Structure of the Wilson disease copper transporter ATP7B

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INTRODUCTION
Ubiquitously found in all kingdoms of life, P-type adenosine triphosphatases (ATPases) use adenosine 5'-triphosphate (ATP) hydrolysis to actively pump substrates across cell membranes (1). In humans, two Cu⁺-transporting P-type ATPases, ATP7A and ATP7B, are key players in maintaining cellular copper homeostasis (2–4). Dysregulation of cellular copper levels is deleterious, as copper functions as an essential cofactor for numerous enzymes, while excess copper is cytotoxic (2–4). Human ATP7A and ATP7B, which share 56% sequence identity, differ in tissue expression, cellular trafficking, and physiological functions (2–4). ATP7A is important for copper uptake in the intestine, and dysfunction causes Menkes disease characterized by systemic copper deficit (5, 6). By contrast, ATP7B is most abundantly expressed in hepatocytes to excrete excess copper into bile, and consequently, missense mutations give rise to Wilson disease that is represented by hepatic copper overload and toxicity (7, 8).

The P-type ATPases, which constitute a large superfamily of membrane pumps that catalyze the transmembrane transport of diverse substrates, all contain a common structural core composed of a transmembrane domain (TMD), an actuator (A) domain, a phosphorylation (P) domain, and a nucleotide-binding (N) domain (1). Substrate transport is actively driven by conformational changes associated with phosphorylation and dephosphorylation of a conserved aspartate residue in the P domain, which result in alternating access of substrate binding sites in the TMD within the membrane (1). The stepwise transport process is described by the Post–Albers cycle with distinct functional states, including high-affinity E1 and low-affinity E2 states for substrate as well as phosphoenzyme intermediate states E1P and E2P (9, 10).

ATP7A and ATP7B belong to the prevalent P1B subfamily of heavy metal–transporting ATPases that are widespread in both prokaryotes and eukaryotes (11). Crystal structures of a prokaryotic homolog from Legionella pneumophila, LpCopA, have delineated the core structure of a P1B ATPase and suggested a potential Cu⁺ transport pathway across cell membranes (12, 13). The structure and ion permeation mechanism appear to differ substantially from those of the prototypical P2 ATPase subfamily exemplified by the best characterized sarcoplasmic reticulum Ca²⁺-ATPase (SERCA1a) (14–17). The LpCopA TMD has two additional P1B-specific transmembrane helices, a long and curved MA helix and a kinked MB helix with an amphipathic helical segment at the cytoplasmic membrane interface, that are not present in the canonical SERCA1a (12, 13). In addition to the common P-type ATPase core pieces comprising the TMD and A, P, and N domains, P1B-ATPases contain one or multiple metal-binding domains (MBDs) in the N terminus (11, 18, 19). The prokaryotic LpCopA acquires a single MBD, which was not resolved in the crystal structures (12, 13). The N terminus is elaborated in higher eukaryotic Cu⁺-ATPases, which contain six consecutive MBDs (MBD1 to MBD6) that receive Cu⁺ from a cytoplasmic Cu⁺ chaperone antioxidant protein 1 (ATOX1) and participate in subsequent Cu⁺ transport across the membrane (19–24). While MBD1 to MBD4 are not strictly required and appear to play a regulatory role, either MBD5 or MBD6 is necessary for Cu⁺ transport (25–27). ATP7A/B additionally contains a conserved C-terminal region absent in LpCopA that appears to communicate with the N-terminal domains to regulate copper-stimulated trafficking (28). These extra segments at both the N and C termini render ATP7A/B much larger in size than the prokaryotic counterparts and confer sophisticated Cu⁺-dependent regulation mechanisms of protein targeting, cellular trafficking, and post-translational modifications (25–28), which remain poorly understood.

To better understand the unique features and physiological and pathological properties of ATP7 proteins, here, we describe a cryo–electron microscopy (cryo-EM) structure of ATP7B from Xenopus tropicalis determined at an overall resolution of ~3.2 Å. The structure reveals a conserved Cu⁺-ATPase core surrounded by the N-terminal...
MBDs and the C-terminal regulatory domain and delineates a plausible Cu⁺ transport pathway lined by a network of amino acids serving as Cu⁺ ligands in the membrane. Meanwhile, the first atomic model of a vertebrate Cu⁺-ATPase provides a molecular blueprint for understanding hundreds of missense mutations associated with devastating Menkes and Wilson diseases and facilitates rational design of therapeutic interventions.

