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Architecture of the human erythrocyte ankyrin-1 complex

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The stability and shape of the erythrocyte membrane is provided by the ankyrin-1 complex, but how it tethers the spectrin-actin cytoskeleton to the lipid bilayer and the nature of its association with the band 3 anion exchanger and the Rhesus glycoproteins remains unknown. Here we present structures of ankyrin-1 complexes purified from human erythrocytes. We reveal the architecture of a core complex of ankyrin-1, the Rhesus proteins RhAG and RhCE, the band 3 anion exchanger, protein 4.2, glycophorin A and glycophorin B. The distinct T-shaped conformation of membrane-bound ankyrin-1 facilitates recognition of RhCE and, unexpectedly, the water channel aquaporin-1. Together, our results uncover the molecular details of ankyrin-1 association with the erythrocyte membrane, and illustrate the mechanism of ankyrin-mediated membrane protein clustering.

A n ordered lattice of spectrin and actin gives mechanical stability and shape to otherwise fragile cellular membranes. The periodic spectrin–actin cytoskeleton was first identified in erythrocytes¹, but has since been found in other cell types, including neurons². The spectrin–actin network is tethered to the membrane by giant, spring-like proteins known as ankyrins, which also serve the purpose of clustering diverse membrane proteins at specific subcellular locations.

Three ankyrin proteins are present in vertebrates: ankyrin-1 (ankyrin-R), ankyrin-2 (ankyrin-B) and ankyrin-3 (ankyrin-G). Ankyrins share a common domain arrangement, consisting of a highly conserved N-terminal membrane binding domain with 24 ankyrin repeats (AR1-24), followed by a spectrin binding module (ZU5-ZU5-UPA; ZZU) and a death domain of unclear function. The characteristic 24-repeat membrane binding domain of ankyrins has a groove on the inner concave surface that mediates recruitment of short peptides from target proteins, as well as mediating autoinhibition by regions of the MBD-ZZU linker and the C-terminal tail. Ankyrin-1 was first identified in erythrocytes³, where it mediates the attachment of the band 3 anion transport protein (AE1; SLC4A1) to spectrin⁴. It is also found in other tissues, including skeletal muscle and brain, where it has recently been shown to mediate clustering of Kv3.1b K⁺ channels in specific neuronal subtypes⁵. Ankyrin-2 and 3 play pivotal roles as dynamic adaptor platforms in cardiomyocytes and striated muscle cells⁶, and in neurons of the central⁷ and peripheral⁸ nervous systems, respectively.

In erythrocytes, ankyrin-1 attaches the cytoskeleton to the membrane via direct interaction with spectrin⁹, and additionally mediates clustering of integral membrane proteins, helping maintain the structure of the membrane and coordinating the spatial organization of transporters and channels involved in the regulation of cellular volume and cytosolic composition. Mutations in components of the erythrocyte ankyrin-1 complex are linked to an inherited anemia known as hereditary spherocytosis, in which the characteristic biconcave disc shape of healthy erythrocytes is lost, resulting in small, fragile, spherical cells known as spherocytes¹⁰. Ankyrin and spectrin mutations are also associated with several other genetic diseases, including neurological and cardiac abnormalities¹¹. The erythrocyte ankyrin-1 complex is known to contain the band 3 chloride/ bicarbonate exchanger, the adaptor molecule protein 4.2, the single transmembrane proteins glycophorin A (GPA) and glycophorin B (GPB), and the Rhesus proteins RhAG and RhCE¹². Other proteins that are believed to associate with the ankyrin-1 complex include GLUT1 (ref. ¹³), LW and CD-47 (ref. ¹⁴). However, the stoichiometry and architecture of the complex remains unknown, making it close to impossible to reconstitute recombinantly.

Here, we purified the ankyrin-1 complex from human erythrocytes, and captured a series of high-resolution structures by single-particle cryogenic-electron microscopy (cryo-EM), complemented by a subtomogram average of the complex in the context of native erythrocyte membrane fragments. These revealed its architecture and composition, shedding light on the mechanism of membrane protein recruitment by ankyrin and protein 4.2, and identifying aquaporin-1 as an unanticipated component of the ankyrin-1 complex.

Results

Structure determination of the ankyrin-1 complex. The ankyrin-1 complex was purified from digitonin-solubilized human erythrocyte ghost membranes by density gradient centrifugation, followed by size-exclusion chromatography (Fig. 1a, Supplementary Fig. 1a,b and Supplementary Video 1). The final sample contained a major 1.2-MDa species as assessed by mass photometry¹⁵ (Supplementary Fig. 1c). The presence of main components of the erythrocyte ankyrin-1 complex, including ankyrin-1, band 3, protein 4.2, GPA and RhAG was assessed by SDS–PAGE and confirmed by mass spectrometry analysis (Supplementary Table 1).

Cryo-EM analysis (Table 1) of the purified sample revealed both a mixture of ankyrin-1 containing complexes, as well as smaller

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Fig. 1 | Architecture of the human erythrocyte ankyrin-1 complex. a, The overall cryo-EM density map of the ankyrin-1 complex (Class 1a, transparent white surface) shown in three different views, with local reconstructions of different regions superposed. Band 3-I, II, III are colored orange, lilac and yellow, respectively; ankyrin is colored red, protein 4.2 green, RhCE purple and RhAG blue and light blue, GPA is colored in magenta and GPB in aquamarine.
b, Model of the ankyrin-1 complex in the plane of the membrane (left panel), viewed from the cytoplasm (central panel) and from the outside the plasma membrane (right panel). Reference colors are the same used in a. c, Schematic of the proposed ankyrin-1 complex in the RBC. Inside the dashed line the connection between ankyrin and spectrin is depicted. d, Subtomogram average of the ankyrin-1 complex from native vesicles with rigid body fit atomic model.

complexes of free band 3 with GPA (Extended Data Fig. 1a,b and Supplementary Fig. 1d).

After repicking using a Topaz model trained on the smaller complexes¹⁶, ab initio reconstruction and refinement of the band 3–GPA class gave a C1 reconstruction at 2.8 Å. Both protomers were in the outward open state, and the transmembrane domains exhibit apparent C2 symmetry. Refinement gave a 2.3 Å reconstruction with C2 symmetry applied, allowing us to build a complete atomic model of the band 3–GPA complex (Extended Data Fig. 1c).

Refinement of all 710,000 ankyrin-1-containing particles against an ab initio reference yielded a 2.4 Å consensus reconstruction (Extended Data Fig. 2). Well-ordered density is observed for the Rh heterotrimer, ankyrin-1, protein 4.2 and the cytoplasmic domains of a single band 3 dimer (denoted subsequently as Band 3-I). Masked refinement and in some cases focused three-dimensional- (3D-) variability analysis was used to improve the density quality for each component (Extended Data Fig. 3).

Subclassification of the ankyrin complexes revealed six distinct classes with variable composition (Extended Data Fig. 4a). The core architecture is conserved in all six classes, but with elaborations. Class 1 had a larger micelle than observed in the consensus refinement, accommodating two additional band 3 dimers (denoted Band 3-II and Band 3-III) that interact directly with ankyrin. Two-dimensional (2D) classes of Class 1 resemble complexes identified in native membrane vesicles using cryo-electron tomography (cryo-ET) (Fig. 1d, Extended Data Fig. 5, Supplementary Fig. 2 and Supplementary Videos 2 and 3). Classes 2 and 5 had an additional component present that proved to be an aquaporin-1 (AQP1) tetramer. Class 4 has an overall architecture similar to Class 1, but with a different orientation of the Band 3-I transmembrane region and the presence of an unknown membrane protein.

While the cytoplasmic domains of Band 3-I were well ordered in the consensus refinement, the transmembrane (TM) domains were very poorly resolved, which we rectified using the following classification scheme. After local refinement gave a poor quality 3.8 Å map, 3D-variability analysis was carried out in cryoSPARC using a mask around the Band 3-I TM. Clustering with a single mode corresponding to an order–disorder transition of the transmembrane region allowed identification of a 226,000 particle subset for which local refinement gave a 2.8 Å map with excellent density quality, and a well-resolved interface with protein 4.2 (Extended Data Fig. 6). A similar approach was applied to improve the density of Band 3-II and Band 3-III starting from Class 1, resulting in 3.3 and 3.0 Å local refinements, respectively.

