A saposin-lipoprotein nanoparticle system for membrane proteins

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A limiting factor in membrane protein research is the ability to solubilize and stabilize such proteins. Detergents are used most often for solubilizing membrane proteins, but they are associated with protein instability and poor compatibility with structural and biophysical studies. Here we present a saposin-lipoprotein nanoparticle system, Salipro, which allows for the reconstitution of membrane proteins in a lipid environment that is stabilized by a scaffold of saposin proteins. We demonstrate the applicability of the method on two purified membrane protein complexes as well as by the direct solubilization and nanoparticle incorporation of a viral membrane protein complex from the virus membrane. Our approach facilitated high-resolution structural studies of the bacterial peptide transporter PeptTso2 by single-particle cryo-electron microscopy (cryo-EM) and allowed us to stabilize the HIV envelope glycoprotein in a functional state.

Membrane proteins are encoded by approximately 30% of all open reading frames1. Membrane proteins have essential roles in many cellular processes, such as signal transduction, cell-to-cell communication, membrane transport and lipid and energy metabolism, and they represent the targets of more than 60% of drugs in clinical use2. However, membrane proteins are difficult to study because of their instability and tendency to aggregate when extracted from their natural lipid-bilayer environment. In order to maintain their integrity, an artificial hydrophobic environment is needed. Detergent micelles are commonly used to solubilize membrane proteins, but they often have adverse effects on protein activity, stability and solubility and can interfere with the experimental setup3–5.

A major challenge is maintaining membrane proteins in a lipid-like environment while keeping them stable and monodisperse in solution, so that they become accessible for biochemical, biophysical and structural studies. Methods for reconstituting membrane proteins in lipid nanoparticles, such as liposomes and high-density lipoprotein particles6,7 based on apolipoproteins, also termed nanodiscs, provide a potential solution. These approaches have been used routinely for biochemical and biophysical studies of membrane proteins and have also been applied for structure determination by single-particle cryo-EM8–10. However, optimization of the technologies for individual membrane proteins is relatively laborious.

Here we present a lipid nanoparticle system that is based on the saposin protein family. Saposins are known to be modulators of lipid membranes11,12, mostly at an acidic pH within lysosomes. Given their lipid-binding properties, we hypothesized that they could be used to generate a nanoparticle system for the incorporation of membrane proteins (Fig. 1). In this paper we report a methodology for reconstituting membrane proteins in saposin-lipoprotein (Salipro) nanoparticles that allowed us to stabilize fragile membrane protein complexes in detergent-free buffer systems for functional and structural studies.

RESULTS
Generation of saposin-lipid nanoparticles
Members of the saposin protein family have membrane-binding and lipid-transport properties11,12. Several crystal structures of saposin proteins have been reported, in both the absence and the presence of lipids or detergents13–16. It was recently demonstrated that saposin A forms lipid complexes at an acidic lysosomal pH when incubated with unilamellar liposomes16. We therefore rationalized that saposins could be used as scaffolding proteins in a versatile lipid nanoparticle system to reconstitute membrane proteins in a lipid environment, without using liposomes and an acidic pH.

First we tested whether it was possible to incorporate lipids into saposin A complexes without using an acidic pH or liposomes. We incubated saposin A (Online Methods) with detergent-solubilized phospholipids at physiological pH 7.4 and then removed the

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Salipro nanoparticles stabilize membrane proteins. Incubation of saposin A with lipids and membrane proteins leads to self-assembly of soluble and stable nanoparticles that adjust to the size of the incorporated molecule. Models of saposin-lipid complexes were adapted from PDB IDs 4DDJ, 4APS, 4NCO, 3DIN and 2DOB; virus envelope structure adapted with permission from ref. 48.

Given the Salipro nanoparticles’ capacity to integrate detergent-solubilized lipids at physiological pH, we focused on the potential for reconstituting membrane proteins. We incubated detergent-purified membrane proteins with lipids and saposin A and then carried out the detergent-removal steps described above. The model proteins we tested included an archaeal mechanosensitive channel17 (T2, 4 predicted transmembrane helices per monomer, 32.9 kDa, putative homopentamer) and a bacterial peptide transporter18,19 (PepTSo2, 14 transmembrane helices per monomer, 56 kDa, homotetramer).

When SEC buffer without detergent is used, these membrane proteins are not soluble but do aggregate and will not elute from the column. In contrast, after incubation with saposin A and lipids, the gel-filtration profile of the archaeal T2 channel showed three symmetric peaks (Fig. 2b); the first peak corresponded to the T2 channel incorporated into Salipro nanoparticles, whereas the second and third peaks originated from lipid-only Salipro nanoparticles and monomeric saposin A, respectively. Thus we concluded that saposin A, lipids and the T2 channel associate in such a way that they form water-soluble particles with incorporated membrane protein.