RESULTS
Structure determination
We expressed and purified the full-length *X. tropicalis* ATP7B protein consisting of 1467 amino acids, which shares 67% sequence identity with the human ortholog (fig. S1). We observed protein degradation likely owing to the flexible linkage of the six N-terminal MBDs. Nuclear magnetic resonance (NMR) spectroscopy studies have indicated that the individual MBDs are loosely connected like beads on a string except for MBD5 and MBD6 forming a compact unit preceding the first TM helix (21, 22, 29). Thus, for structural studies, we removed the first four of six MBD domains (MBD1 to MBD4) that are not essential for Cu⁺ transport (25–27, 30). The truncated construct, which we termed xATP7BEM, consists of amino acids 481 to 1467 with intact MBD5 and MBD6 and C terminus. The resulting construct has a predicted molecular weight of ~107 kDa (Fig. 1A). Consistent with previous studies (25–27), purified xATP7BEM lacking MBD1 to MBD4 retains ATPase activity (Fig. 1B). Furthermore, the ATP turnover rate of xATP7BEM (7.8 min⁻¹) is higher than that of the full length (3.4 min⁻¹) (Fig. 1B). This observation supports the notion that MBD1 to MBD4 negatively regulate ATPase activity before stimulation by cytoplasmic Cu⁺ chaperone ATOX1 (31).

We purified xATP7BEM to homogeneity and determined the cryo-EM structure at an overall resolution of ~3.2 Å (Fig. 1C, fig. S2, and table S1). The cryo-EM densities are of high quality for the TMD and A and P domains, allowing unambiguous model building (Fig. 1C and fig. S3). The densities were less well resolved for the

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Fig. 1. Cryo-EM structure of xATP7B. (A) Membrane topology and domain organization of ATP7B. ATP7B contains six N-terminal MBDs (orange), a TMD with eight TM helices (cyan and green), and four cytoplasmic domains including A (yellow), P (blue), N (magenta), and C-terminal (red) domains. The MBD1 to MBD4 deletion construct used for cryo-EM analysis is indicated by a pair of scissors. The cyan lines mark the approximate membrane boundaries. (B) ATP turnover rates for the full-length wild type (WT; 3.4 ± 0.1 min⁻¹), xATP7BEM (7.8 ± 0.5 min⁻¹), and C-terminal deletion (xATP7BEM ΔC) using the inactive mutant (D1024A) as the baseline for subtraction. Data are means ± SD (three independent measurements). For comparison, the maximum turnover rate of LpCopa was ~11.7 min⁻¹ (12). (C) Cryo-EM density map. The TMD, cytoplasmic domains (MBD5 and MBD6, CTD, and A, P, and N domains), and the MBD6-TMD linker are indicated. (D) Structure of xATP7BEM. The TM helices (MA, MB, and M1 to M6) and cytoplasmic domains are indicated.
cytoplasmic N domain, and thus the crystal structure of the prokaryotic LpCopA [Protein Data Bank identification (PDB ID): 3RFU] was used to guide model building in this region (12). Three-dimensional variability analysis (3DVA) revealed that the densities corresponding to MB5 to MB6, which were relatively weak in the consensus reconstruction, were present in only a subset of the particles. After 3D classification, we obtained two subsets, one in which MB5 and MB6 were ordered (state 1) and one in which the MB5 and MB6 densities were absent (state 2). To model MB5 and MB6, we docked the crystal structure of the isolated human MBD4 (PDB ID: 6A71) into the state 1 density map and then manually adjusted (32). Assignment of MB5 and MB6 was further corroborated by the visible densities for the MB5-MBD6 and MB6-MA linkers (Fig. 1C). The final atomic model, which was refined to good stereochemistry, fits well into the density maps (Fig. 1, C and D, fig. S3, and table S1).

Overall structure of ATP7B

The xATP7B EM structure was determined in the presence of Mg$^{2+}$ and AlF$_4^-$ in the absence of the substrate ions (Cu$^+$), thus presumably representing an occluded, Cu$^+$-released E2-P$_i$ transition state in the Post-Albers cycle. Mg$^{2+}$ and AlF$_4^-$ are bound at the conserved phosphorylation site constituted by the invariant “DKTG” motif from the P domain and the “TGE” loop from the A domain (fig. S4A), which is analogous to the observation in the crystal structure of LpCopA captured in the E2-P$_i$ state (12). The assignment of the functional state is further supported by the global structural agreement between xATP7B EM and LpCopA (fig. S4B) (12).

