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Structure of hepcidin-bound ferroportin reveals iron homeostatic mechanisms

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Author Contributions

C.B.B. purified FPN and Fab fragments, established biochemical approaches to reconstitute a ferroportin-hepcidin complex, established and performed hepcidin binding and calcein transport assays, prepared samples for cryo-EM, screened samples by negative stain and cryo-EM, and determined the X-ray crystal structure of Fab45D8. C.M.A. identified optimal freezing conditions for FPN samples, collected cryo-EM data, and determined high resolution cryo-EM maps by extensive image processing. R.C.K. and A.S.P. performed and analysed molecular dynamics simulations under the guidance of R.O.D. S.G. collected negative stain and cryo-EM data for FPN samples bound to different Fabs, and identified Fab45D8 as a fiducial marker for structure determination. S.S. aided in establishing and performing the calcein transport assay. T.A. provided fluorescently labeled hepcidin and anti-FPN antibodies for structure determination, A.M. built and refined models of FPN and Fab45D8. The manuscript was written by C.B.B., C.M.A., Y.C., and A.M. with revisions provided by all authors. The overall project was supervised by Y.C. and A.M.

Tara Arvedson is employed by Amgen and reports Amgen stock. The other authors declare no competing interests.

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Abstract

The serum iron level in humans is tightly controlled by the action of the hormone hepcidin on the iron efflux transporter ferroportin. Hepcidin regulates iron absorption and recycling by inducing ferroportin internalization and degradation¹. Aberrant ferroportin activity can lead to diseases of iron overload, like hemochromatosis, or iron limitation anemias². Here, we determined cryogenic electron microscopy (cryo-EM) structures of ferroportin in lipid nanodiscs, both in the apo state and in complex with cobalt, an iron mimetic, and hepcidin. These structures and accompanying molecular dynamics simulations identify two metal binding sites within the N- and C-domains of ferroportin. Hepcidin binds ferroportin in an outward-open conformation and completely occludes the iron efflux pathway to inhibit transport. The carboxy-terminus of hepcidin directly contacts the divalent metal in the ferroportin C-domain. We further show that hepcidin binding to ferroportin is coupled to iron binding, with an 80-fold increase in hepcidin affinity in the presence of iron. These results suggest a model for hepcidin regulation of ferroportin, where only iron loaded ferroportin molecules are targeted for degradation. More broadly, our structural and functional insights are likely to enable more targeted manipulation of the hepcidin-ferroportin axis in disorders of iron homeostasis.

Introduction

Iron is essential for life. Despite this central role in biology, free ferrous iron (Fe^{2+}) is toxic. In excess, iron can catalyze the production of free radicals, leading to cellular damage. Iron levels are therefore tightly controlled, both at the cellular and organism level.

In mammals, iron levels are regulated by the action of hepcidin, a peptide hormone, on ferroportin (FPN), the only known iron efflux transporter^{2–4} (Fig. 1a). FPN mediates absorption of dietary iron by transporting ferrous iron across the basolateral surface of intestinal enterocytes. FPN also enables iron recycling from hepatocytes and macrophages⁵. Synthesis of FPN is transcriptionally regulated by cellular hypoxia, iron and heme concentrations, and inflammatory signaling⁶. In settings of elevated serum iron levels, liver-derived hepcidin negatively regulates cell surface FPN by acutely blocking iron transport⁷ and inducing FPN ubiquitination, internalization, and degradation^{1,8–10}. Therefore, hepcidin decreases serum iron levels by suppressing FPN-mediated dietary iron absorption and release of iron from cellular stores.

Iron disorders in humans can result from dysregulation of hepcidin or FPN. Deficits in hepcidin-mediated regulation of FPN, often due to hereditary hemochromatoses, lead to iron overload and widespread tissue damage affecting the liver, heart, pancreas, and joints^{11–13}. By contrast, inappropriate elevation of hepcidin levels yields iron-restricted anemia^{14,15}.

Although several approaches to restore aberrant FPN function have been evaluated in clinical trials ^{16–18}, none have thus far succeeded.

The molecular mechanism of FPN regulation by hepcidin remains incompletely defined. A confluence of human genetics studies and structure-function evaluations have identified key regions of FPN important in hepcidin regulation^{7,19–22}. To understand how FPN transports iron, and how this process is regulated by hepcidin, we used a combination of cryogenic electron-microscopy (cryo-EM), molecular dynamics simulations, and *in vitro* biochemical assays. These studies reveal the molecular recognition of iron and hepcidin by FPN, demonstrate acute inhibition of iron transport by hepcidin, and suggest a new regulatory mechanism enabling hepcidin to selectively target actively transporting ferroportin molecules for degradation.

Structures of apo and hepcidin-bound FPN

We screened the antigen-binding fragments (Fabs) of antibodies previously raised against FPN²³ for use as a fiducial mark to guide image alignment of a small membrane protein embedded in a lipid nanodisc. This approach leverages a strategy we previously proposed for structure determination by single particle cryo-EM²⁴. Fab45D8 was previously determined to be non-competitive with hepcidin and does not induce FPN internalization²³. Indeed, Fab45D8 did not alter the binding properties of hepcidin to nanodisc-reconstituted FPN. (Extended Data Fig. 1).

We determined cryo-EM structures of FPN reconstituted in a NW9 nanodisc 25 and bound to Fab45D8, both in the apo state (3.2 Å, Extended Data Table 1, Extended Data Fig. 3) and bound to hepcidin and Co^{2+} (2.5 Å, Extended Data Fig. 4). We reconstituted FPN metal transport in a liposome-based calcein quenching assay, which yielded a V_{max} of -0.00269 ΔFs^{-1} and K_m of 13.6 μM for FeCl $_2$ (Fig. 1b, Extended Data Fig. 2). Unlike Fe $^{2+}$, Co^{2+} is not readily oxidized and therefore provides a tractable surrogate divalent metal for FPN biochemical and structural studies. $CoCl_2$ was also robustly transported by FPN with a V_{max} of -0.00339 ΔFs^{-1} and K_m of 33.5 μM , supporting its use as a substrate mimetic. Fab45D8 decreased both the V_{max} and K_m of $CoCl_2$ transport (Extended Data Fig. 2), suggesting that Fab45D8 prefers a subset of conformations within the FPN ensemble, but does not inhibit transitions required for metal transport.

The cryo-EM density map of FPN enabled building of an atomic model of FPN regions important for iron transport and hepcidin binding^{2,7}, and a portion of the intracellular loop 3 (ICL3) important in hepcidin-induced FPN internalization^{8–10}(Extended Data Fig. 5). The FPN extracellular loop 5 (ECL5) remains unresolved, likely due to conformational flexibility. We separately obtained an X-ray crystal structure of Fab45D8 at 2.1 Å to enable modeling (Extended Data Table 2).

Both cryo-EM structures reveal a monomeric FPN bound to a single Fab45D8 molecule, which binds FPN extracellular loop 2 (ECL2) (Fig. 1b, Extended Data Fig. 6). Similar to other MFS transporters, FPN contains twelve transmembrane (TM) helices arranged in two domains (Fig. 2a), with a large central cavity open to the extracellular side in both structures

(Fig. 2b). Ferroportin shares structural similarity with a previously characterized bacterial homolog, the bbFPN transporter²⁶ (Extended Data Fig. 6). Unlike most other MFS transporters, the FPN TM7 is interrupted by a short non-helical stretch of six residues. This unique feature, previously posited to be important in iron binding²⁷, is shared between FPN and bbFPN. Similar to bbFPN, FPN metal transport is stimulated by transmembrane pH gradients, but addition of valinomycin had no significant effect (Extended Data Fig. 7).

Several interacting residues define an intracellular gate that keeps the N- and C- domains of FPN in an outward open conformation. Similar to an interaction in bbFPN²⁶, R489 in TM11 of the C-domain forms an ionic interaction with D157 in TM4 of the N-domain (Fig. 2b), which is further supported by an extended ionic and hydrogen-bonding network including residues E486 (TM11) and R88 (TM3). An additional cluster of ionic and hydrogen bonding interactions between TM5 in the N-domain and TM10 in the C-domain of FPN further stabilizes the outward open conformation (Fig. 2c). Mutation of several residues within the intracellular gate leads to FPN loss of function in ferroportin disease, highlighting the importance of the gate in coordinating the conformational steps necessary for iron efflux^{19,28}.

Iron binds to the N and C domains of FPN

Cryo-EM maps from a Co^{2+} -hepcidin-FPN complex revealed two new densities in the central cavity of FPN, corresponding to single metal binding sites in the N and in the C domain respectively (Fig. 3a–c). Within the C domain, Co^{2+} directly interacts with C326 in TM7b and H507 in TM11 while making a water-mediated contact with D325 and the backbone carbonyl of T320 (Fig. 3b). Intriguingly, the tetrahedral coordination geometry for Co^{2+} is fulfilled by the carboxy terminus of hepcidin. Within the N domain, we observe density for Co^{2+} coordinated directly by TM1 residues D39 and H43 (Fig. 3c). Mutation of D39 to alanine or glutamate abolishes iron transport^{22,27}, which was previously ascribed to disruption of a putative calcium binding site within the N domain of FPN²⁷. Our structure, by contrast, suggests that D39 directly binds the substrate. Transport assays with reconstituted FPN, however, revealed clear modulatory effects of calcium (Extended Data Fig. 7). We observed a modest decrease in both K_m and V_{max} for Co^{2+} with intra-liposomal $CaCl_2$, whereas the rate of transport was increased by ~40% for Fe^{2+} .

The identification of two metal binding sites within FPN raises the question of whether both sites are required for iron efflux. Disease causing mutations and experimental mutagenesis studies suggest that the N domain metal site is critical for iron efflux, while the C domain site may not be required for transport. Several hereditary hemochromatosis mutations in FPN directly coordinate Co²⁺ in the C domain, including C326S/F/Y^{13,29–31} and H507R³². These mutations are predicted to disrupt the precise tetrahedral coordination geometry required for metal binding within the C domain. However, these mutants are fully capable of iron transport, and lead to iron overload even in the presence of hepcidin. While mutation of the C domain residue D325 leads to decreased iron efflux²², recent modeling studies of an inward open conformation of FPN suggests that D325 may interact with the N domain in the outward open state; loss of iron efflux in D325 mutants may therefore be caused by disruption of the extracellular gate³³. By contrast, mutation of D39 to alanine in the N

domain metal binding site of FPN completely abolishes iron efflux from HEK293 cells²², suggesting that the N domain may be a primary site for effluxed iron. Because the C domain metal-binding site is important for hepcidin binding, it may primarily serve an important iron-dependent regulatory function in hepcidin control of FPN activity.

