (**Fig. 3b**). Superposition of the globular domains of the bGSDM and mammalian GSDM structures demonstrates that bGSDM forms the sharpest angle with respect to the palm region, rotating 37° inward compared to the hGSDMD conformation (**Fig. 3c**). This rotation compresses the bGSDM α1 thumb into the globular domain, where the Nand C-termini are held in unusually close proximity (**Fig. 3c**). Unlike the mammalian GSDM fingers whose β-strands differ in length from 15–22 aa, the bGSDM fingers are nearly uniform at 17–18 aa each, and the fingertips, which resemble 2–7 aa β-hairpins in mammalian GSDMs, are expanded into 8–11 aa loops (**Fig. 3b,c**). Importantly, multiple other bGSDM sequences, including the *Lysobacter enzymogenes* bGSDM involved in anti-phage defense, are each readily modeled using the *Vitiosangium* bGSDM active-state structure supporting that the conformational changes controlling bGSDM activation are likely conserved across diverse phyla (**Extended Data Fig. 8a**).



Fig. 3 | **Conservation of the GSDM active state structure from bacteria to mammals. a**, Top, DALI Z-scores from searching the active *Vitiosangium* bGSDM structure against the full Protein Data Bank (PDB). PDB depositions with multiple similar chains were manually removed to show only unique hits. Below the graph are representative pore-forming protein structures from the membrane attack complex/perforin superfamily (MACPF) and cholesterol-dependent cytolysins (CDCs), in addition to active mammalian gasdermins. Pneumolysin and perforin structures have been reduced by 60% in size. b, Inactive to active state transition of the *Vitiosangium* bGSDM structure. Left, transition from the crystal structure inactive state (PDB ID 7N51) to the active cryo-EM structure (this study). Right, topology diagrams of the inactive and active structures labeling α -helices and β -sheets based on mammalian GSDM structure superimposed onto the mammalian GSDM structures from (a) and active hGSDMB (PDB ID 8ET2). Structures were superimposed by their globular domains and angles were measured using an axis at the base knuckle of the bGSDM to points at the positively charged residues are indicated, including R13 of mGSDMA3 has been proposed to interact with the cardiolipin headgroup⁶. **d**, Views of *Vitiosangium* bGSDM slinky and pore models. On the far right, the bGSDM pore is shown encompassing the human GSDMD pore model (PDB ID 6VFE).

Comparison of the *Vitiosangium* bGSDM inactive-state crystal structure with the activestate cryo-EM model additionally explains bacteria-specific transitions that direct activation. Whereas the mammalian GSDM fingers form almost exclusively through the elongation of existing β -strands, the bGSDM fingers form via the local refolding of several alpha-helices ($\alpha 2''$, $\alpha 2'''$, and

 α 4) and the β 7' strand, which are largely absent in mammalian GSDM structures (**Fig. 3b** and **Extended Data Fig. 8b,c**). In particular, the α 2" helix contributes multiple residues to the hydrophobic groove housing the palmitoylated N-terminal cysteine residue that stabilizes the bGSDM autoinhibited state. Though no clear electron density could be observed for the cysteine in the active structure, reorientation of the α 1 thumb is predicted to flip the covalently bound palmitoyl for insertion into the surrounding lipid membrane (**Fig. 3b**).

To visualize the *Vitiosangium* bGSDM in its natural closed-pore state, we assembled a 54-mer structure based on the radius of the slinky oligomeric conformation and then realigned protomers into a 52-mer geometric model corresponding to the dominant size observed in 2D classification of intact pores (**Fig. 3d**). The resulting model supports continuous hydrogen bonding around a giant pore formed by 208 β -strands. Notably, the *Vitiosangium* bGSDM pore has an inner diameter of 330 Å that is wide enough to completely encompass the largest known mammalian pore of hGSDMD (**Fig. 3d**). We next introduced mutations to residues along key regions of the *Vitiosangium* bGSDM oligomeric interface and used a cell death assay to analyze bGSDM-mediated pore formation in bacterial cells. We observed that *Vitiosangium* bGSDM single substitution mutants retained the ability to induce cell death, indicating that the bGSDM pore assembly is remarkably stable and robust to minor perturbations across the protein length—a feature distinct from mammalian GSDM counterparts (**Extended Data Fig. 9**)^{6,17}.

Mechanism of bGSDM membrane assembly

The failure of single mutations in conserved regions and at the oligometric interface to inhibit bGSDM-mediated cell death suggests that bGSDM pore formation is a robust and energetically favorable process. Previously, X-ray crystal structures of Vitiosangium and other bGSDMs revealed palmitoylation of a conserved N-terminal cysteine residue that is also required for bGSDM-mediated cell death¹⁵. We therefore hypothesized that palmitovlation may contribute to the mechanism of membrane pore formation. To determine the function of palmitoylation of the Vitiosangium bGSDM, we mutated the conserved cysteine and residues in the surrounding pocket of the inactive structure and tested the function of these bGSDM variants for the ability to induce bacterial cell death. We observed that most mutations of the palmitoyl binding pocket did not compromise bGSDM-mediated cell death, including mutations of V89 and F111 which rearrange into the active state $\beta_3-\beta_5$ fingertip loop and knuckles of the β_5 strand, respectively (Fig. 4a,b and **Extended Data Fig. 8**). However, we found that alanine mutations of conserved L211 and F213 residues located between the $\beta 8$ and $\beta 9$ strands in the active state partially disrupted bGSDM function, and a C4A mutation to the modified cysteine residue itself resulted in bGSDM inactivation and near-complete rescue of bacterial growth (Fig. 4a.b). We next purified C4A mutants of Vitiosangium and Nostoc bGSDMs and analyzed both proteins in vitro with liposome rupture assays. Similar to our previous analysis of the Runella bGSDM¹⁵, the Vitiosangium and Nostoc bGSDMs purified as stable proteins but each were significantly reduced in the ability to form membrane pores and cause liposome leakage (Fig. 4c). These results demonstrate that the N-terminal palmitoyl modification plays a direct role in bGSDM pore formation (Fig. 4c).

