

Conduction pathway for potassium through the *E. coli* pump KdpFABC

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This **valuable** study provides new insights into the movement of ions through the bacterial pump KdpFABC, which regulates intracellular potassium concentration, by solving a 2.1 Å cryo-EM structure of the nanodisc-embedded active wild-type protein, and carrying out mutagenesis and activity assays. Although the structural data and analysis are **solid**, additional information about other structural classes identified in the EM data, as well as a discussion of relevant work done by others, would further strengthen these findings. The description of the activity assays is currently **incomplete** because more information is required to rigorously assess these experiments. This work will be of interest to the membrane transporter and channel communities and to microbiologists interested in osmoregulation and potassium homeostasis.

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Abstract

Under osmotic stress, bacteria express a heterotetrameric protein complex, KdpFABC, which functions as an ATP-dependent K^+ pump to maintain intracellular potassium levels. The subunit KdpA belongs to the Superfamily of K^+ Transporters and adopts a pseudo-tetrameric architecture with a membrane embedded selectivity filter as seen in K^+ channels. KdpB belongs to the superfamily of P-type ATPases with a conserved binding site for ions within the membrane domain and three cytoplasmic domains that orchestrate ATP hydrolysis via an aspartyl phosphate intermediate. Previous work has hypothesized that K^+ moves parallel to the membrane plane through a 40-Å long tunnel that connects the selectivity filter of KdpA with the binding site in KdpB. In the current work, we have reconstituted KdpFABC into lipid nanodiscs and used cryo-EM to image the wild-type pump under turnover conditions. We present a 2.1 Å structure of the E1~P-ADP conformation, which reveals new features of the conduction pathway. This map shows exceedingly strong densities within the selectivity filter and at the canonical binding site, consistent with K^+ bound at each of these sites in this conformation. Many water molecules occupy a vestibule and the proximal end of the tunnel, which becomes markedly hydrophobic and dewetted at the subunit interface. We go on to use ATPase and ion transport assays to assess effects of numerous mutations along this proposed conduction pathway. The results confirm that K^+ ions pass through the tunnel and support the existence of a low affinity site in KdpB for releasing these ions to the cytoplasm. Taken

together, these data shed new light on the unique partnership between a transmembrane channel and an ATP-driven pump in maintaining the large electrochemical K^+ gradient essential for bacterial survival.

Introduction

The Kdp (“K-dependent”) system was discovered in the 1970’s as one of three major systems for maintaining intracellular K^+ levels in *Escherichia coli* (Epstein and Davies, 1970 [↗](#); Rhoads et al., 1976 [↗](#)). The transmembrane gradient of K^+ maintained by these systems represents a fundamental energy source that enables a variety of secondary processes necessary for cell growth and homeostasis (Stumpe et al., 1996 [↗](#)). Further work revealed that the Kdp system consists two overlapping operons with six genes comprising *kdpFABC* and *kdpDE*. The former encodes a hetero-tetrameric pump that binds K^+ with high affinity and uses ATP to transport it into the cell (Gassel et al., 1999 [↗](#); Laimins et al., 1978 [↗](#)). The latter encodes a two-component regulatory system in which KdpD senses osmotic stress (e.g., low extracellular K^+ concentrations) and acts as a histidine kinase to activate KdpE, which in turn induces expression of *kdpFABC* (Walderhaug et al., 1992 [↗](#)). More recently, KdpD was discovered to have a second function as a serine kinase that acts directly on KdpFABC to inhibit its activity when normal K^+ levels have been restored in the growth media (Silberberg et al., 2024 [↗](#)).

KdpA and KdpB subunits descend from two distinct and well-studied membrane transport superfamilies and in KdpFABC, engage in an unprecedented partnership to actively pump K^+ into the cell (Pedersen et al., 2019 [↗](#)). Early studies identified a series of mutants in KdpA affecting the apparent affinity of the Kdp system, thus determining the minimal K^+ concentration at which cells would grow (Buurman et al., 1995 [↗](#); Epstein et al., 1978 [↗](#)). Later, it was recognized that KdpA belongs to the Superfamily of K^+ Transporters (SKT superfamily), which also includes bona fide K^+ channels such as KcsA, TrkH and KtrB (Durell et al., 2000 [↗](#)). As such, KdpA is characterized by pseudo four-fold symmetry and a ‘selectivity filter’ that captures ions from the periplasm (Miller, 2000 [↗](#)). KdpB was recognized to have signature sequences of the P-type ATPase superfamily which are generally known for coupling ATP hydrolysis to transport of substrates across membranes of organisms from all kingdoms of life (Hesse et al., 1984 [↗](#)). This association was confirmed by its formation of an aspartyl phosphate intermediate and characteristic inhibition by vanadate (Puppe et al., 1992 [↗](#)). Function and phylogenetic relationships of KdpF and KdpC are less clear, though they both are suggested to serve as structural chaperones that stabilize the hetero-tetrameric complex (Pedersen et al., 2019 [↗](#)). Like other P-type ATPases, KdpB employs the Post-Albers reaction cycle (**Fig. 1** [↗](#)) involving two main conformations (E1 and E2) and their phosphorylated states (E1~P and E2-P) to drive transport (Albers, 1967 [↗](#); Post et al., 1969 [↗](#)).

A series of X-ray and cryo-EM structures of KdpFABC from *E. coli* have led to proposals of a novel transport mechanism befitting the unprecedented partnership of these two superfamilies within a single protein complex. According to these proposals, K^+ enters the complex from the extracellular side of the membrane through the selectivity filter of KdpA, but is blocked from crossing the membrane. Instead it travels parallel to the membrane plane through a 40-Å long tunnel to reach KdpB. Here the K-ion moves to a binding site in the transmembrane domain of KdpB that resembles canonical sites from other P-type ATPases. Like these other pumps, binding of ions triggers autophosphorylation of a conserved aspartate (Asp307) in the phosphorylation domain (P-domain). Consequent conformational changes in KdpB pinch off the tunnel and induce release of K^+ to the cytoplasm, though the precise location of the low-affinity, release site remains uncertain. In addition, there is uncertainty about the content of the tunnel which has been modeled either as water (Sweet et al., 2021 [↗](#)) or dehydrated K^+ ions (Silberberg et al., 2022 [↗](#), 2021 [↗](#)) in previous work. To shed further light on this unusual transport mechanism, we employ functional and structural approaches to characterize the conduction pathway of K^+ .