Hepcidin binding may influence interactions between FPN and the metal. To understand divalent metal binding to FPN in the absence of hepcidin, we used all-atom molecular dynamics simulations. We first performed six simulations of apo-FPN in a hydrated lipid bilayer with Fe²⁺ ions initially positioned randomly in bulk solvent. In all six independent simulations, Fe²⁺ ions bound spontaneously to the C domain, localizing to the unwound region of TM7 near residues D325, D504, and H507 within hundreds of nanoseconds (Fig. 3d and Extended Data Fig. 8). The Fe²⁺ ion also occasionally moves closer to TM1 to interact with D39 in the N domain. In parallel simulations of apo-FPN without Fe²⁺ ions, we observed mobility of TM7b with significant fluctuations of D325 (Fig. 3e), consistent with comparatively weaker cryo-EM density for TM7b as compared to other transmembrane helices in apo-FPN (Extended Data Fig. 8). By contrast, both D325 and TM7b are less mobile in simulations with iron bound at the C domain site (Fig. 3e). Divalent metal binding to the C domain may therefore stabilize an otherwise dynamic TM7b in a conformation that favors hepcidin binding.

Hepcidin occludes outward open FPN

Hepcidin binds FPN in a central cavity between the N and C domains, acting as a molecular cork to occlude the iron efflux pathway (Fig. 4a). Consistent with acute inhibition of iron transport by hepcidin⁷, we observed a decrease in Co^{2+} transport by reconstituted FPN when hepcidin is included within the liposome. By contrast, hepcidin added to the external solution had minimal effect on metal transport (Fig. 4b), suggesting that the calcein assay primarily measures transport in the forward direction (cellular efflux). Indeed, constitutively inhibited transport of a liposome-reconstituted hepcidin-FPN complex could be rescued by addition of membrane permeable β -mercaptoethanol, which likely denatures internally bound hepcidin by reducing its disulfide bonds (Fig. 4c).

Although the conformations of apo- and Co²⁺-hepcidin-bound FPN are highly similar (RMSD 0.73 Å), the Co²⁺-hepcidin-bound structure shows a ~3 Å rigid body separation of the N and C domains on the extracellular side of FPN (Fig. 4d). Within the N domain, hepcidin binding leads to displacement of TM2 from the central cavity, driven in part by a specific contact between I6 of hepcidin and T61 and Y64. Within the C domain, the largest conformational changes occur around the Co²⁺ binding site, leading to changes in the conformation of TM7b and the extracellular side of TM11.

Hepcidin makes extensive polar and hydrophobic contacts with FPN with a total buried surface area of ~1300 Å². Our structure of hepcidin bound to FPN provides insight into disease-causing mutations associated with FPN gain of function in hereditary hemochromatoses. Several FPN mutations decrease hepcidin binding to FPN, including N144H/D/T^{34–36}, C326S/F/Y^{13,29–31}, Y333H³⁷, Y501C^{7,38}, D504N^{7,39} and H507R³². Of these, C326 and H507 directly coordinate the cobalt ion and mutations therefore indirectly

affect hepcidin affinity or alter the atomic basis of binding specificity (Fig. 4e). Other interactions between FPN and hepcidin are either hydrophobic or depend on hepcidin amide backbone atoms, which is consistent with the relatively high tolerance of amino acid substitutions within hepcidin⁴⁰. For example, Y333 fits into a hydrophobic cavity in hepcidin and hydrogen bonds with the backbone carbonyl of hepcidin residue M21 (Fig. 4e). D504 in FPN coordinates the backbone amide of hepcidin H3 while Y501 π -stacks with the imidazole side chain of H3. A further hydrogen bond between N144 and Y501 further stabilizes this interaction network. In contrast to these mutations, the Y64N/H mutants retain hepcidin binding⁷, but are completely resistant to hepcidin-induced FPN ubiquitination^{41,42}. The outward displacement of TM2 near Y64 induced by hepcidin may therefore be important for FPN ubiquitination (Fig. 4d).

Hepcidin is coupled to iron binding

The direct interaction between hepcidin and Co^{2+} in our structure is unexpected (Fig. 4f), and suggests that divalent metals may be important for hepcidin binding to FPN. We therefore directly tested the effect of Fe^{2+} and Co^{2+} on hepcidin affinity at FPN. A fluorescently tagged version of hepcidin labeled at position 17 (Rhodamine greenhepcidin²³, RhoG-Hep, Fig. 4e) bound to nanodisc-reconstituted ferroportin with an apparent K_D of 210 nM (p K_D = -6.67 ± 0.02) (Fig. 4g). In the presence of 10 μ M FeCl₂, we observed a significantly increased affinity of 2.5 nM (p K_D = -8.61 ± 0.21), an almost 80-fold change in the potency of hepcidin at FPN. Addition of $CoCl_2$ also increased hepcidin affinity. By contrast, $CaCl_2$ did not support high affinity hepcidin binding (Extended Data Fig. 7), potentially due to an inability to form a direct interaction bridging FPN and hepcidin. Consistent with a cooperative effect, the effect of $CoCl_2$ on hepcidin binding was saturable (Extended Data Fig. 7).

The reference range for hepcidin concentration in healthy adults is $\sim 1-30$ nM⁴³. Our *in vitro* binding experiment with purified FPN indicates minimal hepcidin binding to FPN in the absence of a divalent metal, suggesting that metal binding to FPN may regulate hepcidin activity *in vivo*. Our structure of FPN bound to Co²⁺ and hepcidin revealed conformational changes in TM7b associated with hepcidin binding (Fig. 3e) and a direct contact between hepcidin and Co²⁺(Fig. 3b); both could be important for the observed effect of Fe²⁺ and Co²⁺ on hepcidin binding. We therefore tested whether disruption of the C domain iron binding site influences hepcidin binding. Even in the presence of 50 μ M CoCl₂, the D325N, C326S, and H507R mutants bound hepcidin weakly, titrating in a micromolar range similar to wild-type FPN in the absence of divalent metals (Fig. 4h). These results highlight the critical role of the C domain metal site in potent hepcidin binding to FPN, which is likely important in homeostatic control of iron levels in a physiological setting.

Discussion

Ferroportin is a central regulator of iron homeostasis in humans. Our structural and biochemical studies here reveal that hepcidin binding to FPN is greatly potentiated by iron itself, potentially due to the stabilizing effect iron has on the hepcidin-binding site of FPN (Fig. 4i). With iron, the binding affinity of hepcidin falls in a range concordant with the

concentration of hepcidin observed in healthy human adults. In normal iron homeostasis, this may enable hepcidin to selectively bind and regulate FPN molecules actively transporting iron and loaded with Fe²⁺, while sparing FPN molecules located on cells with low transport activity. Hepcidin binding to FPN would both trap the transporter in an outward open state and limit iron egress; both actions acutely decrease iron efflux, as has been recently reported⁷. Elevated hepcidin levels likely inappropriately overcome this regulatory strategy and degrade FPN even in the absence of active iron efflux. The potentiation of hepcidin activity by iron may therefore have immediate consequences for the development of hepcidin mimetics currently in clinical trials¹⁶. Furthermore, hepcidin antagonism by direct targeting of FPN may require molecules with high potency to overcome the nanomolar effect of the hormone in the presence of iron. The structural and functional insights into FPN function presented here therefore provide critical foundations for the discovery of therapeutics for human disorders of iron homeostasis.

METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

Expression and purification of human ferroportin

The wild-type human FPN gene was cloned into a pVL1392 vector containing an expression cassette comprised of a C terminal human rhinovirus 3C (HRV-3C) protease recognition sequence followed by a human protein C epitope tag (EDQVDPRLIDGK) and an 8x polyhistidine tag. Baculovirus was generated using Spodoptera frugiperda Sf9 insect cells (unauthenticated and untested for mycoplasma contamination, Expression Systems 94-001F) and the construct was expressed in Spodoptera frugiperda Sf9 insect cells. Cells were collected 48 h after transduction and stored at -80°C until further use. Frozen cell pellets were thawed and washed with a hypotonic buffer (20 mM HEPES pH 7.50, 1 mM EDTA, supplemented with 20 µg/mL leupeptin, and 160 µg/mL benzamidine) before solubilizing with 50 mM HEPES pH 7.5, 300 mM NaCl, 1% (w/v) n-dodecyl-β-D-maltopyranoside (DDM, Anatrace), 0.1% (w/v) cholesteryl hemisuccinate (CHS, Steraloids), 1 mM EDTA, and supplemented with 20 µg/mL leupeptin, and 160 µg/mL benzamidine for 1 h at 4°C. Following centrifugation, the resulting supernatant was loaded on homemade anti-protein C antibody Sepharose beads and washed extensively in 50 mM HEPES pH 7.50, 300 mM NaCl, 2 mM CaCl₂, 0.1% (w/v) DDM, 0.01% (w/v) CHS. FPN was eluted with 50 mM HEPES pH 7.50, 300 mM NaCl, 0.1% (w/v) DDM, 0.01% (w/v) CHS, 0.2 mg/mL Protein C peptide (Genscript), and 5 mM EDTA. The protein was concentrated with a Vivaspin 100kDa MWCO concentrator and the monomeric FPN fraction was collected after sizeexclusion chromatography (SEC) over a Superdex S200 Increase 10/300 GL column (GE Healthcare) equilibrated with 20 mM HEPES pH 7.50, 100 mM NaCl, 0.1% (w/v) DDM, and 0.01% (w/v) CHS.

FPN mutants used in binding studies were generated using the Q5 site-directed mutagenesis method (NEB) and cloned, along the WT FPN construct, into a pEZT vector containing an

expression cassette comprised of a C terminal human rhinovirus 3C (HRV-3C) protease recognition sequence followed by an 8x polyhistidine tag and a 1D4 epitope tag (TETQSVAPA). The resulting vectors were transfected into Expi293F Human Embryonic Kidney cells (Life Technologies) with the Expifectamine transfection kit (Life Technologies) using 1 µg DNA per mL culture as per the manufacturer's instructions. Cultures were harvested 48 h post transfection and stored at -80° C until further use. FPN mutants were purified as described for WT, except the cleared solubilized supernatant was loaded on Ni-NTA Sepharose beads and washed extensively in 50 mM HEPES pH 7.50, 300 mM NaCl, 0.1% (w/v) DDM, 0.01% (w/v) CHS, 30 mM Imidazole. FPN was eluted with 50 mM HEPES pH 7.50, 300 mM NaCl, 0.1% (w/v) DDM, 0.01% (w/v) CHS, 250 ml Imidazole. The protein was concentrated and purified by SEC as described above.