Peak fractions corresponding to Salipro-T2 were pooled, concentrated and flash-frozen before further analysis via electron microscopy (EM). Salipro-T2 particles that
had been concentrated, frozen, thawed and diluted maintained their assembled state as judged by analytical SEC (Supplementary Fig. 2a). Negative-stain EM showed a homogeneous population of single particles that adopted random orientations on the grid (Fig. 2c,d). The particles were more distinct than the T2 channel embedded in micelles of the detergent n-dodecyl-β-D-maltoside.\(^\text{17}\) In side views, both individual particles and two-dimensional (2D) class averages of Salipro-T2 particles displayed two moieties of different sizes, representing a more compact domain and a bilobed domain (Fig. 2d).

**Cryo-EM of Salipro-POT**

Proton-coupled oligopeptide transporters (POTs) regulate the uptake of di- and tripeptides and are conserved among all kingdoms of life. We tested our ability to integrate the bacterial peptide transporter PeptTSo2, referred to here as POT, into Salipro nanoparticles. We incubated purified POT with saposin A and lipids. The gel-filtration profile of the bacterial peptide transporter showed three symmetric peaks (Supplementary Fig. 2b), with the first peak corresponding to the membrane protein incorporated into Salipro nanoparticles. Generally, the formation of Salipro particles with incorporated membrane proteins such as POT is feasible within a physiological pH range (Supplementary Fig. 2c). Prior to further analysis, peak fractions of Salipro-POT were pooled, concentrated and flash-frozen. Thermal unfolding analysis using dye-free differential scanning fluorimetry\(^\text{20}\) showed that the peptide transporter was significantly more stable when embedded in Salipro nanoparticles (melting temperature \(T_m = 72\ \text{°C}\)) than when in the detergent nonyl-β-D-maltopyranoside \(T_m = 43\ \text{°C}\) (Supplementary Fig. 2d).

We used single-particle cryo-EM to determine the structure of POT embedded in Salipro nanoparticles. POT is composed mostly of transmembrane helices and contains only short loop regions that are not embedded in the membrane. Thus there are no significant features outside of the membrane that can be used to align images of the molecules, which makes POT a difficult target for single-particle cryo-EM. We prepared cryo-EM grids of the Salipro-POT complex by the standard plunge-freezing method\(^\text{21}\) at a moderate concentration \((0.7\ \text{mg}\ \text{ml}^{-1})\). Cryo-EM images showed that the frozen hydrated complexes of Salipro-POT were homogeneous and monodisperse (Fig. 3a).

We collected a cryo-EM data set according to established data-acquisition and motion-correction procedures.\(^\text{22}\) The 2D class averages displayed the features of secondary and tertiary structures, as well as the tetrameric organization of the peptide transporter (Fig. 3b). Using a total of 9,913 particles, we determined a 3D reconstruction following a gold-standard refinement procedure with fourfold symmetry to a resolution of 6.5 Å (estimated with the criterion of Fourier shell correlation (FSC) = 0.143) (Fig. 3c). All secondary structural features were clearly resolved in the final 3D density map (Fig. 3d). The Euler angle distribution showed that particles were evenly distributed in different orientations in vitreous ice (Fig. 3e). The local resolution in the core region of the nanoparticle, representing the transmembrane helices of the peptide transporter, was significantly higher than that in the regions that could be attributed to the more flexible saposin-lipid scaffold (Fig. 3f and Supplementary Fig. 3). The cryo-EM density revealed a square-shaped disc with overall dimensions of \(12\ \text{nm} \times 12\ \text{nm} \times 5\ \text{nm}\) with well-resolved transmembrane helices (Fig. 4a and Supplementary Video 1). Given the size (Fig. 4a) and the apparent stoichiometry (Supplementary Fig. 2c) of Salipro-POT, we can assume that each particle is composed of four POT molecules and four saposin A molecules (Fig. 4a and Supplementary Video 1).

The crystal structure of the peptide transporter\(^\text{19}\) could be directly docked into the cryo-EM density as a rigid body without modification (Fig. 4). Whereas the transmembrane helices were well resolved, the Salipro scaffold of saposin and lipids seemed to be more flexible (Fig. 3f and Supplementary Fig. 3). Fitting of saposin A into the scaffold density was therefore done with a map displayed at a lower isosurface level (Fig. 4a and Supplementary Video 1). The general dimensions of the Salipro scaffold were in agreement with a recently published structure of saposin A.\(^\text{16}\) At the current low resolution for the saposin-lipid belt, we cannot exclude the possibility that saposin might also directly interact with the transmembrane helices of the bacterial transporter. However, we hypothesize that if saposin did indeed
Figure 4 | Cryo-EM structure of Salipro-POT. (a) 3D density map of the tetrameric bacterial transporter, filtered to 6.5 Å, showing each of the subunits and tentative placement of saposin (orange and blue, respectively) in the density. Shown are a top view and a side view cut perpendicularly to the plane of the membrane. (b) Representative views of the cryo-EM density and rigidly fitted X-ray crystallography model (PDB ID 4TPH). (c) Selected side-chain densities of bulky residues such as Arg, Trp, Phe and Tyr.

bind directly to a transmembrane helix, it would be less flexible, and thus in turn it would be possible to resolve the α-helices of saposin. This was not the case in the current study; the flexibility and hence lower resolution of the saposin-lipid scaffold did not allow for resolution of the α-helices of saposin (Fig. 3f and Supplementary Fig. 3).