We hypothesized that the palmitoyl group stimulates intermediate steps during bGSDM pore assembly. To test this hypothesis, we conducted atomistic MD simulations of the early and late steps of pore assembly (**Fig. 4d–f**). Starting from the full-length inactive *Vitiosangium* bGSDM crystal structure (*i*), we removed the 29 aa CTD and placed the inactive Δ CTD structure on a model bacterial membrane (*ii*). Despite lacking the inhibitory CTD, this structure was remarkably stable in two 2 µs long simulations with a Cα root mean squared deviation (r.m.s.d.) of ~2 Å from the initial structure. The palmitoyl remained sheathed inside the inactive state binding pocket. In the inactive state structure, the CTD forms contacts primarily with the bGSDM proto-finger region¹⁵. To mimic disorder in this region that likely occurs over an extended time following loss of inhibitory CTD contacts, we performed additional simulations (10 × 1 µs) for which the α2", α2"'', and α4 helices, the β7' strand, and their connecting loops were remodeled as fully disordered

loops (iii). We observed that the palmitoyl fatty-acid tail gains flexibility and in four of the ten replicate simulations explored regions outside of the pocket. Starting from these structures, we then performed steered MD simulations in which we pulled the palmitoyl fatty-acid tail into the membrane. In three 1 µs long replicate simulations, we observed that once the palmitoyl group entered the membrane it remained stably integrated in the inner leaflet without external bias (iv). In simulations of a single active state protomer and of 2–3-meric bGSDMs in the membrane, the bGSDMs formed stable coiled assemblies. For the trimer, narrow membrane-spanning and waterfilled channels formed, similar to those observed for hGSDMD¹⁹ (Fig. 4e and Extended Data Fig. 10). C4 palmitoylation did not noticeably affect the stability or shape of these small, possibly sublytic, pores. Finally, we simulated a complete 52-mer pore within a membrane. The pore was remarkably stable with r.m.s.d. values of ~3 Å with or without palmitovlation (Fig. 4f and Supplementary Video 1). Taken together, our simulation data suggest that unfolding of the bGSDM proto-finger region precedes palmitoyl membrane insertion. Integration of the covalently attached fatty-acid tail may ensure stable membrane binding of the vulnerable, partially unfolded state and push the system towards pore formation. Our simulations show that once the β-sheet is membrane inserted, the palmitoyl likely plays only a subordinate role.



Fig. 4 | Model for membrane insertion and pore formation of palmitoylated bGSDMs. a, Hydrophobic pocket surrounding the Nterminal palmitoylated cysteine residue of the Vitiosangium bGSDM in the inactive state (PDB ID 7N51). Select hydrophobic residues and the palmitoylated cysteine residue (C4) are indicated. b, The N-terminal cysteine is critical for cell toxicity bacteria, while residues that surround it in the hydrophobic pocket are not. Colony forming units (CFUs) per mL of E. coli grown on LB-agar plates (Extended Data Fig. 9). CFU were determined from bacteria harboring plasmids encoding the full-length bGSDM (full), the N-terminal poreforming domain alone (Δ CTD), or single mutations of the Δ CTD grown in triplicate. c, The N-terminal cysteine is critical for efficient pore formation in vitro. Liposome leakage assay of WT and N-terminal cysteine mutated (C4A) Vitiosangium and Nostoc bGSDMs. Engineered bGSDMs were paired with their matched site-specific protease. Error bars represent the SEM of three technical replicates and statistical significance was determined by one-way ANOVA and Tukey multiple comparison test. n.s. ≥ 0.05; ****P < 0.0001. d, Structural transitions during bGSDM activation. Schematic shows the anchoring process starting from the (i) inactive crystal structure of the bGSDM (PDB ID 7N51) before proteolytic cleavage; followed by snapshots of MD simulations of (ii) the membrane-adhered inactive Δ CTD; (iii) the inactive state with remodeled, disordered residues of the α 2", α 2", α 4 helices and the β 7 strand; and (iv) the inactive state with remodeled disordered residues after 1 µs of unbiased simulation following a steered simulation in which the palmitoyl (green) was inserted into the membrane (iv). e, Schematic of the unresolved intermediate steps of oligomerization and membrane insertion. f, MD simulation of the fully assembled 52-mer pore with and without palmitoylated C4. Snapshot of the pore in top view (left) and front view (bottom right) after 1 µs of simulation. Solvent and membrane lipid tails are omitted for clarity. (top, right) r.m.s.d. fluctuations of the palmitoylated and the non-palmitoylated systems with respect to the average structure of the last 900 ns of the simulation (transparent lines) and smoothed fluctuations (opaque lines; smoothed using a 30 ns window).

