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Dispatched uses Na+ flux to power lipid-modified Hedgehog release

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Abstract

The Dispatched protein, which is related to the NPC1 and PTCH1 cholesterol transporters^{1,2} and to H^+ -driven transporters of the RND family^{3,4}, enables tissue-patterning activity of the lipid-modified Hedgehog protein by releasing it from tightly -localized sites of embryonic expression^{5–10}. Here we determine a cryo-electron microscopy structure of the mouse protein Dispatched homologue 1 (DISP1), revealing three $Na⁺$ ions coordinated within a channel that traverses its transmembrane domain. We find that the rate of Hedgehog export is dependent on the Na⁺ gradient across the plasma membrane. The transmembrane channel and Na⁺ binding are disrupted in DISP1-NNN, a variant with asparagine substitutions for three intramembrane aspartate residues that each coordinate and neutralize the charge of one of the three $Na⁺$ ions.

Competing interests The authors declare no competing interests.

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Author contributions

Q.W. purified and characterized DISP1-A, DISP1-A-NNN and ShhN-DISP1-A complex and SCUBE2 protein, and developed and performed ShhNp binding and release assays. D.E.A. prepared cryo-EM grids, collected and processed cryo-EM data, and performed the 3DVA. R.M. devised and aided the SCUBE2 purification. J.H. performed ShhNp release assays. K.D., D.E.A., Y.Z., and Q.W. built the models. Y.M. provided mouse genetic epistasis analysis while in the laboratory of P.A.B. at Johns Hopkins University School of Medicine. All authors participated in discussion and analysis of the data. P.A.B., Q.W., D.E.A., K.D., Y.C., and Y.Z. prepared the manuscript.

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DISP1-NNN and variants that disrupt single $Na⁺$ sites retain binding to, but are impaired in export of the lipid-modified Hedgehog protein to the SCUBE2 acceptor. Interaction of the amino-terminal signalling domain of the Sonic hedgehog protein (ShhN) with DISP1 occurs via an extensive buried surface area and contacts with an extended furin-cleaved DISP1 arm. Variability analysis reveals that ShhN binding is restricted to one extreme of a continuous series of DISP1 conformations. The bound and unbound DISP1 conformations display distinct $Na⁺$ -site occupancies, which suggests a mechanism by which transmembrane $Na⁺$ flux may power extraction of the lipid-linked Hedgehog signal from the membrane. Na⁺-coordinating residues in DISP1 are conserved in PTCH1 and other metazoan RND family members, suggesting that Na+ flux powers their conformationally driven activities.

> During embryogenesis, the Hedgehog signal projects its tissue-patterning influence many cells beyond its sites of expression in 'organizers' such as the vertebrate notochord, the floor plate of the neural tube and the zone of polarizing activity in the developing $limb^{11-14}$. The mature Sonic hedgehog protein signal (ShhNp) is covalently lipid-modified¹⁵ by autoprocessing-mediated cholesterol attachment at its C terminus¹⁶ and acyl transferasemediated attachment of palmitate at its N terminus^{17,18}. The resulting dual-lipid-modified protein is tightly membrane associated and thus requires dedicated machinery for its packaging and release in long-range patterning. This machinery includes the $DISP1⁵⁻¹⁰$, which is required in Hedgehog-producing cells, and SCUBE, a secreted protein that can be supplied by other cells that do not produce or respond to the Hedgehog signal^{19–21}. Although DISP1 and the related Hedgehog receptor PTCH1 are similar in sequence and transmembrane topology and both interact with the lipid-modified Hedgehog signal, DISP1-mediated Hedgehog release promotes Hedgehog pathway activity, whereas PTCH1 attenuates this activity by inhibiting the essential response component Smoothened (SMO) (Extended Data Fig. 1a, b, Supplementary Discussion).

DISP1 structure determination

We determined the structure of a stable, functional N-terminal truncation of DISP1 (DISP1- A) (Extended Data Figs. 1c–e, 2). Monomeric DISP1-A purified into a mixture of lauryl maltose neopentyl glycol (LMNG), glyco-diosgenin (GDN) and cholesteryl hemisuccinate (CHS) was efficiently cleaved at an essential furin site22 (Extended Data Fig. 1f), and 3D classification of a single-particle cryo-EM dataset revealed high-resolution structures of two major classes—relaxed (R) (2.5 Å resolution); and tense (T) (2.5 Å resolution)—each comprising at least two subclasses (Extended Data Figs. 3, 4, Extended Data Table 1). We further applied 3D variability analysis23 (3DVA; Methods), to systematically model the conformational landscape within our dataset as a series of orthonormal principal components representing specific modes of motion. We found conformations resembling R and T at opposite ends of the second principal component (PC2) of this dataset (Extended Data Fig. 5, Supplementary Video 1), and we present a model of DISP1-A built from the density of R.

DISP1-A structure resembles that of PTCH1, with twelve transmembrane helices (TM1– TM12) derived from apparent duplication of a six-transmembrane-helix unit (TM1–TM6 and TM7–TM12) arranged in pseudodyad symmetry and preceded by amphipathic

transverse helices at the cytoplasmic membrane surface (pre-TM1 and pre-TM7). Each unit contains a large extracellular loop derived from corresponding positions between TM1 and TM2 and between TM7 and TM8 (Fig. 1a, b).

Unlike the extracellular domains (ECDs) of PTCH1, which are juxtaposed to form a conduit for sterol transport1, the mouse DISP1 ECDs are splayed apart in a Y-shaped manner (Fig. 1a), as also recently reported for Drosophila Disp24 and human DISP125. DISP1 also resembles PTCH1 in the presence of ferredoxin-like $\alpha + \beta$ open-faced sandwich folds in the membrane-proximal parts of ECD1 and ECD2 (Extended Data Fig. 1g).

Each ECD accepts secondary structure elements from the other loop, enclosing a bowlshaped space between the two domains. The linker within the TM1–TM2 loop is the site of furin cleavage²² (Arg276-Glu-Arg-Arg279), opening the bowl at a location we designate as 'front'. The ECDs contain seven disulfide bonds, five in ECD1 and two in ECD2 (Fig. 1b, Extended Data Figs. 2, 6), and five N-linked glycosylation sites (Fig. 1b, Extended Data Figs. 1g, 2). Except for pre-TM1 and pre-TM7, most intracellular sequence in DISP1-A is not resolved, including four N-terminal residues (1, 2, 172 and 173), 14 residues preceding pre-TM7 (667–680) and 378 residues at the C terminus (1144–1521). In the ECDs, 19 residues including the furin cleavage site (264–282) and residues 399–409 are not resolved.

We modelled 28 lipid or detergent molecules, mostly at annular positions around the transmembrane domain. Twenty-six of these molecules were CHS, 17 corresponding to the inner leaflet and 9 to the outer leaflet of the membrane, presenting the appearance of a sterol bilayer (Fig. 1c, Supplementary Video 2). Two of the nine outer leaflet sterols were raised, and the front-side lifted sterol may correspond to an intermediate state in ShhNp export (see below)—a nearby lipidic group, modelled as LMNG, levers the lifted sterol upward (Fig. 1c, Supplementary Video 2).We also modelled a non-proteogenic density within a hydrophobic cavity at the distal tip of ECD2 as an LMNG molecule (Fig. 1c, Supplementary Video 2).