The cryo-EM density allowed for clear identification and positioning of the POT transmembrane helices (Fig. 4b). Moreover, the EM density revealed loop and helix densities that could not be modeled in the crystal structure of this peptide transporter19. The map also exhibited bulky side-chain densities from amino acids, the positions of which were in good agreement with the rigid-body fitting of the crystal structure of the peptide transporter (Fig. 4c).

Stabilization of the functional HIV-1 spike protein

Membrane proteins of enveloped viruses often represent potent antigens that can be used for vaccination24. The trimeric HIV-1 envelope glycoprotein (HIV-1 spike), which mediates viral entry into host target cells, is a prominent example, as it is the only viral protein exposed to the immune system during infection24.

Initially we screened six detergents with high critical micelle concentrations. HEGA-10, MEGA-10 and C-HEGA-11 preserved the trimeric HIV-1 spike during solubilization on ice for 10 min, as judged by blue native (BN)-PAGE gel shift assay (Supplementary Fig. 4a). The viral spikes in detergent dissociated completely into monomers within 30 min at 37 °C (Supplementary Fig. 4b). To stabilize the trimeric association, we reconstituted the HIV-1 spike in Salipro nanoparticles. We solubilized purified VLPs with 9 mM HEGA-10 in the presence of 0.1 mg ml−1 saposin A (for optimization, see Supplementary Fig. 4b) and then removed HEGA-10 using spin SEC columns (Supplementary Fig. 5). BN-PAGE analysis showed that HIV-1 spikes eluted as soluble trimers only in the presence of saposin A, indicating reconstitution in Salipro–HIV spike particles (Fig. 5c). When saposin A was omitted, HIV-1 spikes could not be resolved because of protein aggregation (Fig. 5c).

Figure 5 | Stabilization of soluble and functional HIV-1 spike proteins in Salipro nanoparticles. (a) The HIV-1 spike protein in the viral membrane. The HIV-1 envelope glycoprotein (gp) (i.e., the spike) consists of two subunits, the peripheral gp120 subunit and the transmembrane gp41 subunit, forming a heterotrimer. (b) Salipro–HIV spike reconstitution. Purified VLPs containing HIV-1 spikes (brown) were solubilized with 9 mM HEGA-10 in the presence of 0.1 mg ml−1 saposin A (for optimization, see Supplementary Fig. 4b). After detergent removal, the soluble trimers were resolved on spin SEC columns. (c) Lectin purification of Salipro–HIV spikes. Particle purity was assayed using SDS-PAGE and SYPRO Ruby protein staining. The gels show efficient removal of contaminating viral internal proteins. CA, capsid; MA, matrix. (e) The HIV-1 spike has a native fold and preserves its function in Salipro nanoparticles. Shown are representative gels from BN-PAGE analysis of Salipro–HIV spike particles incubated at 37 °C for 16 and 90 h and then at 37 °C for 2 h with 10 μg ml−1 HIV-1 spike ligands, PG16 (150 kDa) (lane 3), sCD4 (50 kDa) (lane 4), sCD4 and 17b Fab (50 kDa) together (lane 5), 17b Fab alone (lane 6) or no ligand (lane 2). Lane 1 shows results for a control sample without any ligand that was kept on ice. Binding of the ligands can be followed by the reduced migration of the Salipro–HIV spike particles.
A second purification step using lectin affinity chromatography successfully removed viral matrix and capsid proteins, as well as excess unbound saposin A, leading to a pure Salipro–HIV spike sample as determined by SDS-PAGE (Fig. 5d and Supplementary Fig. 5).

We investigated the stability and functionality of radiolabeled lectin-purified Salipro–HIV spikes by BN-PAGE. Whereas detergent-solubilized HIV-1 spikes dissociated completely within 30 min at 37 °C (Supplementary Fig. 4), Salipro–HIV spikes remained intact for 16 h and 90 h at 37 °C as judged by BN-PAGE (Fig. 5e). Subsequent to the 16-h and 90-h incubations at 37 °C, we assessed the functionality and native features of the Salipro–HIV spikes by binding them to several protein ligands for additional 2 h at 37 °C. Binding of these ligands to the Salipro–HIV spike increased its apparent molecular weight and therefore decreased its mobility in BN-PAGE correspondingly, as shown before with detergent–HIV-1 spike complexes.27

The broadly neutralizing antibody PG16 binds to variable loops V1-V2 and V3 at the top of preferentially intact and functional trimeric HIV-1 spikes27–29. We found that PG16 bound efficiently to the Salipro–HIV spikes even after 16-h and 90-h incubations at 37 °C (Fig. 5e). In addition, a soluble version of the primary HIV-1 receptor CD4 (sCD4) bound stoichiometrically to the Salipro–HIV spikes (Fig. 5e). Binding of CD4 induces a conformational change in the spike that is essential for binding of the coreceptor site-directed monoclonal antibody 17b Fab30,31. We analyzed the binding of 17b Fab to Salipro–HIV spike–sCD4 complexes via BN-PAGE, which revealed efficient binding (Fig. 5e). We noted that 17b Fab did not bind to the Salipro–HIV spikes in the absence of sCD4 (Fig. 5e). This confirmed that sCD4 binding induced proper conformational changes to the reconstituted Salipro–HIV spikes. Together, our data demonstrate that the HIV-1 spike can be properly reconstituted in Salipro nanoparticles, preserving the fragile trimeric HIV-1 spike structure in a native, stable and functional state for at least 90 h at 37 °C.