Our study collectively defines an ancient structural mechanism of GSDM pore formation. Comparison of the cryo-EM structure of the Vitiosangium bGSDM active-state with a previouslydetermined crystal structure¹⁵, allowed us to model how bacteria-specific domains in the inactive state transform into membrane-spanning β -sheets to form huge ~1.3 MDa pores. Reconstitution of pore formation using five diverse bGSDMs demonstrates that pore size is heterogeneous in nature and intrinsically determined by protein sequence. Mammalian GSDM pores have welldefined roles in secretion of interleukin signals¹⁷. Though interleukins are not conserved in bacteria, the divergent sizes observed for bGSDM pores suggest other molecules may be released to provide an additional layer of complexity to bGSDM-mediated anti-phage defense²¹. Whereas mammalian GSDMs and pore-forming toxins (PFTs) recognize specific lipids in target membranes^{4,5,22}, a remarkable feature of bGSDMs is the ability to form pores on membranes with simple lipid compositions¹⁵. This lack of specificity may reflect the absence of selective pressures on bGSDMs to distinguish the plasma membrane from the organelle membranes present in eukaryotes, or self from non-self membranes required for PFTs. Consistent with this observation, the α 1 thumb, which in mGSDMA3 mediates interactions with the cardiolipin head group via R13⁶. is reoriented in the active-state bGSDM (Fig. 3c). This finding suggests that cardiolipin-specific interactions are not conserved features of GSDM pores, despite the prevalence of cardiolipin in bacterial membranes. Nevertheless, future studies should investigate whether phylogenetically diverse bGSDMs are customized to the complex protein and lipid features of native bacterial membrane membranes²³. Besides extrinsic membrane features, bGSDMs appear to broadly utilize intrinsic properties such as palmitoylation to overcome energy barriers for pore formation on target membranes. Two recent studies describe the surprising requirement of enzymedependent palmitoylation of human and mouse GSDMD for pore formation in cells^{24,25}. Though the palmitoylated cysteine (C191 in humans) is located within the fingertip region instead of on the thumb in bGSDMs, we suggest that our model for palmitovl membrane insertion may explain why the modification is conserved across the tree of life.

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Competing Interests None declared.

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Methods

Bacterial growth assays

pBAD plasmids encoding full-length and Δ CTD GSDMs were constructed using gene synthesis (IDT) or PCR amplification from existing plasmids followed by Gibson assembly. Constructs expressing single amino acid mutations were generated by "around the horn" (ATH) mutagenesis or gene synthesis (IDT). The following GSDM sequences were used in this study: human GSDMD (Uniprot ID P57764), Runella bGSDM (IMG ID 2525253496), Unclassified Bacteroidetes metagenomic isolate bGSDM (simplified to "Bacteroidetes bGSDM", IMG ID 2806880301), Nostoc bGSDM (IMG ID 2631173059), Vitiosangium bGSDM (IMG ID 2831770670), and Ideonella bGSDM (IMG ID 2684147428). C-terminal truncations for the \triangle CTD GSDMs were based on experimentally demonstrated or predicted cleavage sites. For cleavage site predictions. proteolysis was assumed to occur within exposed loops of X-ray crystal or AlphaFold predicted structures, and before a small P1' residue (Gly, Ala, or Ser). The ∆CTD GSDMs had the following sequence lengths: 1–275 (hGSDMD), 1–247 (Runella bGSDM), 1–247 (Bacteroidetes bGSDM), 1-246 (Nostoc bGSDM), 1-237 (Vitiosangium bGSDM), and 1-244 (Ideonella bGSDM). Sequence-verified plasmids were transformed into E. coli TOP10 cells and plated on LB agar plates with 100 µg mL⁻¹ ampicillin and 1% glucose. Single colonies were used in triplicate to inoculate 5 mL liquid cultures in LB with 100 µg mL⁻¹ ampicillin and 1% glucose and were incubated with shaking overnight at 37°C. The next morning, cultures were 10-fold serially diluted in 1× phosphate-buffered saline (PBS) from 10° to 10^{-7} and 5 µL of each dilution was spot onto LB agar plates with 100 µg mL⁻¹ ampicillin and 1% glucose (to repress expression) or 0.2% arabinose (to induce expression from the arabinose promoter). Spots were allowed to dry at room temperature and plates were then grown overnight at 37°C. The plates were imaged using a Chemidoc system (Bio-Rad) and colony forming units (CFUs) were determined by counting colonies at appropriate dilutions. For conditions in which no colonies appeared in the 10^{-1} and the 10[°] dilution was uncountable, it was assumed that 1 colony was present in 10[°] and therefore 200 CFU mL⁻¹ total.

Recombinant protein expression and purification

bGSDM sequences were engineered for controlled activation by inserting site-specific protease cleavage motifs into predicted or verified cleavage sites using the target cleavage sequences for Tobacco Etch Virus (TEV) protease (ENLYFQ/G), the 3C-like protease (LEVLFQ/GP), the thrombin protease (LVPR/GS), or the WELQut protease (WELQ/G) where the backslash indicates the cleaved bond. bGSDM-encoding sequences were codon-optimized for E. coli and constructed by DNA synthesis (IDT). Gene fragments were cloned by Gibson assembly into a custom pET vector for expression fused to a 6×His-tagged SUMO2 tag. The N-terminal methionine residue was removed from each sequence and replaced by a single serine residue left as a scar following SUMO2 tag removal by the human SENP2 protease. For select constructs, alanine mutants of N-terminal cysteines were prepared by ATH mutagenesis or DNA synthesis (IDT). 40 total engineered bGSDM constructs were screened, and the best identified constructs were for the *Bacteroidetes* bGSDM with the TEV site replacing residues E242–G248, the *Nostoc* bGSDM with the WELQut site replacing residues V243–G247, the *Vitiosangium* bGSDM with the thrombin site replacing residues P234–M239, and the *Ideonella* bGSDM with the WELQut site replacing residues A241–G245.