Table 1 Cryo-EM data collection, r	efinement and valid	ation statistics					
				Ankyrin-1 compl	ex consensus refinemei	Ħ	
	Band3-GPA (PDB 7UZ3) (EMD-26874)	Composite (PDB 7V0K) (EMD-26943)	Rh-AR1-5 (PDB 7UZQ) (EMD-26916)	Protein 4.2 (PDB 7UZS) (EMD-26917)	Ankyrin N-term (PDB 7UZU) (EMD-26918)	Band3-I cytosolic domains (PDB 7UZV) (EMD-26919)	Band3-I TM domains (PDB 7V07) (EMD-26940)
Data collection and processing							
Magnification	105,000×	105,000×	105,000×	105,000×	105,000×	105,000×	105,000x
Voltage (kV)	300	300	300	300	300	300	300
Electron exposure (e ⁻ /Ų)	58	58	58	58	58	58	58
Defocus range (µm)	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5
Pixel size (Å)	0.83	0.83	0.83	0.83	0.83	0.83	0.83
Symmetry imposed	C1	C1	C1	C1	CI	C	C1
Initial micrographs	14,926	14,926	14,926	14,926	14,926	14,926	14,926
Final micrographs	14,464	14,464	14,464	14,464	14,464	14,464	14,464
Initial particle images	1,781,329	1,781,329	1,781,329	1,781,329	1,781,329	1,781,329	1,781,329
Final particle images	166,626	710,437	710,437	710,437	710,437	710,437	226,351
Map resolution (Å)	2.33	2.39	2.17	2.20	2.34	2.47	2.82
FSC threshold	0.143	0.143	0.143	0.143	0.143	0.143	0.143
Map sharpening <i>B</i> factor ($Å^2$)	-47.7	-42.5	-52.5	-56.5	-65.5	-70.1	-80.1
Model composition							
Non-hydrogen atoms	9,315	32,037	10,403	5,367	3,744	4,682	9,162
Protein residues	1,114	3,997	1,315	658	483	584	1,114
Ligand molecules	20	14	4	0	0	0	20
R.m.s. deviations							
Bond lengths (Å)	0.0054	0.0039	0.0062	0.0043	0.0029	0.0027	0.0059
Bond angles ($^{\circ}$)	0.76	0.68	0.91	0.72	0.57	0.58	0.90
Validation							
Clashscore	2.87	7.29	4.49	2.70	3.14	3.69	8.34
Poor rotamers (%)	0.84	2.26	2.27	1.79	0.52	2.84	3.89
Ramachandran plot							
Favored (%)	97.82	98.07	97.99	98.92	96.87	98.26	97.91
Allowed (%)	2.18	1.80	1.78	1.08	3.13	1.39	1.81
Disallowed (%)	0	0.13	0.23	0	0	0.35	0.27
							Continued

						Class	1					
	Composite (PDB 8CS9) (EMD-26960)	Rh-AR1-5 (PDB 7V0S) (EMD- 26949)	Protein 4.2 (PDB 7V0Q) (EMD- 26948)	Ankyrin N-term (PDB 7VOM) (EMD- 26944)	Ankyrin C-term (PDB 7VOX) (EMD- 26952)	Band3-I cytosolic domain (PDB 7V0T) (EMD- 26950)	Band3-II cytosolic domain (PDB 7VOU) (EMD- 26951)	Band3-III cytosolic domain (PDB 7V0Y) (EMD- 26953)	Band3-I TM (PDB 8CRQ) (EMD- 26955)	Band3-II TM (PDB 7V19) (EMD- 26954)	Band3-III TM (PDB 8CRR) (EMD- 26956)	Rh-GPB (PDB 8CRT) (EMD- 26958)
Data collection and process	sing											
Magnification	105,000×	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x	105,000x
Voltage (kV)	300	300	300	300	300	300	300	300	300	300	300	300
Electron exposure (e ⁻ /Ų)	58	58	58	58	58	58	58	58	58	58	58	58
Defocus range (μm)	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5	-0.5 to -1.5
Pixel size (Å)	0.83	0.83	0.83	0.83	0.83	0.83	0.83	0.83	0.83	0.83	0.83	0.83
Symmetry imposed	C1	CJ	C	C1	CJ	CI	CI	CI	CI	CJ	C1	C
Initial micrographs	14,926	14,926	14,926	14,926	14,926	14,926	14,926	14,926	14,926	14,926	14,926	14,926
Final micrographs	14,464	14,464	14,464	14,464	14,464	14,464	14,464	14,464	14,464	14,464	14,464	14,464
Initial particle images	710,437	710,437	710,437	710,437	710,437	710,437	710,437	710,437	710,437	710,437	710,437	710,437
Final particle images	126,197	126,197	126,197	126,197	126,197	126,197	126,197	126,197	85,645	44,624	96,966	22,953
Map resolution (Å)	2.75	2.44	2.44	2.65	2.97	2.71	3.0	2.95	3.14	3.33	2.96	3.0
FSC threshold	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143	0.143
Map sharpening B factor (Å ²)	-35.1	-57.5	-58.7	-65.9	-84.7	-79.2	-101.2	-104.3	-83.1	-79.7	-77.4	-39.9
Model composition												
Non-hydrogen atoms	60,944	10,303	5,301	3,724	2,818	4,658	4,453	4,454	8,936	8,936	8,936	17,925
Protein residues	7,677	1,315	658	483	375	584	558	558	1,114	1,114	1,114	2,305
Ligand molecules	27	c	0	0	0	0	0	0	80	8	80	9
R.m.s. deviations												
Bond lengths (Å)	0.0063	0.0042	0.0036	0.0034	0.0023	0.0037	0.0050	0.0031	0.0035	0.0022	0.0041	0.0032
Bond angles (°)	1.16	0.72	0.61	0.56	0.63	0.63	0.75	0.63	0.56	0.49	0.57	0.58
Validation												
Clashscore	7.77	5.04	3.48	3.55	6.52	6.19	7.40	5.27	4.15	4.97	4.26	5.01
Poor rotamers (%)	С	2.55	2.15	1.29	5.05	3.45	2.50	1.46	1.37	1.16	0.95	2.36
Ramachandran plot												
Favored (%)	97.36	98.22	98.92	96.66	95.96	98.26	98.54	98.54	98.46	98.37	97.91	98.50
Allowed (%)	2.42	1.70	1.08	3.34	4.04	1.57	1.46	1.46	1.54	1.63	2.09	1.45
Disallowed (%)	0.22	0.08	0	0	0	0.17	0	0	0	0	0	0.04
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Fig. 2 | Interactions between ankyrin-1 and the proteins in the ankyrin-1 complex. a, Overview of ankyrin-1 interactions with binding partners in the ankyrin-1 complex. The proteins are depicted in cartoon representation. Transmembrane domains of band 3, GPA and GPB are omitted for clarity. Ankyrin (red) interacts with RhCE (purple) and AQP1 (salmon), and both interact with the peptide binding groove of AR1-5; protein 4.2 (green) interacts with the exterior surface of AR1 and AR6-13; Band 3-II (B3-II) (lilac) and Band 3-III (B3-III) (yellow) interact predominantly with the exterior surface of AR1-3; Band 3-II (B3-II) (lilac) and Band 3-III (B3-III) (yellow) interact predominantly with the exterior surface of AR17-24, the N terminus domain of Band 3-III also interacts with the peptide binding groove of AR6-10. Insets refer to regions highlighted in **b-f**. On the top left of **a**, a small thumbnail is included to show how the figure is positioned compare to Fig. 1a. **b-f**, Close-up views of the interactions between protein 4.2 (**c**); ankyrin and RhCE (**d**); ankyrin and Band 3-II (**e**) and ankyrin and Band 3-III (**f**). The key residues that mediate the interactions in the interfaces are shown as sticks. Key interactions are indicated with black dotted lines.

By applying this clustering approach to Class 2, a particle subset was identified with improved AQP1 density, resulting in a 3.0 Å C1 map that facilitated unambiguous fitting of the AQP1 crystal structure, and identification of the regions of AQP1 that interact with ankyrin-1 and protein 4.2. Application of local C4 symmetry yielded a 2.4 Å map (Supplementary Video 4).

We have used the reconstructions described above to build atomic models for class 1a (Fig. 1b), which we believe represents an intact complex, and class 2, in which the interaction with AQP1 is best defined, as well as for the consensus refinement of all classes, which had the best resolved density at the core of the complex. AQP1 is present in classes 2 and 5. The remaining classes may represent genuine diversity in the composition of complexes present in the membrane, or partial dissociation of a single native complex (Extended Data Fig. 4a). The association of AQP1 in classes 2 and 5 appears compatible with the arrangement of band 3 dimers in Class 1, despite a substantial rigid body rearrangement of the transmembrane domains of Band 3-I in the presence of AQP1 (Extended Data Fig. 4b). We therefore assess that in vivo, the three band 3 dimers identified here, as well as AQP1, are likely to be part of the same complex. This conclusion is supported by analysis of tomograms obtained from vesicles generated by extrusion of native erythrocyte

membranes. The appearance of particles in the tomograms is very similar to the 2D classes obtained by single-particle analysis (Supplementary Video 2), and subtomogram averaging of particles extracted from the tomograms identifies a class with near identical architecture to Class 1/1a (Fig. 1d, Supplementary Fig. 2a,b and Supplementary Video 3). The absence of CD-47 and LW from the complex may indicate that they are loosely associated, and lost during digitonin extraction, or that they are associated with a distinct complex that we have not isolated.

Architecture of the human erythrocyte ankyrin-1 complex. The architecture of Class 1a is presented in Fig. 1 and Supplementary Video 1. The membrane-embedded core of the complex is formed by the (RhAG)₂(RhCE) heterotrimer. Ankyrin binds to the N and C termini of RhCE, via the first five ankyrin repeats (AR1-5), which are oriented parallel to the membrane. The subsequent ankyrin repeats (AR6-24) emerge perpendicularly downward from a T interface with AR1-5, and gradually recurve to run almost parallel to the membrane.