Na⁺ ions in a transmembrane tunnel

The transmembrane domain contained three non-proteogenic densities (sites I, II and III), each associated with one of the three acidic residues in TM4 and TM10 that are conserved in PTCH1 (Fig. 2a, Extended Data Fig. 7a–c). These densities appear to be metal cations, well-coordinated by oxygens from side-chain hydroxyls and main-chain carbonyls, and at least one oxygen from water (site III) (Fig. 2a). Site I and II cations lie on the axis of pseudodyad symmetry between TM1–TM6 and TM7–TM12, each receiving coordination bonds from symmetry-related oxygens in both opposing six-transmembrane helix units. The site III cation is asymmetrically positioned, its ligands deriving entirely from TM4 and TM5.

The coordination number is 5 for sites I and II and 6 for site III, the centre-to-centre distances between ions and their associated oxygens averaging $2.50 \text{ Å}, 2.51 \text{ Å},$ and 2.47 Å, respectively. Each density in these three sites is associated with a single carboxylate and with oxygens from hydroxyls and main chain carbonyls, ruling out alkaline earth metal ions such as Ca^{2+} or Mg^{2+} , which require additional charge neutralization, display a more rigid geometry and a tighter coordination sphere, and are rarely coordinated by main-chain

carbonyls. The average bond distances match those characteristic²⁶ of Na⁺ (2.4–2.5 Å) rather than K^+ (2.7–3.2 Å). We therefore conclude that the densities at sites I, II and III derive from Na⁺, as further validated by the CheckMyMetal web server²⁶.

To assess the role of Na+ in DISP1 activity, we modified a quantitative assay for release of ShhNp⁶ by: (1) replacing luciferase between amino acid residues 131 and 132 of Shh with nanoluciferase (Extended Data Fig. 8a); (2) use of DISP1-A–mEGFP (Extended Data Fig. 1c); and (3) use of purified SCUBE2 protein in defined medium as the acceptor (Extended Data Fig. 8b, c). This assay provides a rapid, quantitative measure of the rate of nanoluciferase (nanoLuc)–ShhNp release into the medium over a period of minutes (Extended Data Fig. 8d). Under reduced Na^+ , with the organic cation N-methyl-d-glucamine $(NMDG⁺)$ used to maintain osmolarity²⁷, we found that release by DISP1-A activity ranged from near zero at 10 mM extracellular $Na⁺$ to partial and full restoration of release at extracellular $Na⁺$ concentrations of 40 and 155 mM (Fig. 2b, Extended Data Fig. 8d); a similar observation was reported during revision of our manuscript²⁸.

Mutation of DISP1-A D571, D572 and D1049—the charge-neutralizing residues in TM4 and TM10—to asparagine, resulting in DISP1-A-NNN, caused disruption of ShhNp release (Fig. 2c) in vitro, as previously established in vivo⁶. Disruption of sites I, II and III individually also disrupted release of ShhNp (Fig. 2c, Extended Data Fig. 8e). Given that all of these variant proteins are stable and retain the ability to bind ShhNp (Extended Data Fig. 8f), the results indicate that $Na⁺$ coordination is critical for ShhNp release.

Utilization of a Na⁺ gradient to power DISP1 activity implies Na⁺ flux across the membrane. We found a path extending from the extracellular surface of the DISP1 transmembrane domain to the intracellular surfaces²⁹; most water molecules within the transmembrane domain were clustered near this channel (Fig. 3a, Supplementary Video 3). We determined the structure of DISP1-A-NNN at 3.2 Å resolution (Extended Data Fig. 3, Extended Data Table 1), revealing that the $Na⁺$ ion densities at sites I, II and III are indeed absent (Fig. 3b), and the side chains of the asparagine residues substituted for aspartates and some of the remaining liganding side chains are rearranged (Fig. 3b) compared with the wild-type protein. We compared water-excluded internal cavities within the transmembrane domains of these proteins (Extended Data Fig. 7d, Supplementary Discussion), and found that a large cavity within DISP1-A containing the three $Na⁺$ ions is substantially disrupted in DISP1-A-NNN by a series of side-chain rotamer swaps, including interactions across the pseudodyad axis between the altered residues D571N and D1049N, with concomitant straightening and movement of TM4 and TM10 towards each other (Fig. 3c, Extended Data Fig. 7d, bottom, Supplementary Discussion). These movements, associated with Na+ loss in DISP1-A-NNN, bring into closer proximity the pseudodyad halves of the transmembrane domain (TM1–TM6 and TM7–TM12), suggesting possible modes of allosteric communication from the nexus of $Na⁺$ coordination outward towards ECD1, ECD2 and other Hedgehog-interacting regions of DISP1.

Structure of a ShhN–DISP1-A complex

We determined the structure of DISP1-A in complex with ShhN, a truncated Shh containing residues 26–189 and lacking lipid modifications (2.7 Å overall resolution) (Extended Data Figs. 4, 6, 9, Extended Data Table 1). The largest of the two major density classes contained substantial additional density corresponding to ShhN perched near the top front side of DISP1-A, between ECD1 and ECD2 (Fig. 4a). In the model of this ShhN–DISP1-A complex (Fig. 4b), the ShhN portion of the map resolved residues 39–189, as well as the previously described Zn2+ ion³⁰ and the di-atomic Ca²⁺ cluster³¹, along with all major secondary structure elements and most side chains (Fig. 4a, b, Extended Data Figs. 6, 9).

In the complex, ECD1 and ECD2 of DISP1-A bind ShhN in a pincer-like grasp (Fig. 4a, b), including hydrophobic contacts and possible ionic interactions with total buried surface area of 989.3 Å2 (−8.6 kcal mole−1; Methods). The position of ShhN relative to DISP1-A in our complex structure is distinct from the position recently proposed by Cannac et al. 24 for Drosophila Hedgehog (Extended Data Fig. 9c). Notably, the furin-cleaved front-side linker of DISP1-A, which is considerably sharper than in the DISP1-A apo structure, extends from ECD2 upwards and around ShhN in a one-armed molecular embrace (Fig. 4a, b). We extended the model of this linker by an additional 13 residues preceding the furin cleavage site (Extended Data Fig. 2). Improved ECD resolution in the ShhN–DISP1-A map also enabled us to extend the DISP1-A model by an additional 11 residues (399–409) in distal ECD1 (Fig. 4b, Extended Data Fig. 2), revealing a hydrophobic cavity that could interact with an extended hydrophobic moiety (Extended Data Fig. 6f), and reminiscent of the hydrophobic cavity at distal ECD2 (Fig. 1c, Extended Data Fig. 6g).

Conformational dynamics of DISP1

We applied 3DVA23 to this dataset and found the first principal component (PC1) to be anchored at opposite ends by DISP1-A molecules either complexed with or lacking the ShhN ligand (Extended Data Fig. 10a, Supplementary Video 4). A similar range of DISP1-A conformations is found within PC1 of 3DVA of the DISP1-A apoprotein dataset (Extended Data Fig. 10b), suggesting that ShhN binds preferentially to a restricted subset of common DISP1-A conformations.

Close examination revealed that Na+ sites are differentially occupied at opposite extremes of the PC1 reaction coordinates for datasets with or without ShhN (Extended Data Fig. 10a, Supplementary Video 4), with densities corresponding to all three $Na⁺$ ions present at the end anchored by unbound DISP1-A, and only $Na⁺$ site I occupied at the opposite end, anchored by the ShhN-bound complex. $Na⁺$ occupancy thus drives, or is permissive of, alternative conformational states of DISP1, suggesting that chemiosmotically driven flux through the DISP1 transmembrane tunnel could induce conformational changes that cause Hedgehog release after initial binding.