Plasmids were transformed into BL21(DE3)-RIL *E. coli* and plated on MDG (0.5% glucose, 25 mM Na₂HPO4, 25 mM KH₂PO₄, 50 mM NH₄Cl, 5 mM Na₂SO₄, 2 mM MgSO₄, 0.25% aspartic acid, 100 mg mL⁻¹ ampicillin, 34 mg mL⁻¹ chloramphenicol, and trace metals) agar plates. 3–5 of the resulting colonies were used to inoculate starter cultures of MDG media for overnight growth at 37°C. MDG starter cultures were used to inoculate 2×1 L of M9ZB media (0.5% glycerol, 1% Cas-amino Acids, 47.8 mM Na₂HPO₄, 22 mM KH₂PO₄, 18.7 mM NH₄Cl, 85.6 mM NaCl, 2 mM MgSO₄, 100 mg mL⁻¹ ampicillin, 34 mg mL⁻¹ chloramphenicol, and trace metals) and grown at

 37° C with shaking at 230 rpm until the culture reached an OD₆₀₀ of ~2.5. Cultures were then chilled on ice for 20 min and induced for expression with 0.5 mM IPTG before shaking overnight at 16°C and 230 rpm. The next morning, expression cultures were pelleted by centrifugation, washed with PBS, re-pelleted, and pellets were flash frozen on liquid nitrogen and stored at ~80°C until purification.

Protein purification was performed at 4°C with all buffers containing 20 mM HEPES-KOH (pH 7.5) and other components as described below. Cell pellets were thawed and lysed by sonication in buffer containing 400 mM NaCl, 30 mM imidazole, and 1 mM DTT. Lysate was clarified by centrifugation and passing through glass wool, bound to NiNTA agarose beads (QIAGEN), which were subsequently washed with buffer containing 1 M NaCl, 30 mM imidazole, and 1 mM DTT. Protein was eluted with buffer containing 400 mM NaCl, 300 mM imidazole, and 1 mM DTT. Protein was eluted with buffer containing 400 mM NaCl, 300 mM imidazole, and 1 mM DTT. Human SENP2 protease was added to the eluate to cleave off the SUMO2 tag before dialyzing overnight into buffer containing 125–250 mM KCl and 1 mM DTT. For initial screening of candidates, the dialyzed protein was directly purified by size-exclusion chromatography (SEC) using a 16/600 Superdex 75 column equilibrated with 250 mM KCl and 1 mM TCEP. For select candidates, SEC was preceded by passing dialyzed protein through a 5 mL HiTrap Q ion exchange column (Cytiva) in 125 mM and 1 mM DTT. Sized proteins were brought to concentrations >20 mg mL⁻¹, using 10 kDa molecular weight cut-off concentrators (Millipore), flash frozen on liquid nitrogen, and stored at -80° C until use.

Liposome assays

Liposomes were prepared from 8 mg lipid total from chloroform dissolved stocks (Avanti) of 1,2dioleoyl-sn-glycero-3-phosphocholine (DOPC), 1-palmitoyl-2-oleoyl-glycero-3-phosphocholine (POPC), or a 3:1 w/w mixture of DOPC or POPC and cardiolipin (CL). Lipids were dried by transferring to a glass vial and evaporating chloroform under a stream of nitrogen gas and held under vacuum overnight in the dark. Liposomes for electron microscopy were prepared by resuspending the dried lipids with 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCl. To prepare liposomes for terbium-based liposome rupture assays, and the resuspension buffer was supplemented with 15 mL TbCl₃ and 50 mM sodium citrate. Liposomes were resuspended in a 800 μ L volume (10 mg mL⁻¹) by vortexing for 2 min, transferring to a 1.7 mL microfuge tube, and freeze thawing 3× with liquid nitrogen and a warm water bath. The crude liposome suspension was transferred to a glass tube, pulled into a glass syringe (Hamilton), and then extruded 21× with a mini-extruder (Avanti) using 19 mm nucleopore membranes with 0.2 μ M pores (Whatman) and 10 mm filter supports (Avanti). Extruded liposomes were purified by SEC with a 10/300 Superdex 200 column equilibrated with 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCl. Fractions containing liposomes were combined and stored at 4°C until use.

Liposome leakage assays were carried out with purified DOPC liposomes containing TbCl₃. 40 µL liposome reactions were assembled in a clear well 96-well PCR plate (BioRad) from 24 µL liposomes and 16 µL dipicolonic acid (DPA) dilution buffer at a final concentration of 20 µM DPA. Cleavage reactions were assembled in triplicate for the addition of 10 µL per well with final concentrations of 50 µM bGSDM with or without paired proteases in a buffer containing 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCl. Site-specific proteases were used at the following final concentrations: 5 µM homemade TEV protease, 0.28 units µL⁻¹ PreScission protease (Genscript), 0.25 units mL⁻¹ thrombin protease (MP Bioscience), or 0.1 units µL⁻¹ WELQut protease (ThermoFisher). After adding proteases, cleavage reactions were immediately added to wells containing liposomes and plate contents were spun to the bottom of wells by brief centrifugation. The plate was sealed and transferred to a Synergy HI microplate reader (BioTek) to monitor fluorescence resulting from Tb³⁺–DPA complex formation for 6 h with measurements every 1 min by excitation/emission at 276/545 nm.