Protein 4.2 is oriented vertically, such that the myristoylated N terminus contacts the membrane, with one flat surface bound to AR1 and AR6-13 of ankyrin, while the distal face recruits band 3.



Fig. 3 | Protein 4.2 is an adaptor that mediates binding of Band 3-I. a, The 2.4 Å cryo-EM density map of protein 4.2 from local refinement of the consensus reconstruction. **b**, Structure of protein 4.2 colored by domain from the N (blue) to the C (red) terminus. The four domains are labeled. The proteins are depicted as cartoon representations. **c**, Model of protein 4.2 (green) and band 3-I (orange), PIP₂ is colored in yellow. The proteins are shown as cartoon representations, and the membrane is represented in gray. **d**, The outer face of protein 4.2 forms the primary binding site for Band 3-I. Protein 4.2 is displayed as a molecular surface colored by domain from the N (blue) to the C (red) terminus, band 3-I cytosolic domain is displayed as ribbon and the parts of Band 3-I that are not interacting with protein 4.2 are shown in transparency.

Three band 3–GPA dimers (Band 3-I, Band 3-II and Band 3-III) form part of the complex. Band 3-I binds to the distal face of protein 4.2 via the band 3 cytoplasmic domain and N-terminal peptide. Band 3-II and Band 3-III, by contrast, both interact directly with ankyrin. Band 3-II interacts with the outer face of AR17-19 via the cytoplasmic domain, while the cytoplasmic domain of Band 3-III binds to the outer face of AR21-24 and the Band 3-III N-terminal peptide runs back along the inner groove of ankyrin.

Membrane association and target recruitment by ankyrin-1. We observe simultaneous engagement of the ankyrin-1 membrane binding domain with multiple targets (Fig. 2a) that fall into three distinct categories: membrane embedded (RhCE and AQP1), extramembrane (cytosolic domains of band 3) and peripheral adapters (protein 4.2). The integral membrane targets, RhCE and AQP1, both interact with the peptide binding groove of AR1-5, the adaptor (protein 4.2) interacts with the exterior surface of AR1-13 and the two extramembrane targets (Band 3-II and Band 3-III) interact predominantly with the exterior surface of AR17-24, supplemented in the case of Band 3-III by interactions of the N terminus with the peptide binding groove of AR6-10.

The Rh heterotrimer forms the primary membrane attachment site for ankyrin-1, mediated via an interaction between the N and

C termini of RhCE and the first five repeats of ankyrin-1 (Fig. 2d). The structure of ankyrin-1 in the complex (Extended Data Fig. 7a) adopts a T-shaped arrangement in which AR1-5 are orthogonal to AR6-24, with the peptide binding groove of AR1-5 oriented toward the membrane. This is distinct from the extended conformation observed for ankyrin-2 bound to autoinhibitory motifs¹⁷ (Extended Data Fig. 7b). Comparison of the structures of AR1-9 of ankyrin-2 bound to a peptide from the NaV1.2 channel¹⁸, an ankyrin membrane target, with structures bound to the autoinhibitory peptide, shows substantial flexibility at the AR5-6 interface (Extended Data Fig. 7b,c), albeit not to the extent seen here in ankyrin-1. A sequence alignment of the 24 repeats of ankyrin-1 (Extended Data Fig. 7e) shows that, while AR5 and AR6 are highly conserved when compared to the equivalent regions of ankyrin-2 and ankyrin-3 (Extended Data Fig. 7f), they are the most divergent of the ankyrin repeats when compared with the other repeats in the ankyrin-1 MBD. The rearrangement of the AR1-5 module orients the canonical protein binding groove to directly face the membrane, to bind the membrane-embedded targets RhCE (in all classes) and AQP1 (classes 2 and 5) (Extended Data Fig. 4a).

Protein 4.2 forms multiple sites of attachment to ankyrin-1, all of which are outside the canonical peptide binding groove (Fig. 2c). Protein 4.2 acts as an adaptor molecule and a stabilizer for ankyrin,

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Fig. 4 | Aquaporin is a component of the ankyrin-1 complex. a, The 2.8 Å cryo-EM density map of the aquaporin in Class 2. Densities corresponding to the four subunits are colored in light orange, deep pink, medium purple and hot pink. **b**, Close up of AQP1 palmitoylation of Cys 87. **c**, Aquaporin-1 is present in classes 2 and 5 (Extended Data Fig. 5). The cryo-EM density map of Class 2 shows inside the membrane the density and atomic models corresponding to aquaporin (salmon), Rh (purple, blue and light blue) and Band 3-I (orange). **d**, The model of Class 2 shows that AQP1 (salmon color) is located at the mutual interface of protein 4.2 (green), ankyrin (red), RhCE (purple) and Band 3-I (orange). **e**, Close up of the interaction region between AQP1, protein 4.2 and ankyrin. The membrane is represented in gray. The key residues that mediate the interactions in the interfaces are shown as sticks. Unmodeled density of putative lipid headgroups is shown as an orange surface.

with the outer face of protein 4.2 forming the attachment site for Band 3-I (Fig. 2b). Interaction of domain 1 of protein 4.2 with the exposed 'edge' of AR1 appears to further stabilize the T-shaped conformation of the ankyrin-1 membrane binding domain (Fig. 2c).

In Class 1, Band 3-II and Band 3-III directly bind to ankyrin, interacting with repeats 17–19 and 21–24, respectively (Fig. 2e, f). The N terminus of Band 3-III runs back along the inner ankyrin groove, with residues 2–24 forming an ordered interaction with AR6-10 and the AR5-6 linker (Extended Data Fig. 8a–c and Supplementary Video 5). This peptide is reversed when compared to the binding mode of ankyrin autoinhibitory motifs (Extended Data Fig. 8d–f), but in the same direction as the NaV 1.2 N-terminal peptide bound to ankyrin-2 (ref. ¹⁸) (Extended Data Fig. 7d). The orientation of the Band 3-I cytoplasmic domains is inverted with respect to the membrane when compared with Band 3-II and Band 3-III, which are both directly bound by ankyrin (Extended Data Fig. 9a). Interacting surfaces of band 3 cytoplasmic domains are distinct and nonoverlapping (Extended Data Fig. 9b).

Structure of protein 4.2 and interactions with band 3. The 2.2 Å structure of protein 4.2 is solved here in the context of the ankyrin-1 complex (Fig. 3a,b). As previously suggested^{19,20}, the fold of protein 4.2, is very similar to that of the closed conformation

of tissue transglutaminase (TG2). The central papain-like domain (domain 2) is surrounded by three Ig-like domains (domains 1, 3 and 4). The N-terminal glycine of protein 4.2 is known to be myristoylated²¹, and here we see that the N terminus directly contacts the membrane. The inner face of protein 4.2 interacts with ankyrin-1, while the outer face forms the primary binding site for Band 3-I (Fig. 3c,d). The Band 3-I binding mode shares some broad similarities with the binding mode of Band 3-II to ankyrin (Extended Data Fig. 9b), insofar as it involves recognition of a 3D epitope via an extensive interface and is augmented by interaction of the free N terminus of band 3 with a peptide recognition groove. The interaction with Band 3-I is mediated by interactions with D4 and D1 of protein 4.2 (Fig. 2b), and the N terminus of Band 3-I is nestled in a groove formed by D1, D2 and D4. In Class 2, D1 of protein 4.2 also appears to be forming interactions with AQP1. D1 is also forming an interface with loops near the transmembrane dimerization interface of Band 3-I, in a region that also forms cytoplasmic-TM contacts in Band 3-II (Extended Data Fig. 9c).

Aquaporin is a component of the ankyrin-1 complex. A final resolution of 2.5 Å after local refinement with C4 symmetry allows unambiguous fitting of the crystal structure of AQP1 to the density map (Fig. 4a). Clear density is seen for waters in the pore (Extended Data Fig. 10a), as well as a lipid-like density directly attached to Cys-87, which we have assigned as a palmitoylation



Fig. 5 | (RhAG)₂(**RhCE) trimer. a**, The structure of the (RhAG)₂(RhCE) trimer as viewed from the cytoplasm (right panel), and as viewed in the plane membrane (central and right panels). RhCE is colored in purple, RhAG1 in light blue and RhAG2 in dark blue. Two molecules of cholesterol (yellow) are bound to each RhAG. b, The 2.4 Å cryo-EM density map of the (RhAG)₂(RhCE) complex. Densities corresponding to RhCE, RhAG1 and RhAG2 and cholesterol are colored as in **a. c**, Refined atomic *B* factors, averaged per residue. RhCE has higher *B* factors than RhAG, likely indicating more flexibility/ mobility. **d**, Cryo-EM density map of the Band 3-III-GPA, (RhAG)₂(RhCE) and GPB that mediates the interaction between the two. **e**, Close up of the interaction region between GPB and RhAG1. The side chain of M71 is located in a hydrophobic pocket in the surface of RhAG1, and the key residues that mediate the interactions are shown as sticks.

site (Fig. 4b). AQP1 is located at the mutual interface of Band 3-I, protein 4.2 and AR1-2 (Fig. 4c,d). The interaction appears to be largely mediated by interactions of the N terminus of AQP1 with AR1-2, and interactions of the C terminus of AQP1 with domain 1 of protein 4.2 (Fig. 4e). In addition, strong, unmodeled density is present in the cleft between AQP1 and RhCE, underneath a patch of positively charged residues contributed by ankyrin, suggesting the possibility of a lipid-mediated interaction with ankyrin-1 and the Rh heterotrimer (Extended Data Fig. 10b). The position of the Band 3-I transmembrane domains is altered substantially in the presence of AQP1 as compared to Class 1 or the consensus refinement, with a translation of roughly 16 Å putting it in close contact with AQP1 and breaking the interaction of Band 3-I with protein 4.2.