**DISCUSSION**

Here we demonstrate that it is possible to incorporate membrane proteins into saposin-based lipid nanoparticles that allow for structural and functional studies of membrane proteins in detergent-free buffer systems. The formation of Salipro nanoparticles is comparably straightforward: lipids are mixed with membrane proteins and saposin A, which leads to a rapid self-assembly reconstitution process. Moreover, it is possible to incorporate viral membrane proteins by mixing saposin A with virus particles, leading to a fully functional viral membrane protein embedded in Salipro nanoparticles. The nanoparticles are composed of lipids and saposin A, a member of the saposin-like protein family.11–16 This class of proteins is composed of amphipathic helices and six invariable cysteine residues that form disulfide bridges, leading to extraordinary stability of the domains.11 We assume that the intrinsic stability of the saposin proteins, which surround the lipid core of the nanoparticle, in turn renders the nanoscale protein-lipid complexes remarkably stable and uniform in a wide temperature range (Supplementary Fig. 1c).

Various methodologies have been developed to stabilize membrane proteins in an aqueous environment7,32–39 (Supplementary Table 1). In efforts to achieve high resolution, both amphipols and maltose-neopentyl glycol-3 analogs have been shown to facilitate structural studies of membrane proteins,40,41 indicating their stabilizing effect on membrane proteins and their tight binding to the membrane protein. However, these systems do not provide a lipid environment, which is known to be of great importance for membrane protein structure and function.

Discoidal high-density lipoprotein particles, also termed nanodiscs42, provide a lipid environment, but they do not bind tightly to the transmembrane helices of the membrane protein. Additional apolipoprotein-based nanoparticle systems have been developed that may vary in diameter with respect to the classical nanodisc.32,35,39 Generally, the size of a nanodisc is dictated by the length of the scaffolding apolipoprotein belt at an optimum lipid content.42 The classical nanodisc technology was established more than 15 years ago, but so far the application of nanodiscs for single-particle cryo-EM studies has been limited to relatively large targets at subnanometer resolution9,10.

The Salipro nanoparticle system combines the advantages of adapting to the size of the incorporated membrane proteins (such as detergents or amphipols) with those of a lipid environment. Moreover, adjustment of the methodology for individual membrane proteins is comparably straightforward. Using the Salipro system, in this study we determined a subnanometer-resolution structural picture of the POT complex, which is a challenging target for structural studies by single-particle cryo-EM because it contains only short loop regions and has almost no structural features that stick out of the plane of the membrane region. Therefore, our image alignment was driven entirely by the structural features of the transmembrane domain. This may have been facilitated by the relatively small size of the saposin-lipid scaffold, which does not significantly affect the alignment process but enhances the contrast in cryo-conditions (Figs. 3 and 4 and Supplementary Fig. 3).

It has been shown previously that two saposin A molecules can encapsulate a lipid core.16 Here, in the presence of a large membrane protein complex, we observed that several saposin A proteins may encircle a large hydrophobic lipid-protein assembly in the case of the homotetrameric POT transporter (Fig. 4a and Supplementary Video 1). This suggests significant flexibility of saposin A, allowing it to assemble into homogeneous and stable lipoprotein complexes by adapting to the size of the incorporated molecules. This intrinsic flexibility of saposin A–lipid complexes is likely to facilitate the formation of nanoparticles that can accommodate membrane proteins of different sizes and, at the same time, assist in the formation of homogeneous lipid nanoparticles (Figs. 2–4).

Moreover, we have also presented a method to extract and stabilize fragile membrane proteins from the virus membrane. The HIV-1 spike is the only viral protein that is exposed to the immune system by intact virus particles and therefore is an interesting target for HIV vaccine development24,43,44. Antigen candidates that could preserve or sufficiently mimic the native conformation of the HIV-1 spike including the important membrane domain have long been sought after.43,44 It has been shown that regions proximal to the membrane as well as the membrane domain of the HIV-1 spike are important for the generation of broadly neutralizing antibodies to this virus45–47.

To our knowledge, the HIV-1 spike protein preparation presented here using the Salipro system represents the first approach
that allows for stabilization of the HIV-1 spike, including of the important membrane domains, in a soluble and functional state. Thus, the strategy applied here may represent a novel approach for the generation of immunogen or vaccine candidates that might induce broadly neutralizing antibodies to HIV-1. We envisage that this approach may be applicable to other viral envelope proteins such as influenza virus hemagglutinin, Ebola virus G protein and hepatitis C virus E protein.