The TMD comprises eight TM helices that closely resemble those of LpCopA, including the two P1B-specific helices (MA and MB) preceding the canonical six TM helices (M1 to M6) shared by P-type ATPases (Fig. 1, C and D). The first TM helix, MA, is connected to MB6 via a partially ordered linker that is ~20 Å in length. The MB helix contains a sharp kink, enabled by two consecutive glycine residues that are conserved among Cu$^+$-ATPases (fig. S1). Consequently, the second helical segment in MB runs approximately parallel to the membrane. A short linker follows MB and connects to the common core architecture of a P-type ATPase, which sequentially consists of M1, M2, the A domain, M3, M4, the P and N domains, M5, and M6.

Unique to eukaryotic Cu$^+$-ATPases, the cytoplasmic N terminus contains six MBDs, which confer exquisite regulation mechanisms (19). The cryo-EM structure reveals that the last two MBDs (MB5 and MB6) are distantly positioned from the lipid bilayer and pack against both the A and P domains in one state and uncoupled from the transporter in the other (Fig. 1, C and D, and fig. S2). The locations of MB5 and MB6, clearly resolved in the density map of state 1, differ substantially from the previously suggested positions of the single N-terminal MBD in LpCopA (12, 33). In addition, in contrast to LpCopA, eukaryotic ATP7 transporters contain a C-terminal tail following the TMD, which comprises ~90 amino acids. In the xATP7B EM structure, the C terminus emerging from M6 wraps around the P domain and reaches MB5 (Fig. 1, C and D), thus forming an integral structural unit specific for eukaryotic ATP7 proteins. Together, it appears that both the extra N- and C-terminal regions constitute peripheral structural domains that surround the P and A domains, presumably providing further regulatory mechanisms through direct interdomain interactions (Fig. 1D).

Transmembrane Cu$^+$ pathway

The overall arrangement of the cytoplasmic domains and, in particular, the positions of the obligatory MBDs (MBD5 or MBD6) indicate...
that the Cu⁺ entryway is most likely constituted by the cytoplasmic interface created by MA, MB, and M2 (Figs. 1, C and D, and 2A), which aligns with the proposed entry site of LpCopA (12, 13). The Cu⁺-free structure, together with the distal placement of MBD5 and MBD6 above the lipid bilayer, prevents visualization of how Cu⁺ is transferred from the MBDs to the membrane entrance. Notably, the poorly ordered linker connecting MBD6 and MA (Fig. 1C) makes no contacts with other cytoplasmic domains. Together with the MBDs being disordered in state 2, it appears that the flexible linker may allow the long-range movement of MBDs necessary for Cu⁺ delivery to this entry site.

To illuminate the potential Cu⁺ transport pathway across the membrane, we mapped the conserved Cu⁺-binding residues, including cysteine, methionine, and histidine as well as charged amino acids in the transmembrane region (Fig. 2A). These residues delineate a virtually complete route allowing Cu⁺ movement across the lipid bilayer from the cytoplasmic entry to the luminal exit. The presumed cytoplasmic Cu⁺ entry site comprises multiple conserved charged residues (Fig. 2B). A ring of negatively charged residues, including E643, D727, E778, and D915, appears to form a cation-binding site. Notably, two conserved histidine residues in the vicinity, H721 and H779, likely facilitate Cu⁺ entry by serving as additional Cu⁺ ligands. In the absence of Cu⁺ ions, a conserved arginine residue, R775, occupies the center of the acidic ring and seals the cytoplasmic side of the entryway. The negative electrostatic potential of the acidic ring is partially balanced by nearby lysine residues K717 and K782. Therefore, Cu⁺ entering this acidic binding site would likely displace the side chain of R775, which appears to be critical in stabilizing the occluded-to-the-cytoplasm conformation and thus functions as a cytoplasmic gate facilitating Cu⁺ occlusion. Consistent with this proposed role of R775, a single missense mutation of the corresponding residue in human ATP7B (R778L) represents one of the most