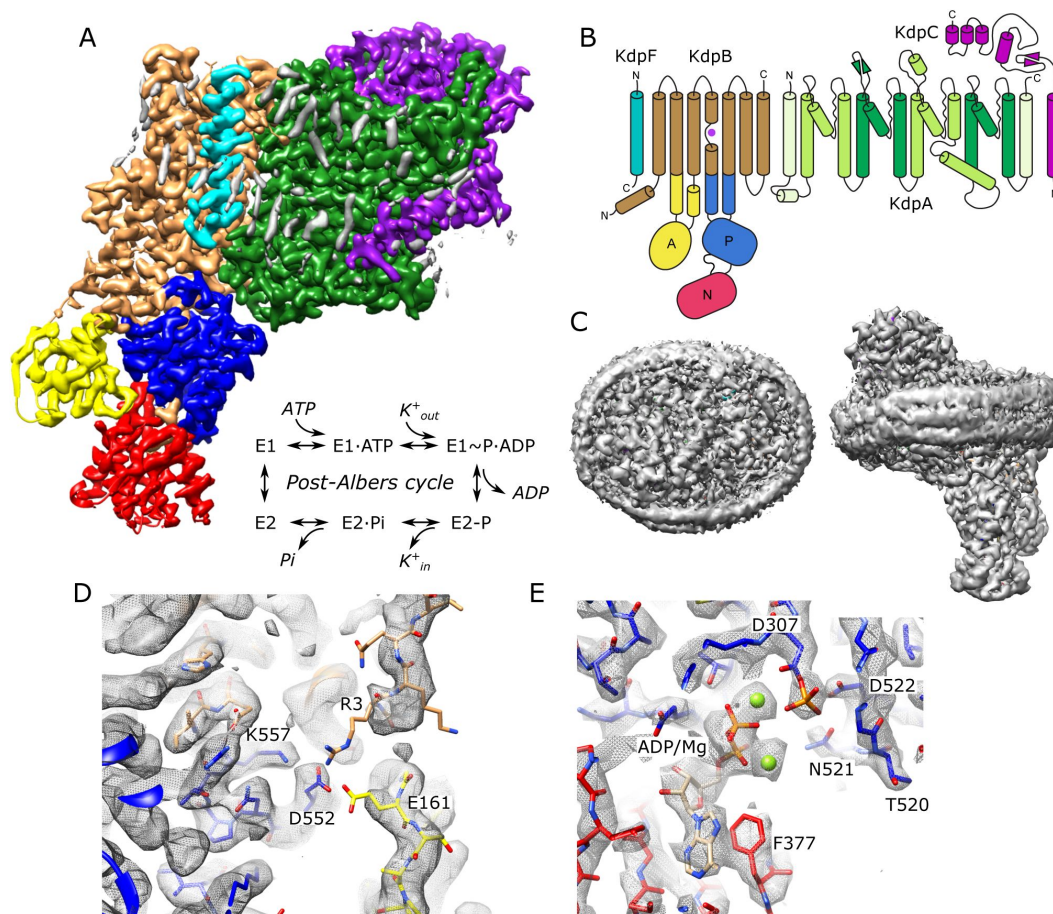


Figure 1.

Cryo-EM structure of KdpFABC in nanodiscs.

(A) Overview of the density map with subunits rendered in conventional colors corresponding to those in panel B. The Post-Albers reaction scheme is shown in the inset. (B) Diagram illustrating the transmembrane topology of KdpFABC. (C) Rendering of the unsharpened map at a lower density threshold illustrates the presence of the membrane scaffolding protein surrounding the nanodisc. (D) Detail of the newly resolved N-terminus of KdpB revealing a potential interaction between Glu161, Arg3, Asp552 and Lys557. (E) Detail of the nucleotide binding site of KdpB showing ADP associated with two Mg^{2+} ions and the phosphorylated catalytic residue: Asp307. Panels A, C and D show the unsharpened map at thresholds of 7, 3 and 4 σ ; panel E shows the sharpened map at 3.9 σ .

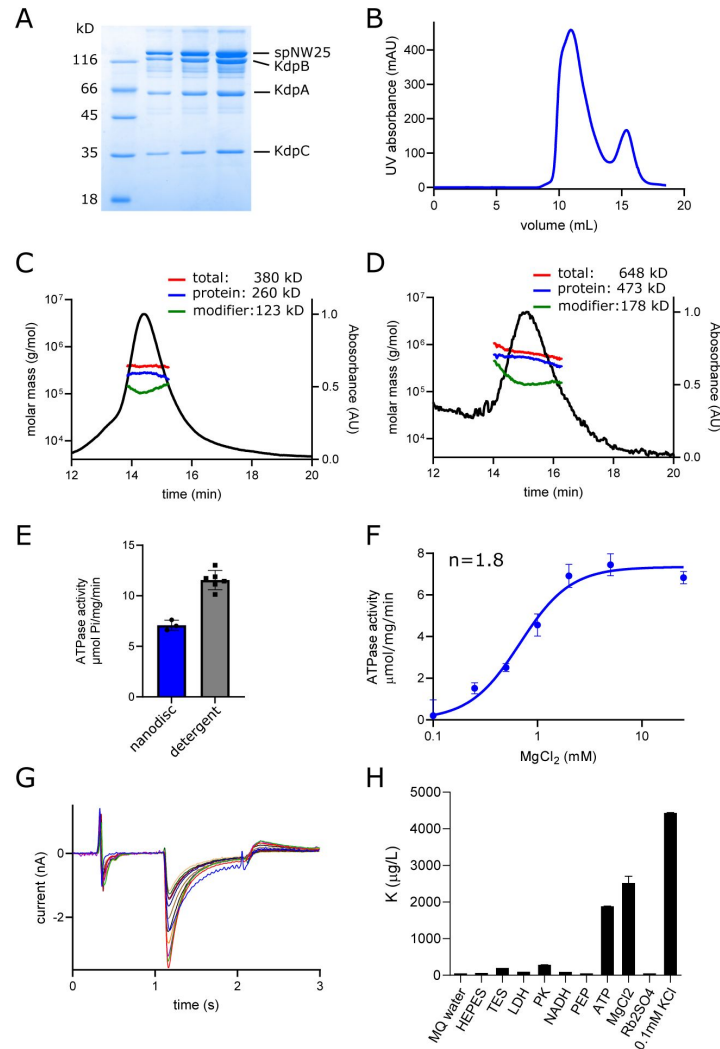


Figure 1 - figure supplement 1.