Membrane extraction of Hedgehog lipid adducts

The release of mature Hedgehog signal requires extraction of its covalently linked cholesteryl and palmitoyl adducts from the membrane; thus, the lifted sterol on the front

side of DISP1-A (Fig. 1c) may correspond to an intermediate state of the cholesteryl adduct as it is being removed from its hydrophobic environment. The hydrophobic cavities at the distal tips of ECD1 and ECD2 (Extended Data Fig. 6f, g) are large enough to accommodate the covalently linked Hedgehog lipid adducts, suggesting that they may act as waystations for the ShhNp lipid adducts after membrane extraction and before complex formation with SCUBE2.

Discussion

We propose that DISP1-A extraction and release of the lipid-modified Hedgehog signal proceeds via a state activated for recruitment of SCUBE2 (Fig. 4c). We suggest that activation by DISP1-A binding to Shh at a location approximately 25 Å above the membrane aids in extraction of the ShhNp-linked lipids, and the lifted sterol at the front side of DISP1-A may represent an intermediate in cholesteryl adduct extraction (Fig. 1c). Three prominent features characterize this activated state: (1) a pincer-like action of the split ECDs; (2) a molecular embrace of Shh by the furin-cleaved linker arm; and (3) the possible accommodation of lipid adducts in distal ECD1 and ECD2. This activated state is correlated with loss of $Na⁺$ from sites II and III (3DVA, ShhN-bound state) (Extended Data Fig. 10, Supplementary Video 4), followed by Na^+ flux down its chemiosmotic gradient to re-occupy sites II and III and cause release of ShhNp to SCUBE2—a process incorporating opening of the ECD pincers, expulsion of the ECD1 and ECD2 lipid adducts and withdrawal of the linker embrace, followed by resetting to permit another cycle.

We note from structure-based alignment that each of 15 protein-derived $Na⁺$ -liganding oxygens in DISP1 (10 from side-chains, 5 from main-chain carbonyls) is conserved in PTCH1, as are the three charge-neutralizing intramembrane acidic residues (Extended Data Fig. 11). This degree of conservation exceeds that of the overall protein sequence (19% identity overall), and strongly supports the functional role of $Na⁺ flux$ in PTCH1 suppression of SMO activity, consistent with a previous demonstration of PTCH1 dependence on a transmembrane $Na⁺$ gradient for its function²⁷. All of these residues are also conserved by sequence alignment in PTCH2, which also functions to suppress $SMO³²$. By contrast, DISP2, which does not rescue disp function in Drosophila⁶ and probably lacks Hedgehogreleasing activity, has only one of three charge-neutralizing intramembrane acidic residues and six of ten side-chain oxygens. Prokaryotic RND transporters harbour distinct residues at these Na+-coordinating sites (Extended Data Fig. 11), but most retain interactions between TM4 and TM10 that are conditional on cation binding, pointing to a common mechanism for harnessing chemiosmotic gradients (Supplementary Discussion).

How do DISP1 and PTCH1 produce such distinct functional effects on Hedgehog-pathway activation? Our analysis suggests the use of chemiosmotic force from the metazoan transmembrane Na+ gradient to power the release of lipids from membranes. For PTCH1, cholesterol removal via an enclosed sterol conduit in the PTCH1 ECD regulates SMO by depriving it of sterol that must be bound within the interior of its seven-transmembrane helix bundle for activity^{33,34}. DISP1 also appears to extract lipids from the membrane, but these lipids are covalently attached to the Hedgehog protein signal, necessitating a split of the DISP1 ECD to accommodate the protein moiety. Evolutionary acquisition of the furin

cleavage site that permits this split, and of the linker arm that contacts ShhNp, apparently occurred by insertion into the Disp1 gene of a single small exon that encodes both features (Extended Data Fig. 2). A fundamental mechanistic similarity is indicated by a lifted sterol in both DISP1 and PTCH1, at corresponding positions within the 12-transmembrane helix bundle35. The unusual positions of these sterols at corresponding locations in both proteins suggest partial extraction from the membrane, perhaps representing a common intermediate in the lipid extraction activities of both proteins.

Methods

No statistical methods were used to predetermine sample size. All proteins, including DISP1 variants, ShhN, ShhNp and SCUBE2, were of mouse origin (Mus musculus). Cell lines were not tested for mycoplasma contamination. All cell lines grew normally.

DISP1 expression and purification

All constructs were cloned using Gibson Assembly. In DISP1-A, DISP1-A-NNN and the ion-site mutants, amino acids 3 to 171 were deleted from the full-length $(1-1521)$ DISP1. For protein expression, DISP1-A–ShhNp binding assays and ShhNp release assays, different DISP1 variants were cloned into the pcDNA3.1 vector¹. For detection of protein expression levels in Extended Data Fig. 1d, HEK 293T cells (ATCC, CRL-3216) were seeded into a six-well plate. Plasmids encoding DISP1 or DISP1-A with the SBP and HA tags at the C terminus were transiently transfected into the cells using Lipofectamine 3000 (ThermoFisher). Two days after transfection, the cells were lysed and incubated for 1 h at 4 °C in buffer containing 20 mM HEPES pH 7.5 (Teknova), 300 mM NaCl (Teknova), 2 mg ml−1 iodoacetamide (Sigma), 1% n-dodecyl-β-d-maltoside (DDM) (Anatrace), 0.2% CHS (Anatrace) supplemented with protease inhibitors (ThermoFisher). Protein extracts were cleared by a subsequent centrifugation (100,000g, 30 min, 4 °C). Western blotting was performed using Criterion TGX stain-Free precast gels (Bio-Rad) and Immobilon-P PVDF membrane (Millipore). PVDF membranes were immunoblotted overnight at 4 °C with anti-HA rabbit polyclonal (1:1,000, Cell Signaling) or anti-β-actin rabbit polyclonal (1:1,000, Cell Signaling) antibodies followed by a 1-h incubation with the corresponding horseradish peroxidase-conjugated secondary antibodies (1:2,000, Promega) at room temperature. The blots were developed using SuperSignal West Pico PLUS substrate (ThermoFisher).

For large-scale protein purification, C-terminally streptavidin-binding peptide (SBP)-tagged DISP1-A was cloned into a BacMam expression vector, PVLAD6. Baculoviruses were produced in Sf9 cells (ATCC, CRL-1711) as previously described^{1,27}. HEK 293F cells (ThermoFisher, R79007) were cultured in suspension using Free-Style medium (ThermoFisher) until the density reached $1.5-2 \times 10^6$ cells per ml, and cells were transduced with DISP1-A-SBP baculoviruses in the presence of 10 mM sodium butyrate (Sigma). Forty-eight hours after transfection, cells were collected and stored at −80 °C. For DISP1-A-NNN, plasmids encoding DISP1-A-NNN-SBP were transfected into the HEK 293F cells using linear polyethylenimines (Polysciences). Both DISP1-A and DISP1-A-NNN were purified essentially as previously described1. Cells were disrupted using a hypotonic buffer (20 mM HEPES pH 7.5, 10 mM MgCl₂ (Teknova), 10 mM KCl (Teknova), 0.25 M