Expression and purification of MSP

Constructs encoding MSP-NW9 or MSP-NW11²⁵ in the pET28b vector (Addgene #133442) were transformed into BL21(DE3) Rosetta Escherichia coli, and grown in terrific broth medium supplemented with 2 mM MgCl₂ and 0.1% (w/v) glucose at 37°C. At OD₆₀₀ of ~0.6, expression was induced by addition of 400 μM isopropyl β-d-1-thiogalactopyranoside (IPTG) and lowering the temperature to 20°C. Cells were harvested after 16 hours and resuspended into 5 mL lysis buffer (200 mM Tris pH 8.0, 500 mM NaCl, 1% (v/v) Triton X-100 (Sigma), 0.02 mg/mL leupeptin, 0.16 mg/mL benzamidine, and benzonase) per gram pellet. After stirring for 30 min at 4°C, cells were lysed by pulsed sonication on ice. The lysate was cleared by centrifugation at 15,000 × g for 25 min at 4°C and loaded on Ni-NTA Sepharose. Ni-NTA beads were washed with 50 mM Tris pH 8.0, 500 mM NaCl, 1% (v/v) Triton, then 50 mM Tris pH 8.0, 500 mM NaCl, 50 mM sodium cholate, then 50 mM Tris pH 8.0, 500 mM NaCl, and finally with 50 mM Tris pH 8.0, 500 mM NaCl, 30 mM Imidazole. MSP was eluted with 50 mM Tris-HCl pH 8.0, 500 mM NaCl, 400 mM Imidazole and dialyzed into 50 mM Tris-HCl, pH 8.0, 100 mM NaCl, 1 mM EDTA, 0.1 mM TCEP at 4°C. The following day, MSP was concentrated on a Vivaspin 10-kDa MWCO concentrator, aliquots were flash frozen in liquid nitrogen and stored at -80°C for reconstitution.

Isolation, expression and purification of Fab45D8

The heavy and light chain sequences of mAb45D8²³ were separately cloned into pcDNA3.4 and the resulting vectors were transfected into Expi293F Human Embryonic Kidney cells (unauthenticated and untested for mycoplasma contamination, Life Technologies) using a 1:2 mass ratio of light and heavy chain DNA with the Expifectamine transfection kit (Life Technologies) as per the manufacturer's instructions. Supernatant containing mAb45 was harvested 136 h after transfection and loaded on homemade Protein G Sepharose beads and extensively washed with a buffer comprising 20 mM HEPES pH 7.50, and 100 mM NaCl. mAb45D8 was eluted with 100 mM glycine (pH 3.0) and fractions were immediately neutralized with 200 mM HEPES pH 7.50. To generate the Fab fragment, 10 mg of purified mAb45D8.1 was diluted into 9.5 ml freshly prepared cleavage buffer (20 mM sodium phosphate pH 7.00, 10 mM EDTA, and 10 mM cysteine) and treated with 0.5 ml agarose immobilized papain (Thermo Scientific) at 37°C. After 16 h the cleaved Fab45D8 fragment was purified by reverse Protein A affinity chromatography, followed by SEC into buffer

comprising 20 mM HEPES pH 7.50 and 100 mM NaCl. Fab45D8 was concentrated on a Vivaspin 10-kDa MWCO concentrator, and aliquots were flash frozen in liquid nitrogen and stored at -80° C for later use.

Reconstitution of FPN into lipidic nanodisc

Purified FPN (0.2–0.5 mg) was mixed with purified MSP and a lipid mixture containing a 2:3 weight ratio of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC, Avanti) and 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(1'-rac-glycerol) (POPG, Avanti). For reconstitution into NW9 nanodiscs, an FPN:MSP:Lipid molar ratio of 1:20:1100 was used. For reconstitution into NW11 nanodiscs, an FPN:MSP:Lipid molar ratio of 1:20:800 was used. The reconstitution sample was nutated for 1 h at 4°C before addition of 0.2 g/mL SM2-BioBeads (BioRad), and the reconstitution sample was further nutated overnight at 4°C before removal of the biobeads. FPN containing nanodiscs for cryo-EM studies were purified by loading the reconstitution sample on anti-protein C antibody Sepharose beads and washing extensively with 20 mM HEPES pH 7.50, 100 mM NaCl, and 1 mM CaCl₂ to remove empty nanodiscs. FPN containing nanodiscs were eluted with 20 mM HEPES pH 7.50, 100 mM NaCl, 0.25 mM EDTA, and 0.2 mg/mL Protein C peptide (Genscript) and concentrated on a Vivaspin 50-kDa concentrator. FPN containing nanodiscs for binding studies were purified by SEC over a Superdex S200 Increase 10/300 GL column (GE Healthcare) equilibrated with 20 mM HEPES pH 7.50, 100 mM NaCl.

Crystallization and structure determination of Fab45D8

Purified Fab45D8 was diluted to 13.0 mg/mL in 20 mM HEPES pH 7.5, 100 mM NaCl. Fab45D8 crystals were obtained in 0.3 M trimethylamine-N-oxide (TMAO), 0.1 M Tris pH 8.5, and 30% (w/v) PEG 2000 MME at 20 °C. Individual crystals were flash frozen in liquid nitrogen after a 30 s soak in 0.3 M trimethylamine-N-oxide (TMAO), 0.1 M Tris pH 8.5, and 30% (w/v) PEG 2000, and 20% v/v ethylene glycol. A full diffraction dataset was collected at the Advanced Photon Source GM/CA-CAT beamline 23ID-B, and processed using xia2dials⁴⁴ implementation of XDS⁴⁵. The structure of the Fab was solved by molecular replacement using Phaser⁴⁶, with a search model of a closely related germline mouse monoclonal antibody (PDB ID: 6BZV⁴⁷) lacking complementarity determining regions (CDRs). The model was iteratively improved by refinement in Coot⁴⁸ and Phenix⁴⁹ Data collection and refinement statistics are summarized in Extended Data Table 2. The final model contained 96.77%, 2.23% and 0% in the favored, allowed and outlier regions of the Ramachandran plot, respectively as assessed by MolProbity⁵⁰.

Calcein transport assay for divalent cations

FPN was reconstituted into liposomes for divalent cation transport assays. Empty liposomes were prepared as a 3:1 mass ratio of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (POPE, Avanti) to POPG dissolved in chloroform, followed by gentle evaporation of the chloroform under a stream of nitrogen gas, and overnight desiccation. The lipids were dissolved in 20 mM HEPES pH 7.40, 100 mM KCl to a final concentration of 12.5 mg/mL, sonicated until optically clear, subjected to multiple freeze-thaw cycles, and extruded through a 400 nm polycarbonate filter (Avestin) to generate unilamellar vesicles. Subsequently 0.13% (w/v) Triton-X100 (Sigma) was added to destabilize liposomes,

corresponding to the onset of solubilization as determined by turbidity measurements at 540 nm. Purified FPN was added at a 1:50 protein to lipid mass ratio and incubated for 15 min at 4°C. The FPN-Hepcidin complex was reconstituted by incubating 1µM purified FPN with 3 μM Hepcidin (Bachem) and 3μM CoCl₂ for 30 min on ice before addition of liposomes. The FPN-Fab complex was reconstituted by incubating 1μM purified FPN with 2 μM purified Fab45D8 for 30 min on ice before addition of liposomes. Control liposomes devoid of FPN were prepared in parallel using the same concentration of DDM. To remove excess detergent, 0.05 g/mL of SM2-BioBeads were added to the sample and nutated for 1 hr at 4°C, then 0.05 g/mL SM2-BioBeads were added followed by incubation overnight at 4°C, and finally addition of 0.08 g/mL SM2-BioBeads followed by incubation for 2 hr at 4°C. Liposomes were harvested by ultracentrifugation at 264,500 × g for 30 min and resuspended at a concentration of 4.0 mg/mL lipids in internal buffer comprised of 100 mM KCl and 20 mM HEPES pH 7.40 or pH 7.50, before flash freezing in liquid nitrogen and storage at -80°C. On the day of the transport assay, liposomes were thawed and diluted to 2 mg/mL with internal buffer containing 250 μM calcein (Sigma), and where indicated 5 μM hepcidin (Bachem) or 1.25 mM CaCl₂ was added. The samples were then subjected to three freezethaw cycles, and extruded through a 400 nm polycarbonate filter. The calcein containing liposomes were concentrated and washed four times by repeated ultracentrifugation and resuspension in external buffer containing 100 mM NaCl and (i) 20 mM HEPES pH 6.80, pH 7.40, pH 7.50, or pH 8.20, or (ii) 20 mM Tris/MES pH 6.00, pH 7.50 or pH 9.00. Reconstitution yields determined by the Micro BCA protein assay kit (Pierce) were 205–222 pmol/mg lipid for FPN samples, and 275 pmol/mg lipid for the FPN-Fab45D8 complex assuming 1:1 stoichiometric incorporation. Presence of FPN and Fab were further verified by SDS-PAGE. Immediately prior to the assay, liposomes were diluted to 0.125 – 0.25 mg/mL lipid in external buffer. Time-course fluorescence traces were recorded as 1-2 s integrations using a FluoroMax-4 (Horiba) or a FluoroLog (Spex) with λ_{ex} of 490 nm and λ_{em} of 520 nm. Steady state fluorescence was recorded for at least 2 min, before addition of small aliquots of freshly prepared stocks of either FeCl2, CoCl2, MnCl2, NiCl2, CuCl2, or ZnCl₂. To stabilize the ferrous (Fe²⁺) state, we prepared iron as a 1:10 ratio of sodium ascorbate:FeCl₂ immediately prior to the experiment. A negative inside membrane potential was generated by incubating liposomes with 100 nM of the K⁺-selective ionophore valinomycin (Sigma) for 1 min prior to the experiment. To determine the full extent of the calcein quenching response, 10 µM of the divalent cation ionophore calcimycin (Sigma) was added at the end of the experiment. Transport data was normalized to the mean baseline fluorescence intensity prior to addition of ion, and initial rates of transport were obtained from linear regression and fit to the Michaelis-Menten function in GraphPad Prism 8 (GraphPad Software). Statistical significance was determined using student's t-test. For clarity, mixing artifacts ± 5 sec from the time of addition have been removed from representative fluorescence traces.