Fig. 3. CTD and the N-terminal MBDs. (A) Overall structure highlighting positions of the CTD and the N-terminal MBD5 and MBD6. AlF₄⁻ bound in the P domain and the presumed Cu⁺-binding motifs in MBD5 and MBD6 are highlighted. (B) MBD5 interacts with both the A and P domains and contacts the CTD. (C) Interface between MBD6 and the A domain. (D) Orthogonal view as in (A) to illustrate domain organization of MBD5 and MBD6, CTD, and the A and P domains. (E) The α helix in the C-terminal tail αCTD interacts with the P domain. (F) The C-terminal β strand βCTD extends the β sheet of the P domain. (G) ATP turnover rates for xATP7B EM and mutants weakening inter-domain interactions. Data are means ± SD (three independent measurements).
and I870 of the A domain, and L1250 of the P domain. In addition, E526 of MBD5 and R824 of the A domain form salt bridges. In contrast to MBD5 making extensive interdomain interactions, MBD6 appears to be rather detached from the core pieces, only contacting the A domain via salt bridge interactions involving R613, E815, and D826 (Fig. 3, C and D).

In the P-type ATPases, conformational changes in the cytoplasmic domains (A, P, and N domains) driven by ATP hydrolysis are coupled with ion movement in the TMD (1). In the cryo-EM structure, MBD5 interacts with multiple cytoplasmic domains via its Cu\(^{2+}\)-binding motif and nearby residues. This observation indicates that MBD5 could engage Cu\(^{2+}\)-dependent regulation of structural rearrangements of the cytoplasmic domains and, consequently, Cu\(^{2+}\) movement in the TMD. Both Cu\(^{2+}\)-binding motifs in MBD5 and MBD6 are distant from the cytoplasmic Cu\(^{2+}\)-ATPase entry point as estimated by the Ca-Ca distances between R775 and C496 (~50 Å) and between R775 and C572 (~56 Å). Therefore, if MBD5 or MBD6 participates in delivering Cu\(^{2+}\) to the membrane entry site, long-range domain movements in MBD5 or MBD6 are expected to occur.

**N-terminal MBDS**

Cu\(^{2+}\)-ATPases from different organisms share the core catalytic pieces but have a varying number of MBDS in the N terminus (11). The prokaryotic Cu\(^{2+}\)-ATPases contain a single MBD, and the lower eukaryotic ones have two or four, whereas the higher eukaryotic transporters comprise six consecutive MBDS. NMR studies of individual MBDS have revealed a conserved ferredoxin-like \(\beta\alpha\beta\beta\alpha\beta\) fold consisting of ~70 amino acids with an invariant “MxCxxC” Cu\(^{2+}\)-binding motif (11, 21, 41–43). The ATP\(_7\) transporters engage the MBDS to receive Cu\(^{2+}\) ions from cytosolic Cu\(^{2+}\) chaperones such as ATOX1, which shares the same \(\beta\alpha\beta\beta\alpha\beta\) structural fold, and to relay Cu\(^{2+}\) ions to the intramembranous transport pathway (19–24). However, overall structural arrangement of the MBDS in the intact transporters and the mechanisms of Cu\(^{2+}\) delivery and Cu\(^{2+}\)-dependent regulation of transport activity via these MBDS remain unknown. Mutagenesis studies of ATP\(_7\)B have demonstrated that either MBD5 or MBD6, but not MBD1 to MBD4, is required for Cu\(^{2+}\) transport activity (25–27), suggesting that MBD1 to MBD4 play a regulatory role, whereas MBD5 or MBD6 is essential for Cu\(^{2+}\) relay to the membrane entry site. Consistently, purified xATP\(_7\)B\(_{EM}\) lacking MBD1 to MBD4 showed increased ATP\(_\text{ase}\) activity compared with the full-length enzyme (Fig. 1B).

In the xATP\(_7\)B\(_{EM}\) structure, MBD5 simultaneously contacts the A, P, and C-terminal domains (Fig. 3A). Notably, the Cu\(^{2+}\)-binding motif, which consists of M494, C496, and C499, is precisely located at the multidomain junction (Fig. 3B). In the vicinity of the Cu\(^{2+}\)-binding site, MBD5 and the A and P domains are held together by a hydrophobic interface comprising I497 and L520 of MBD5, V861

prevalent of the more than 700 Wilson disease mutations that have been identified (8, 34).