Detergent screens of bGSDM pores

4 mL bGSDM cleavage reactions were prepared in glass tubes with a buffer containing 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCl. Reactions contained final concentration of liposomes at ~0.8 mg mL⁻¹, bGSDMs at 20 μM, and proteases as described above. Screens were initially performed using liposome compositions found to be most ideal by negative-stain EM analysis. Reactions were carried out at room temperature overnight. The next day, liposomes were harvested by centrifuging 150 μL portions at 60,000 × g at 4°C for 30 min with a TLA100 rotor. The supernatant containing uncleaved bGSDM and protease was removed, and pellets were resuspended using 1/5 or 1/10 dilutions of detergents stocks from the JBScreen (Jena Bioscience) or Detergent Screen (Hampton). Detergent resuspensions were incubated for 30 min at room temperature, centrifuged at 60,000 × g at 4°C for 30 min with a TLA100 rotor, and the supernatants were analyzed by SDS-PAGE. Select candidate extractions were analyzed by negative-stain EM and cryo-EM. All detergent-extractions were stored at 4°C until use. Of the detergents screened, the *Bacteroidetes* bGSDM pores were found to extract in 0.34 mM DDM, the *Vitiosangium* bGSDM pores were found to extract in 5.16 mM DDMAB.

Negative-stain electron microscopy

Select bGSDMs were cleaved with paired proteases in the presence of liposomes in a volume of 50 µL in a buffer containing 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCI. Reactions contained liposomes at concentrations of ~4 mg mL⁻¹ and all bGSDMs at 25 μ M with proteases concentrations as described above. Samples were incubated at room temperature overnight in glass tubes. To confirm cleavage, samples made in parallel without liposomes were analyzed by 15% SDS-PAGE and stained by Coomassie blue (Extended Data Fig. 2). To identify optimal liposomes for imaging with each bGSDM pore sample, reactions were prepared using liposomes composed of DOPC or POPC lipids with or without CL. Reactions were diluted 1/10 in 20 mM HEPES-KOH (pH 7.5) and 150 mM NaCl buffer and 5 µL was applied to glow-discharged 400 mesh copper grids coated with formvar/carbon film (Electron Microscopy Sciences). Grids were blotted after 1 min, stained with 2% uranyl formate for 2 s, blotted immediately, and air dried. Imaging was performed at 80 kV using the JEOL JEM 1400plus using AMT Image Capture Engine Software version 7.0.0.255. Select samples were imaged at 40,000× magnification to acquire enough micrographs for measuring the inner diameters of >50 pores. Bacteroidetes, Nostoc, and Ideonella bGSDM pores were all imaged in DOPC liposomes, while the Vitiosangium bGSDM was imaged from POPC-CL liposomes. Pore dimensions were measured manually using ImageJ version 2.9.0. Select detergent-extracted bGSDM pore samples were applied to grids and imaged and measured in the same manner as liposomes.

Cryo-EM sample preparation and data collection

For all bGSDM samples, 3 µL of sample was vitrified on grids using a Mark IV Vitrobot (ThermoFisher). 30 min prior to sample vitrification, grids were glow discharged using an easiGlow[™] (Pelco). Grid type and blotting time was optimized for each sample using a double-sided blot with a constant force of 0, in a 100% relative humidity chamber at 4°C, and a 10 s wait prior to plunging in liquid ethane before storing in liquid nitrogen. For the *Bacteroidetes* pores embedded in liposomes, the 1.2/1.3 Carbon Quantifoil[™] grids were used with a blotting time of 10 s. For the DDM extracted *Bacteroidetes* and *Vitiosangium* bGSDM pores, lacey carbon grids with carbon thin film support were used with a blotting time of 9 s. For the *Vitiosangium* bGSDM pores and pore-slinky mixtures extracted in HECAMEG detergent, lacey carbon grids with an additional 2 nm layer of carbon thin film were used with a blotting time of 9 s. The pore-slinky mixture represents the same sample as the pore-only sample, after leaving it for one week at 4°C. For the *Vitiosangium* bGSDM oligomeric slinkies extracted in DDMAB detergent, 2/1 Carbon

Quantifoil[™] grids with an additional 2 nm layer of carbon thin film were used with a blotting time of 9 s.

Cryo-EM data were collected either using a Talos Arctica (ThermoFisher) operating at 200 kV or a Titan Krios (ThermoFisher) microscope operating at 300 kV. Both microscopes were equipped with a K3 direct electron detector (Gatan). SerialEM software version 3.8.6 was used for all collections. Data collection on grids of the Bacteroidetes pores embedded in liposomes was performed using a Titan Krios microscope and a total of 4.340 movies were acquired at a pixel size of 0.83 Å, a total dose of 54.5 e⁻ /Å², dose per frame of 1.1e⁻ /Å² at a defocus range of -1 to -2 µm. Grids of the DDM extracted Bacteroidetes bGSDM pores were imaged using a Talos Arctica microscope and a total of 5,364 movies were acquired at a pixel size of 1.1 Å, a total dose of 51.7 e- $/Å^2$, dose per frame of 1.1e- $/Å^2$ at a defocus range of -1.0 to -2.0 µm. Data collection on grids of the Vitiosangium bGSDM pores was performed using a Titan Krios and a total of 8,930 movies were acquired at a pixel size of 0.83 Å, a total dose of 48.35 e- /Å², dose per frame of 1.1– $/Å^2$ at a defocus range of -1.0 to -2.0 μ m. Data collection on grids of the Vitiosangium bGSDM slinky-like oligomers extracted in DDMAB detergent was performed using a Titan Krios microscope and a total of 8,156 movies were acquired at a pixel size of 1.06 Å, a total dose of 51.8 e⁻ /Å², dose per frame of 1.04 e⁻ /Å² at a defocus range of -0.7 to -2.0 μ m. Data collection on grids of the Vitiosangium bGSDM pore-slinky mixture extracted in HECAMEG was performed using a Titan Krios microscope and a total of 4,818 movies were acquired at a pixel size of 0.83 Å, a total dose of 52.68 e- $/Å^2$, dose per frame of 1.1e- $/Å^2$ at a defocus range of -1.0 to -2.0 um. Micrographs shown in Extended Data Figure 4 and 6 were prepared using IMOD version 4.11.3.