Hepcidin binding assays

Fluorescence polarization measurements were performed using rhodamine-green labeled hepcidin labeled at position 17 (RhoG-hepcidin) 23 . For FPN saturation binding experiments, samples were prepared in a black 384-well plate (Greiner) containing 0–1 μ M of nanodisc reconstituted NW11-FPN and 5 nM RhoG-hepcidin in sample buffer comprising 20 mM

HEPES pH 7.50, 100 mM NaCl, and supplemented with FeCl₂, CoCl₂ or MnCl₂ as indicated. For ion stimulation experiments, 100 nM NW11-FPN and 5 nM RhoG-Hepcidin was mixed with 0–600 μ M of CoCl₂. For Fab binding experiments, 100 nM NW11-FPN and 5 nM RhoG-Hepcidin was mixed with 0 – 3 μ M of Fab45D8 in sample buffer containing 10 μ M CoCl₂. Binding reactions were equilibrated for 60 min at RT, and fluorescence polarization was recorded on a Biotek Synergy H4 (Agilent) in polarization mode using fixed bandpass filters with λ_{ex} of 484 nm and λ_{em} of 520 nm. Binding data was processed in GraphPad Prism 8 (GraphPad Software) using 3- or 4-parameter curve fitting.

Analytical fluorescence size exclusion chromatography (FSEC) was performed by mixing 25 µg of NW11-FPN with 2 × fold molar excess of RhoG-Hepcidin in sample buffer comprised of 20 mM HEPES (pH 7.50), 100 mM NaCl and 10 µM CoCl2. Samples were incubated for 20 min on ice and 1.5 × molar excess of Fab45D8, or sample buffer, was added followed by incubation for 30 min on ice. For homologous competition, 1 µM NW11-FPN was mixed with 30 µM unlabelled hepcidin (Bachem) in sample buffer comprised of 20 mM HEPES (pH 7.50), 100 mM NaCl and 10 µM CoCl, and incubated for 30 min on ice. Then 2 µM RhoG-Hepcidin was added and the sample incubated for 30 min on ice. Samples were injected on a Superdex 200 Increase 10/300 GL column (GE Lifesciences) preequilibrated in 20 mM HEPES pH 7.50, 100 mM NaCl, and 10 µM CoCl2. RhoG-Hepcidin fluorescence was recorded using an FP-1520 Intelligent Fluorescence Detector (Jasco) with λ_{ex} of 493 nm and λ_{em} of 524 nm.

Cryo-EM Sample Preparation and Data Collection

Nanodisc-reconstituted apo-FPN was mixed with 1.15 molar excess of Fab45D8 and incubated on ice for 30 min. The complex was purified by size-exclusion chromatography over a Superdex S200 Increase 10/300 GL column (GE Healthcare) equilibrated with 20 mM HEPES pH 7.50, 100 mM NaCl. For Co^{2+} /hepcidin samples, 600 μ M CoCl₂ and 30 μ M hepcidin (Bachem) were added to nanodisc-reconstituted FPN and incubated for 20 minutes on ice prior to addition of Fab45D8. The resulting complex was purified over size-exclusion chromatography as for the apo sample but with the addition of 100 μ M CoCl₂ in the chromatography buffer. Collected fractions were supplemented with fresh hepcidin to 30 μ M. For both preparations, fractions containing the nanodisc-FPN-Fab45D8 complex were concentrated to ~3 mg/ml on a Vivaspin 50-kDa MWCO concentrator and freshly used for electron microscopy.

For high-resolution cryo-EM, the apo-FPN-Fab45D8 complex was diluted to 0.0375~mg/mL in 20 mM HEPES pH 7.5, 100~mM NaCl directly prior to vitrification, and $2~\mu\text{L}$ sample was applied to glow-discharged gold holey carbon 1.2/1.3~300-mesh grids (Quantifoil) coated inhouse with graphene oxide 51-53. Grids were blotted for 2-4 seconds at 0~force and 10~seconds wait time before being plunge vitrified in liquid ethane using a MarkIV Vitrobot (ThermoFisher). The blotting chamber was maintained at 22°C and 100% humidity during freezing.

 $Co^{2+}/hepcidin$ samples were diluted to 1.5 mg/mL in gel filtration buffer (20 mM HEPES pH 7.5, 100 mM NaCl, 100 μM CoCl $_2$) before vitrification on UltrAufoil 1.2/1.3 300 mesh grids. Grids were blotted for 3 seconds at 0 force and 5 seconds wait time before plunge

vitrification in liquid ethane using a MarkIV Vitrobot (ThermoFisher). The blotting chamber was maintained at 22°C and 100% humidity during freezing.

FPN-Fab45D8 and Co2+/hepcidin-FPN-Fab45D8 movies were collected using a Titan Krios (ThermoFisher) outfitted with a K3 camera and Bioquantum energy filter (Gatan). The K3 detector was operated in superresolution mode and the energy filter slit width was set to 20 eV. Movies were collected at a nominal magnification of 105,000x, physical pixel size 0.834Å/pix, with a 70 μ m C2 aperture and 100 μ m objective aperture at a dose rate of 8 e⁻/pixel/second. A total dose of 66 e⁻/Ų was collected as a 120-frame movie, resulting in a 6-second movie with 0.55 e⁻/frame. Data were collected using semi-automated imaging scripts in SerialEM⁵⁴. For FPN-Fab45D8, 5009 movies were collected using a 3×3 image shift pattern at 0° tilt and 406 movies were collected on-axis with a 30° stage tilt in two separate data collection sessions. For Co2+/hepcidin-FPN-Fab45D8, 4395 movies were collected at 0° tilt in a single data collection session.

Cryo-EM Image Processing

For FPN-Fab45D8, data were motion corrected and 2x binned on-the-fly using MotionCor2⁵⁵ in the SCIPION pipeline⁵⁶. Motion corrected micrographs were imported into cryoSPARC⁵⁷ and RELION⁵⁸ and contrast transfer function parameters were calculated using CTFFIND4⁵⁹. CTF information for tilted images were estimated using patch CTF estimation in cryoSPARC. 138,314 particles were selected from 497 micrographs using the Blob picker in cryoSPARC. 2D class averages were generated after extracting the putative particles with a 300-pixel box and binning to 64 pixels. Six of these averages were used as templates for further particle picking. Template picking yielded 4,737,795 particles. These were split into 6 groups to increase speed of processing, extracted in a 300-pixel box, and binned to 64 pixels. 2D classification was run in cryoSPARC with default settings except: number of 2D classes 200, initial classification uncertainty factor 4, number of online-EM iterations 40, batch size per class 300. Objectively "good" (showing clear Fab and receptor density) class averages were selected and exported to RELION format using csparc.py⁶⁰. Class averages that were not classified as "good", but were not clearly ice contamination or graphene oxide edges, were run through a second round of 2D classification with default settings except: number of 2D classes 200, number of online-EM iterations 40, batch size per class 300. All "saved" class averages from the second rounds of 2D classification in cryoSPARC were selected and exported to RELION format using csparc.py. Particles were extracted from CTF-corrected images in RELION at a box size of 300 pixels and binned to 128 pixels. 1,326,130 particles, in three groups (detailed in Extended Data Fig. 3) were classified in 3D with image alignment in RELION using: an initial model generated in cryoSPARC from 80,000 particles collected on a Talos Arctica filtered to 40 Å, C1 symmetry, a regularization parameter of 4, for 30-35 iterations with no mask. Particles from classes with resolved transmembrane (TM) helices were selected, extracted in a 300-pixel box, and imported back into cryoSPARC. Non-uniform refinement was run with default settings and no resolution limit, resulting in angle and shift assignments for 850,000 particles. These particles were subsequently exported to RELION format using csparc.py and run through 3D classification without image alignment. Four of the 12 classes were selected, imported into cryoSPARC, run through non-uniform alignment with an

automatically generated mask, and refined to a reported global resolution of 3.2 Å. The resulting map showed signs of mild preferred orientation (Extended Data Fig. 3). The particles were exported into RELION format using csparc.py, converted into an image stack, and imported into cisTEM⁶¹ as a refinement package. The particles were reconstructed and half-maps were generated using the "generate 3D" command. These half-maps, as well as the half-maps from cryoSPARC were run through our lab's directional Fourier shell correlation (dFSC) program⁶² clearly showing a more evenly weighted range of views in the map generated by cisTEM. Maps were sharpened in RELION. Average resolutions are reported using the FSC = 0.143 cut-off63 and were estimated in cryoSPARC and cisTEM. Local resolution estimates were performed through cryoSPARC, using a similar algorithm to blocres⁶⁴, with an automatically generated kernel size of 24 voxels.

For Co²⁺/hepcidin-FPN-Fab45D8, data were motion corrected and 2x binned on-the-fly using MotionCor2⁵⁵ in the SCIPION pipeline⁵⁶. Motion corrected micrographs were imported into cryoSPARC⁵⁷ and contrast transfer function parameters were calculated using patch CTF estimation through cryoSPARC⁵⁹. 3,753,516 particles were selected from 4395 micrographs using the template picker in cryoSPARC with 2D averages from the apo-FPN dataset as templates. These particles were extracted in a 360-pixel box, and binned to 64 pixels. 2D classification was run in cryoSPARC with default settings except: number of 2D classes 200, initial classification uncertainty factor 4, number of online-EM iterations 40, batch size per class 300. Objectively "good" (showing clear Fab and receptor density) class averages were selected for 3D classification. Class averages that were not classified as "good", but were clearly not ice contamination, were run through a second round of 2D classification with default settings except: number of 2D classes 200, number of online-EM iterations 40, batch size per class 300. All "saved" class averages from the second rounds of 2D classification in cryoSPARC were sent to 3D classification. These particles were subjected to two rounds of heterogeneous refinement in cryoSPARC that serves as a "trash collector". Four initial models were used, three generated from an early round of ab initio model generation and our final apo-FPN-Fab45D8 structure. All initial models were filtered to 30Å before refinement. Particles were unbinned and a final heterogeneous refinement was performed with three good initial models of apo-FPN-Fab45D8. Non-uniform refinement was run with default settings and no resolution limit, on the most populated class, showing clear density for hepcidin, resulting in angle and shift assignments for 310,647 particles. The particles were exported into RELION format using csparc.py, converted into an image stack, and imported into cisTEM⁶¹ as a refinement package. The particles were reconstructed and half-maps were generated using the "generate 3D" command. These half-maps, as well as the half-maps from cryoSPARC were run through our lab's directional Fourier shell correlation (dFSC) program⁶² clearly showing a more distributed range of views in the map generated by cisTEM. Maps were sharpened in cisTEM. Average resolutions are reported using the FSC = 0.143 cut-off⁶³ and were estimated in cryoSPARC and cisTEM. Local resolution estimates were performed through cryoSPARC with an automatically generated kernel size of 18 voxels. Another job was run with a kernel size of 20 voxels and results were similar.