M726, immediately beneath R775 in the proposed transport pathway, demarcates the cytoplasmic and intramembranous binding sites and may facilitate Cu\(^{2+}\) movement further into the lipid bilayer (Fig. 2). In the cryo-EM structure, the Ca-Ca distances between M726 in M1 and cysteine residues (C980 and C982) in the invariant “CPC” motif in M4 are ~10 Å. Presumably, conformational changes in M1 and M4 associated with ATP hydrolysis could bring M726, C980, and C982 together to bind Cu\(^{2+}\) through Cu-S coordination with trigonal geometry that is selective for Cu\(^{2+}\) over Cu\(^{+}\) ions (35–37). A network of Cu\(^{2+}\)-binding residues, including M1365, M766, D762, M665, M1360, and M668, lines the transmembrane ion path reaching the luminal side. These Cu\(^{2+}\) ligands seem to be meticulously arranged to relay the bound Cu\(^{2+}\) ions through a series of ligand exchange reactions to deliver Cu\(^{2+}\) to the lumen. An analogous mechanism of Cu\(^{2+}\) translocation has been suggested for the structurally unrelated Ca\(^{2+}\)-ATPase, which shares the same \(\alpha\beta\delta\alpha\beta\) structural fold, and to relay Cu\(^{2+}\) ions from cytosolic Cu\(^{2+}\) chaperones such as ATOX1, which shares the same \(\alpha\beta\delta\alpha\beta\) structural fold, and to relay Cu\(^{2+}\) ions to the intramembranous transport pathway (19–24). However, overall structural arrangement of the MBDS in the intact transporters and the mechanisms of Cu\(^{2+}\) delivery and Cu\(^{2+}\)-dependent regulation of transport activity via these MBDS remain unknown. Mutagenesis studies of ATP\(_7\)B have demonstrated that either MBD5 or MBD6, but not MBD1 to MBD4, is required for Cu\(^{2+}\) transport activity (25–27), suggesting that MBD1 to MBD4 play a regulatory role, whereas MBD5 or MBD6 is essential for Cu\(^{2+}\) relay to the membrane entry site. Consistently, purified xATP\(_7\)B\(_{EM}\) lacking MBD1 to MBD4 showed increased ATP\(_\text{ase}\) activity compared with the full-length enzyme (Fig. 1B).

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C-terminal tail

Following the TMD, the eukaryotic ATP7 transporters contain a C-terminal tail that is absent in prokaryotic homologs (fig. S1). The C-terminal tail forms intimate interactions with the P domain (Fig. 3, D and E). The short C-terminal helix αCTD makes multiple contacts with the P domain (Fig. 3E). Furthermore, the C-terminal tail donates an extra β strand (βCTD) that complements the β sheet of the P domain (Fig. 3F). In addition, the distal C terminus continues toward the N-terminal MBD5, joining the multidomain interface involving MBD5 and the A and P domains (Fig. 3F). Together, the C-terminal tail appears to be an integral structural component extending the cytoplasmic headpieces. Presumably, the C-terminal tail could impart additional regulatory roles specific for the eukaryotic Cu\(^{+}\)-ATPases through direct interactions with MBD5 and the P domain.

Yeast complementation assays with ATP7B have demonstrated that the C terminus is critical for function and communicates with the N terminus as assessed by phosphoproteomics studies (28, 44), in which Cu\(^{+}\) binding at the N terminus resulted in hyperphosphorylation.
of the C-terminal sites. The xATP7B<sub>EM</sub> structure provides direct
evidence that the N and C termini physically interact and thus
offers a straightforward explanation for the observed long-range
cross-talk between the N and C termini. To further probe the
functional role of the C-terminal tail, we removed this portion
beginning at K1384, and the resulting construct (xATP7B<sub>ΔC</sub>)
essentially eliminated the ATPase activity (Fig. 1B). In accordance
with previous studies (44), our structural and functional data
demonstrated the obligatory role of the C-terminal tail that is
unique in eukaryotic Cu<sup>+</sup>-ATPases. In contrast, in P4-ATPases,
analogous C-terminal tail wraps around the cytosolic headpieces
to confer an autoinhibitory role by blocking ATP binding and
constricting domain movement (45–47).

In addition to a conserved Cu<sup>+</sup>-ATPase core, our vertebrate
ATP7B structure reveals previously unknown structural elements at
both the N and C termini (Fig. 3). To further validate these structural
observations, we introduced mutations in the N-terminal MBD5
(L520A/M521A) and MBD6 (R613E) domains and the C-terminal
tail (K1384E) on the xATP7B<sub>EM</sub> background, which presumably
weaken domain-domain interactions with the catalytic core, and
measured the ATPase activity (Fig. 3G). Consistently, all these mutants
demonstrated a reduced ATP turnover rate, thus indicating the involve-
ment of these structural elements in regulation of transport activity
through interactions with the cytosolic headpieces. Notably, the
charge reversal mutation (K1384E) in the C-terminal tail resulted in
the most pronounced reduction of the ATP turnover rate.