sucrose (Sigma)) in the presence of benzonase (Sigma) and protease inhibitors. Following centrifugation (100,000g, 30 min, 4° C), the pellets were solubilized using a glass Dounce tissue homogenizer in lysis buffer (20 mM HEPES pH 7.5, 300 mM NaCl, 2 mg ml−1 iodoacetamide, 1% DDM, 0.2% CHS) supplemented with protease inhibitors. The sample was incubated for 2 h at 4° C with gentle rotation, then centrifuged (100,000g, 30 min, 4 °C) for collection of the supernatant. The C-terminally SBP-tagged DISP1-A or DISP1- A-NNN was captured using streptavidin–agarose affinity resin (ThermoFisher). After a 3-h incubation period at $4 \degree C$, the resin was packed into a disposable column by gravity flow. The resin was first washed with 30 column volumes (CVs) of wash buffer (buffer W, 20 mM HEPES pH 7.5 and 300 mM NaCl) containing 0.03% DDM, 0.006% CHS; then with 30 CVs of buffer W containing 0.12% LMNG (Anatrace), 0.024% CHS and 0.025% GDN (Anatrace) to exchange detergents; and finally with 30 CVs of buffer W containing 0.012% LMNG, 0.0024% CHS and 0.0025% GDN. Protein was eluted in buffer W containing 0.012% LMNG, 0.0024% CHS, 0.0025% GDN and 2.5 mM biotin (Sigma). The eluted protein was concentrated with an Amicon Ultra filter (100 kDa cut-off, Millipore) and loaded onto a Superose 6 Increase 10/300 GL column (GE Healthcare) in buffer W containing 0.012% LMNG, 0.0024% CHS and 0.0025% GDN plus 5 mM CaCl₂ (Sigma).

ShhN expression and purification

A DNA fragment encoding mouse ShhN residues 26–189 was cloned into a pHTSHP vector with N-terminal His8 tags³¹. Protein was expressed in Escherichia coli strain BL21(DE3) and purified as previously described¹. The cell lysate was first centrifuged at $8,000$ g for 15 min (4 \degree C), then the filtered supernatant was applied to the Ni-NTA resin. After elution from the Ni-NTA column, the N-terminal tags were cleaved with HRV3C protease. The protein was dialysed against 20 mM HEPES, 100 mM NaCl and 7 mM β-mercaptoethanol, and further purified by SP sepharose cation-exchange chromatography.

SCUBE2 expression and purification

Mouse SCUBE2 lacking amino acids 30–281 (to mimic the effect of furin cleavage at an internal site) was 3×Flag-tagged at the C terminus and expressed in Flp-In T-REx 293 cells (ThermoFisher, R78007), as previously described²⁰. Cultures were induced with doxycycline and transitioned to serum-free medium (DMEM (ThermoFisher) supplemented with 1x insulin–transferrin–selenium (ThermoFisher), 10 mM HEPES, 1× GlutaMAX and 5 μg ml−1 doxycycline). After 3–4 days, SCUBE2-conditioned medium was collected and clarified by centrifugation and membrane filtration. After concentrating by ultrafiltration (Vivaflow 200/30 kDa, Sartorius), SCUBE2 was purified by heparin and anti-Flag M2 affinity chromatography. Eluted protein was concentrated to 9.13 mg ml−1 by ultrafiltration (Amicon Ultra 30 kDa, Millipore). The eluting 3×Flag peptide was washed out during the final concentration step.

Cryo-EM sample preparation and data acquisition

For DISP1-A, 2.5 µl of purified and concentrated sample at a concentration of 1.56 mg ml⁻¹ was applied to glow-discharged gold grids covered with holey carbon film (Quantifoil, 300 mesh 1.2/1.3) and plunge frozen using a Vitrobot Mark IV, with blotting time of 4 s and blotting force 0, at 8 °C and 100% humidity. DISP1-A-NNN cryo-EM grids were prepared similarly, but using gold-grids covered by a holey gold film (Quantifoil UltrAufoil, 300 mesh 1.2/1.3) and a sample concentration of 0.3 mg ml⁻¹. For ShhN–DISP1-A complex, purified DISP1-A was incubated with ShhN at a 2.7:1 molar ratio (27 μM ShhN, 10 μM DISP1-A) for 3 h (4 °C) before cryo-sample preparation. DISP1-A–ShhN grids were prepared in the same way as that for DISP1-A.

Cryo-EM data for DISP1-A and ShhN–DISP1-A were collected with a Titan Krios microscope operated at 300 keV and equipped with a Bio Quantum post-column energy filter with zero-loss energy selection slit set to 20 eV. Using SerialEM37, images were recorded automatically at a nominal magnification of $105,000\times$ (true magnification 59,880×), with defocus range set to −0.5 to −1.5 μm. Images were recorded with a post-GIF Gatan K3 camera in super-resolution single-electron counting mode with a calibrated pixel size of 0.4175 Å (physical pixel size of 0.835 Å). The DISP1-A-NNN samples were imaged on a different Titan Krios with identical parameters, but a slightly different calibrated pixel size of 0.417 Å. Data collection statistics are shown in Extended Data Table 1.

Image processing and map calculation

As part of an on-the-fly image-processing pipeline, which processes micrographs immediately after acquisition, we used MotionCor 2^{38} to correct motion of each dose fractionated movie stack, sum motion-corrected frames with and without dose weighting, and bin images in Fourier space to the physical pixel size $(\sim 0.835 \text{ Å})$ for further image processing. Motion-corrected, dose-weighted sums were used for contrast transfer function determination and resolution estimation in cryoSPARC³⁹. Particles were picked in cryoSPARC first using a Gaussian template, and then re-picked with low-resolution templates generated from an initial structure. Subsequent image processing was carried out in RELION 40 and cryoSPARC. The flow chart of image processing is illustrated in Extended Data Fig. 3 (DISP1-A and DISP1-A-NNN) and Extended Data Fig. 9 (ShhN–DISP1-A). Conversion of data from cryoSPARC to RELION was performed using UCSF pyem 41 . Directional Fourier shell correlation curves were determined as previously described 42 .

3DVA23 of DISP1-A alone was conducted based on the early non-uniform refinement shown in Extended Data Fig. 3 (2.4 Å nominal resolution, 974,614 particles), using a five-component decomposition solved over 40 iterations. During 3DVA, the data were filtered by an 8th-order Butterworth high-pass filter with a cut-off frequency of 40 Å and a 1.5-order Butterworth low-pass filter with a cut-off frequency of 3.5 Å, as well as a soft mask containing the entire protein but excluding the detergent micelle. Trajectories representing each variability component were computed using 'simple' mode (that is, each step is calculated by weighted addition of variability component maps to the mean structure) with 20 frames and no filters. 3DVA of ShhN–DISP1-A was performed similarly, based

on non-uniform refinement of all apo and complex particles combined (2.7 Å nominal resolution, 356,610 particles), but with no high-pass filter owing to the need to capture the appearance and disappearance of ShhN.

Model building and refinement

The de novo atomic models of DISP1-A were built in $Coot^{43}$, refined in PHENIX⁴⁴ and finalized in ISOLDE⁴⁵. Starting from a predicted model from $PHYRE2^{46}$ and a combined density of R and T conformers, we manually fitted transmembrane helices into the corresponding density and then traced the backbone of ECDs and linkers in between those transmembrane helices in Coot. The preliminary model was then mutated based on Mouse DISP1 sequence and refined using real space refinement in PHENIX. Based on residual unmodelled density near the protein, ligands were added and refined in real space using Coot and PHENIX, respectively. Starting from this 'common model' built referring to the combined density of R and T, we obtained models for each conformer (R, T and NNN) by refining this 'common model' against the final density of each conformer individually, removing atoms unresolved in each density and then finalizing the models with ISOLDE (largely to improve steric clashes). The ShhN–DISP1-A complex was treated in the same manner, but using several crystal structures of mammalian Shh as initial models (Protein Data Bank (PDB) IDs: 1VHH, 4C4N and 6PJV). All models were validated by wwPDB validation server47 and no major issues were reported in our three models.