Model building and refinement

For apo-FPN, a homology model of human FPN in the outward open state was built using Modeller⁶⁵, with a previously determined X-ray crystal structure of outward open bbFPN (PDB ID: 5AYN)²⁶ as a template. After truncating putatively flexible regions (N-terminus, ICL3, ECL5, and C-terminus), the resulting model was fit into the 3.2 Å cryo-EM map of FPN:Fab45D8 using Chimera⁶⁶. The initial template was manually rebuilt in Coot⁴⁸ and iteratively refined with real space refinement implemented in Phenix⁴⁹ Model geometry was assessed using MolProbity⁵⁰. Modeling of FPN-Co²⁺-hepcidin complex used the apo-FPN structure as a starting point, with iterative improvements in Coot and Phenix. We used a previously determined X-ray crystal structure of hepcidin bound to a neutralizing Fab as a starting model²¹ for the peptide. Further validation was performed with EMRinger⁶⁷ to compare the map and final model. Map-to-model FSCs were calculated within Phenix. Figures were prepared in Chimera⁶⁶ and PyMol.

Molecular dynamics simulations

The structure of the outward-open apo conformation of human FPN was used as the starting coordinates for all simulations. Three different conditions were simulated (Supplementary Table 1): (1) the iron-absent condition, where no iron was added; (2) the iron-bound condition, where an Fe²⁺ ion was placed in the proposed iron binding site 5.6 Å from D325 α -carbon, 6.9 Å from D504 α -carbon, and 7.8 Å from H507 α -carbon; (3) the iron-in-bulk-solvent condition, where 15 Fe²⁺ ions were placed randomly in the water box outside the protein using Dabble⁶⁸.

Simulation coordinates were prepared by removing non-FPN molecules from the initial structure. Prime (Schrödinger) was used to model missing side chains, and neutral acetyl and methylamide groups were added to cap protein termini. The unresolved loops between TM6-TM7 and TM9-TM10 (residues 239–290 and 394–451 respectively, inclusively) were not modeled. The termini surrounding these loops were capped. PropKa was used to determine the dominant protonation state of all titratable residues at pH 7^{69,70}. The structure was internally hydrated using Dowser⁷¹. Dabble was used to additionally fill the extracellular cavity⁶⁸. The structure was aligned using the Orientation of Proteins in Membranes (OPM) server⁷².

Using Dabble, the protein was inserted into a pre-equilibrated 1-palmitoyl-2-oleoylphosphatidylcholine (POPC) membrane bilayer. For all simulations except condition three (iron in bulk solvent), sodium and chloride ions were added at 150 mM to neutralize the system. For condition three, chloride ions were added to neutralize the system resulting in a concentration of 108 mM. A periodic box was used with dimensions 90×90 Å in the x-y plane and a water buffer of 10 Å above and below the protein to the periodic boundary. We used the CHARMM36m parameters for lipids, proteins, sodium and chloride ions, and the TIP3P model for waters $^{73-75}$. The Fe²⁺ Lennard-Jones parameters were obtained from Li *et al.* 's compromise model ⁷⁶.

All simulations were run on a single Graphical Processing Unit (GPU) using the Amber18 Compute Unified Device Architecture (CUDA) version of particle-mesh Ewald molecular

dynamics (PMEMD)^{77,78}. For each condition, 6 independent simulations were run. For each independent simulation, the system was minimized with 500 steps of steepest descent followed by 500 steps of conjugate gradient descent three times. 10 and 5 kcal mol⁻¹ -Å² harmonic restraints were used on the protein, lipid, and Fe²⁺ ions for the first and second minimization, respectively. 1 kcal mol $^{-1}$ Å $^{-2}$ harmonic restraints were used on the protein and Fe²⁺ ions for the final minimization. The system was then heated from 0 K to 100 K over 12.5 ps in the NVP ensemble with a Langevin thermostat and harmonically restraining the protein heavy atoms and Fe²⁺ ions with a restraint of 10 kcal mol⁻¹ \mathring{A}^{-2} . The system was further heated with the same restraints from 100 K to 310 K in the NPT ensemble over 125 ps. The system was equilibrated with harmonic restraints on protein heavy atoms and Fe²⁺ ions for 30 ns. The restraint strength started at 5 kcal mol⁻¹ Å⁻² and was reduced by 1 kcal $\text{mol}^{-1} \text{ Å}^{-2} \text{ every 2 ns for the first 10 ns and then by 0.1 kcal } \text{mol}^{-1} \text{ Å}^{-2} \text{ every 2 ns for the}$ final 20 ns. Production simulations were performed at 310 K and 1 bar using the NPT ensemble, a Langevin thermostat and a Monte Carlo barostat. Every 200 ps snapshots were saved. All simulations were run for at least 2.2 µs. These simulations used a 4-fs time step with hydrogen mass repartitioning⁷⁹. Bond lengths to hydrogen atoms were constrained using SHAKE^{79,80}. Non-bonded interactions were cut off at 9 Å.

Simulation Analysis Methods

Molecular dynamics simulation snapshots were reimaged every 1 ns and centered using CPPTRAJ package in AmberTools18⁸¹. Simulations were visualized using Visual Molecular Dynamics and figures prepared in PyMOL⁸². Time traces from simulation were smoothed using a moving average with a window size of 15 ns unless otherwise indicated and visualized with the PyPlot package from Matplotlib. For all analysis in the manuscript that required structural alignment, we aligned to the initial Ferroportin structure using the backbone atoms of residues 26–116, 127–228, 308–483, and 492–543.

To investigate the localization of Fe²⁺ ions, the iron-in-bulk-solvent simulations (condition 3) were analyzed. To visualize the density of Fe²⁺ ions under a given simulation condition, the position of Fe²⁺ ions was recorded every 10 ns for each of the 6 simulations, each 2 μ s in length. Each Fe²⁺ ion position was then drawn as a point superimposed on the starting structure (Fig. 3d). To quantify the binding events, the distance between iron and the closest side chain oxygen atom on D325 was measured. This distance was graphed over 1.2 μ s, including the equilibration time (Fig. 3d, Extended Data Fig. 8b).

To investigate the dynamics and conformation of the TM7b region, the iron-absent (condition 1) and iron-bound (condition 2) simulations were compared. The TM1-TM7b distance was measured using distance between the $C\alpha$ of V51 and the $C\alpha$ of Y333. For each simulation, we calculated the average of the distance over 2.2 μ s, excluding equilibration. The average over the simulations for each condition was plotted with error bars representing the standard error of the mean (s.e.m.) (Extended Data Fig. 8g). The dynamics of D325 were also investigated. The dynamics were visualized by overlaying representative frames showing the movement of D325, the binding site, and TM7b. For iron-absent simulations, frames from a single replicate at 200, 350, 500, and 550 ns were overlaid (Extended Data Fig. 8d). For iron-bound simulations, frames from a single simulation at 200, 500, 725, 1000

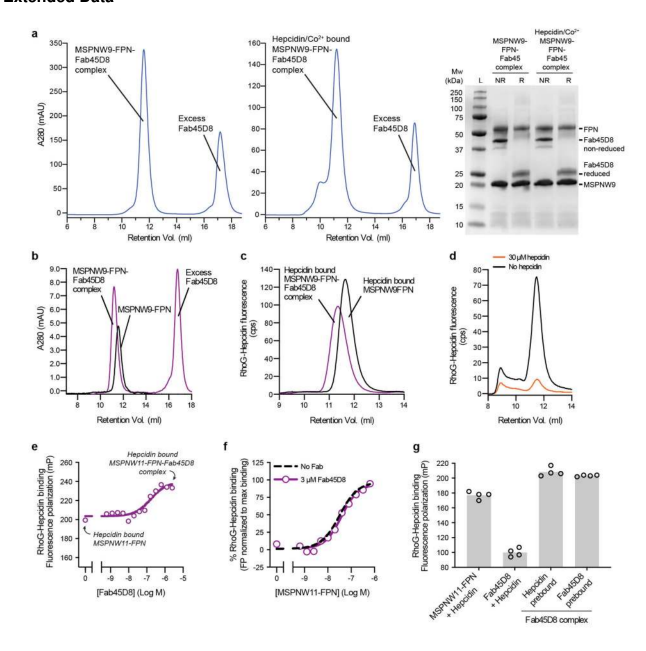
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ns were overlaid (Extended Data Fig. 8e). The conformational range of D325 was quantified by measuring the distance between $C\gamma$ of D325 and $C\beta$ of S47. This was visualized for one simulation for each condition over a time of 1 μs inclusive of equilibration (Extended Data Fig. 8g). For each independent simulation, the mean of the $C\gamma$ D325 - $C\beta$ S47 distance was calculated over 2.2 μs . For each condition, the average over the simulations was plotted with error bars representing the s.e.m. (Extended Data Fig. 8g). The flexibility of D325 was quantified by calculating the root-mean-square fluctuation (RMSF) of the side-chain atoms of D325 using an in-house script (Extended Data Fig. 8g). Statistical significance was determined using the Mann-Whitney U test (two-tailed).

Data Availability

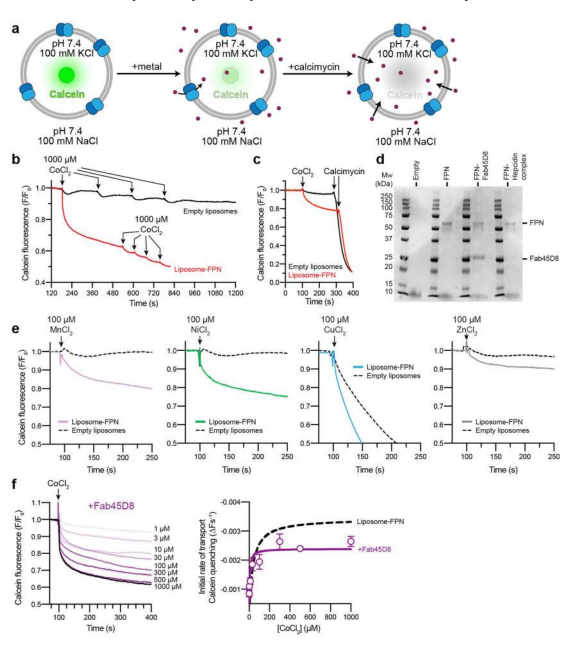
All data generated or analyzed during this study are included in this published article and its Supplementary Information. Crystallographic coordinates and structure factors for the Fab45D8 complex have been deposited in the Protein Data Bank under accession code 6W4V. Coordinates for Fab45D8-FPN complex have been deposited in the Protein Data Bank under accession code 6W4S and the maps have been deposited in the Electron Microscopy Data Bank under accession code 21539. Coordinates for the FPN-Co²⁺-hepcidin-Fab45D8 complex have been deposited in the Protein Data Bank under accession code 6WBV and the maps have been deposited in the Electron Microscopy Data Bank under accession code 21599.