**Wilson disease mutations**

Our cryo-EM structure provides a molecular blueprint for under-
standing Wilson disease mutations. We mapped known missense
human mutations of Wilson disease onto the cryo-EM structure of
frog ATP7B and labeled these disease mutations according to the
human ATP7B sequence for consistency with the Wilson disease
database (Fig. 4) (48). Most of the known disease mutations are located
within the structurally resolved region of the full-length protein,
including the TMD and cytosolic A, P, and N domains. This reinforces
the engagement of each of these domains during the transport cycle
driven by ATP hydrolysis. Notably, several mutations occur in struc-
tural elements located at the N-terminal MBDs and the obligatory
C-terminal tail that interact with the cytosolic headpieces. In particu-
lar, the human mutation R616Q/W (corresponding to R613 in frog
ATP7B) would disrupt salt bridge interactions between MBD6
and the A domain (Fig. 3C). Another disease mutation Q1399R
(equivalent to Q1405 in frog ATP7B) is located at the interface
between the C-terminal tail and the P domain. These observations
further strengthen the critical roles of these peripheral regulatory
components.

**DISCUSSION**

This work now reveals the first high-resolution structure of a verte-
brate Cu<sup>+</sup>-ATPase, which delineates the shared and distinct features
among Cu<sup>+</sup>-ATPases and provides a molecular blueprint for fur-
ther understanding numerous Wilson disease mutations. The Cu<sup>+</sup>
ATPases from eukaryotes and prokaryotes share a core structure
consisting of the A, P, and N domains and a P1B-specific TMD with
eight TM helices (Fig. 5). In contrast to prokaryotic Cu<sup>+</sup>-ATPases,
the eukaryotic counterparts contain a greater number of N-terminal
MBDs and an extended C-terminal tail. These extra structural
components surround the conserved cytosolic headpieces of the
Cu<sup>+</sup>-ATPase apparatus and form extensive interdomain packing
interactions. Both the N-terminal MBDs and C-terminal tail appear to
be strategically positioned to exert additional regulatory roles
by interfering with conformational changes in cytosolic headpieces,
which are necessary for coupling ATP hydrolysis to TMD move-
ments that facilitate active Cu<sup>+</sup> transport across the membrane.
Furthermore, structural organization of these domains indicates
that the regulatory roles are poised to be modulated by Cu<sup>+</sup> bind-
ing to MBDs.

The arrangement of TM helices is conserved between prokaryotic
and eukaryotic Cu<sup>+</sup>-ATPases (root mean square deviation of the Cu
atoms between eight TM helices is ~1.6 Å). Mapping of potential
Cu<sup>+</sup> ligands in the TMD, including cysteine, methionine, and
charged amino acids, suggests a similar stepwise shuttle mechanism
for Cu<sup>+</sup> transport as well as an analogous “platform” entry site to
accommodate the N-terminal MBDs or a cytosolic chaperone for
Cu<sup>+</sup> transfer (Fig. 5 and fig. S5) (12). Presumably, structural
rearrangements, associated with Cu<sup>+</sup> loading to the MBDs and
ATPase activity in the cytosolic domains, drive unidirectional Cu
movement from the cytosolic entry site through the intramembrana-
rous binding sites and to the luminal side. Notably, the wide exit
tunnel to the lumen observed in the presumed E2-Pi state supports
a unique ion-release mechanism for Cu<sup>+</sup>-ATPases as previously
suggested (13). The Cu<sup>+</sup>-coordinating amino acids seem to be
delicately arranged within the membrane to relay Cu<sup>+</sup> via ligand
exchange reactions. In comparison with the prokaryotic Cu<sup>+</sup>-ATPases,
the eukaryotic counterparts constitute a more refined transmem-
brane Cu<sup>+</sup> path that involves additional Cu<sup>+</sup>-binding amino acids
such as M766 and M668 (Fig. 5). Notably, the eukaryotic ATP7B
transporters have evolved to engage a conserved basic residue R775,
which corresponds to glutamine at the equivalent position in LpCopA
(fig. S1), as the cytoplasmic gatekeeper for Cu<sup>+</sup> entry. Coincidently,
an amino acid substitution at this position in human ATP7B (R778L)
gives rise to a prevalent Wilson disease mutation (8, 34).