Visualizations of the atomic models (figures and videos) were made using UCSF Chimera48, ChimeraX49 and PyMOL (The PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC.). The ion conduction path was mapped using Caver 3^{29} and ion coordination geometries were analysed with the CheckMyMetal web server²⁶. DISP1– AShhN interaction interface area and the solvation free-energy gain upon the formation of the interfaces were calculated using $PISA^{50}$. Clustal Omega⁵¹ and the ESPript server⁵² was used in preparation of the protein sequence alignment (Extended Data Fig. 2).

Rapid, quantitative ShhNp release rate assay

Disp−/− mouse embryonic fibroblasts (MEFs)6,20, were cultured on 24-well plates in DMEM supplemented with 10 % FBS and 1% penicillin-streptomycin-glutamine (ThermoFisher). DISP1–mEGFP variants were transfected into the cells along with a NanoLuc–Shh fusion in which the nanoluciferase coding sequence was inserted between E131 and D132 in Shh. The mass ratio of plasmids expressing DISP1 variants and NanoLuc–ShhNp was 1 to 2. An EGFP expression construct was used as a negative control for conditions lacking DISP1 variants. Two days after transfection, following a quick wash with DMEM, cells were incubated with cycloheximide (100 μg ml⁻¹, Sigma) in DMEM for 50 min. Cells were then incubated in the desired medium either with or without SCUBE2 protein (1 μM (Fig. 2b) or 150 nM (Fig. 2c)) at 37 °C with gentle shaking (125 rpm). Small aliquots of supernatant were removed sequentially at multiple time points, and collected supernatants were cleared by a centrifugation at 3,000g for 30 min at 4 °C. NanoLuc-ShhNp luminescence level was measured using Nano-Glo Luciferase Assay System (Promega), following the manufacturer's protocol, and the measured luminescence was corrected to

account for the decreased supernatant volume through time. At the end of the experiment, cells were washed with PBS and lysed with the passive lysis buffer (Promega). Released NanoLuc–ShhNp luminescence at each time point was normalized to the GFP level, the total protein in cell lysates (measured using Pierce BCA protein assay kit, ThermoFisher), and to the total luminescence of ShhNp (luminescence level in cell lysates plus total NanoLuc–ShhNp released into supernatant by the final time point). The linear portion of the release curves was used to calculate the ShhNp release rate. ShhNp release endpoint assay Disp^{-/−} MEFs^{6,20}, were cultured on six-well plates in DMEM supplemented with 10 % FBS and 1% penicillin-streptomycin-glutamine. All cell transfections were performed using Lipofectamine 3000. Plasmids encoding DISP1 variants were transfected into the cells along with full-length Shh constructs. The ratio of plasmids expressing DISP1 variants to Shh was 1 to 2. For conditions that did not have DISP1 variants or Shh, a GFP construct was used as a negative control to replace the missing partner. Two days after transfection, following a quick wash with DMEM, the cells were shifted to $SCUBE2$ ($30-281$) (mouse SCUBE2, with an N-terminal truncation)-conditioned medium or control medium (600 μl per well). After 16 h of incubation, the medium was collected and centrifuged at 18,000g for 30 min (4 °C). The supernatant was probed by immunoblotting for ShhNp (anti-Shh rabbit monoclonal, 1:1,000, Cell Signaling), 3×Flag-tagged SCUBE2 (anti-Flag m2 mouse monoclonal, 1:1,000; Sigma). Cells were lysed using a buffer containing 20 mM HEPES pH 7.5, 300 mM NaCl, 2 mg ml⁻¹ iodoacetamide, 1% DDM, 0.2% CHS supplemented with protease inhibitors (400 μl per well, 1 h at 4 °C). After a centrifugation of 100,000g for 30 min at 4 °C, cell lysate was probed by immunoblotting for ShhNp (anti-Shh rabbit monoclonal, 1:1,000), DISP1–HA or DISP1–mEGFP (anti-HA rabbit polyclonal, 1:1,000 or anti-GFP rabbit polyclonal, 1:2,000, Invitrogen), and β-actin (rabbit polyclonal, 1:1,000). SCUBE2-conditioned medium was prepared in the same way as described above in SCUBE2 expression and purification. Control medium was prepared as for SCUBE2 conditioned medium but without doxycycline induction.

DISP1-ShhNp co-immunoprecipitation

The DISP1-ShhNp binding assay (Extended Data Fig. 8f) was conducted using the HEK 293 Flp-In T-REX ShhNp-producing cell line²⁰. ShhNp-producing cells were maintained in DMEM with 10% FBS and 1% penicillin-streptomycin-glutamine. Plasmids of C-terminally SBP and HA-tagged DISP1-A variants were transiently transfected into the cells using Lipofectamine 3000. One day after transfection, full-length Shh expression was induced by 5 μg ml−1 Doxycycline (Sigma) in DMEM F-12 (ThermoFisher) supplemented with 1% insulin-transferrin–selenium and 1% penicillin-streptomycin-glutamine. Another day after ShhNp expression was induced, cells were lysed and incubated for 1 h at 4 °C in the same buffer as that in ShhNp release endpoint assay. Following a subsequent centrifugation (100,000g, 30 min, 4 °C), Streptavidin-agarose affinity resin was used to pull down theSBPtagged DISP1-A variants, together with any bound ShhNp, from the supernatant. After a 2-h incubation period at 4 °C, the resin was washed with buffer W containing 0.03% DDM and 0.006% CHS. Protein was eluted in the same buffer supplemented with 2.5 mM biotin. Immunoprecipitates were analysed by Western blot as described above using the following

antibodies: anti-HA rabbit polyclonal (1:1,000), anti-Shh rabbit monoclonal (1:1,000) and anti-β-actin rabbit polyclonal (1:1,000).

Reporting summary

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

Extended Data

Extended Data Fig. 1 |. Structurally related transporters Dispatched and Patched in Hedgehog signalling.

a, b, Opposing functions of Dispatched and Patched in Hedgehog signalling. **a**, The HH protein signal, covalently modified by cholesterol and palmitate, requires the action