Extended Data



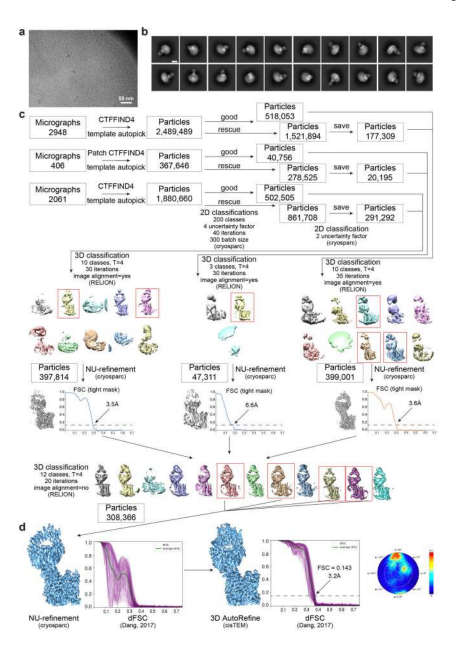
Extended Data Fig. 1 | **. Biochemistry of purified human ferroportin and Fab45D8 complex. a,** Size exclusion chromatography of purified apo and hepcidin/Co²⁺ bound FPN-Fab45D8 complex. SDS-PAGE gel of purified complex under non-reducing (NR) and reducing (R) conditions. SEC and SDS-PAGE analyses are representative and were performed four and two times for apo and hepcidin/Co²⁺ bound FPN-Fab45D8 samples respectively. **b,** Fab45D8 binds nanodisc-reconstituted FPN (nanodisc-FPN), as assessed by size-exclusion chromatography. **c,** Fluorescence size exclusion chromatography (FSEC) of rhodamine green hepcidin (RhoG-hepcidin). RhoG-hepcidin co-elutes with nanodisc-FPN and with a nanodisc-FPN:Fab45D8 complex in the presence of CoCl₂. **d,** FSEC shows RhoG-hepcidin binding to nanodisc-FPN is competed by excess unlabelled hepcidin used for structural

studies. **e**, Fluorescence polarization of RhoG-hepcidin increases further with Fab45D8, consistent with formation of a larger complex. Importantly, Fab45D8 does not decrease RhoG-hepcidin binding. Data points are means from n = 3 technical replicates. **f**, RhoG-hepcidin binding to nanodisc-FPN in the presence of $10 \,\mu\text{M}$ CoCl₂ measured by fluorescence polarization in the absence ($K_D = 31 \,\text{nM}$) and presence of $3 \,\mu\text{M}$ Fab45D8 ($K_D = 39 \,\text{nM}$). Data points are means from n = 3 technical replicates. **g**, RhoG-hepcidin binds to FPN (FPN+hepcidin) but not to Fab45D8 alone (Fab45D8+hepcidin). Order of hepcidin and Fab45D8 addition does not influence the increase in fluorescence polarization for a Fab45D8-FPN-hepcidin complex. Data points are means from n = 4 technical replicates.



Extended Data Fig. $2\mid$. Calcein quenching assay for FPN metal transport.

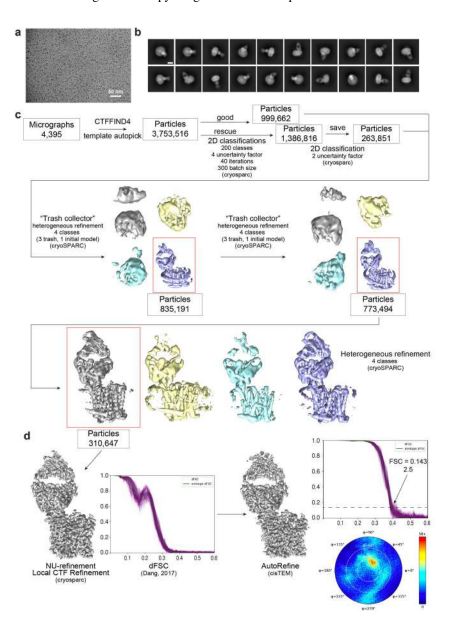
a, Schematic of calcein-based assay to measure transport of divalent cations. Purified FPN is reconstituted into proteoliposomes containing calcein. Transport of divalent cations leads to quenching of calcein fluorescence, which can also occur through the divalent cation selective ionophore calcimycin. b and c, Specificity of the calcein transport assay. FPN reconstituted liposomes facilitate uptake of 1000 µM Co²⁺ added (arrows, red trace) to the external bath solution, producing a calcein quenching response that follows a single exponential. Further additions of Co²⁺ demonstrates the transport process has reached saturation. Calcein fluorescence from empty liposomes (black trace) returns to baseline after multiple additions of 1000 μM Co²⁺. c, Additions of Co²⁺ followed by calcimycin results in rapid and almost complete quenching. Fluorescence traces are representative experiments. d, Liposome reconstitution samples analyzed by reducing SDS-PAGE, the analysis is representative and was performed three times for FPN samples and twice for FPN samples containing FPN-Hepcidin or FPN-Fab45D8. e, Calcein transport assay queried against a panel of putative transition metal ion substrates at 100 µM. Notably, the larger ions Ni²⁺ and Zn²⁺ are also reported FPN substrates⁸³. The smaller ion Mn²⁺ was previously found not to be effluxed by FPN in oocytes⁸³, but appears to be transported in liposomes. Cu²⁺ interferes with empty and FPN-containing liposomes, leading to non-specific quenching of calcein. Fluorescence traces are representative experiments. f, Fab45D8 decreases cobalt transport Vmax $(2.40 \times 10^{-3} \pm 0.075 \times 10^{-3} \Delta F s^{-1})$ and Km $(4.68 \pm 0.54 \mu M)$. Data points are means \pm s.e.m. from n = 3 independent experiments.



Extended Data Fig. 3 |. Cryo-EM data processing for FPN-Fab45D8.

a, Representative motion-corrected micrograph collected on the Titan Krios showing monodisperse FPN-Fab45D8 nanodisc particles. Four different FPN-Fab45D8 samples were imaged on GO-coated Quantifoil grids times with similar results. **b,** Examples of "good" 2D class averages that were used in 3D classification. Similar quality class averages were produced when processing six unique subsets of the final particles used. Scale bar is 50 Å. **c,** Flowchart showing image processing pipeline for FPN-Fab45D8. Initial processing, through 2D classification, was performed in cryoSPARC. Particles were then transferred, using csparc2star.py, to RELION for 3D classification, then to cryoSPARC for a nonuniform refinement, and finally to cisTEM for refinement. The number of particles moving into each

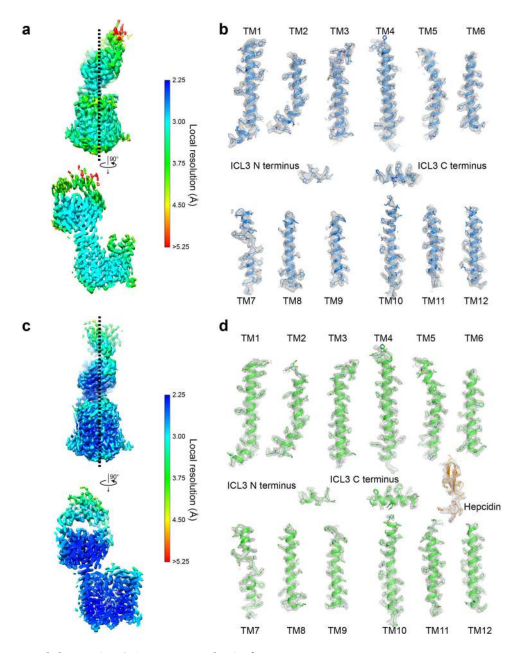
step are noted. **d**, Final refinements from cryosparc and cisTEM beside their directional FSC curves calculated using dfsc.0.0.1.py. Angular distribution plot from cisTEM is shown.



Extended Data Fig. 4 |. Cryo-EM data processing for Co²⁺-hepcidin-FPN-Fab45D8.

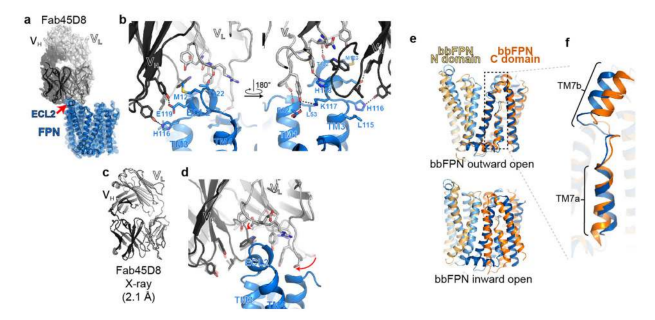
a, Representative motion-corrected micrograph collected on the Titan Krios showing monodisperse Co²⁺-hepcidin-FPN-Fab45D8 nanodisc particles. Two indepedent Co²⁺-hepcidin-FPN-Fab45D8 samples were frozen on UltrAufoil grids with similar micrograph quality. **b,** Examples of "good" 2D class averages that were used in 3D classification. **c,** Flowchart showing image processing pipeline for Co²⁺-hepcidinF-PN-Fab45D8. Initial processing was performed in cryoSPARC. Local CTF refinement, with a conservative high-resolution limit of 4 Å, was performed before nonuniform refinement using default

parameters. A final subset of particles was transferred to cisTEM for refinement. The number of particles moving into each step are noted. **d,** Final refinements from cryosparc and cisTEM beside their directional FSC curves calculated using dfsc.0.0.1.py. Angular distribution plot from cisTEM is also shown.