**MATERIALS AND METHODS**

**Cloning, expression, and purification of ATP7B**

DNA encoding *X. tropicalis* ATP7B was codon-optimized, synthesized
(Bio Basic), and served as the template for subcloning. Deletion of
the N-terminal 480 amino acids corresponding to MBD1 to MBD4
(xATP7B<sub>EM</sub>) improved biochemical stability and decreased protein
degradation and thus was suitable for structural studies. DNA
was polymerase chain reaction–amplified and ligated into the 5’
Xho I and 3’ Eco RI cloning sites of a modified pPICZ-B vector
(Invitrogen) under a methanol-inducible promoter. The expres-
sion construct contains a C-terminal PreScission protease cleavage
site linked to a green fluorescent protein (GFP)–His<sub>10</sub> tag. The plasmid
was linearized by Pme I and transformed into *Pichia pastoris* for
expression.

Yeast cells were disrupted by milling (Retsch MM400) and re-
suspended in lysis buffer containing 50 mM Hepes (pH 7.0), 150 mM
NaCl, 5 mM MgCl<sub>2</sub>, 0.5 mM adenosine 5’-diphosphate (ADP),
10 mM NaF, and 1 mM AlCl<sub>3</sub> supplemented with protease inhibi-
tors [leupeptin (2.5 μg/ml), pepstatin A (1 μg/ml), 4-(2-aminoethyl)
benzenesulfonyl fluoride hydrochloride (100 μg/ml), aprotinin
(3 μg/ml), 1 mM benzamidine, and 200 μM phenylmethylsulfonyl
fluoride] and deoxyribonuclease I. Cell membranes were solubilized

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in 1% (w/v) lauryl maltose neopentyl glycol (Anatrace) and 0.1% cholesteryl hemisuccinate for 2 hours with stirring at 4°C. Solubilized ATP7B protein was separated from the insoluble fraction by centrifugation for 1 hour at 30,000g and incubated with 3 ml of TALON metal affinity resin (Clontech) for 3 hours at 4°C. Resin was then washed with 10 column volumes of 20 mM Hepes (pH 7.0), 150 mM NaCl, 5 mM MgCl₂, 0.5 mM ADP, 10 mM NaF, 1 mM AlCl₃, 10 mM imidazole, and 85 μM glyco-diosgenin (GDN; Anatrace). Bound protein was eluted with buffer containing 200 mM imidazole and further incubated with anti-GFP nanobody bound to Glutathione Sepharose 4B resin (GE Healthcare) for another 2 hours at 4°C. Unbound protein was washed away with 10 column volumes of wash buffer containing 20 mM Hepes (pH 7.0), 150 mM NaCl, 5 mM MgCl₂, 0.5 mM ADP, 10 mM NaF, 1 mM AlCl₃, and 85 μM GDN. The C-terminal GFP-His₁₀ tag was removed by digestion with PreScission protease on the column at 4°C overnight. The flowthrough from the column was then concentrated using an Amicon Ultra-15 centrifugal filter unit (molecular weight cutoff 100 kDa) and injected into a Superose 6 Increase 10/300 gel filtration column (GE Healthcare) at 4°C overnight. The flowthrough from the column was then concentrated using an Amicon Ultra-15 centrifugal filter unit (molecular weight cutoff 100 kDa) and injected to density modification in Phenix using a soft mask (55). The refined consensus model was docked into the state 2 structure, and the missing MBDS were removed from the model before real-space refinement in Phenix.

**Cry-EM sample preparation and imaging**

Four microliters of purified xATP7BEM protein at ~7 mg/ml was added to Au 400 mesh Quantifoil R1.2/1.3 holey carbon grids (Quantifoil) that had been glow-discharged for 60 s. After a 20-s incubation, the grids were blotted for 2 s with a blot force of 0 and rapidly plunged into liquid nitrogen-cooled liquid ethane using a FEI Vitrobot Mark IV (FEI Thermo Fisher Scientific). A total of 3439 images were recorded in super-resolution mode at a nominal magnification of 22,500x on a Titan Krios (Thermo Fisher Scientific) operating at 300 kV equipped with a K2 Summit (Gatan). The calibrated super-resolution pixel size was 0.544 Å/pixel. The dose rate was eight electrons per pixel per second for 8 s using 0.2-s frames, and images were acquired with a defocus range of ~1.2 to ~2.5 μm using SerialEM (49).