Structure and composition of the Rh channel. In the structure of the erythrocyte ankyrin-1 complex presented here, we have resolved the structure of the Rh heterotrimer at 2.2 Å (Fig. 5a,b and

Extended Data Fig. 3) revealing that it contains two molecules of RhAG and one of RhCE (Extended Data Fig. 10c,d), with the N and C termini of RhCE interacting with ankyrin-1 (Fig. 2d). RhAG also helps stabilize this interaction, as the RhCE termini pack on top of the adjacent RhAG protomers. Several cholesterol and digitonin molecules are bound to the transmembrane regions of both RhAG and RhCE on both leaflets of the membrane (Fig. 5a,b). The pore of RhAG shows the expected twin-histidine motif in the center of the pore (Extended Data Fig. 10e), with a hydrogen bonded network of water molecules similar to that seen in the structures of AmtB and RhCG. The pore of RhCE is open to both the cytosolic and extracellular sides, with only a short central occlusion (Fig. 5c and Extended Data Fig. 10f), raising the possibility that RhCE may have a role in membrane transport. The density for RhCE is also weaker than for RhAG, despite the fact that RhCE is the subunit that directly engages with ankyrin. This weaker density (and correlated higher atomic B factors) may indicate flexibility, possibly consistent with an unresolved conformational change corresponding to pore

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Fig. 6 | Band 3 in complex with GPA. a, The structure of band 3 in complex with GPA as viewed in the plane of the membrane, as viewed from the cytoplasm (central panel) and as viewed from inside the cell. Band 3 protomers are colored aqua green and gray, GPA in magenta and molecules of cholesterol (yellow) PIP₂ (orange) and POPC (blue) are shown. Glycans are colored green. **b**, The 2.3 Å cryo-EM density map of the band 3-GPA complex. Density corresponding to band 3, glycophorin, cholesterol (bound to glycophorin) and PIP₂ are colored as in **a**. The three orientations of the map are the same as **a**. **c**, Close up of the interactions between band 3 (aqua green), glycophorin (magenta) and a cholesterol molecule (yellow). The key residues that mediate the interactions are shown as sticks. **d**, Close up of the band 3 transmembrane dimer interface. PIP₂ (orange) sits in the middle of the site. The key residues that mediate the interactions in the interfaces are shown as sticks. **e**, Cut through the TM of ankyrin-1 complex shows the position of GPA bound to band 3 in the context of the complex. **f**, PIP₂ (orange) sits in the middle of the dimerization site of Band 3-I and close to the interaction site between Band 3-I and protein 4.2. The key residues that mediate the interactions in the interaction site between Band 3-I and protein 4.2. The key residues that mediate the interactions in the interactions in the interfaces and between Band 3-I TM and protein 4.2 are shown as sticks.

opening (Fig. 5c). Inspection of the map at sites of RhCE/D variation strongly suggests that RhCE is predominantly incorporated (Extended Data Fig. 10h), consistent with previous findings suggesting preferential association of RhCE with the band 3 complex²².

Local refinement of the Rh trimer in Class 1 revealed an additional transmembrane helix bound to RhAG1 in a groove formed by TM1, TM3 and TM5, with the C terminal end extending toward Band 3-III. Further classification, using a mask around the Rh trimer and Band 3-III, identified a class of 22k particles in which this density was much better defined, allowing assignment as GPB (residues 59–91) (Fig. 5d); however, given the sequence similarity with GPA in the TM region some ambiguity remains in this assignment. The N-terminal half of the helix forms the best-defined interactions with RhAG1, with the side chain of Met71 buried in a hydrophobic pocket formed by Trp93, Ile146, Met16, Leu19 and Leu12 (Fig. 5e). The C-terminal end of the GPB helix (residues 86–91) appears to be forming a weak interaction with the N-terminal juxtamembrane helix of Band 3-III.

Structure of the band 3 dimer complexed with GPA. In the structure of the free band 3 dimer (Fig. 6a,b), a GPA monomer is bound

at each end of the band 3 dimer, with the GPA transmembrane helix interacting primarily with TM4 of band 3 and the extracellular N-terminal peptide interacting with the TM7-8 loop (Fig. 6c). A tightly bound cholesterol molecule is located near the extracellular side, wedged between the transmembrane helix of GPA and TM8 of band 3 (Fig. 6c). A prominent lipid headgroup density is located at the intracellular dimer interface of band 3, where it binds to a patch of positively charged residues. Based on the distinctive shape of the density, and previous biochemical data showing enrichment of phosphoinositides in purified band 3 (ref. ²³) we have assigned this density as PI(4,5)P₂ (Fig. 6d and Supplementary Video 6). This density is also observed in classes of the intact ankyrin complex, where it is located very close to the interface of Band 3-I with protein 4.2 (Fig. 6e,f).

Discussion

Ankyrins form part of an intricate submembrane lattice that provides mechanical stability, shape and local functional organization to cellular membranes in many cell types including erythrocytes and neurons. In erythrocytes, ankyrin-1 forms the nucleus of a

macrocomplex that acts to tether spectrin to the membrane³ and organizes several highly abundant membrane proteins involved in gas exchange, pH regulation and control of cellular volume²⁴ (Supplementary Fig. 3). Mutations in elements of this complex lead to defects in erythrocyte size, shape and stability¹⁰. However, the architecture and composition of the complex, as well as the mechanism of ankyrin-1 recruitment to the membrane, have so far remained unresolved. Here, we used single-particle cryo-EM to determine structures of an erythrocyte ankyrin-1 complex that includes both the RhAG/CE heterotrimer and band 3. We provide a mechanistic hypothesis for ankyrin recruitment and autoinhibition, showing how ankyrin-1 can recruit three band 3 dimers at distinct sites, clarifying the role of protein 4.2 and identifying aquaporin-1 as an unanticipated component of the ankyrin-1 complex.

Membrane-bound ankyrin-1 adopts a T-shaped conformation. The discontinuous, T-shaped conformation of ankyrin-1 we observe in the membrane-bound state is strongly suggestive of a rearrangement on membrane binding, which we propose is linked to the mechanistic basis of both ankyrin membrane association and autoinhibition. The only previous structure of a full-length ankyrin membrane binding domain is that of ankyrin-2 fused to the autoinhibitory C-terminal tail of ankyrin-1 (ref. ¹⁷), which binds in a continuous groove that traverses the entire length of the molecule. Constructs lacking the autoinhibitory peptide failed to crystallize, likely because they were too conformationally heterogeneous. In contrast, in our structure (Fig. 1a), the ankyrin-1 repeats AR1-5 are oriented parallel to the membrane and orthogonal to the downstream ankyrin repeats. While there is no autoinhibited structure of ankyrin-1 available, the membrane binding domain and the AR5-6 region in particular are highly conserved in all three ankyrin proteins, suggesting that this is unlikely to be an isoform-specific difference. In the ankyrin-2 AR1-24 structure, the autoinhibitory peptide occupies a groove that crosses the AR5-6 boundary, stabilizing them in the 'extended' state. By contrast, in our structure, the N-terminal peptide of Band 3-III occupies this same groove, but ends at the AR5-6 interface, with the very N-terminal 11 residues, including the Syk phosphorylation site Y8, appearing to stabilize it in the T-shaped, 'hammerhead' configuration (Extended Data Fig. 8d and Supplementary Video 5). Deletion of these 11 residues in the band 3 Neapolis mutant causes a severe hemolytic anemia, without compromising insertion into the membrane²⁵. Phosphorylation of Y8 also causes alterations in erythrocyte size and shape²⁶, consistent with a role for this segment in stabilizing the membrane-associated configuration of ankyrin. We hypothesize that the Band 3-III N-terminal peptide displaces the autoinhibitory motif from the inner groove of ankyrin, freeing AR1-5 to adopt the membrane-associated configuration and leaving the AR1-5 groove unoccupied, facilitating recognition of RhCE and AQP1. Band 3-III is also the band 3 dimer that has a GPB-mediated interaction with RhAG1 in Class 1a. It is possible that pre-association of RhAG/CE and Band 3 via GPB could facilitate recruitment of ankyrin-1, with band 3 first displacing the autoinhibition motif, and then AR1-5 binding to RhCE (Supplementary Fig. 4). Binding to protein 4.2 further stabilizes the T-shaped configuration of ankyrin, resulting in the core complex. In this model, the autoinhibited state is not competent to bind to the membrane not only because the inner groove is occupied, but also because AR1-5 are unable to access membrane-embedded targets in the 'extended' state.