of DISP1 and SCUBE for release from the membrane of producing cells. HH then uses its palmitoyl adduct to clog the sterol transport conduit and block the function of its receptor PTCH1 in responding cells. The loss of PTCH1 sterol transport activity permits accumulation of cholesterol within the inner leaflet to levels that activate SMO by binding within its seven transmembrane helix bundle, resulting in activation of the GLI transcriptional effector of Hedgehog signalling. **b**, As Ptch1 is a target for GLI activation, the X-Gal staining of a $PtchI^{LacZ}$ knock-in allele⁵³ provides an indication of Hedgehog pathway activity (leftmost embryo). Homozygous mutation of $Disp1^{6-10}$ causes a loss of nearly all embryonic Hedgehog pathway activity $(2nd$ embryo from left)⁵⁴, whereas homozygous disruption of Ptch1 leads to unregulated ectopic pathway activity, regardless of the functional status of *Disp1* (rightmost two embryos)⁵³. **c–f**, Truncated DISP1 protein. As murine full-length DISP1 protein was poorly expressed in HEK293 cells, we tested a variety of constructs, and ultimately settled on an N-terminal truncation. Although its export activity was partially reduced, truncated DISP1-A protein nevertheless mediated efficient release of ShhNp (autoprocessed, lipid-modified Shh protein) into cell culture medium containing SCUBE2 (mouse SCUBE2, lacking amino acids 30–281) upon transfection into Disp−/− mouse embryonic fibroblasts6,20 (MEFs), indicating preservation of its function. **c**, DISP1 mEGFP and DISP1-A-mEGFP in HEK293T cells with magnified insets showing expression on the cell membrane. **d**, Western blot showing high level of DISP1-A expression relative to DISP1 (both proteins were SBP and HA-tagged at the C-terminus). **e**, Functional assay (Methods, ShhNp release endpoint assay) in Disp−/− MEFs. Culture media and cell lysates from transiently transfected $\text{Disp}^{-/-}$ MEFs were probed by immunoblotting for expression of DISP-mEGFP (SBP, HA and mEGFP tagged at the C-terminus, see panel **c**), ShhNp, and SCUBE2. DISP1 and DISP1-A both released ShhNp in the presence of SCUBE2. β-ACTIN, loading control. **f**, Size exclusion chromatography of purified DISP1-A, together with the SDS-PAGE of the indicated fraction, corresponding to monomeric DISP1-A. Panels **c–f**, show representative results ($n = 4$ biologically independent replicates). **g**, Structural comparison between DISP1-A and mouse PTCH155 (PDB ID: 7K65). The ECDs, closely apposed in PTCH1 to form a conduit for sterol transport, in DISP1-A are splayed apart. Conserved ferredoxin-like $\alpha + \beta$ open-faced sandwich folds are highlighted (lime for ECD1) and pink for ECD2). Magnified views in spectral sequence from N to C termini of the $\alpha + \beta$ open-faced sandwich folds in DISP1-A ECD1 (bottom left) and ECD2 (bottom right). Distal structures inserted into the peripheral loops of the ferredoxin-like folds are structurally unrelated to each other or to distal PTCH1 ECD structures. Red symbols indicate the five N-linked glycosylation sites, three in ECD1 (N362, N390 and N475) and two in ECD2 (N834 and N915), which can be inferred from additional densities that extend from N-X-S/T sequences in the extracellular loops. Panels **d, e, f**, see Supplementary Fig. 1 for gel source data.

Extended Data Fig. 3 |. Cryo-EM data and image processing flow for DISP1-A and DISP1-A-NNN.

a, A representative cryo-EM micrograph ($n = 4487$ for DISP1-A, $n = 4467$ for DISP1-A-NNN) and several highly-populated, reference-free 2D class averages are shown for DISP1-A (left) and DISP1-A-NNN (right). The micrograph for DISP1-A-NNN has been contrast-stretched for display in order to account for the presence of a gold edge in the upper left corner of the image (the DISP1-A-NNN particle distribution on this grid necessitated targeting of the gold edge). **b**, Schematic flow-chart representing the image processing approach for DISP1-A. Thumbnail images of each 3D class or refinement are shown along with global GS-FSC resolution in black, particle counts in red, and dashed black boxes to

indicate selected 3D classes. After separation of the R and T conformations in the first round of 3D classification, the identical processing flows for the two conformations are shown in parallel. Cryo-EM map (Red box) and atomic model of R conformation are used in main figures to present DISP1-A features. Subclasses of R and T conformations, R_1 , R_2 , T_1 and T3, are labeled.

Extended Data Fig. 4 |. Cryo-EM density and atomic model quality.

Fourier shell correlation curves (**a**), particle orientation distributions (**b**), and local resolution maps (**c, d**) are shown for R and T conformations of DISP1-A, DISP1-A-NNN, and for ShhN–DISP1-A complex. The 'gold-standard' independent half-map FSC curves and

orientation distributions were determined during refinement in cryoSPARC, map-to-model FSC curves were calculated in PHENIX using protein chains only, and directional FSC curves were estimated as in ref.⁴². The orientation distributions are plotted such that an elevation angle of 0° corresponds to a 'side-view' perpendicular to the transmembrane helices; in each case the predominant views are 'side-views' at a wide range of azimuthal angles. Local resolution estimates were computed using the BLOCRES algorithm as implemented in cryoSPARC.

Extended Data Fig. 5 |. Coincident modes of DISP1 conformational flexibility revealed by 3D classification and 3D variability analysis (3DVA).

a, Overlay comparing front view of R and T conformations (khaki and salmon, respectively) from 3D classification. Major conformational changes are localized to the extracellular domains. **b**, Cut-away view showing the formation of a 'kink' in the back-side linker of the T conformation, with an accompanying shift of about one helix turn that breaks a hydrogen bond between linker residue H777 and the backbone carbonyl of K767. **c**, Extracellular view of the R and T conformations, highlighting the movement of secondary structure elements in ECD1 (> 5 Å) and ECD2 (\sim 2 Å). The shift of ECD1 and the formation of the inter-ECD linker 'kink' appear intimately related. Numbers indicate distances (Å) between the Cα of F772 in R and T (marked by \blacktriangle in **b**) and R382 and S898 (marked by \blacktriangleright and \bigcirc , respectively, in **c**). **d**, Comparison of the most extreme R and T conformations from 3D classification (right) and the two extreme ends of PC2, PC2+ (R-like) and PC2- (T-like), from 3DVA left) shows that a nearly identical mode of motion is captured by both techniques. **e**, Distributions of DISP1-A particles stratified by their 3D class along the PC2 axis of 3DVA, demonstrating colinearity between the 3DVA trajectory and the 3D subclasses ordered by

Extended Data Fig. 6 |. Representative cryo-EM densities from selected structural features. a–d, Representative cryo-EM densities from 3D reconstruction of DISP1-A, conformation R. a, Densities of all transmembrane helices. **b**, Representative densities of beta-sheets from ECD1 and ECD2. **c**, Cryo-EM densities of three representative disulfide bonds. **d**, Cryo-EM densities of five representative CHS molecules. **e**, Representative cryo-EM densities in ShhN from 3D reconstruction of ShhN–DISP1-A complex. **f, g**, Representation of surface

hydrophobicity, viewed from top (**f**, left) or a top-front position of DISP1-A (**g**, left). Close-up views of hydrophobic cavities in both ECD1 (**f**, right) and ECD2 (**g**, right). A hydrophobic track beginning near the front-side lifted sterol (cyan) extends outward from the membrane (dotted shape). The enclosed hydrophobic conduit employed by the PTCH1 protein for sterol movement away from the membrane is formed by the juxtaposition of ECD1 and ECD2. In DISP1, although ECD1 and ECD2 are split apart in a manner that would bisect this conduit, the ECD2 portion retains a series of hydrophobic residues that line its inner surface, which could perhaps form a partial hydrophobic conduit to the hydrophobic cavities near the distal tips of ECD1 and ECD2, analogous to the sterol conduit within the center of the conjoined ECDs of PTCH1.

Extended Data Fig. 7 |. Na+-coordinating amino acid residues in DISP1, and conformational rearrangements within the transmembrane Na+ pathway in DISP1-A-NNN. a, b, Side (**a**) and extracellular (**b**) views of three Na^+ ion binding sites within the transmembrane domain, each labeled with its associated charge-neutralizing intra-membrane Asp. **c**, Residues in DISP1 with carbonyl oxygens or side-chain residues that coordinate $Na⁺$ and carboxylate residues that neutralize $Na⁺$ charge are conserved in PTCH1 (see also Extended Data Fig. 11). **d**, Solvent excluded surfaces (1.4 Å probe radius) reveal

transmembrane cavities within DISP1-A (left) that are altered in DISP1-A-NNN (right). Two extracellular branches (upper insets) that provide access to the central channel can be individually opened or closed depending on alternate conformations of M557 ('A' and 'B' in DISP1-A) and alternate conformations of L1035 ('A' and 'B' in DISP1-A-NNN). Significant rearrangements also take place around a mid-membrane water-filled cavity containing the three ion coordination sites (lower insets). In DISP1-A, short kinks in TM4 and TM10 position I568 above this cavity. In DISP1-A-NNN, however, TM4 and TM10 straighten, rotating I568 and D571N and D1049N into this cavity, dramatically reducing its volume and isolating the channel from its intracellular exit (see Supplementary Discussion).