In summary, Salipro particles provide a versatile nanoparticle platform for membrane proteins and may have a wide range of potential applications, from structural biology to the discovery of new pharmaceutical agents and the therapeutic delivery of protein-based therapeutics and vaccines.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. The 3D cryo-EM density map of Salipro-POT has been deposited in the Electron Microscopy Data Bank under accession number 3302.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

J.F. developed the concept, performed experiments, analyzed data and wrote the manuscript. R.L. developed, performed and analyzed HIV-related experiments. J.-P.A. carried out all cryo-EM experiments, including data acquisition, processing and data interpretation. F.G., P.M., A.F. and C.L. purified membrane proteins. A.F.-P.S. prepared grids, collected negative-stain data and conducted negative-stain data on lipid-only Salipro nanoparticles. J.A.G.B., H.G., Y.C. and P.N. contributed project feedback and comments on the manuscript. All authors contributed to data interpretation and preparation of the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare competing financial interests: details are available in the online version of the paper.
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Expression and purification of saposin A. Protein expression was carried out using a vector with the coding region for saposin A inserted into a pNIC-Bsa4 plasmid. The cloning vector pNIC28-Bsa4 adds an N-terminal hexahistidine tag with an integrated TEV protease cleavage site. The cleaved target protein represents the saposin A polypeptide with three additional N-terminal Ser-Met-Gly residues as a result of the cloning strategy, with the protein sequence MHHHHHHSSGVDLGTENLYFQSMGSLP CDICKDVVTAAAGMDLKDANEELIVYELKTCDWLPKPMN SASCKEIVD6LYVLD1IKGEM5RPGVCSALNCLSES. The protein was expressed using E. coli Rosetta gam-2 (DE3) (Novagen). Cells were grown at 37 °C in Terrific broth supplemented with tetracycline, chloramphenicol and kanamycin and induced with 1.0 mM IPTG. Three hours after induction we collected the cells by centrifugation at 12,000 g for 15 min; we discarded the supernatant. The cell pellet was resuspended in lysis buffer (20 mM HEPES, pH 7.5, 150 mM NaCl, 20 mM imidazole) and disrupted by sonication. Lysates were subjected to centrifugation at 26,000 g for 30 min, and the supernatant was heated to 85 °C for 10 min and then subjected to an additional centrifugation step at 26,000 g for 30 min. Preparative immobilized metal affinity chromatography (IMAC) purification consisted of batch-adsorption of the supernatant by end-over-end rotation with Ni Sepharose 6 Fast Flow medium for 60 min. After binding of saposin A to the IMAC resin, the chromatography medium was packed in a 10-mm inner-diameter open gravity flow column, and unbound proteins were removed by washing with 15 bed volumes of lysis buffer. The resin was washed with 15 bed volumes of wash buffer WB2 (20 mM HEPES, pH 7.5, 150 mM NaCl, 40 mM imidazole). Saposin was eluted by the addition of five bed volumes of elution buffer EB (20 mM HEPES, pH 7.5, 150 mM NaCl, 400 mM imidazole). The eluate was dialyzed overnight against gel filtration buffer GF (20 mM HEPES, pH 7.5, 150 mM NaCl) supplemented with recombinant TEV protease. We removed TEV protease containing an uncleavable His-tag from the eluate by passing the eluate over 2 ml of IMAC resin. Cleaved target protein was concentrated to a volume of 5 ml using centrifugal filter units and loaded onto a HiLoad Superdex 200 16/60 GL column using an ÄKTAexplorer 10 chromatography system. Peak fractions were pooled and concentrated to 1.2 mg/ml protein. Typically, a 1–1 bacterial culture yields 5–10 mg of purified saposin. The protein sample was flash-frozen in liquid nitrogen and stored at −80 °C.

Expression and purification of POT and T2. POT was expressed and purified according to a recently described protocol18,19. T2 was purified as described recently17.