Band 3 clustering is mediated by ankyrin-1 and protein 4.2. The ankyrin-1 complex structure reveals four distinct sites of target recruitment (Fig. 2a). We term these sites the juxtamembrane (AR1-5 binding of RhCE and AQP1), the adaptor-mediated (protein 4.2 recruitment of Band 3-I), the central (AR16-18 binding of Band 3-II) and the distal site (Band 3-III binding, including

interactions of N-terminal peptide). The observation that band 3 associates directly with ankyrin at the distal and central sites is consistent with studies of AR13-24 binding to the cytoplasmic domain of band 3, which identified a high affinity and a low affinity site²⁷. Band 3 has variously been characterized as binding to ankyrin as a bona fide tetramer, or a 'pseudotetramer', in which ankyrin links two band 3 dimers. Our structure is consistent with the latter model. The central site is broadly in agreement with the EPR-based structural model of Kim et al.²⁸, as well as studies showing the importance of loops 175–185 and 63–73 of the band 3 cytoplasmic domain for interaction with ankyrin²⁹. In contrast, the distal site uses an entirely different interface on both ankyrin and the band 3 cytoplasmic domain, and is further stabilized by interactions of the N-terminal peptide of band 3 with the peptide binding groove of ankyrin.

Monomeric GPA binds to the band 3 dimer. One GPA monomer binds to each band 3 protomer, with the GPA transmembrane helix packing with TM3 and TM4 of band 3, and the extracellular N-terminal extension interacting with the TM7-8 loop, which includes E658, the critical residue for formation of the Wright blood group antigen³⁰. H85 of GPA appears to be forming a salt bridge with E658, likely representing the interaction that is variously present and absent in the different Wright blood types. This had previously been proposed to originate from a salt bridge with R61 of GPA³⁰, which is disordered in our structure. A cholesterol-mediated interaction with TM8 of band 3 is also observed, consistent with the known importance of TM8 in the band 3-GPA interaction³¹. Only monomeric GPA is observed binding to band 3, despite the known propensity of GPA to dimerize. This is consistent with studies showing that mutations that inhibit GPA dimerization do not inhibit band 3 dependent effects of GPA³², but still leaves a puzzle: why does the GPA dimer not interact with band 3? Structural alignment of the crystal structure of the GPA dimer³³ with band 3-bound GPA shows that the second molecule of GPA would have to be embedded entirely within the membrane, in an almost horizontal orientation, which is highly energetically unfavorable. We speculate that in the membrane, GPA exists in a monomer-dimer equilibrium and band 3 selectively binds monomeric GPA. In the context of the Class 1 model of the ankyrin-1 complex, the GPA molecules bound to Band 3-II and Band 3-III are in close proximity, although we cannot be certain they are interacting. The possibility of GPA mediating end-to-end attachment of two band 3 dimers would be consistent with both previous molecular dynamics simulations of band 3 in the presence of modeled GPA³⁴, as well as the known propensity of detergent-solubilized band 3 purified from erythrocytes to form 'tetramers' in solution³⁵.

Protein 4.2 facilitates recruitment of Band 3-I and AQP1. The structure of the ankyrin-1 complex also establishes a role for protein 4.2, a highly abundant component of the erythrocyte. Protein 4.2 is known to interact with both band 3 and ankyrin³⁶, but the implication of these interactions was unclear. We observe that protein 4.2 plays a role as an adaptor, stabilizing the interaction of ankyrin with the membrane and facilitating recruitment of an additional dimer of the band 3 anion exchanger (Band 3-I) via interactions with both the cytoplasmic and TM regions, the latter of which appears to be at least partially mediated by a PIP₂ lipid bound at the band 3 transmembrane dimer interface. Furthermore, protein 4.2 appears to play an additional role in facilitating the recruitment of AQP1, the C terminus of which directly interacts with domain 1 of protein 4.2.

Aquaporin-1 is a component of the ankyrin-1 complex. The identification of aquaporin-1 as a component of the ankyrin-1 complex (Fig. 4) was an unexpected finding, although consistent with fluorescence resonance energy transfer measurements indicating that AQP1 is located within 8 nm of band 3 dimers³⁷. The observation

that AQP1 expression is decreased, and glycosylation pattern altered in both murine and human erythrocytes lacking band 3 (ref. ²⁴) is also compatible with a role for AQP1 in the ankyrin-1 complex. It has been proposed that the cluster of membrane proteins that ankyrin-1 organizes in the red blood cell (RBC) membrane forms a CO₂ metabolon³⁸, linking facilitated diffusion of CO₂ across the membrane, anion exchange of Cl⁻/HCO₃⁻ via band 3, and interconversion of dissolved CO₂ and carbonic acid in the cytosol via carbonic anhydrase, which binds to the C-terminal tail of band 3 (ref. 39) and catalyzes conversion of CO₂ and H₂O to HCO₃⁻ and H⁺. A role for AQP1 in the CO₂ metabolon would be consistent with the known capacity of AQP1 to conduct both CO₂ and H₂O (ref. ⁴⁰), both of which are substrates for carbonic anhydrase. RhAG may also participate indirectly in this process, by allowing facilitated diffusion of NH₃ and consequent modulation of [H⁺], via the equilibrium of $NH_3 + H^+$ and NH_4^+ .

RhCE recruits ankyrin-1 to the membrane. The heterotrimeric Rh ammonia channel plays an important role in ammonium homeostasis in the blood⁴¹, and in maintaining the stability of the RBC membrane¹². Humans express five Rh family proteins-RhAG, RhCE and RhD in erythrocytes, and RhBG and RhCG in the kidney epithelium. The structure of the RhCG homotrimer⁴² showed that the eukaryotic Rh family shares a great deal of similarity to the bacterial AmtB family of ammonium transporters, including a conserved twin-histidine motif in the central pore. It has been suggested that RhAG mediates ammonia or possibly carbon dioxide transport^{43,44} across the erythrocyte membrane⁴⁵. The functional role of RhCE and RhD in the structure and function of the RBC membrane has been much debated, as has the stoichiometry of the heterotrimeric Rh complex. The 2.2 Å resolution structure of the (RhAG)₂RhCE heterotrimer (Fig. 5a,b) in the context of the ankyrin-1 complex resolves the stoichiometry question, and suggests that the primary role of RhCE might be as a recruitment site for ankyrin-1, via binding of the N and C termini to AR1-5 of ankyrin-1. Other putative roles for RhCE remain speculative. It has been proposed, based on homology modeling⁴⁶ and the absence in RhCE/D of the characteristic twin-histidine motif, that RhCE and RhD lack a membrane transport function completely. Our work, while not conclusively demonstrating that RhCE retains a transport function, shows that RhCE has a pore that extends almost completely across the membrane, impeded by only by a small cluster of hydrophobic residues and that it seems to be more dynamic than RhAG. We therefore suggest that the possibility of an auxiliary transport function for RhCE in the erythrocyte membrane may be worth investigating.

Concluding remarks. In summary, we have reported the structure of an erythrocyte ankyrin-1 complex, purified from human erythrocytes. We have structurally characterized two direct binding sites for the band 3 anion exchanger on ankyrin-1, as well as one band 3 binding site mediated by protein 4.2, identified aquaporin-1 as a component of the complex, determined the stoichiometry of the heterotrimeric Rhesus ammonia channel and identified RhCE as the ankyrin-1 binding subunit of the Rhesus channel. Taken together these findings offer insights into the molecular architecture of the RBC membrane, and show how ankyrins can facilitate the simultaneous recruitment of multiple classes of membrane proteins, to enable functional organization of membrane transport processes.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/ s41594-022-00792-w. Received: 22 December 2021; Accepted: 24 May 2022; Published online: 14 July 2022

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Methods

Preparation of erythrocyte ghost membranes. Ghost membranes were prepared as described in Niggli et al.⁴⁷. Briefly, RBCs from healthy blood donors were provided by the Transfusion Center of the Hospital of Padua (Italy). Samples were obtained from voluntary and informed blood donation for transfusions. Erythrocytes from fresh human blood samples were washed twice in 5 volumes of 130 mM KCl and 10 mM Tris-HCl, pH 7.4. The cells were hemolyzed in 5 volumes of 1 mM EDTA and 10 mM Tris-HCl, pH 7.4, and centrifuged at 18,000g for 10 min. The ghost membranes were then washed five times in the hemolysis buffer, and four additional times in 10 mM HEPES, pH 7.4. The hemoglobin-free ghost membranes were fi ally resuspended in 130 mM NaCl, 20 mM HEPES, pH 7.4, 0.5 mM MgCl₂, 0.05 mM CaCl₂ and 2 mM DTT, and stored at -80 °C.