Extended Data Fig. 8 |. Quantitative luciferase-based assay of ShhNp release.

a, Inserting Nanoluciferase coding sequence between E131 and D132 of Shh does not affect Shh autoprocessing and associated lipid modification. Immunoblotting of ShhNp detects both precursor and processed forms of the protein. Most of the expressed nanoluciferase-inserted Shh is in the processed form. **b**, Coomassie blue staining of purified SCUBE2 ($30-281$) protein. **c**, Composition of buffers used in this study(see Fig. 2b, c). **d**, Time course of NanoLuc-ShhNp release at different Na⁺ concentrations (see panel **c**) with or without SCUBE2 ($30-281$, 1μM). One representative set of normalized data (see Methods) with linear regression (dashed lines) is shown for each condition $(n = 4$ biologically independent experiments). The release rate determined as the slope of the linear regression line is presented in Fig. 2b. **e**, Western blot-based ShhNp release assay (Methods, ShhNp release endpoint assay). Culture media and cell lysates from transiently transfected Disp−/− MEFs were probed by immunoblotting for DISP

HA (SBP and HA-tagged at the C-terminus), ShhNp, and SCUBE2. β-ACTIN, loading control. **f**, DISP-ShhNp binding assay using HEK293 with stably integrated constructs for doxycycline-inducible expression of full-length Shh20. DISP1-A variants tagged with SBP and HA at the C-terminus (DISP-HA) were immunoprecipitated with Streptavidin resin, and ShhNp detected by Western blot. Alterations in ion site residues as follows: DISP1- A-NNN, D571N/D572N/D1049N; DISP1-A-VVVVVA, T613V/T614V/T1087V/T1088V/ T610V/S611A; Site I (NVV), D1049N/T613V/T614V; Site II (NVV), D571N/T1087V/ T1088V; Site III (NVA), D572N/T610V/S611A; Site I (LVV), D1049L/T613V/T614V; Site II (LVV), D571L/T1087V/T1088V; Site III (LVA), D572L/T610V/S611A. Panels **a, b, e, f**, show representative results ($n = 3$ biologically independent replicates, see Supplementary Fig. 1 for gel source data).

Extended Data Fig. 9 |. Cryo-EM data and image processing flow for ShhN–DISP1-A complex, and comparison to Drosophila HhNDisp complex.

a, A representative cryo-EM micrograph ($n = 1687$) and highly populated reference-free 2D class averages for ShhN–DISP1-A. **b**, Schematic flow-chart illustrating the image processing used for ShhN–DISP1-A data. Thumbnail images are shown for reference-based 3D classes and high resolution refinements. Dashed black boxes indicate 3D classes selected for the next processing step, with class particle counts in red and refinement GS-FSC resolutions in black. The label, '3D classification with references' indicates that explicit 'apo' and 'complex'references were used to seed the classification. **c**, Left, the

cryo-EM density from our ShhN–DISP1-A complex (transparent grey), overlaid on a ribbon diagram of ShhN (goldenrod). Right, cryo-EM density from the Drosophila HhNDisp complex in Cannac et al²⁴. (transparent grey, EMD: 10464), overlaid on the ribbon diagram of HhN (orchid, model extracted from PDB ID: 6TD6). Middle, superimposition of the ShhN–DISP1-A and HhNDisp complex models reported here and in Cannac et al.24, with mouse DISP1-A and ShhN in khaki and goldenrod, and Drosophila Disp and HhN in pink and orchid. The model of Drosophila HhN was based on docking of a Drosophila HhN structure⁵⁶ within a 4.8 Å density map. Relative to Drosophila HhN, murine ShhN is translated upwards, away from the membrane, and rotated towards the right. Asterisks indicate corresponding positions near the N termini of ShhN and HhN proteins. This difference is somewhat puzzling in light of the ability of mammalian DISP1 to rescue Drosophila disp mutant function⁶, and we cannot definitively account for it. One functional difference is that mammalian DISP1 cooperates with SCUBE2 for its Hedgehogreleasing activity, whereas Drosophila lacks a Scube orthologue. The ECD domain split and the Hedgehog interaction with the furin-cleaved linker arm together help explain the requirement for furin cleavage²² in DISP1 function.

Extended Data Fig. 10 |. Conformational dynamics link intramembrane Na+ site occupancy to Hedgehog release.

a, Three-dimensional variability analysis (3DVA) of the ShhN–DISP1-A complex dataset reveals a conformational series with ShhN bound or absent at opposite ends of the first principal component (PC1). Front (first line) and top (second line) views of reconstructed densities from the extremes of PC1 (unbound, khaki; bound, salmon) are shown, with a superimposed view of these extremes in the centre. The overall changes in ECD position are illustrated by lines drawn atop the reconstructed densities and by schematized diagrams to the left. Reconstructed densities at the $Na⁺$ coordination sites in the transmembrane domain indicate that these sites change from fully occupied in the unbound state to site I only

occupied in the bound state. See also Supplementary Video 4. b, 3DVA analysis of the apoprotein preparation shows a similar conformational series along PC1, along with similar shifts in Na⁺ site occupancy.

Extended Data Fig. 11 |. Na+ ion utilization by DISP1, PTCH1, and other members of the RND transporter family.

a, Close-up views of Na⁺ ion sites I, II, and III in DISP1, showing the locations of liganding oxygens from amino acid side chains and main-chain carbonyls (see main text), and of the corresponding locations in PTCH135 (PDB ID: 6RMG), based on structural alignment of the two proteins. Note the presence in both proteins of a charge-

neutralizing acidic residue at each site, and the conservation of side-chain oxygens as ligands. **b**, Structural alignment, using Chimera Matchmaker, of RND family members, including DISP1, PTCH1, NPC1, and several prokaryotic RND transporters. **c**, Tabulated conservation of charge-neutralizing residues (3 total) or oxygen ligands from amino acid side-chains (10 total) in the indicated proteins, aligned from structure (**b**, see PDB IDs from the second right column) or, if possible without ambiguity, aligned from sequence. References to specific sequences, structures, or background information are given within the table, including refs $.1,2,6,24,27,32,35,57-71$. Note the close conservation of Na⁺-liganding side-chains and charge-neutralizing residues in PTCH1, known to require Na^+ for its activity, and in Disp from Drosophila melanogaster. The prokaryotic Na^+ -utilizing SecD1/ SecF1 peptide translocator from Vibrio alginolyticus (encoded as two peptides; aligned by homology to Thermus thermophilus SecDF), in contrast, appears to have evolved a distinct mode of Na⁺ interaction. See Supplementary Discussion. Abbreviations: Mmus, Mus musculus; Dmel, Drosophila melanogaster; Hsap, Homo sapiens; Ecol, Escherichia coli; Paer, Pseudomonas aeruginosa; Cjej, Campylobacter jejuni; Abau, Acinetobacter baumannii; Msme, Mycolicibacterium smegmatis; Bmul, Burkholderia multivorans; Tthe2, Thermus thermophilus; Valg, Vibrio alginolyticus.

Extended Data Table 1.

Cryo-EM data collection, refinement and validation statistics.