Biochemical preparation of KdpFABC.

(A) SDS gel of nanodisc sample used for cryo-EM showing the presence of KdpA, KdpB, KdpC and the membrane scaffolding protein, spNW25. KdpF is not visible due to its small size (3 kD); typically, it stains poorly and runs very near the die front of the gel. (B) Elution profile from FPLC size-exclusion column of the sample used for making cryo-EM grids; the main peak is composed of nanodiscs containing KdpFABC and the lower peak contains empty nanodiscs. Slight asymmetry of the main peak is consistent with the presence of single- and double-occupancy nanodiscs seen during image processing. (C-D) MALS analysis during HPLC elution of nanodiscs reconstituted with POPC/DOPA lipids and *E. coli* polar lipids, respectively. This analysis indicates DOPC/DOPA lipids produce smaller nanodiscs containing ~1 KdpFABC molecule (panel c), whereas *E. coli* polar lipids produce larger, more heterogeneous nanodiscs (panel d). (E) ATPase activity is preserved after nanodisc reconstitution, though at a reduced rate attributable to the lipid environment. (F) Mg^{2+} dependence of ATPase activity indicates a Hill coefficient (n) of ~1.8, which is consistent with the two Mg^{2+} ions associated with ADP in the structure. (G) Example of transport current measured by solid supported membrane electrophysiology on WT KdpFABC. (H) Inductively coupled plasma mass spectrometry analysis of stock solutions used for ATPase assay indicate substantial K^+ contamination from ATP and $MgCl_2$ solutions. These data represent the mean and std. dev. of triplicate measurements.

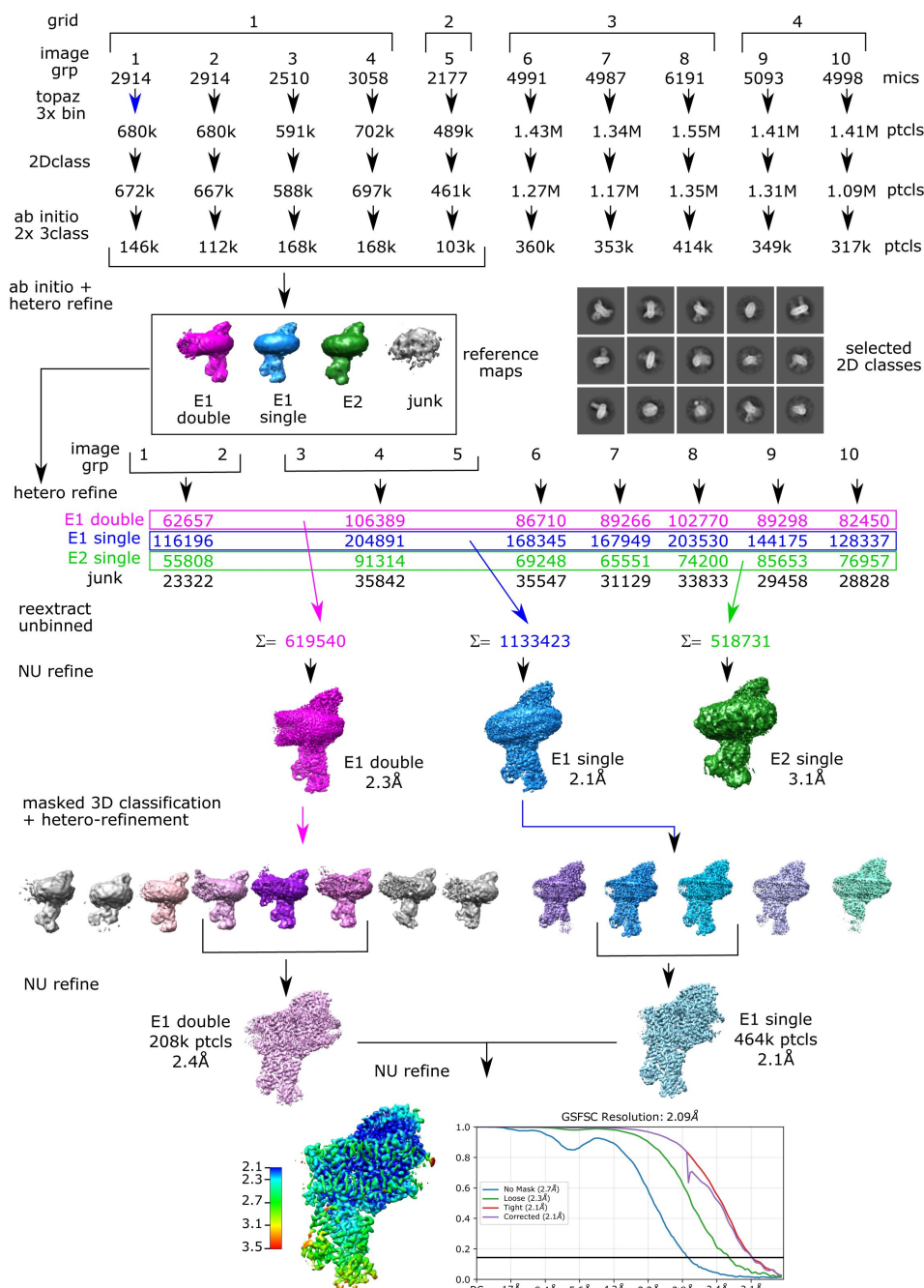


Figure 1 - figure supplement 2.

Cryo-EM processing pipeline.