Cryo-EM image processing and model building

For the *Bacteroidetes* bGSDM pores and the *Vitiosangium* bGSDM pores and pore-slinky mixtures, movies frames were imported to cryoSPARC²⁶ for patch-based motion correction, patch-based CTF estimation, and 2D classification. For the *Vitiosangium* bGSDM oligomeric slinkies, movies frames were pre-processed using the on-the-fly-processing scheme of RELION (version 3.1)²⁷. Motion correction was performed using MotionCor2²⁸ and the motion corrected movies were imported to cryoSPARC for patch-based CTF estimation, 2D and 3D particle classification, non-uniform refinement, and masked local refinement. The resulting cryo-EM density was post-processed using DeepEMhancer²⁹ prior to model building. Local resolution was estimated using cryoSPARC.

The globular domain of the inactive-state *Vitiosangium* bGSDM (7N51) was used as a starting model after docking it into a single protomer of the slinky-like oligomer using Coot³⁰. A complete protomer model was manual built in Coot and iterative corrections were made based on Phenix real-space refine³¹. The protomer model was validated in Phenix with MolProbity³². The 54-mer slinky model in Figure 3d was constructed by extending from a docked protomer by continuously fitting new protomers within the cryo-EM map density. To make the 52-mer pore model in Figure 3d, the protomer models were realigned using a custom script and a geometric model based on the radius of a low-quality 3D reconstruction of anisotropic cryo-EM data of the 52-mer pore. Protomers were realigned to preserve the inter-subunit hydrogen bonding pattern of the slinky-like oligomer. Structure figure panels were generated using UCSF ChimeraX³³ and PyMOL version 2.4.0.

Molecular dynamics simulations

Membrane. We set up the systems for molecular dynamics (MD) simulations using Charmm-GUI³⁴. As membrane template, we used 15.2 × 15.2 nm² patch mimicking the bacterial inner membrane (PVPE:PVPG:cardiolipin, 15:4:1) surrounded with 150 mM NaCl solution. We minimized the potential energy of this patch in 5,000 steepest descent steps and performed 250 ps of NVT and 1.625 ns of NPT equilibration while stepwise decreasing the initial positional

restraints of the phosphate z positions and the lipid tail dihedral angles. We used this membrane to set up simulation systems of 1–3 subunits of the *Vitiosangium* bGSDM. For simulations of the full 52-mer pore we multiplied the membrane patch along the xy plane to create a $62.3 \times 62.3 \text{ mm}^2$ large membrane patch.

bGSDM. We set up 1, 2, 3 and 52-mer *Vitiosangium* bGSDM systems in pore conformation by extracting the respective oligomer structures from the full 52-mer pore, with and without palmitoylated C4, inserting them into the membranes and removing all lipids with atoms overlapping with any protein (or C4 palmitoyl) atom, using Charmm-GUI to create simulation topologies. We alleviated membrane asymmetries by removing lipids from the overfilled leaflet. Subsequently, we resolvated the system with TIP3P water³⁵, removed water molecules in the membrane, and then added NaCl to a sodium concentration of 150 mM and chloride atoms to neutralize the charge of the system.

For simulations of the inactive conformation of *Vitiosangium* bGSDM, we removed the 29 aa CTD from the inactive crystal structure (PDB ID 7N51¹⁵) and either directly placed the inactive Δ CTD on the membrane patch or first remodeled α 2", α 2"', α 4 and β 7', and their connecting loops as fully disordered loops using swiss-model³⁶.

All MD simulations were performed using Gromacs version 2022.4 (Abrahams Gromacs reference) and the CHARMM36m force-field³⁷. We used a standard integration timestep of 2 fs. Electrostatic interactions were computed using the particle-mesh Ewald (PME) algorithm³⁸. Real-space electrostatic and van-der-Waals interactions were cut-off at 1.2 nm. We constrained the distance of bonds with hydrogen atoms using LINCS³⁹. We maintained a constant temperature of 37°C using the velocity-rescale algorithm⁴⁰ with a time constant of 1 ps, controlling protein, membrane and solvent (water and ions) individually. During system equilibration, we used the Berendsen barostat⁴¹ for semi-isotropic pressure coupling (x and y dimensions coupled together) with a time constant of 5 ps, a reference pressure of 1 bar, and a compressibility of 4.5×10^{-5} bar⁻¹. After the equilibration phase, we switched to the Parrinello-Rahman algorithm⁴² for pressure control, while keeping all other control parameters the same.

Each system was first energy minimized for 5,000 steepest descent steps. Then, equilibration simulations of 50 ns length each were performed. During these equilibration simulations, the initial positional restraints of the heavy atoms of the protein (and the phospholipids) were gradually lifted (Supplementary Table 1). No positional restraints were used during the production simulations. Simulations were run for times indicated in Supplementary Table 2 and in the case of replicate simulations were started with randomly drawn initial velocities according to the Maxwell-Boltzmann distribution. The two simulations of the membrane bound inactive structure were not started from the exact same system, but from two systems with initially slightly different conformations of the palmitoyl fatty acid tail.