Purification of the erythrocyte ankyrin-1 complex. The general purification workflow is presented in Supplementary Fig 1. Ghost membranes were solubilized at a protein concentration of 4 mg ml-1 in 130 mM KCl and 10 mM HEPES, pH 7.4, protease inhibitor tablet (cOmplete, EDTA-free Protease Inhibitor Cocktail, Millipore Sigma), 1% (w/v) digitonin (Carbosynth), 1 mM ATP, 1 mM MgCl₂ and 1 mM PMSF, at 4°C for 1 h. Unsolubilized material was removed by centrifugation at 26,000g for 30 min. The supernatant was applied to a PD10 column (to reduce the detergent concentration) equilibrated with 0.05% (w/v) digitonin, 130 mM KCI, 20 mM HEPES, pH 7.4, 1 mM ATP, 1 mM MgCl, and 1 mM PMSF. The sample was than applied on a glycerol step gradient (30-12% glycerol) and centrifuged for 15h at 25,000 r.p.m. (SW 32 rotor, Beckman). The separation and distribution of proteins was confirmed by SDS-PAGE gel (4-20% Mini-PROTEAN TGX Precast Protein Gels, Biorad). Fractions containing high molecular weight species were pooled together and concentrated to <500 µl using a 100-kDa cutoff concentrator. The sample was loaded on a Superose 6 10/300 Increase column (Cytiva) equilibrated in 0.05% (w/v) digitonin, 130 mM KCI, 20 mM HEPES, pH 7.4, 1 mM ATP, 1 mM MgCl₂ and 1 mM PMSF. Fractions of a peak around an 11 ml elution volume were analyzed using a mass photometer (Refeyn OneMP), pooled and concentrated to 8 mg ml⁻¹.

Vesicle preparation. Ghost membranes (roughly 100 mg) were pelleted (6,000g, 10 min), washed five times in 130 mM KCl and 10 mM HEPES, pH7.4 and then homogenized with the same buffer. The sample was sonicated using microprobe (amplitude 30, pulse on 5s, rest time 10s, total pulse on 50s). Large fragments are removed by centrifugation (6,000g, 10 min) and the supernatant was extruded (Avanti Polar Lipid) ten times using a 400 nm filter. After extrusion, vesicles were collected by ultra-centrifugation at 35,000 r.p.m. for 30 min (S120-AT3, Sorvall). The pellet, which contains vesicles, was resuspended at a final concentration of 5 mg ml⁻¹. All steps were performed at room temperature to facilitate vesicle formation.

Mass photometry (iSCAMS). Mass photometry experiments were performed using a Refeyn OneMP (Refeyn Ltd). Data acquisition was accomplished using AcquireMP (Refeyn Ltd 172 v.2.3). Samples were evaluated with microscope coverslips. The coverslips (cover glass thickness 1.5 24 × 50 mm, Corning) were washed with ddH₂O and isopropanol. Silicone templates were placed on top of the coverslip to form reaction chambers immediately before measurement. The instrument was calibrated using NativeMark Protein Standards (NativeMark Unstained Protein Standard, ThermoFisher). Then 10 µl of fresh room temperature buffer was pipetted into a well, the focal position was identified and locked. For each acquisition, 1 µl of the protein (at a final concentration after dilution of 200 nM) was added to the well and thoroughly mixed. Mass photometry signals were recorded for 60 s to allow detection of at least $2 \times 10^{\circ}$ individual binding events. Data analysis was carried out using the DiscoverMP software.

Cryo-EM and Cryo-ET grid preparation and data collection. Here, 3 µl of purified ankyrin complex at 8 mg ml⁻¹, with a surface-active additive (0.01% (w/v) glycyrrhizic acid) was added to a glow discharged (PELCO easiGlow) holey gold grid (Quantifoil UltrAuFoil grids (R0.6/1, Au 300-mesh gold)) and blotted for 4 to 6 s at 4°C and 100% humidity using the Vitrobot system (ThermoFisher), before plunging immediately into liquid ethane for vitrification. Images were collected on a Titan Krios electron microscope (FEI) equipped with a K3 direct electron detector (Gatan) operating at 0.83 Å per pixel in counting mode using Leginon automated data collection software. Data collection was performed using a dose of $58 e^{-}/Å^2$ across 50 frames (50 ms per frame) at a dose rate of $16 e^{-}$ pixel⁻¹s⁻¹, using a set defocus range of -0.5 to -1.5 µm. A 100-µm objective aperture was used. A total of 14,464 micrographs were collected.

Vesicles for cryo-ET were prepared similarly using 1.2/1.3-µm holey gold grids (Quantifoil UltrAuFoil) and blotted for 10 s using a Vitrobot Mark IV (ThermoFisher), before plunging immediately into liquid ethane for vitrification. About 100 tilt series were collected on a Titan Krios electron microscope (FEI) equipped with a K3 direct electron detector (Gatan), energy filter (Gatan) and Cs-corrector, operating at 2.077 Å per pixel in counting mode using Leginon automated data collection software⁴⁸. Tilt series were collected bidirectionally from [0:+51]° then [0:-51]° in 3° increments, with a dose of $3.13 e^{-/Å^2}$ across eight frames (57 ms per frame) total dose of $112 e^{-/Å^2}$ across 36 total images per tilt series, using a nominal defocus range of -3.5 to -4.5 µm.

Cryo-EM data processing. The final cryo-EM data processing workflow is summarized in Supplementary Fig. 3. Orientation distributions, Fourier shell correlation (FSC) plots and validation statistics are presented in Extended Data Fig. 1c and Table 1. Maps, masks and raw videos have been deposited at EMDB (Table 1) and EMPIAR (IDs EMPIAR-11043). Subsequent steps were performed in cryoSPARC v.3.2–v.3.3 unless otherwise indicated⁴⁹.

Motion correction, contrast transfer function (CTF) estimation and micrograph curation. Patch-based motion correction and dose weighting of 14,464 cryo-EM videos was carried out in cryoSPARC using the Patch Motion job type. Patch-based CTF estimation was performed on the aligned averages using Patch CTF.

Particle picking. An initial round of exploratory processing was performed using Blob Picker. Ranking the micrographs by the number of particles remaining after 2D classification, 58 exemplary micrographs were selected for manual picking to create a training dataset for the Topaz neural network-based picker¹⁶. Then 1,510 particles corresponding to large complexes were manually picked from 58 micrographs. Topaz was run in training mode using a downsampling factor of 6 and an estimated number of particles per micrograph of 200. The resulting model was used to pick particles using Topaz Extract. Initially, 1.8 million particles were extracted in a box of 600 pixels and Fourier cropped to 150 pixels for initial cleanup (2D and initial 3D classification).

2D classification. Multiple rounds of 2D classification were performed to isolate homogeneous subsets of particles to use for ab initio reconstruction of compositionally distinct classes. The 48 most occupied 2D classes from the initial round of 2D classification are presented in Extended Data Fig. 1b.

Initial model generation. Heterogeneous ab initio reconstruction was performed on a variety of subsets of particles selected by 2D classification, to generate a diverse range of initial models representing distinct 3D classes in the data for further classification.

Initial 3D classification. Heterogeneous refinement was performed using the Topaz picked initial particle stack of 1.8 million particles, with 12 representative initial reconstructions as map inputs. Four well-defined, high-resolution classes of the ankyrin complex were identified, as well as several well-defined classes of smaller particles, most of which were identified as free band 3 dimers. Then 710,000 particles corresponding to ankyrin complexes were re-extracted with recentering in a 450 pixel box.

Consensus refinement. An initial refinement of all 710,000 ankyrin complex particles was performed using nonuniform refinement with on-the-fly refinement of per particle defocus, and global refinement of beam tilt and trefoil aberrations, giving an initial consensus refinement with a resolution of 2.3 Å. The use of dynamic masking and real-space windowing of particle images during refinement was disabled. Masked refinements of the Rh trimer, ankyrin-1 membrane binding domain, protein 4.2, the cytoplasmic domains of Band 3-I and the transmembrane domains of Band 3-I were performed starting from the global consensus refinement with local angular searches.

Subclassification of ankyrin complex particles. Based on subclassification of each of the original four classes, six nonredundant classes were identified. The 710,000 stack of clean ankyrin complex particles from the consensus refinement was subjected to heterogenous refinement using these six classes as inputs, as well as five decoy classes, using an initial low-pass filter of 50 Å and a batch size per class of 5,000 particles. The composition and occupancy of the resulting classes is summarized in Supplementary Fig. 5.

Subclassification using 3D-variability analysis. In the case of Band 3-I, Band 3-II, Band 3-III and AQP1, another approach was taken to isolate a set of particles with improved density for the mobile component (Extended Data Fig. 6). First, a local refinement with a mask around the mobile component was performed, yielding a relatively poor resolution, noisy local reconstruction (3.8 Å in the case of Band 3-I starting from the consensus refinement). Then, 3D-variability analysis was carried out as implemented in cryoSPARC 3.2, selecting five modes, with a filter resolution of 3-4.5 Å. Reconstructions were calculated along each mode, and a mode corresponding to an order-disorder transition of the mobile subregion was identified. This mode was then used (using the 3D Variability Analysis Display job type) to split the particles into 20 clusters, and a reconstruction calculated for each. Particles belonging to clusters with well-defined, high-resolution features were combined, realigned globally by nonuniform refinement and subjected to another round of local refinement, In the case of Band 3-I local refinement from the consensus refinement of all ankyrin-1 complexes, this yielded a map with a resolution of 2.8 Å.

Processing of free band 3–GPA dimer particles. In a separate processing pathway, after identification of the free band 3 dimer class using multi class ab initio reconstruction, several successive rounds of heterogeneous refinement were

performed using a single band 3 dimer class and eight random density decoys, against the original 1.8 million particle stack. This allowed isolation of a subset of 37,000 particles that were re-extracted in a 320 pixel box and refined with nonuniform regularization to obtain a reconstruction at 2.8 Å resolution. Training a Topaz model and repicking the dataset with a model biased toward band 3–GPA dimer particles, and resoluted a substantial increase in the number of high-quality particles, and resulted in a 2.3 Å reconstruction from 166,000 particles.