Four grids from a single sample were imaged to produce a total of ~50,000 micrographs. These were divided into ten groups of 2000-6000 micrographs for initial processing. After motion correction and determination of defocus parameters, particles were picked using TOPAZ and extracted with a box size of 335 Å and 3-fold binning. Multiple rounds of 2D classification (a few examples of 2D classes are shown) and ab initio reconstruction were used to remove false positive picks and empty nanodiscs. A subset of these particles were further classified to generate four main classes representing single occupancy nanodiscs with E1 and E2 conformations, double occupancy nanodiscs which were mainly E1 conformation, and junk. These low-resolution structures were used as references for hetero-refinement of particles partitioned in seven groups. Particles conforming to each conformation were combined and used for masked 3D classification, where the mask encompassed the cytoplasmic domains of KdpB. Particles representing the E1~P-ADP conformation, which represented the largest class, were combined for a final non-uniform refinement job.

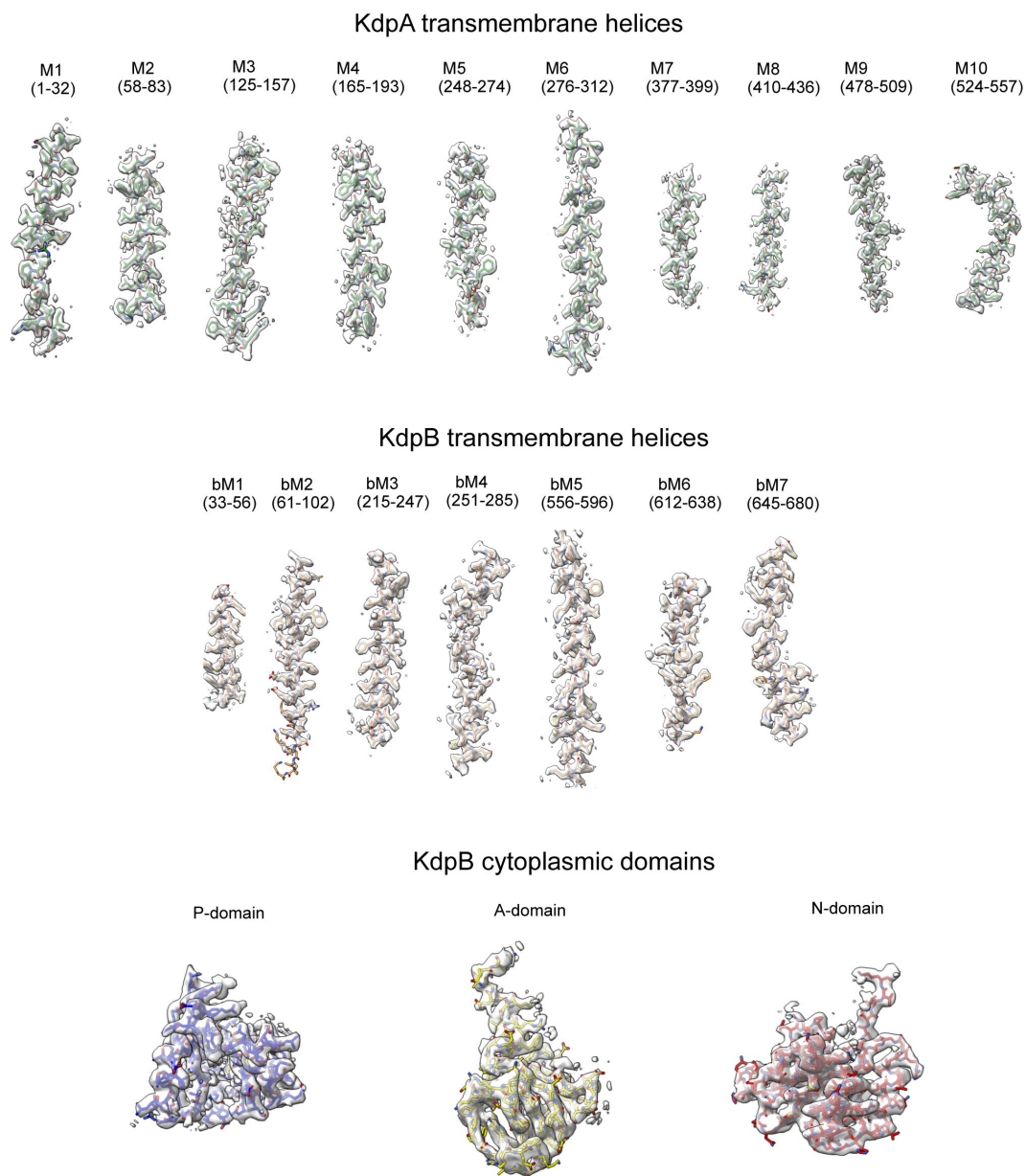


Figure 1 - figure supplement 3.

Cryo-EM densities.

Correlation between map densities and model for individual transmembrane helices from KdpA as well as transmembrane helices and cytoplasmic domains of KdpB. The sharpened map was used for helices and the P-domain with a threshold of 5.7 σ , whereas the unsharpened map was used for the more flexible A- and N-domains at 2.5 σ .

We mutated a series of residues along this pathway to assess their functional impact using both ATPase and transport activities. We also determined a cryo-EM structure at substantially higher resolution (2.1 Å) compared to previous structural studies. In addition, the structure was obtained under the most native conditions to date, namely with uninhibited, wild-type (WT) protein embedded in lipid bilayer nanodiscs under active turnover conditions induced by the addition of K^+ and ATP. The structure reveals new details along the transport pathway, including clear evidence of tunnel hydration as well as K^+ occupancy in the selectivity filter in KdpA and at the canonical binding site in KdpB. Map densities in the catalytic domains of KdpB unambiguously show phosphorylation of Asp307 in the P-domain and binding of ADP together with two Mg^{2+} ions in the nucleotide binding domain (N-domain), confirming the conformation to be E1~P-ADP. Effects of mutation at the entrance of the selectivity filter and within the tunnel confirm the transport pathway. In addition, mutations at the canonical ion site in KdpB as well as at a proposed release site provide experimental evidence for the allosteric changes that drive transport.