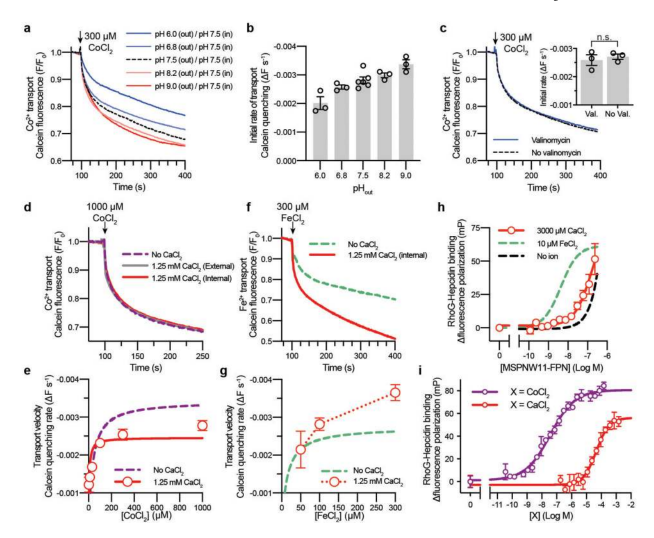
Extended Data Fig. 5 |. Cryo-EM map density for FPN.

Local resolution estimation in cryoSPARC for the apo-FPN-Fab45D8 complex (**a**) and for the Co2+-hepcidin-FPN-Fab45D8 complex (**c**). Two views of each complex are shown. Shown in grey mesh is cryo-EM map density for individual FPN transmembrane helices for the apo-FPN-Fab45D8 complex (**b**) and for the Co²⁺-hepcidin-FPN-Fab45D8 complex (**d**). Mesh depicts density within a 2.5 Å radius of any modeled atom.



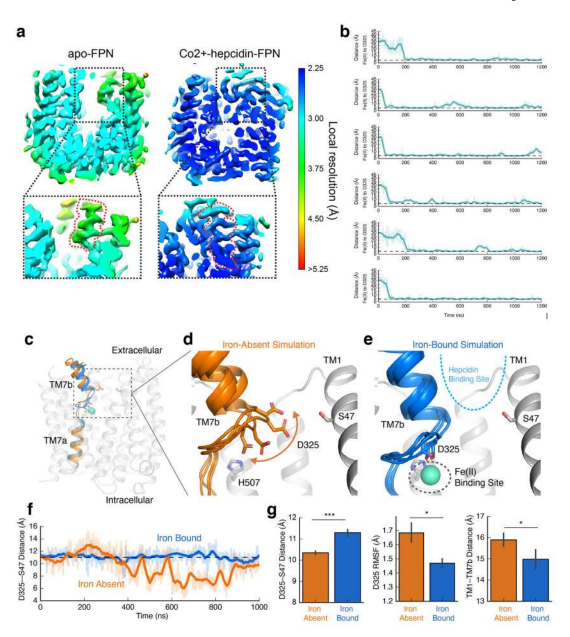
Extended Data Fig. 6 |. Structure of Fab45D8 and comparison of FPN with bbFPN.

a, Cryo-EM density of FPN and Fab45D8. Although density is observed for the constant regions of Fab45D8, it is not of sufficient quality to unambiguously model. **b,** Fab45D8 makes extensive contacts with FPN extracellular loop 2 (ECL2) with both the heavy (V_H) and light (V_L) chains. **c,** Crystal structure of Fab45D8 at 2.1 Å. **d,** Comparison of Fab45D8 alone (transparent cartoon and sticks) and bound to FPN. The binding site residues of Fab45D8 change minimally upon binding FPN. **e,** Human FPN aligned to the outward-open (PDB: 5AYN) and inward-open (PDB: 5AYO) conformations of bbFPN. **f,** Unique architecture of TM7 shared between human FPN and bbFPN.



Extended Data Fig. 7 |. Effect of pH and ions on metal transport and binding. a and b, Co²⁺ transport by FPN is potentiated by an opposing H⁺-gradient. a, Time-course calcein fluorescence quenching response after addition of 300 µM CoCl₂ (arrow) for FPNliposomes containing 100 mM KCl and HEPES, pH 7.50 that were diluted into buffer comprised of 100 mM NaCl and (i) HEPES pH 6.8, (ii) HEPES pH 7.5, (iii) HEPES pH 8.2, (iv) Tris/MES pH 6.0, or (v) Tris/MES pH 9.0. b, initial rates of transport obtained from the linear phase of time-course fluorescence quenching experiments at the indicated pH values. Data points are means \pm s.e.m. from n = 3 experiments, c. Calcein fluorescence responses for FPN-liposomes containing 100 mM KCl and HEPES, pH 7.50 diluted into 100 mM NaCl and HEPES, pH 7.50. A membrane potential ($\Delta \Psi$) was generated by addition of the K + selective ionophore valinomycin prior to the experiment, which creates an outward K+ gradient. Not significant (p = 0.5986, Student's unpaired t-test, two-tailed) difference was observed for the rate of transport by valinomycin addition. Bars are means \pm s.e.m with corresponding data points from n=3 independent experiments. **d-g** Calcium modulates the FPN transport mechanism. d, Time-course calcein fluorescence quenching response after addition of 1000 µM CoCl₂ (arrow) for FPN-liposomes preloaded with 1.25 mM CaCl₂ internally (red) or 1.25 mM CaCl₂ added to the external solution (grey). e, Initial rates of

Co²⁺ transport in the presence of internal 1.25 mM CaCl₂ (red circles). Data points are means \pm s.e.m from n=3 independent experiments. Quantified transport rates in Supplementary Table 2. **f**, Calcein fluorescence response for FeCl₂. **g**, Initial rates of Fe²⁺ transport. Data points are means \pm s.e.m from n=3 independent experiments. **h-i**, Calcium does not support high affinity hepcidin binding to FPN. **h**, Binding of 5 nM RhoG-hepcidin to nanodisc-FPN in the presence of 3 mM CaCl₂ was determined by fluorescence polarization (EC₅₀ >100 nM). Data points are means \pm s.e.m from n=3 independent experiments. **i**, Ca²⁺ stimulates the partial binding of 5 nM RhoG-hepcidin to 100 nM nanodisc-FPN with an EC₅₀ of 47.1 \pm 4.1 μ M, confirming that the Ca²⁺ binding site in FPN titrates in the physiological range (~1.25 mM) of free ionized Ca²⁺. In contrast, Co²⁺ stimulates binding of 5 nM RhoG-hepcidin to 100 nM nanodisc-FPN with an EC₅₀ of 35.8 \pm 3.6 nM. However, depletion of RhoG-hepcidin leads to a n_{Hill} = 0.52, and the true affinity of cobalt may be higher than this value. Importantly, these results indicate that binding of physiologically relevant concentrations of hepcidin would be exceedingly ineffective to apo and Ca²⁺-bound FPN. Data points are means \pm s.e.m from n=3 independent experiments.



Extended Data Fig. $8\mid$. Molecular dynamics simulations of iron binding to apo FPN.

a, Local resolution of FPN cryo-EM density. Insets show local resolution around TM7b. **b**, Each graph corresponds to an independent simulation where Fe²⁺ ions start in solution and bind spontaneously to the proposed iron binding site. Distance shown is from the ion to the nearest oxygen atom of the D325 side chain. Thick traces represent a 15-ns sliding mean and thin traces represent unsmoothed values. Timetraces include 30 ns of equilibration. The average time to bind (the time from start of production simulation to when the measured distance is less) was 71 ns. **c**, Representative conformations of TM7 from simulations with iron bound (blue) or absent (orange). **d**, In the absence of bound iron, D325 is mobile and can move into the cavity between TM7b and TM1. The top of TM7b can also tilt away from TM1. Four representative frames from simulation are overlaid. **e**, In simulations with Fe²⁺

bound, interaction with the ion restricts the mobility of D325 and, in turn, TM7b. **f**, Ironbound and iron-absent simulations show differences in the dynamics and position of D325, as measured by the distance between D325 $C\gamma$ and S47 $C\beta$. **g**, Comparison of conformation and dynamics with and without iron bound. With iron bound, D325 moves away from TM1 into the iron binding site (left), the root-mean-square fluctuation (RMSF) of D325 decreases (middle), and the extracellular end of TM7b moves closer to TM1 (right). For these comparisons, 6 simulations for each condition were used, each 2.0 μ s in length. Error bars are s.e.m. and p-values were calculated using Mann-Whitney U test (two-tailed); the p=0.001 (left plot), 0.046 (middle plot), 0.046 (right plot).

Extended Data Table 1:

Cryo-EM statistics

	FPN-Fab45D8 (EMDB-21539) (PDB 6W4S)	FPN-Co ²⁺ -hepcidin-Fab45D8 (EMDB-21599) (PDB 6WBV)
Data collection and processing	,	
Microscope/Detector	Titan Krios/Gatan K3 with Gatan Bioquantum Energy Filter	Titan Krios/Gatan K3 with Gatan Bioquantum Energy Filter
Imaging software and collection	SerialEM, 3×3 image shift	Serial EM, 3×3 image shift
Magnification	105,000	105,000
Voltage (kV)	300	300
Electron exposure (e–/Å ²)	66	66
Dose rate (e-/pix/sec)	8	8
Frame exposure (e-/Å ²)	0.55	0.55
Defocus range (µm)	-0.8 to -2.0	−0.8 to −2.0
Pixel size (Å)	0.834 (physical)	0.834 (physical)
Micrographs	5,415	4,395
Reconstruction		
Autopicked particles (template-based in cryosparc)	4,737,795	3,753,516
Particles in 3D classification	1,326,130 (RELION)	1,263,513 (cryosparc)
Particles in final refinement	308,386 (cisTEM)	310,647 (cisTEM)
Symmetry imposed	Cl	Cl
Map sharpening B factor (Å ²)	-130	-90
Map resolution, global FSC (Å)		
FSC 0.5, unmasked/masked	7.1/3.4	4.0/2.7
FSC.143, unmasked/masked	4.0/3.2	3.3/2.5
Refinement		
Initial model used (PDB code)	5AYN	5AYN, 3H0T
Model resolution (Å)		
FSC 0.5, unmasked/masked	3.6/3.4	3.0/2.8
Model composition		
Non-hydrogen atoms	4968	6869

	FPN-Fab45D8	FPN-Fab45D8 FPN-Co ²⁺ -hepcidin-Fab45D8
	(EMDB-21539)	(EMDB-21599)
	(PDB 6W4S)	(PDB 6WBV)
Protein residues	650	892
B factors ($Å^2$)		
Protein	67.31	27.30
Ligand	N/A	55.15
Water	N/A	31.32
R.m.s. deviations		
Bond lengths (Å)	0.004	0.006
Bond angles (°)	0.563	0.872
Validation		
MolProbity score	2.31	2.10
Clashscore	7.90	7.10
Poor rotamers (%)	4.45	4.26
EMRinger score	2.37	4.32
CaBLAM score	2.06	0.69
Ramachandran plot		
Favored (%)	94.38	96.47
Allowed (%)	5.62	3.53
Disallowed (%)	0	0