Generation of Salipro lipid particles. We prepared detergent-solubilized lipids by dissolving the respective lyophilized lipid powder (brain lipid mix from bovine brain from Sigma-Aldrich; phosphatidylserine, phosphatidylcholine, phosphatidylglycerol, phosphatidylethanolamine and E. coli total lipid extract from Avanti Polar Lipids) in buffer D (50 mM HEPES, pH 7.5, 150 mM NaCl, 1% n-dodecyl-β-D-maltoside (DDM)) to obtain a stock solution that contained 25 mg/ml lipids and then incubated it for 1 h at 37 °C. If not otherwise stated, a brain lipid extract from bovine brain, Type I (Sigma-Aldrich) was used for nanoparticle generation. Prior to Salipro particle reconstitution, the lipid stock solutions were diluted to 5 mg/ml using buffer E (50 mM HEPES, pH 7.5, 150 mM NaCl, 0.03% DDM). For the reconstitution of Salipro particles, 10 µl of purified saposin A and 5 µl of the respective detergent-solubilized lipids were incubated for 10 min at 37 °C. Subsequently, 50 µl of GF buffer was added to the mixture and incubated for 10 min at room temperature. The reconstitution mixture was subjected to a gel filtration analysis on a Superdex S200 5/150 analytical gel filtration column (GE Healthcare) and eluted with GF buffer, giving rise to a monodisperse peak of Salipro particles. For a medium-scale preparation, 60 µl of a brain lipid solution (5 mg/ml brain lipids (Sigma-Aldrich), 50 mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) were incubated with 100 µl of purified saposin A (1.2 mg/ml, 20 mM HEPES, pH 7.5, 150 mM NaCl) and incubated for 5 min at 37 °C, after which 160 µl of GF buffer was added and the mixture was incubated for 10 min at room temperature. Then 230 µl of GF buffer were added and the sample was subjected to a gel filtration step on a Superdex200 10/300 GL column using an ÄKTAexplorer 10 chromatography system equilibrated with GF buffer. Fractions containing Salipro particles were pooled and concentrated to 1 mg/ml using Amicon Centricron filter devices at 3,000 g.

In one experiment, Salipro particles were generated after incubation with an increasing excess of lipids. Here varying amounts of lipid (Lipids 5, 5 µg; Lipids 12.5, 12.5 µg; Lipids 25, 25 µg; Lipids 50, 50 µg; Lipids 100, 100 µg) from a brain lipid solution (5 mg/ml brain lipids (Sigma-Aldrich), 50 mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) were incubated for 10 min at 37 °C. 10 µl of purified saposin A (1.2 mg/ml, 20 mM HEPES, pH 7.5, 150 mM NaCl) were added and incubated for 1 min at 37 °C, and GF buffer was added to a final volume of 41 µl. After 10 min of incubation at room temperature, 24 µl of GF buffer (pH 7.5) were added and the samples were subjected to a gel filtration step on a SuperdexTM 200 5/150 GL analytical gel filtration column using an ÄKTAmicro chromatography system (GE Healthcare) equipped with the A-905 autosampler, which automatically injected 25 µl of protein-containing sample. Analytical gel filtration runs were performed at 4 °C at a flow rate of 0.2 ml/min in GF buffer.
Incorporation of membrane proteins into Salipro particles. To reconstitute the archaean mechanosensitive channel T2 in Salipro nanoparticles, we incubated 5μl of a brain lipid solution (5mg/ml brain lipids (Sigma-Aldrich), 50mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) for 5 min at 37 °C, after which we added 2μl of purified membrane protein T2 (10mg/ml, 20mM Tris-HCl, pH 7.5, 150 mM NaCl, 5% glycerol, 0.03% DDM) and incubated the sample to a second incubation step at 37 °C. Subsequently, we added 20μl of purified saposin A (1.2mg/ml, 20mM HEPES, pH 7.5, 150 mM NaCl) and incubated the samples for 5 min at 37 °C. Then we added 24μl of 1× PBS (pH 7.4) and incubated the mixture for 5 min at room temperature, after which we added 39μl of 1× PBS (pH 7.4) to the mixture. The sample was subjected to a gel-filtration step on a Superdex S200 5/150 analytical gel filtration column, equilibrated and eluted with 1× PBS (pH 7.4).

For a medium-scale preparation of Salipro-T2, we incubated 40μl of a brain lipid solution (5mg/ml brain lipids (Sigma-Aldrich), 50mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) and 5μl of protease inhibitor stock solution (1×Complete EDTA-free tablet in 1ml H2O) for 5 min at 37 °C. Then we added 24μl of 1× PBS (pH 7.4) and incubated the mixture for 5 min at room temperature, after which we added 39μl of 1× PBS (pH 7.4) to the mixture. The sample was subjected to a second incubation step at 37 °C. Subsequently, we added 140μl of purified saposin A (1.2mg/ml, 20mM HEPES, pH 7.5, 150 mM NaCl) and incubated the mixture for 5 min at 37 °C. Then we added 145μl of 1× PBS (pH 7.4) and incubated the mixture for 5 min at room temperature, after which we added 150μl 1× PBS. The sample was subjected to a gel filtration step on a Superdex200 10/300 GL column using an ÄKTAexplorer 10 chromatography system equilibrated with 1× PBS (pH 7.4). Fractions containing Salipro-T2 were pooled and concentrated to 0.7mg/ml using Amicon Centricon filter devices at 3,000g. The protein sample was flash-frozen in liquid nitrogen and stored at −80 °C until further use.