Results

Structure of KdpFABC in the E1~P conformation

For expression, we used an *E. coli* strain lacking the dual-function kinase KdpD that is responsible for post-translational inhibition of KdpFABC depending on the growth conditions. In particular, phosphorylation of Ser162 by KdpD confounded previous attempts to survey mutations along the putative transport pathway (Sweet et al., 2020) and produces off-cycle, inhibitory conformations (Huang et al., 2017; Silberberg et al., 2022). As a result, many previous structures (Huang et al., 2017; Silberberg et al., 2021; Stock et al., 2018; Sweet et al., 2021) feature the S162A mutation to avoid inhibition rather than the fully WT protein used for the current work. Our WT complex displayed high levels of K^+ -dependent ATPase activity and generate robust transport currents (Fig. 1 – figure suppl. 1). Unlike previous structural analyses, which relied on detergent-solubilized preparations, we reconstituted the purified complex into lipid nanodiscs to provide a more native lipid environment. Rather than use inhibitors to stabilize a particular state for cryo-EM imaging, we added ATP and froze the sample under active turnover. We collected a large data set (~50,000 micrographs) and used 3D classification to segregate various conformational states (Fig. 1 – figure suppl. 2). In the paper, we report on the most highly populated state (44% of particles), which resulted in the highest resolution reported for KdpFABC to date (2.1 Å, Table 1) in which all transmembrane helices and cytoplasmic domains are well resolved (Fig. 1 – figure suppl. 3). Fig. 1 shows an overview of the structure, including the belt formed by the nanodisc scaffolding protein which is visible at a lower density threshold. Densities at the catalytic site indicate that Asp307 in the P-domain is phosphorylated and that ADP is bound within the N-domain (Fig. 1e), consistent with the E1~P-ADP conformation. Two Mg^{2+} ions are associated with ADP, consistent with a Hill coefficient of 1.8 determined from Mg^{2+} titration of ATPase activity (Fig 1 – figure suppl. 1f). The overall conformation is quite similar to previous structures of the E1 state in detergent (RMSD for Cα atoms of 0.8-1.5 Å) including E1~P states captured under turnover conditions (Silberberg et al., 2022) and the E1-ATP state stabilized with AMPPCP (Sweet et al., 2021). Unlike previous structures, our map resolves most of the N-terminus of KdpB, revealing an interaction involving Glu161 from the A-domain, Arg3 from the N-terminus as well as Asp552 and Lys557 from the P-domain (Fig. 1d). This interaction is notable given the role of Glu161 in hydrolysis of the aspartyl phosphate in the subsequent E2-P state. Furthermore, post-translational regulation of KdpFABC involves phosphorylation of the neighboring residue, Ser162, which leads to inhibition of this dephosphorylation step and thus cessation of transport (Sweet et al., 2020).

Deposition	
PDB	9OC4
EMDB	EMD-70308
Data collection and processing	
Magnification	130kx
Voltage (kV)	300
Electron exposure (e ⁻ /Å ²)	50
Defocus range (μm)	1.0-3.0
Pixel size (Å)	0.93
Symmetry imposed	C1
Initial particle images (no.)	10,382,000
Final particle images (no.)	672,405
Map resolution	2.09
FSC threshold (Å)	0.143
B factor (Å ²)	59.2
Resolution range (Å)	2-3.5
Model Refinement	
Model composition	
Non-hydrogen atoms	11068
Protein residues	1455
Ligands	7
RMS deviations	
Bond lengths (Å)	0.005
Bond angles (°)	1.026
Validation	
MolProbity score	1.45
Clashscore	5.91
Rotamer outliers (%)	1.23
CaBLAM outliers (%)	1.46
Rama-Z score	0.63
Ramachandran plot	
Favored (%)	97.71
Allowed (%)	2.29
Disallowed (%)	0.00
Model vs. Data CC (mask)	0.86

Table 1.

Structure determination of KdpFABC.

Passage of K⁺ through the selectivity filter of KdpA

Selectivity filters are a defining feature of the SKT superfamily that includes KdpA. The selectivity filter from KcsA has four binding sites denoted S1-S4 (Morais-Cabral et al., 2001 [DOI](#)) and previous structures of KdpFABC have consistently shown strong density only at the equivalent S3 site. In the current map, however, clear densities are visible at all four sites, likely due to increased resolution (Fig. 2a [DOI](#)), but density at S3 remains much stronger. In fact, this S3 site contains the strongest densities in the entire map, measuring 7.9x higher than the threshold used for Fig. 2a [DOI](#) (Fig. 2 - figure suppl. 1a [DOI](#)). This suggests that S3 is occupied by a well-ordered K⁺ ion. The higher resolution also allows accurate placement of carbonyl atoms in the selectivity filter and the resulting coordination geometry (Fig. 2 - figure suppl. 1b [DOI](#)) is consistent with S3 acting as the primary binding site for K⁺. S3 is tightly coordinated by the eight surrounding carbonyl atoms at an average distance of 2.8 Å (s.d. 0.2 Å), which is consistent with the expected center-to-center distance for K⁺ and oxygen atoms of its hydration shell (Naranjo et al., 2016 [DOI](#)). At S2, coordination is noticeably looser with the four lower carbonyls at an average of 3.0 Å (s.d. 0.9 Å) and the four upper carbonyls at 3.6 Å (s.d. 0.1 Å). The S1 position is only coordinated from below at an average distance of 3.1 Å (s.d. 0.3 Å) consistent with its role in dehydrating the incoming K⁺ ion. Similarly, the S4 position is well coordinated from above at 3.1 Å (s.d. 0.2 Å) with only 2 oxygen ligands below, both of which come from side chain oxygen atoms instead of main-chain carbonyls. Additional densities provide coordination at both S1 and S4, which we have modeled as water given that their density levels are similar to the surrounding protein.

The functional role of this selectivity filter has previously been probed by mutagenesis of Gln116 at the entrance and Gly232 in the middle (Buurman et al., 1995 [DOI](#)). We have expanded studies of these sites by comparing rates of ATP hydrolysis and K⁺ transport to assess coupling, and by substituting a variety of amino acids to explore the role of electrostatics on ion binding.