Extended Data Table 2:

Fab45D8 crystal structure statistics

	Fab45D8 (PDB 6W4V)
Data collection	
Space group	$P2_1$
Cell dimensions	
<i>a, b, c</i> (Å)	72.19, 36.43, 84.95
a , β , γ (°)	90.0, 112.26, 90.0
Resolution (Å)	39.31 – 2.09 (2.14 – 2.09) ^a
R_{sym} or R_{merge}	0.087 (1.385)
$I/\sigma I$	7.9 (1.2)
Completeness (%)	99.9 (99.9)
Redundancy	3.3 (3.5)
CC (1/2) (%)	99.6 (30.0)
Refinement	
Resolution (Å)	39.31 – 2.09
No. reflections	24736
$R_{\text{work}} / R_{\text{free}} (\%)$	21.6 / 24.7
No. atoms	
Protein	3356

	Fab45D8 (PDB 6W4V)
Ligand/ion	20
Water	125
B-factors	
Protein	57.87
Ligand/ion	86.17
Water	51.43
R.m.s. deviations	
Bond lengths (Å)	0.01
Bond angles (°)	0.92

^aValues for parentheses are for highest-resolution shell.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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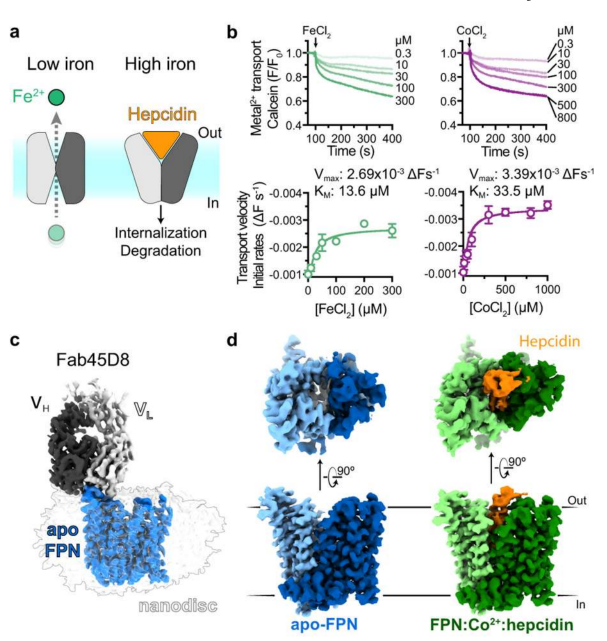


Fig. 1 |. Structures of human ferroportin.

a, Ferroportin effluxes cellular iron (Fe²⁺) by an alternating access mechanism. Hepcidin binds to outward-open ferroportin and induces ubiquitination and degradation. **b,** Transport of Fe²⁺ and Co²⁺ by human ferroportin reconstituted into calcein containing liposomes. Time-course experiments were performed by addition of ions (arrow) and the steady-state kinetic analysis was performed by fitting initial rates, obtained from the linear phase of transport, to the Michaelis-Menten function. Time-course data are representative experiments, and data points are means +/- s.e.m. from n = 3 independent experiments. **c,** Cryo-EM map of apo-FPN-Fab45D8 complex in lipid nanodisc. **d,** Cryo-EM density of apo and Co²⁺/hepcidin bound FPN. The N and C domains are colored in different shades of blue

for a po-FPN and green for $\mathrm{Co^{2+}/hepcidin}$ bound FPN. Hepcidin (orange) binds to an extracellular facing cavity in FPN.

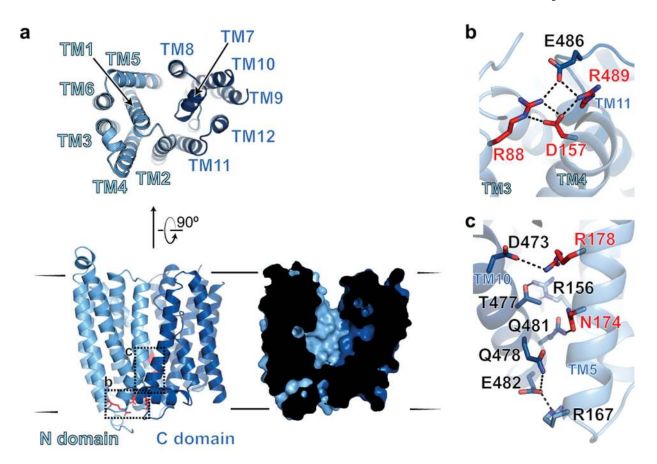


Fig. 2 |. Structure of apo-FPN.

a, Ribbon diagram of FPN reveals 12 transmembrane helices. The N- and C-domains are colored in different shades of blue. Cutaway surface view (right) shows outward open conformation. **b**, Intracellular gating residues are shown as sticks. **c**, TM10 and TM5 form an extensive network of interactions, further stabilizing the outward open conformation. Residues highlighted in red in (**b**) and (**c**) are known loss-of-function mutations that lead to ferroportin disease in humans.

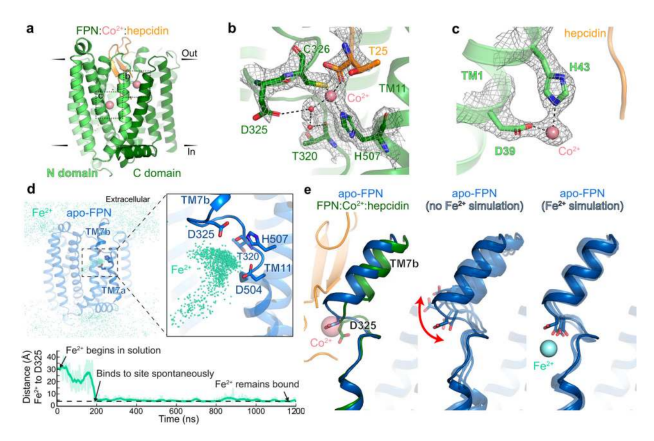


Fig. 3 |. Iron binds to the N and C domains of FPN.

a, Ribbon diagram of FPN-Co²⁺-hepcidin complex. Closeup view of cryo-EM density for Co^{2+} ion (pink) in the FPN C domain (**b**) and in the FPN N domain (**c**). **d,** Top: In molecular dynamics simulations with Fe^{2+} initially positioned randomly in bulk water surrounding FPN, the Fe^{2+} ions spontaneously bind to a region near H507, D325, and D504. The aggregated position of Fe^{2+} ions from six simulations, each 2 μ s in length, is shown superimposed with apo FPN. Bottom: In one representative simulation, an Fe^{2+} ion binds within 200 ns and remains localized at this site for >1000 ns. Distance shown is from the ion to the nearest oxygen atom of D325. Thick trace is a 15-ns sliding mean. **e,** Comparison of TM7b conformation in apo-FPN and FPN bound to Co^{2+} and hepcidin. In simulations without Fe^{2+} , TM7b is dynamic, with significant fluctuation of D325. D325 coordinates Fe^{2+} in simulations and leads to decreased TM7b motion.

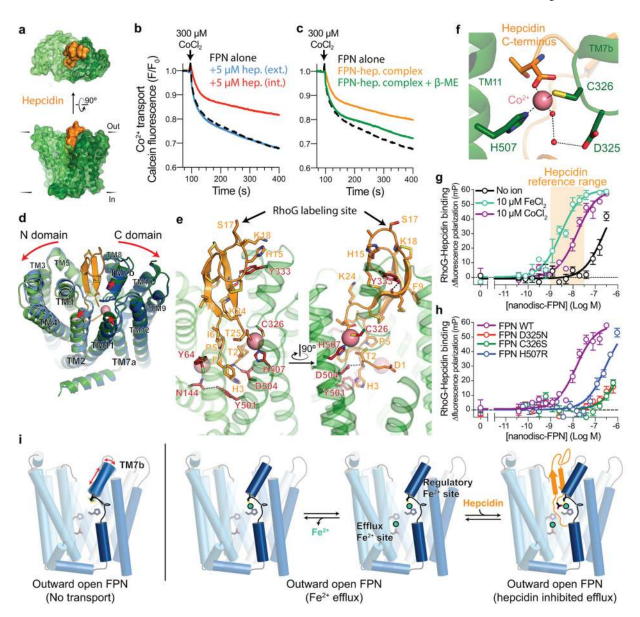


Fig. 4 |. Hepcidin binding to FPN requires iron.

a, Surface representation of the FPN-Co²⁺-hepcidin complex. **b,** Hepcidin (hep.) added internally (int.) to calcein-loaded liposomes inhibits FPN transport. External (ext.) hepcidin has no effect on transport. **c,** Reconstitution of a ferroportin-hepcidin (FPN-hep.) complex shows inhibited transport. Addition of 1 mM β-mercaptoethanol (β-ME) rescues transport activity. All transport traces are averaged values from n = 3 independent experiments. **d,** Ribbon diagrams of apo-FPN (blue) aligned to FPN-Co²⁺-hepcidin (green, orange, and pink spheres. Red arrows highlight structural differences. **e,** Closeup views of the hepcidin binding site. Residues in red are known hepcidin resistance mutations. Rhodamine green (RhoG) - hepcidin labeling site on position 17 is highlighted. **f,** Closeup of C domain metal binding site. **g,** Fluorescence polarization increase in rhodamine green-labeled hepcidin (RhoG-hepcidin) as nanodisc-reconstituted FPN is titrated with a K_D of 210 nM. Addition of 10 μM FeCl₂ or CoCl₂ increases the affinity of hepcidin to 2.5 nM and 7.7 nM (pK_D = -8.11

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 \pm 0.16), respectively. Hepcidin concentration range in healthy human adults is shown in orange. Data points are means \pm s.e.m. from n=3 independent experiments. **h**, C domain metal binding site mutants decrease RhoG-hepcidin binding affinity at FPN, even in the presence of 50 μ M CoCl₂. Data points are means \pm s.e.m. from n=3 independent experiments. **i**, Model for iron-coupled hepcidin regulation of FPN function. In settings of iron efflux, TM7b is conformationally stabilized by iron coordination in the C domain regulatory site. High affinity hepcidin binding to outward open FPN depends on the direct coordination of iron in the C domain.