To reconstitute the bacterial peptide transporter POT in Salipro nanoparticles, we incubated 5μl of a brain lipid solution (5mg/ml brain lipids (Sigma-Aldrich), 50mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) and 5μl of protease inhibitor stock solution (1×Complete EDTA-free tablet in 1ml H2O) for 5 min at 37 °C. Then we added 20μl of purified membrane protein POT (10mg/ml, 20mM Tris-HCl, pH 7.5, 150 mM NaCl, 5% glycerol, 0.03% DDM) and subjected the mixture to a second incubation step at 37 °C. Subsequently, we added 140μl of purified saposin A (1.2mg/ml, 20mM HEPES, pH 7.5, 150 mM NaCl) and incubated the mixture for 5 min at 37 °C. Then we added 145μl of 1× PBS (pH 7.4) and incubated the mixture for 5 min at room temperature, after which we added 150μl 1× PBS. The sample was subjected to a gel filtration step on a Superdex S200 5/150 analytical gel filtration column, equilibrated and eluted with 1× PBS (pH 7.4). For a medium-scale preparation of Salipro-POT, we incubated 40μl of a brain lipid solution (5mg/ml brain lipids (Sigma-Aldrich), 50mM HEPES, pH 7.5, 150 mM NaCl, 0.28% DDM) for 5 min at 37 °C, added 20μl of purified membrane protein POT (10mg/ml, 20mM Tris-HCl, pH 7.5, 150 mM NaCl, 5% glycerol, 0.03% DDM) and subjected the mixture to a second incubation step at 37 °C. Subsequently, we added 80μl of purified saposin A (1.2mg/ml, 20mM HEPES, pH 7.5, 150 mM NaCl) and incubated the mixture for 5 min at 37 °C. We transferred this mixture into individual tubes (23μl in each tube) and added 50μl of buffers with various pH levels to the tubes (pH 4.5: 20mM trisodium citrate, 100mM NaCl; pH 6: 20mM trisodium citrate, 100mM NaCl; pH 7.5: 20mM Tris-HCl, 100mM NaCl; pH 9: 20mM Tris-HCl, 100mM NaCl). After 5 min of incubation, the samples were subjected to a gel filtration step on a Superdex S200 5/150 analytical gel filtration column, equilibrated and eluted with buffers at the corresponding pH (4.5–9).

Negative-stain electron microscopy. For Salipro lipid-only discs, carbon-coated copper grids (400 mesh) were glow discharged in low-pressure air to render them hydrophilic for 20 s before application of 4μl sample for 30 s. Washing was done with three drops of water before staining with 4μl of uranyl formate (1%, Polysciences, USA) for 30 s. The grids were air dried after the removal of excess stain solution with torn filter paper. Negative-stain image data were collected with a JEMOL2100F field emission gun transmission electron microscope at an accelerating voltage of 200 keV and a final magnification of 69,500× on a CCD (charge-coupled device) camera (4K × 4K, Tietz Video and Image Processing System, GmbH Gauting, Germany) with 15-μm pixel size (corresponding to 2.16 Å on the specimen level).

Salipro-T2 samples were stained with uranyl acetate and imaged on an FEI Tecnai Polara TEM at 100 kV at a magnification of 59,000× and a pixel size of 1.91 Å on an UltraScan 4000 CCD camera (Gatan). Individual particles were windowed semi-automatically with the swarm option of e2boxer (EMAN2 package). After normalization, 33,000 particles were subjected to reference-free 2D classification and class averaging using the EMAN2 routines (e2refine2d).

Thermal unfolding analysis. Thermal unfolding analysis was performed using dye-free differential scanning fluorimetry using two samples each for POT in detergent (1× PBS, pH 7.4, 0.4% NM) and Salipro-POT (1× PBS, pH 7.4) with a sample volume of 10μl per capillary at a concentration of 0.2 mg/ml. The temperature was increased by 1 °C/min from 20 °C to 95 °C using a Prometheus NT.48 instrument (Nanotemper).

Cryo-EM. The POT data were collected and processed as described. We applied 2.5μl of purified sample at 0.7 mg/ml concentration to a glow discharged Quantifoil holey carbon grid.
(400 mesh, 1.2-µm hole size) and blotted it at 100% humidity using the FEI Vitrobot Mark I for 5.5 s. A total of 288 images of the vitrified sample were subsequently collected on an FEI TF30 Polara electron microscope operated at 300 kV, equipped with a Gatan K2 Summit direct electron detector camera. The images were recorded with a defocus in a range from 1.5 to 2.5 µm at a nominal magnification of 31,000×, corresponding to a calibrated physical pixel size of 1.2156 Å/pixel on the specimen at a dose rate on the camera set to 8.2 counts per physical pixel per second.

The sample was exposed for a total time of 6 s, with each image fractionated into 30 subframes, each with an accumulation time of 0.2 s per frame, with a total accumulated dose of 41e-/Å² on the specimen.

**Image processing.** Dose-fractionated super-resolution image frames were drift-corrected using whole-frame motion correction; a sum of all subframes in each image stack was used for further processing. The particles embedded in vitreous ice were monodisperse and exhibited good contrast (Fig. 3a), even at low defocus.