**Image processing and map calculation**

The super-resolution movies were gain-corrected, Fourier-cropped to a pixel size of 1.088 Å/pixel, and aligned using whole frame and local motion correction using MotionCor2 (50). Contrast transfer function (CTF) parameters were estimated using CTFFIND 4.1.10 (51). Particles were automatically selected using Laplacian of Gaussian autopicking in Relion 3.0 (52). False-positive selections were eliminated using iterative heterogeneous classification in cryoSPARC v3 (53) using models generated from the ab initio algorithm in cryoSPARC v3. After Bayesian polishing in Relion and global and local CTF refinement in cryoSPARC v3, the consensus refinement from 257,208 particles achieved a resolution of 3.19 Å using nonunimodal refinement in cryoSPARC v3 (54). DVA in cryoSPARC revealed the presence of two distinct states, one in which MBDB5 and MBDB6 were resolved (state 1) and one in which the density for MBDB5 and MBDB6 was absent (state 2). Heterogeneous classification using the maps generated by D3VA yielded a stack of 138,790 particles for state 1 and 118,418 particles for state 2, which achieved resolutions of 3.32 and 3.49 Å, respectively. All reconstructions were subsequently subjected to density modification in Phenix using a soft mask (55).

**Model building and coordinate refinement**

The structure of LpCopA (PDB: 3RFU) was manually docked into the density modified consensus map in UCSF Chimera (56) and then rebuilt in Coot (57). The structure of human MBDB4 (PDB: 6A71) was docked in the densities corresponding to MBDB5 and MBDB6 in state 1 using UCSF Chimera and then rigid-body fit and manually rebuilt in Coot (32). The manually built models were refined in ISOLDE (58) and by using real-space refinement in Phenix with geometric and Ramachandran restraints (59). The refined consensus model was docked into the state 2 structure, and the missing MBDS were removed from the model before real space refinement in Phenix.

**ATPase assay**

All xATP7B constructs used in the ATPase assay were expressed and purified following the procedure described for xATP7BEM, except that NaF and AlCl₃ were removed in the lysis and was buffer, and the gel filtration buffer contained 20 mM Hepes (pH 7.0), 150 mM NaCl, and 40 μM GDN. Peak fractions were collected and concentrated to ~1 to 2 mg ml⁻¹ for the ATPase assay. Concentrated protein was immediately flash-frozen and stored in −80°C until further usage. ATPase activity was measured using Bismuth detection method (12, 60). Measurements were conducted with equal amounts of protein (1 μM) for each construct. In a total reaction volume of 50 μl, protein was mixed with the ATPase assay buffer [40 mM Mops-KOH (pH 6.8), 150 mM NaCl, 5 mM KCl, 5 mM MgCl₂, 20 mM (NH₄)₂SO₄, 20 mM cysteine, 5 mM Na₂S, 0.25 mM Na₂MoO₄, and 80 μM GDN]. Ammonium tetrathiomolybdate (TTM; 10 μM) was added to each reaction mix to eliminate free copper. An increasing amount of ATP was added to initiate the reaction, and the mixture was incubated at 37°C for 15 min. Reactions were stopped by addition of 75 μl of freshly made quench solution [2.86% ascorbic acid, 1 M HCl, 0.48% (NH₄)₂MoO₄, and 2.86% SDS] and then incubated at room temperature for 8 min. Last, 125 μl of stabilizing solutions (3.5% bismuth citrate, 1 M HCl, and 3.5% sodium citrate) was added to the mixture and incubated for 30 min at room temperature. Absorbance was measured at 710 nm using Cytation 5 (BioTek) plate reader. The background from blank (assay buffer with TTM) was subtracted from each data point to retrieve the actual reads. The amount of inorganic phosphate released was determined by using the K₂HPO₄ standard as described previously (60). All measurements were performed in three replicates. The kinetic parameters were deduced from the Michaelis-Menten equation using Prism (www.graphpad.com). We have not been able to establish in vitro Cu⁺-stimulated ATPase activity, because the purified frog ATP7B protein was prone to aggregation with the addition of Cu⁺.  

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abb5508

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Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. The cryo-EM maps of xATP7BEM have been deposited to the Electron Microscopy Data Bank with accession codes EMD-25137 (consensus), EMD-25138 (state 1), and EMD-25139 (state 2). Atomic coordinates of the xATP7BEM structure have been deposited to the Protein Data Bank with accession codes 7SI3 (consensus), 7SI6 (state 1), and 7SI7 (state 2). The expression plasmids of ATP7B can be provided by Washington University pending scientific review and a completed material transfer agreement. Requests for the expression plasmids should be submitted to P.Y. (yuanp@wustl.edu).

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