Local refinements and generation of composite maps. For the consensus refinement, class 1 and class 2, local refinements using several different masks were performed to generate local reconstructions with improved density. Maps were post-processed by density modification, soft masking outside the refinement mask and cropping to a minimal box using phenix.resolve_cryo_em (without using an atomic model in the analysis) and resampled to 0.415 Å per pixel using relion_image_handler to best visualize high-resolution structural details. For deposition of each local refinement, the density modified, cropped and resampled map has been designated as the main map, with nondensity modified half maps that have been cropped and resampled to match the map used for model building. Raw half maps, and an optimally B factor sharpened full reconstruction, are provided as additional maps for each deposition. To aid model building of the complex, composite maps of the consensus (all classes combined), Classes 1 and 2 were generated by aligning all the relevant local maps to the matching global reconstruction, and then combining them by taking the maximum value at each voxel using UCSF Chimera⁵⁰. Composite maps were created from both the density modified maps, and the unsharpened, unmasked local maps, and both have been provided in each composite map deposition, along with the global reconstruction (and matching half maps) as additional maps.

Cryo-EM data validation. Map resolution was estimated using the gold-standard FSC = 0.143 criterion, calculated from half maps using a soft mask in Relion v.3.1 (ref. ⁵¹). Detailed validation statistics are provided in Table 1.

Atomic model building and refinement. An initial model for protein 4.2 was generated using trROSETTA⁵². Ankyrin-1 (ref. ⁵³) (Protein Data Bank (PDB) ID 1N11), Band 3 (refs. 54,55) (PDB IDs 4YZF and 1HYN) and aquaporin crystal structures $^{\rm 56}$ (PDB 4CSK) were used as initial models for building the corresponding proteins. Initial models for RhCE and RhAG were generated by threading using CHAINSAW57 as implemented in the CCP4 package58, using the structure of the RhCG homotrimer as a template⁴² (PDB 3HD6). Initial models were placed in corresponding local refinements and and fit as rigid bodies using UCSF Chimera⁵⁰. Each submodel was then manually extended and completed in $\mathrm{COOT}^{\mathrm{59-61}}$, waters and ligands were added where justified by the density and chemical environment, and each final model was refined against the local map using phenix.real_space_refine62. Finally, models were fit to the composite map, merged, then flexibly fit using phenix.real_space_refine. The combined model of class 1a was rigid body fit to the ankyrin-1 complex subtomogram average, after which ligands, waters and side chains were stripped, leaving a polyalanine model. Figures were prepared using UCSF ChimeraX⁶³. Refinement and validation statistics are provided in Table 1.

Cryo-ET data processing. Tilt images were frame aligned with MotionCor2 (ref. ⁶⁴) without patches or dose weighting. Frame aligned tilt images were used for fiducial-less tilt-series alignment in Appion-Protomo^{65–67}. Aligned tilt series were dose weighted in Appion-Protomo using equation 3 in Grant and Grigorieff⁰⁸ reconstructed for initial visualization with Tomo3D (refs. ^{69,70}) SIRT and separately without dose weighting with Warp⁷¹. Seven binned by four Warp tomograms (pixel size 8.308 Å) were used to train an IsoNet⁷² model (where 300 subtomograms were extracted at 963 voxels after masking with *z* axis cropping to remove void volumes, then trained over 35 iterations), which was then applied to each tomogram for visualization with IMOD⁷³.

Cryo-ET subtomogram averaging. Tilt images were frame aligned with MotionCor2 (ref. 64) without patches or dose weighting. The unaligned tilt images were imported into Warp71 after CTF preprocessing using Warp from the frames. The tilt series were aligned with AreTomo⁷⁴ and the alignment files were imported back into Warp, where CTF correction was performed. Tilt series that had poorly modeled CTFs were manually removed. Tomogram reconstruction was performed in Warp at 8.308 Å per pixel. The missing wedge was corrected using IsoNet (as described above) to enhance visualization for particle picking. Twenty missing wedge corrected tomograms were manually picked with crYOLO75 to obtain 10,000 particle coordinates. The particle coordinates were used to train crYOLO on uncorrected tomograms. Prediction was performed on 60 tomograms where a total of 60,029 particles were obtained. The particles were extracted in Warp at 4 Å per pixel with a box size of 80 pixels. Initial models were generated in Relion v.4 using stochastic gradient descent, with eight classes requested. The resulting initial map that most closely resembled the ankyrin-1 complex was low-pass filtered to 60 Å and used for multiple rounds of 3D classification in Relion v.4 (refs. 76,77) with a spherical mask of 300 Å at an angular sampling interval of 7.5°. After 3D classification, 1,596 particles were re-extracted in a 140 pixel box and used for

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refinement in Relion v.4 with a soft spherical mask, giving a final map with a nominal resolution of 25 Å.

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

Data availability

All raw video files, aligned micrographs and particle coordinates (in two STAR files, one for band 3–GPA and one for the ankyrin complexes) are deposited in the EMPIAR (EMPIAR-11043). The cryo-EM density maps and models have been deposited in the Electron Microscopy Data Bank and the PDB. PDB IDs are: 7UZ3, 7UZE, 7UZQ, 7UZS, 7UZU, 7UZV, 7V07, 7V0K, 7V0M, 7V0Q, 7V0S, 7V0T, 7V0U, 7V0X, 7V0Y, 7V19, 8CRQ, 8CRR, 8CRT, 8CS9, 8CSV, 8CSW, 8CSX, 8CSY, 8CT2, 8CT3, 8CSL and 8CTE. EMDB IDs are: EMD-26874, EMD-26886, EMD-26916, EMD-26917, EMD-26918, EMD-26919, EMD-26940, EMD-26943, EMD-26944, EMD-26948, EMD-26949, EMD-26950, EMD-26951, EMD-26952, EMD-26953, EMD-26974, EMD-26955, EMD-26956, EMD-26958, EMD-26960, EMD-26972, EMD-26973, EMD-26974, EMD-26975, EMD-26978, EMD-26979, EMD-26988.

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Author contributions

EV., with the assistance of L.Y.Y., performed protein preparation for structural analysis. E.V. and T.C. prepared membranes from human erythrocytes. F.V. screened and optimized sample vitrification, and generated cryo-EM data. J.D.J. and A.J.N. carried out tomography data analysis. Single-particle cryo-EM data analysis and model building was conducted by O.B.C. The manuscript was written by O.B.C. and F.V. with input from T.C. Figures were prepared by K.K and F.V.

Competing interests

The authors declare no competing interests.

Additional information

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С

0.2 0.0 DC

10A

5Å

3.3A

2.5Å

2Å



b

Extended Data Fig. 1 | Single particle cryo-EM structure determination. a, A representative micrograph out of 14.464 micrographs. **b**, 2D class averages of particles initially picked with Topaz (~1.8 M), ordered by population. 2D class averages of band 3–GPA (red box) and ankyrin-1 complexes (green box) are highlighted. **c**, Fourier shell correlation (FSC) curves. Maps shown as inset, with refinement mask in the case of local refinements.

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Extended Data Fig. 2 | Cryo-EM workflow and analysis of the ankyrin-1 complex. Flowchart outlining cryo-EM image acquisition and processing performed to obtain the structure of ankyrin-1 complex. All processing was performed using CryoSPARC v.3.3 (see Methods for details).

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RhCE (Ser73-Leu98)



RhAG1 (Asp234-Leu260)



RhAG2 (Ser137-Phe163)



Free Band 3 (TM)

(Ser517-Pro548)



Glycophorin A (Glu89-Gly113)



Band 3-I (TM) (Thr570-Lys592)



Band 3-I (Cyt) (Leu128-Glu142)



Band 3-II (Cyt) (Asp277-Tyr299)



Band 3-III (Cyt) (Arg304-Thr324)



Glycophorin B (Ala59-Ala91)



Protein 4.2 (Asn151-Phe184)



Ankyrin-1 (lle69-Tyr166)



Aquaporin-1 (His209-Phe229)

Extended Data Fig. 3 | See next page for caption.

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Extended Data Fig. 3 | Model/map fit. Cryo-EM densities (transparent gray surface) are shown with corresponding segments of the atomic model; sidechains are rendered in stick representation and colored as in Fig. 1. The densities for GPA and free band 3 are derived from the free band3 class. Density for band 3-I TM derives from local refinement of Class 3. For all the other images we used the density from local refinement of the consensus refinement.

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Extended Data Fig. 4 | See next page for caption.