Consistent with previously work, K⁺ titrations combined with measurement of ATPase (Fig. 2b [DOI](#)) and transport (Fig. 2c [DOI](#)) activities show that WT has affinity in the micromolar range which is severely reduced by mutations of either Gln116 or Gly232. Background activity in the absence of added K⁺ was consistently observed for WT KdpFABC which we attribute to K⁺ contamination found in commercially available sources of ATP and MgCl₂ (Fig. 1 - figure suppl. 1h [DOI](#)).

Nevertheless, the overall correspondence between ATPase and transport titrations indicate that energy coupling is maintained despite differences in apparent affinity. Structural work has shown that G232D is more disruptive to the structure of the selectivity filter (Silberberg et al., 2021 [DOI](#)), which may explain the lower V_{max} seen for this mutant. Indeed, detergent solubilized preparations proved to be relatively unstable during extended ATPase assay titrations. Previous studies also showed a reduction of V_{max} for Q116R, but that work employed the S162A mutation on KdpB to prevent post-translational phosphorylation of that residue and consequent formation of an inhibited complex (Sweet et al., 2020 [DOI](#)). Our use of the KdpD-knockout strain to produce unphosphorylated protein shows that Q116R can achieve WT levels of activity when Ser162 is retained.

Next, we tested the hypothesis that lower apparent affinity of the Q116R substitution was simply due to charge repulsion from an added positive charge at the entrance of the selectivity filter, whereas the G232D substitution results in a more severe disruption of the selectivity filter architecture. Indeed, mapping electrostatic charge on the surface of the WT protein shows an abundance of negative charge at the entrance to the selectivity filter, which would effectively attract substrate cations (Fig. 2d [DOI](#)). Consistent with such an electrostatic effect, we found that Q116E behaves comparably to WT, with substantial background activity and K_m < 10 μM (Fig. 2e [DOI](#)). Furthermore, the K_m for Q116H was pH dependent with decreased apparent affinity (higher K_m) as the His became protonated and thus positively charged (Fig. 2f [DOI](#)). For G232D, we

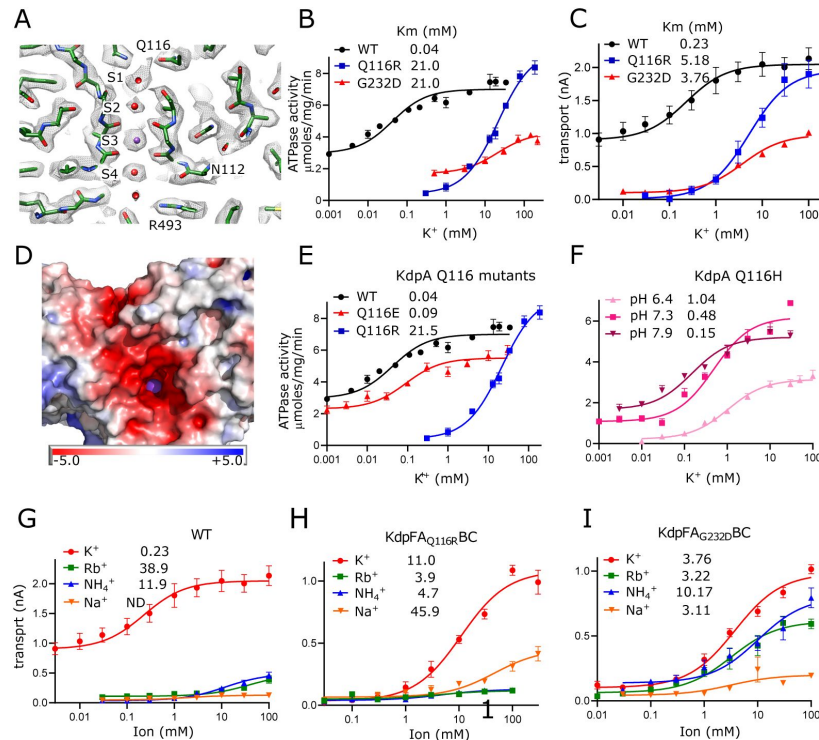


Figure 2.

Selectivity filter of KdpA.

(A) The map reveals densities at all sites (S1-S4) of the selectivity filter. Because the density at the S3 site is the highest in the entire map (**Fig. 2 - figure suppl. 1a**), it has been assigned as a K^+ ion (purple sphere). Grey mesh corresponds to the sharpened map at 6.5σ . (B-C) ATPase and transport assays (respectively) illustrate that Q116R and G232D mutations lower the apparent affinity of the pump (K_m values in mM shown in the legend). Background activity is consistently observed for the WT, which is likely due to K^+ contamination in stock solutions (**Fig. 1 - figure suppl. 1h**). The location of Gln116 is shown in panel a, G232 is identified in **Fig. 2 - figure suppl. 1b**. (D) Electrostatic charge at the entrance to the selectivity filter of the WT pump shows strong negative charge. A K^+ ion is visible in the selectivity filter as a purple sphere. (E-F). Substitution of Glu116 has differing effects that reflect changes in net charge at the entrance to the selectivity filter. (G-I) Transport assays with various monovalent cations show that the strong selectivity of the WT pump is compromised by the G232D mutation but not by the Q116R mutation. **Table 2** shows that measurements from transport assays were broadly consistent with these ATPase data.

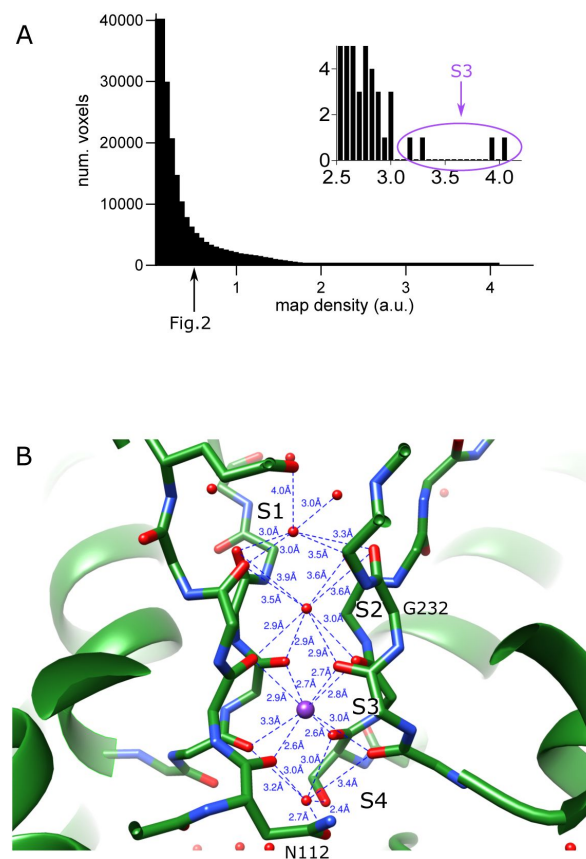


Figure 2 - figure supplement 1.