For all 2D image display and particle picking, we used SamViewer, an interactive image-analysis program written in Python. Defocus was determined using CTFFIND4 (ref. 52). Particle extraction and class averaging were performed in RELION53. Using 12,200 particles, we performed 2D classification (Fig. 3b) and obtained classes with a large number of inner-membrane details. Selected classes were extracted, and a probabilistic initial 3D-model generation procedure (PRIME) implemented in the SIMPLE54 package was used for ab initio 3D-reconstruction generation; this served as a reference model for the maximum likelihood-based 3D classification and auto-refinement procedure implemented in RELION. For the final 3D reconstruction (Fig. 4a,b), 9,913 particles were used and C4 symmetry was imposed; resolutions were estimated using the FSC 0.143 criterion on corrected FSC curves to obtain the map at 6.48 Å (Fig. 3e). ResMap was used to estimate local resolution from the unburned and unsharpened raw density map. Figures were prepared using UCSF Chimera.

**HIV virus-like particle production.** VLPs carrying HIV-1 JR-FL spikes with a deleted cytoplasmic tail (truncated at residue 708), disulfide-linked gp120 and gp41 (Cys501-Cys605), and E168K and N189A mutations (which facilitate binding of PG16 to the JR-FL spike) were produced in HEK 293T cells by calcium phosphate precipitation-mediated cotransfection. For production of [35S]Cys-labeled VLPs, we transfected one 150-cm² culture flask of semiconfluent 293T cells using 10 µg of pCAGGS JRFL gp160SOS,ΔCT,E168K,N189A and 10 µg of pNLA-3Luc.R-E DNA56,57. [35S]Cys was incorporated into the VLPs through metabolic labeling in Cys-free Dulbecco’s modified Eagle’s medium (National Veterinary Institute, Uppsala, Sweden) supplemented with 10 µM unlabeled Cys and 100 µCi of [35S]Cys/ml (Fisher Scientific, Gothenburg, Sweden) from 24 to 48 h after the transfection. Cells debris was removed from the virus-contain media using low-speed centrifugation at 1,000 r.p.m. for 10 min. The cleared supernatant was layered on top of a step gradient composed of 1 ml of 60% sucrose and 4.5 ml of 20% sucrose (wt/wt) in HN buffer (50 mM HEPES, 100 mM NaCl, pH 7.4) and centrifuged at 4 °C for 2 h at 93,000g (22,000 r.p.m.) in a Beckman SW28.1 rotor. Radiolabeled virus was collected from the 20/60% sucrose interface. For unlabeled VLP production, 15 culture flasks (150 cm²) with 293T cells were transfected, and the virus was collected and concentrated as described above. To increase VLP concentration, we diluted the virus to 15–20% sucrose in HN buffer and added it on top of a 1-ml 60% (wt/wt) sucrose cushion, after which we subjected it to a second ultracentrifugation step at 4 °C for 6 h at 22,000 r.p.m. using the SW28.1 rotor. The virus was collected by fractionation from the bottom, and fractions were analyzed using SDS-PAGE and SYPRO Ruby protein stain to visualize the VLP peak for collection.

**Incorporation of the trimeric HIV-1 spike into Salipro nanoparticles.** VLPs were lysed for 10 min on ice in HN buffer supplemented with 9 mM HEGA-10 in the presence of saposin A, and in some cases brain lipids. HEGA-10 was then removed by two successive spin SEC steps (Zeba spin desalting columns, 7K MWCO). The columns were washed and centrifuged according to the manufacturer’s instructions (Pierce). The column removed the HEGA-10 detergent but not saposin A, which with viral lipids replace the detergent in the spike-detergent complexes. Salipro–HIV spike particles eluted from the column void volume together with the other soluble virus proteins. The sample was then incubated for 15 min at 37 °C before lectin purification of the Salipro–HIV spike particles.

**Purification of the Salipro–HIV spike particles.** Salipro–HIV spikes were purified using Galanthus Nivalis lectin (Sigma-Aldrich) affinity chromatography. The lectin binds to high-mannose sugar moieties on the spike protein. The column was washed with 50 column volumes of HN buffer, and then the Salipro–HIV spike particles were eluted using 0.3 M methyl-α-D-mannopyranoside. The methyl-α-D-mannopyranoside was removed from the sample using a spin SEC column (Zeba spin desalting columns, 7K MWCO) and replaced with HN buffer. The Salipro–HIV spikes were concentrated by ultrafiltration using an Amicon Ultra-4 50K MWCO centrifugation filter (Millipore), and sample purity was assayed with SDS-PAGE using SYPRO Ruby protein staining (Lonza Group) and BN-PAGE using silver staining (Thermo Scientific).

**Functional assays of the Salipro–HIV spikes.** Salipro–HIV spike particles were incubated with 10 µg/ml of PG16, scCD4 or 17b Fab in HN buffer for 2 h at 37 °C. The formation of ligand spike complexes was followed by BN-PAGE and phosphorimaging. One volume of two times–concentrated BN-PAGE sample buffer was added, and the samples were incubated for 10 min at room temperature and analyzed by gradient BN-PAGE with 3–8% total acrylamide as described previously58. The dried gels were exposed to phosphorimaging screens (BAS MS2025; Fujifilm, Science Imaging, Nacka, Sweden), and the radiolabeled bands were visualized using a Molecular Imager FX and the Quantity One software (Bio-Rad, Hercules, CA).

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