Supplementary Material

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Data availability

Coordinates for the DISP1-A R conformation, DISP1-A T conformation, DISP1-A-NNN, and ShhN–DISP1-A complex models reported in this paper have been deposited in the Protein Data Bank under accessions 7RPH, 7RPI, 7RPJ and 7RPK, respectively. The maps of DISP1-A R conformation, DISP1-A T conformation, DISP1-A-NNN and ShhN– DISP1-A complex have been deposited in the Electron Microscopy Data Bank under accession codes EMD-24614, EMD-24615, EMD-24616 and EMD-24617, respectively. Further information and requests for data should be directed to the corresponding authors. PDB IDs of cited structures are provided either in the related figure legends or in the Methods.

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Fig. 1 |. Structure and lipid binding sites of DISP1.

a, b, Corresponding views of the cryo-EM map (top) and atomic model (bottom) of DISP1-A (the R conformation, as seen by 3D classification and 3DVA) (Extended Data Figs. 3–5). Domains are coloured as follows: pre-TM1 helix and TM1, blue; TM1–TM2 loop, lime; TM2–TM6, yellow; pre-TM7 and TM7, orange; TM7–TM8 loop, pink; TM8– TM12, lavender. **b**, Topology diagram of DISP1-A showing secondary structure elements (same colours as in **a**), disulfide bonds (gold lines), glycosylation sites (red symbols) and unresolved regions (black dashed lines, amino acid residues 1, 2, 172, 173, 264–282, 399– 409, 667–680 and 1144–1521); (Extended Data Fig. 2). Secondary elements exchanged between extracellular loops to form ECD1 and ECD2 are highlighted by oval shadows. **c**, Front view of DISP1-A (pale ribbon) with resolved LMNG (magenta) and CHS (blue) molecules (see also Supplementary Video 2). A sterol molecule (cyan) located at the front of DISP1 is positioned above other sterols in the outer leaflet and is shown in the magnified inset (right) as a stick model with corresponding density and an associated, tilted LMNG molecule (magenta); this lifted sterol may correspond to an intermediate in the extraction of Hedgehog-associated lipids from the membrane.

Fig. 2 |. Three Na+ densities in the transmembrane domain.

a, Close-up views of the full set of coordinating ligands for each of the three Na+ ions are shown with protein-derived densities superimposed on the atomic model of DISP1-A (overview, top left; dashed red line showing the pseudodyad axis between TM1–TM6 and TM7–TM12). Dashed lines in magnified ion sites denote coordination bonds and corresponding centre-to-centre distances between Na⁺ ions and associated oxygens. Contour levels are 6 σ for sites I and II, and 5 σ for site III. b, c, Na⁺-dependent ShhNp release by DISP1. **b**, NanoLuc-ShhNp released from Disp−/− mouse embryonic fibroblasts (MEFs) transfected to express DISP1-A–mEGFP and NanoLuc–ShhNp, and incubated in defined

medium at different $Na⁺$ concentrations, with or without (control) purified SCUBE2 protein (mouse SCUBE2 lacking amino acids 30–281). NMDG+ was added to maintain osmolarity for Na+ concentrations below 155 mM (Extended Data Fig. 8c). Slopes from linear regression applied to the linear portions of normalized NanoLuc–ShhNp release data were used to calculate release rates (Methods, Extended Data Fig. 8d). NanoLuc– ShhNp release was critically dependent on the $Na⁺$ gradient. NanoLuc–ShhNp release was normalized to DISP1, ShhNp and total protein. RLU, relative luminescence units. **c**, Impaired NanoLuc–ShhNp release (measured as in **b**) for DISP1-A (normalized to 1.0) and variants with alterations disrupting Na+ binding at all three sites (DISP1-A-NNN, D571N/ D572N/D1049N) or individually, at site I (LVV, D1049L/T613V/T614V), site II (LVV, D571L/T1087V/T1088V) or site III (LVA, D572L/T610V/S611A). SCUBE2 concentrations were 1 μM in **b** (30 min time course), and 0.15 μM in **c** (75 min time course). Data in b and c are mean \pm s.d., n = 3 (c) or 4 (b) biologically independent experiments; statistical analysis for SCUBE2+ conditions (datasets normally distributed) used one-way analysis of variance (ANOVA) with Tukey's test (**b**) or Dunnett's test (**c**, with DISP1-A as comparison).

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Fig. 3 |. A transmembrane Na+ permeation channel is disrupted in the structure of DISP1-A-NNN.

a, A continuous pathway through the transmembrane domain of DISP1-A, shown as a purple surface (left), is identified by the Caver 3^{29} PyMOL plugin with a probe of radius 1.0 Å, appropriate for the passage of Na^+ . This channel is partly occupied by resolved water molecules, shown as red spheres (right; superimposition with Caver channel in centre), and provides access to the transmembrane cavity that contains the $Na⁺$ ions and water (Supplementary Video 3). **b**, Close-up views of the three Na^+ ion sites in the DISP1-A-NNN structure (3.2 Å resolution) show no discernible Na⁺ densities, even when contoured at 3σ (Na⁺ densities in DISP1-A were clearly seen at 5σ for all three sites). Site III in particular has collapsed entirely, with TM4 and TM5 moving close enough to permit D572N to form hydrogen bonds with the side-chains of T610 and S611. **c**, In DISP1-A, D571 and D1049 side chains coordinate and neutralize their respective $Na⁺$ ions, maintaining a minimum distance of 5.3 Å, whereas in DISP1-A-NNN, the absence of $Na⁺$ permits D571N and D1049N side chains to rotate toward and interact across the pseudodyad axis at a reduced distance of 3.3 Å. The motion of D571N together with an upward tilting of H1053 disrupt the connection between the channel and the intracellular exit (Supplementary Discussion).

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Fig. 4 |. DISP1-A binds ShhN between its distal ECDs.

a, Cryo-EM density map of a ShhN–DISP1-A complex at 2.7 Å resolution enabled de novo modelling of ShhN at a location 26 Å above the transmembrane domain of DISP1-A. ShhN is in orange; DISP1-A colours are as in Fig. 1a. The furin-cleaved linker, although not resolved in the DISP1-A apo structure, is stabilized within the complex and seen to extend from ECD2 in a one-armed molecular embrace of ShhN. The region of ShhN contacted by this linker corresponds to a polybasic region that interacts with heparin sulfate or other sulfated glycosaminoglycans $(GAGs)$ ³⁶, which would be displaced by the DISP1 molecular embrace. **b**, Three major interaction interfaces between ShhN and DISP1-A. The ECD1 and ECD2 interfaces feature primarily van der Waals interactions, with the extensive ECD2 interface additionally including a hydrogen bond between ShhN K75 and the backbone carbonyl of DISP1-A R241 (located just below the expanded view (not shown)). Near the tip of the furin-cleaved linker arm, R276 of DISP1-A forms an ionic interaction with a sulfate ion that also interacts with basic residues in ShhN (R124, R154 and K88). **c**,

A model for Na+-driven conformational dynamics in the DISP1 activity cycle. Membraneanchored, dual-lipid-modified Hedgehog protein (HH) is activated for release to SCUBE2 by interaction with DISP1 ECDs. Hedgehog activation (left) entails (1) a pincer-like action of the DISP1 ECDs; (2) interaction with and stabilization of the DISP1 furin-cleaved linker (dark blue); and (3) accommodation of Hedgehog-linked lipids in hydrophobic distal cavities of DISP1 ECDs. From this activated state, associated with restricted Na+ occupancy (site I only, from 3DVA, ShhN-bound state), SCUBE2 availability and the chemiosmotic force of the transmembrane Na+ gradient together drive Hedgehog release to SCUBE2 and resetting of DISP1 for resumption of the cycle (right) (see Supplementary Video 4).