In case of the simulations of membrane-bound bGSDM with remodeled proto-finger region, for each of the 10 unbiased replicate simulations, we extracted a frame after 50 ns of production and started a steered MD simulation in which we pulled the palmitoyl fatty acid tail into the membrane. The pull group comprised the carbon atoms of only the fatty acid tail. A harmonic spring (force constant 100 kJ mol⁻¹ nm⁻²) connected its center of mass to a dummy atom that moved into the membrane along the membrane normal at a constant velocity of 0.01 nm ns⁻¹. At 305 ns, after tail insertion, we extracted a frame from one of the simulations and started 3 independent replicate simulations without any external potential applied.

To calculate the C α root mean squared deviation (r.m.s.d.) of the pore, we first calculated the average root mean squared fluctuation (r.m.s.f.) of each residue's C α and excluded the highly flexible protein regions (termini and fingertip regions with r.m.s.f. > 3.5 Å) from the subsequent r.m.s.d. calculation. The r.m.s.d. was calculated with respect to the average structure of the last 900 ns of the production simulation.

Visual analysis as well as image and movie rendering were performed using PyMOL, VMD^{43} , and UCSF ChimeraX³³.

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Data Availability Statement: The coordinates and density map of the active-state *Vitiosangium* bGSDM are being deposited with the PDB and the EMDB under the accession codes NNN and NNN. All other data are available in the manuscript or the supplementary materials.

Extended Data Figures



Extended Data Fig. 1 | Expression of bGSDM NTDs induces potent cellular toxicity. a, Crystal structure of an inactive bGSDM from a *Vitiosangium* species (PDB ID 7N51) and indication of the disordered loop that was targeted for cleavage site engineering. **b**, Colony forming units (CFU) per mL of *E. coli* derived from spot assays shown in panel (b). **c**, *E. coli* harboring plasmids encoding full-length GSDMs (full) or the N-terminal pore-forming domain alone (Δ CTD) were grown on LB-agar plates in triplicate. LB-agar contained either 1% glucose or 0.2% arabinose to repress or induce expression, respectively. Cells were serially diluted and plated out from left (10⁰) to right (10⁻⁷) with 5 µL per spot. Though the *Ideonella* Δ CTD construct does not drastically

reduce the CFUs compared to the full construct, colonies grow more slowly and appear fainter in agreement with toxicity from pore formation.



Extended Data Fig. 2 | Cleavage site engineering enables analysis of diverse bGSDM pores. a, Engineered bGSDMs were treated with or without paired site-specific proteases for 18 h at

room temperature and analyzed by 15% SDS-PAGE and visualized by Coomassie staining. Cleaved bGSDM proteins are indicated with a yellow asterisk. **b**, Full time-course of liposome leakage assays related to Fig. 1b. Error bars represent the SEM of three technical replicates.



Extended Data Fig. 3 | Negative-stain EM micrographs of engineered bGSDM poreliposome samples. Left panels represent larger view fields of micrographs shown in Fig. 1d, while right panels represent a second example used to measure pore sizes for Fig. 1c. Scale bar = 50 nm.



Extended Data Fig. 4 | Cryo-EM data processing of *Bacteroidetes* **bGSDM pores. a,** Representative cryo-EM micrograph and select 2D class averages of bGSDM pores from poreliposome samples. **b,** Representative cryo-EM micrograph and select 2D class averages of DDM-

extracted bGSDM pores. Numbers in the upper left-hand corner of 2D classes represent the number of bGSDM protomers observed in that class. Scale bars = 20 nm.



Extended Data Fig. 5 | Negative-stain EM micrographs of detergent-extracted Vitiosangium bGSDM pores and slinkies. a, HECAMEG detergent-extracted bGSDM pores. Scale bar = 50 nm. **b**, Comparison of inner diameters measured from pore-liposome samples (Fig. 1 and Extended Data Fig. 2) and HECAMEG detergent-extracted bGSDM pores. The number of pores measured (n) from each sample is as follows: pore-liposome (n = 56), extracted pore (n = 189). The black bar represents the average inner diameter of measured pores. **c**, DDMAB detergent-extracted bGSDM slinkies. Scale bars = 50 nm.



Extended Data Fig. 6 | Cryo-EM data processing of Vitiosangium bGSDM pores and slinkies. a, Representative cryo-EM micrograph and 2D class averages of HECAMEG detergent-extracted bGSDM pores. **b**, Single-particle processing schematic of DDMAB detergent-extracted bGSDM slinkies. From top to bottom: representative cryo-EM micrograph and 2D class averages (as in Fig. 2b), particle classification and map refinement, and local resolution estimate of final

map. **c**, Fourier shell correlation (FSC) curves versus resolution of bGSDM slinky map. Resolution was estimated at an FSC of 0.143. Scale bars = 50 nm. **d**, Representative cryo-EM micrograph and 2D class averages of HECAMEG detergent-extracted bGSDM pore-slinky mixture (as in Fig. 2a).



Extended Data Fig. 7 | Cryo-EM model to map fitting of bGSDM slinky and pore. a, 54-mer model of the *Vitiosangium* bGSDM in a slinky-like oligomerization. **b**, Examples regions of model to map fit quality for a single *Vitiosangium* bGSDM protomer. The map surface has been contoured to 16σ .