Selectivity filter of KdpA.

(A) Distribution of raw densities for the entire map. The arrow indicates the threshold used for rendering the map in [Fig. 2](#) (0.5 which corresponds to 6.5σ). The inset shows the extreme tail in the density distribution; the highest four densities (circled) correspond to the K^+ ion modeled at the S3 site. (B) Coordination of densities in the selectivity filter in the refined model. The regular network of bonds at the S3 site suggest that this is the high affinity binding site for K^+ , whereas less regular coordination at other sites is consistent with water.

confirmed previous observations (Schrader et al., 2000 [↗](#); van der Laan et al., 2002 [↗](#)) that this mutation not only reduced affinity, but also ion selectivity relative to WT and Q116R. Whereas the latter are high selectivity for K^+ , G232D also displays robust activity with Rb^+ and NH_4^+ (**Fig. 2g-i** [↗](#)). Notably, very little activity was seen with Na^+ . In general, measurements of ATPase and transport were comparable, indicating that these mutants remain energetically coupled (**Table 2** [↗](#)).

Vestibule in KdpA

Immediately after the selectivity filter there is a large cavity analogous to the vestibule seen in K^+ channels (Doyle et al., 1998 [↗](#)). However, unlike channels where it leads directly to the cytosol, the vestibule of KdpFABC leads to the inter-subunit tunnel. There are many side chain oxygen atoms in this region that produce a generally hydrophilic environment and our new structure reveals numerous discrete densities within this cavity (**Fig. 3a** [↗](#)). Because their magnitude is comparable to or lower than the surrounding protein side chains we model them as water molecules, thus producing a highly hydrated vestibule. Glu370 and Arg493 are the only charged residues within the membrane core of KdpA and are positioned on either side of the S4 site. Glu370 lies at the entrance of the tunnel whereas Arg493 lines the back wall of the vestibule and we used mutagenesis in conjunction with ATPase and transport assays to probe their roles (**Fig. 3b-c** [↗](#), **Table 3** [↗](#)). Changes to Arg493 generally increase K_m (lower apparent affinity) without affecting V_{max} , with Met and Gln substitutions having greater effects than charge reversal (R493E). Neutralization of Glu370 with a Gln or His substitution has a similar effect, but reversing the charge (E370K) severely reduces both apparent affinity and V_{max} , suggesting an important functional role for negative charge in this region. The E370K/R493E double mutant partially restores V_{max} but the apparent affinity is still severely compromised, suggesting that location of charge is essential for efficient movement of K^+ into the tunnel.

Inter-subunit tunnel

A continuous tunnel has been documented in KdpFABC structures in the E1 conformation (Huang et al., 2017 [↗](#)) and has been postulated to provide the conduction pathway for K^+ ions as they move from the selectivity filter in KdpA to the canonical ion binding sites in KdpB (Stock et al., 2018 [↗](#); Sweet et al., 2021 [↗](#)). In our new structure, the discrete densities seen in the vestibule continue into this tunnel, forming a well coordinated network with spacings between 2.5 and 3.5 Å. Whereas previous structures show 5-7 discrete densities extending from the vestibule into the tunnel (Silberberg et al., 2022 [↗](#), 2021 [↗](#); Sweet et al., 2021 [↗](#)), our new structure reveals >30 spherical densities in this region, all of comparable strength to the surrounding protein density. These densities, which we have modeled as water, are most prevalent near the vestibule, which is the wider part of the tunnel, but then disappear completely at the subunit interface, which is the narrowest part of the tunnel and also distinctly hydrophobic (**Fig. 4** [↗](#)).

In order to provide experimental evidence for K^+ transport through the tunnel, we made a series of substitutions to Val496 in KdpA. This residue resides near the widest part of the tunnel and is fully exposed to its interior (**Fig. 4a** [↗](#)). We made substitutions to increase its bulk (V496M and V496W) and to introduce charge (V496E, V496R and V496H). We used the AlphaFold-3 artificial intelligence structure prediction program (Jumper et al., 2021 [↗](#)) to predict structures of these mutants and to evaluate their impact on tunnel dimensions. This analysis predicts that V496W and V496R reduce the radius to well below the 1.4 Å threshold required for passage of K^+ or water (**Fig. 4c** [↗](#)); V496E and V496M also constrict the tunnel, but to a lesser extent.

Measurements of ATPase and transport activity show that negative charge (V496E) has no effect. The longer side chain of V496M reduces transport modestly with no effect on ATPase activity. V496R, which introduces positive charge, completely abolishes activity. V496W and V496H reduce

mutants	ATPase ^a Km (mM)	transport ^b Km (mM)
WT		
K ⁺	0.04 ± 0.01	0.23 ± 0.10
Rb ⁺	30.71 ± 13.45	38.88 ± 22.45
NH ₄ ⁺	- ^c	- ^c
Na ⁺	- ^c	- ^c
G232D		
K ⁺	21.17 ± 2.66	3.76 ± 0.58
Rb ⁺	- ^d	3.22 ± 1.06
NH ₄ ⁺	- ^d	10.17 ± 4.16
Na ⁺	- ^c	- ^c
Q116R	21.45 ± 2.66	5.19 ± 0.12
Q116E	0.09 ± 0.02	0.11 ± 0.46

^a average of N=3 independent measurements with SEM

^b average of N=4 independent measurements with SEM

^c little activity above baseline prevented Km determination

^d protein instability in detergent solution interfered with accurate Km determination

Table 2.