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Extended Data Fig. 4 | Ankyrin-1 complex classes. a, Different views of six main classes of the ankyrin-1 complex. Class 1 contains an (RhAG)₂(RhCE) heterotrimer, ankyrin-1, protein 4.2, and three band 3 dimers (I, II, III). Class 2 exhibits a smaller micelle, retaining the core (RhAG)₂(RhCE)(Ank1)(P4.2) architecture, but including only one band 3 dimer (B3-I), bound to protein 4.2, and aquaporin. Class 3 contains an (RhAG)₂(RhCE) heterotrimer, and a single band 3 (B3-I). Class 4 has a bigger micelle and contains an (RhAG)₂(RhCE) heterotrimer, ankyrin-1, protein 4.2, three band 3 dimers and an unidentified protein "X'. Class 5 contains an (RhAG)₂(RhCE) heterotrimer, ankyrin-1, protein 4.2, aquaporin and Band 3-II and Band 3-III. Class 6 contains an (RhAG)₂(RhCE) heterotrimer, ankyrin-1, protein 4.2, aquaporin and Band 3-II and Band 3-III. Class 6 contains an (RhAG)₂(RhCE) heterotrimer, ankyrin-1, protein 4.2, and Band 3-II. In all the six classes the cytosolic domain of Band 3-II is present. The transmembrane domains of B3-II in classes 2, 3 and 6 appear to be associated with a separate micelle, likely due to either partial dissociation during extraction and/or re-association with free band 3 dimers post-extraction. **b**, Comparison of Class 1 & Class 2. Rh is colored in blue, Band 3-I in orange, Band 3-II in lilac, Band 3-III in yellow and aquaporin in pink. Superposition of Class 1 and Class 2 shows that presence of the additional two band 3 dimers does not exclude the presence of aquaporin.

b



100 nm

2D Classes

Multi-complex 2D slices

С

Tomogram 2D slices

Extended Data Fig. 5 | Cryo-ET of native membrane vesicles. a, A composite slice from the tomogram shown in Supplementary Video 2 with several densities resembling ankyrin-1 complex side-views boxed out. **b**, SPA 2D class side-views for comparison with the tomogram slices. **c**, Magnified tomogram slices from the boxed-out regions in A. The multi-complex 2D slice on the right shows one of many visible strings of ankyrin-1 complex-like densities suggesting potential higher order assemblies. Tomogram slices are 8.3 Å thick. During the process of vesicle formation, vesicles can form in either the native or inverted configurations, with the cytosolic side of the membrane oriented in either direction.

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Extended Data Fig. 6 | Cryo-EM workflow for sub-classification of Band 3-I from the consensus refinement. Flowchart outlining cryo-EM processing performed to improve the structure of Band 3-I (orange) and the bound GPA proteins (magenta). All processing was performed using CryoSPARC v.3.3. A similar procedure was used to improve the density of AQP1 starting from Class 2.

а

b





С













е





Extended Data Fig. 7 | See next page for caption.

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Extended Data Fig. 7 | Conformation of membrane-associated ankyrin-1. a, Ankyrin structure shown as a molecular surface with the inner groove depicted in pink, the convex outer surface in red, in yellow the N-terminal peptide of Band 3-III and in magenta the C-terminal linker of ankyrin-1. **b**, Structural alignment between ankyrin-1 structure (colored by RMSD) and ankyrin-2 (in gray). The first 5 repeats of Ank1 are dramatically rearranged compared to their position in the Ank2 structure. Arrows indicate AR5 and 6. **c**, Structural alignment between ankyrin-2 (gray) and ankyrin-2 bound to Nav1.2 peptide (colored by RMSD), shows that flexibility at this interface also occurs in ankyrin-2. **d**, Hydrophobic surface calculated for the first 5 ankyrin repeats (AR1-5) of Ank1 (top) and Ank2 (bottom). RhCE N-terminal (magenta) and C-terminal fragments (purple) are displayed in stick representation. In the lower figure Ank2 is bound to Nav1.2 peptide (blue). **e**, Sequence alignment of all the 24 ankyrin repeats of Ank1 using MUSCLE⁸² and visualized and colored using Jalview with the ClustalX color scheme. The sequences for AR5 and 6 are outlined in red. **f**, Sequence of AR5 and AR6 from Ank1, Ank2 and Ank3, aligned using MUSCLE⁸² and visualized and colored using Jalview with the clustalX colored using Jalview with the clustalX colored using Jalview with the clustalX colored using Jalview in the region that rearranges in Ank1. AR5 is shorter than the other repeats by 4 residues, and AR6 lacks both the "S/TP' motif at positions 13-14 is replaced by two aspartates.



Extended Data Fig. 8 | See next page for caption.

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Extended Data Fig. 8 | Ankyrin interaction with Band 3-III N-terminal peptide. a, Map at low density threshold allows tracing of the complete Band 3-III N-terminal peptide (dotted yellow line), that runs back along the inner ankyrin groove. **b-c**, Molecular surface of the ankyrin repeats, colored by hydrophobicity, bound to the N-terminal peptide of Band 3-III (yellow). Residues 2-24 of Band 3-III form an ordered interaction with AR6-10 and the AR5-6 linker. Key residues are displayed as sticks. Band 3-III is shown as yellow ribbons. **d**, Structural comparison between ankyrin-1 (top) and ankyrin-2 structures (PDB: 4Y4D) (bottom). Structure of ankyrin-1 (dark red AR1-5 and red AR6-12) is associated with N-terminal peptide of Band 3-I (yellow) and ankyrin-2 (dark blue AR1-6 and light blue AR6-12) is associated with the autoinhibitory domain (AI) of ankyrin-1. **e**, Close up of the AR5-6 of ankyrin-1 in panel S8d. The Band 3-I part colored in green corresponds to the first 11 N-terminal amino acids. **f**, Structural alignment between the first five ankyrin repeats (AR1-5) of ankyrin-1 (dark red) bound to RhCE peptide and ankyrin-2 (dark blue) bound to the autoinhibitory peptide of ankyrin-1. The key conserved residues that mediate the interactions between ANK1 and B3-III and ANK2 and the autoinhibitory region are shown as sticks.

Band 3-I

Protein 4.2

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Extended Data Fig. 9 | Flexibility of band 3 cytosolic domain. a, Models of the three Band 3 dimers (I, II, III). The three cytosolic domains of Band 3 proteins in the complex have different orientations. The Band 3-I cytoplasmic domain is inverted with respect to those of Band 3-II and Band 3-III. The TM part of Band 3 is colored gray. **b**, Two different views of band 3 cytosolic domain represented as molecular surface. Interacting surfaces of band 3 are distinct and non-overlapping. Regions of the band 3 cytosolic domain that interact with proteins in the complex are shown in different colors: green for the region that interact with protein 4.2, red for ankyrin repeats 22-24 and purple for ankyrin repeats 17-19. **c**, Comparison between Band 3-I (yellow) associated with protein 4.2 (green) and Band 3-II (lilac). The model is displayed as ribbon and the surface is shown in transparency. In the right panel a close view of the interactions between Band 3-II cytosolic domain (lilac) and protein 4.2 (green); the two share the same interaction site with the TM part. Key residues for the interaction are displayed as sticks.

Band 3-II

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RhCE-RhAG2 interface

d

f

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RhAG

g



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h













Extended Data Fig. 10 | See next page for caption.

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Extended Data Fig. 10 | (RhAG)₂(**RhCE) trimer interactions. a**, Model of AQP1 with waters (red dots fit inside the map). In the bottom inset a close view of the waters in the central cavity. **b**, Unmodeled density at the interface of aquaporin-1 (salmon) with RhCE (purple) and RhAG (light blue). Potential interacting sidechains are depicted in stick representation. **c**, The structure of (RhAG)₂(RhCE) trimer as viewed from the cytoplasm, RhCE is colored in purple, RhAG1 in light blue and to RhAG2 in dark blue. The cryo-EM density map of the (RhAG)₂(RhCE) trimer is shown in transparency. The 12 TM helixes are numbered (0-11) for each Rh molecule. Gray circles shown the pore position in each subunit. **d**, Interfaces between the three different subunits, with key residues represented as sticks. **e**, Interface between the three Rh protomers. The key residues that mediate the interactions in the interfaces are shown as sticks. Three different parts of the trimerization interface are show in the right figure viewed from the top. **f**, Analysis of the pores of RhCE and RhAG. Figure of the pore in green generated by HOLE. 2D graph of pore radius for RhCE (top graph) and RhAG (bottom graph). **h**, Close up of RhAG2, the map shows an undefine density. The key residues that interact with the undefined density are shown as sticks. **g**, Structure of (RhAG)₂(RhCE) complex and the first five repeats of ankyrin. The amino acids depicted in yellow sticks highlight the sites of variation between RhCE and RhD. Three sites of variation between RhD and RhCE are shown in the right panels, with the density map overlayed. The density map is consistent with the presence of RhCE, not RhD.

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Data collection	Leginon 3.5
Data analysis	For Cryo-EM data processing the following software were used: Appion 3.4, cryoSPARC 3.2-3.3, Topaz 0.2.4 For model building , the following software were used: Coot 0.9.6, Phenix 1.19, Chimera 1.15 , ChimeraX 1.3, HOLE v2.2.005 For sequence alignment the following software were used: Muscle 3.8, Jalview 2.11.1.4 For mass photometry: AcquireMP v2.3, DiscoverMP v2.3 For cryoET data collection and processing: MotionCor2, Appion-Protomo v1.2.2, Tomo3D v2.0, Warp v1.0.9, IMOD v4.11, AreTomo v1.0.12, crYOLO v1.8, Relion v4.0

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