Extended Data Fig. 8 | Functional conservation of the GSDM active state. a, Predicted bGSDM structures are organized from left to right based on percent sequence identity to the Vitiosangium bGSDM: Runella bGSDM (18%), Bradyrhizobium bGSDM (19%), and Lysobacter bGSDM (28%). Phyre homology model threading utilized the active human GSDMD structure (PDB ID 6VFE, left model) or the active Vitiosangium bGSDM structure (this study, right model). Each bGSDM structure was also predicted using AlphaFold after deleting ~20 amino acids from the C-termini and yielded inactive-like structures (center). The modeled sequences are as follows: Runella bGSDM (1–247), Bradyrhizobium bGSDM (1–237), and Lysobacter bGSDM (1–240). b, Structure-based alignment of bacterial and mammalian gasdermins. The Vitiosangium bGSDM active structure was aligned to the active mouse GSDMA3 (PDB ID 6CB8) and human GSDMD (PDB ID 6VFE). Secondary structures are indicated below the sequences for each structure, in addition to secondary structures from the Vitiosangium bGSDM inactive state crystal structure. Residues of the Vitiosangium bGSDM that surround the palmitoyl in the inactive state are boxed in black, and asterisks indicate residues that have been mutated to test their effect on bGSDMmediated bacterial cell death. c, Topology diagrams showing the transitions from inactive to active structures of the Vitiosangium bGSDM (left) and mouse GSDMA3 (right). Conserved α-helices



and β -strands are outlined in black, and the positions of residues universally conserved in charge, identity, or aromaticity are indicated in active state topology diagrams.

Extended Data Fig. 9 | Cell death induced by the Vitiosangium bGSDMs is robust to single mutations across the protein length. a, Colony forming units (CFU) per mL of *E. coli* derived from spot assays shown in panel (c). Growth assays test single charge-swap mutations to residues at select interfaces in the active model. b, Colony forming units (CFU) per mL of *E. coli* derived from spot assays shown in panel (d). **c–e**, *E. coli* harboring plasmids encoding full-length GSDMs (full) or the N-terminal pore-forming domain alone (Δ CTD) were grown on LB-agar plates in triplicate. LB-agar contained either 1% glucose or 0.2% arabinose to repress or induce expression, respectively. Cells were serially diluted and plated out from left (10⁰) to right (10⁻⁷) with 5 µL per spot.



Extended Data Fig. 10 | MD simulations of small membrane pores formed by active 1, 2, and 3-meric bGSDM assemblies. Snapshots of palmitoylated (left) and non-palmitoylated (right) mono- and oligomers after 3.3 µs of simulation in a bacterial membrane in top view and side view. Protein shown in purple cartoon representation, membrane phosphates shown as tan spheres, palmitoylated C4 shown in cyan opaque licorice and transparent surface representation. In the side views, water inside the small pores is shown in blue transparent surface representation. Otherwise, solvent molecules and lipid fatty acid tails are omitted for clarity.

	Vitiosangium bGSDM slinky-like oligomer (EMD-NNNN) (PDB NNNN)
Data collection and processing	
Magnification	81.000
Voltage (kV)	300
Electron exposure $(e - / Å^2)$	51.8
Defocus range (um)	-0.7 to -2.0
Pixel size (Å)	1.06
Symmetry imposed	C1
Initial particle images (no.)	228.905
Final particle images (no.)	132.403
Map resolution (Å)	3.3
FSC threshold	0.143
Map resolution range (Å)	3.44–18.85
Refinement	
Initial model used (PDB code)	/N51
Model resolution (A)	3.52
FSC threshold	0.143
Model resolution range (A)	3.82–3.52
Map sharpening <i>B</i> factor (A ²)	-62.2
Model composition	(=0=
Non-hydrogen atoms	1/3/
Protein residues	228
B factors (A ²)	
Protein	142.94
R.m.s. deviations	
Bond lengths (A)	0.003
Bond angles (°)	0.621
Validation	4.05
MolProbity score	1.95
Clashscore	7.96
Poor rotamers (%)	0.53
Ramachandran plot	o. / / =
Favored (%)	91.15
Allowed (%)	8.41
Disallowed (%)	0.44

Cryo-EM data collection, refinement and validation statistics

Extended Data Fig. 11 | Cryogenic-electron microscopy data summary table. Table contains details of all cryo-EM data collection, processing, and refinement statistics used in this study.

Supplementary Table 1 | Restraints used during energy minimization (EM) and stepwise

equilibration	(EQ)) in kJ	mol ⁻¹	nm ⁻² .
oquinoration		,		

Step	Time [ns]	Backbone	Sidechains	Lipid headgroup (z)	Lipid dihedrals
EM		4000	2000	1000	1000
EQ 1	50	2000	1000	50	0
EQ 2	50	500	100	0	0

Supplementary Table 2 | Overview over bGSDM production simulations.

Conformation	Number of	C4	Simulated	Relaxed	Number
	subunits	palmitoylation	time [µs]	membrane	of atoms
				size [nm²]	
pore	52	yes	1	59.6 × 59.6	4547198
pore	52	no	1	59.7 × 59.7	4549291
pore	1	yes	3.3	12.3 × 12.3	197342
pore	2	yes	3.3	11.9 × 11.9	193401
pore	3	yes	3.3	11.9 × 11.9	192685
pore	1	no	3.3	12.0 × 12.0	194776
pore	2	no	3.3	11.9 × 11.9	193385
pore	3	no	3.3	11.9 × 11.9	192631
Membrane-bound	1	yes	2×2	12.2 × 12.2	203086 /
inactive ∆CTD					203095
Membrane-bound	1	yes	10 × 1	12.2 × 12.2	202541
Δ CTD with					
remodeled proto-					
finger region					
Steered palmitoyl	1	yes	10 × 0.35	12.2 × 12.2	202541
insertion					
Unbiased after	1	yes	3 × 1	12.2 × 12.2	202541
steered palmitoyl					
insertion					