Summary of apparent affinities of the selectivity filter mutations measured by ATPase and transport assays.

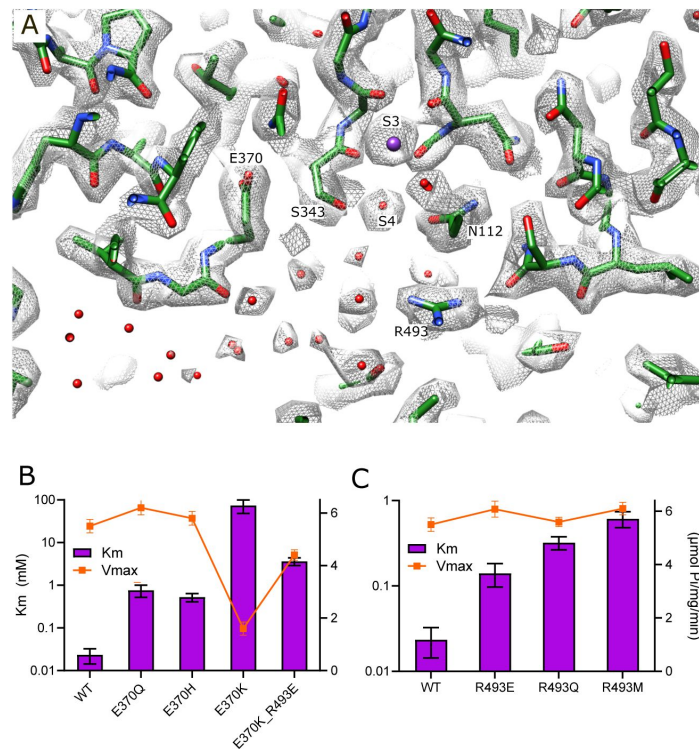


Figure 3.

Vestibule of KdpA.

(A) The map reveals numerous spherical densities immediately below the selectivity filter and leading into the tunnel that connects KdpA with KdpB. Glu370 and Arg493 are the only charged residues in the transmembrane domain of KdpA and reside on either side of this vestibule. Grey mesh corresponds to the sharpened map at 6.5 Å. (B) ATPase assays of E370Q and E370H show that neutralization of charge on Glu370 reduces the apparent affinity without affecting Vmax, but that reversing the charge with E370K almost completely abolishes activity. The double mutant E370K/R493E partially restores activity, but with much lower apparent affinity and Vmax. (C) ATPase assays show that mutation of Arg493 has relatively modest effects on apparent affinity without affecting Vmax. Note that the axis with Km is logarithmic. **Table 3** shows that measurements from transport assays were broadly consistent with these ATPase data.

mutants	ATPase ^a Km (mM)	transport ^b Km (mM)
WT	0.023 ± 0.01	0.23 ± 0.10
E370Q	0.76 ± 0.24	1.32 ± 0.58
E370H	0.53 ± 0.11	0.35 ± 0.12
E350K	74.95 ± 25.8	- ^c
E350K_R493E	3.63 ± 0.74	0.43 ± 0.08
R493E	0.14 ± 0.04	0.50 ± 0.23
R493Q	0.32 ± 0.06	1.52 ± 0.68
R493M	0.61 ± 0.13	0.47 ± 0.08

^a average of N=3 independent measurements with SEM

^b average of N=4 independent measurements with SEM

^c little activity above baseline prevents Km determination

Table 3.

Summary of apparent affinities of the vestibule mutations measured by ATPase transport assays.

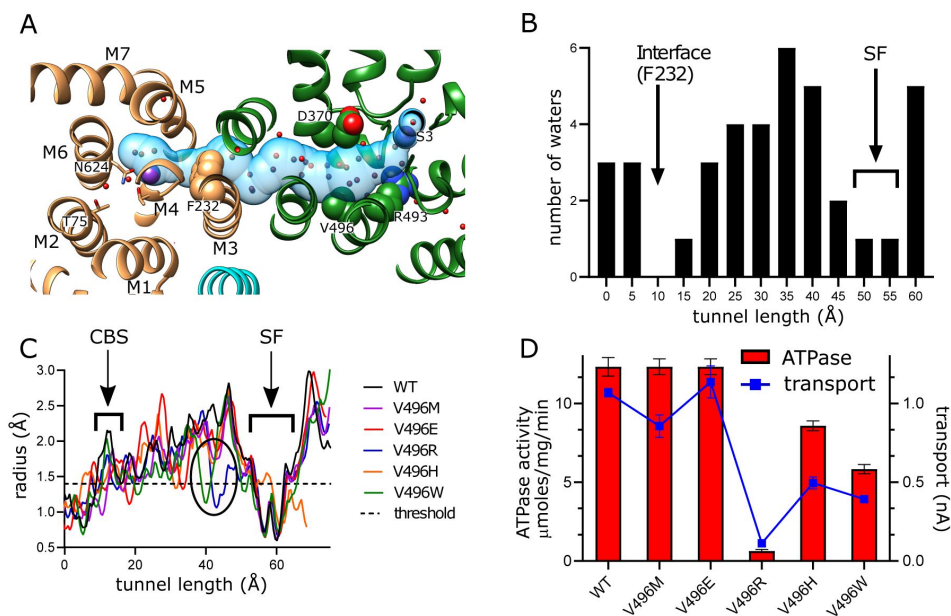


Figure 4.

Tunnel connecting KdpA and KdpB.

(A) The intersubunit tunnel is shown as a transparent blue surface running from the selectivity filter on the right (S3) to the canonical binding site on the left (near N624). Non-protein densities within the tunnel have been modeled as water molecules (red spheres). Asp370, Arg493 in the vestibule and Phe232 at the subunit interface are shown as space-filling models and K^+ as purple spheres. As in other figures, KdpA helices are green and KdpB helices are brown; KdpF is cyan. (B) The distribution of water molecules within the tunnel shows that the vestibule is well populated whereas the subunit interface is quite hydrophobic. Values along the x-axis correspond to the contour length along the tunnel. (C) Tunnel profiles for the experimentally determined model (WT) and predicted models for the Val496 mutants. The horizontal dotted line at a radius of 1.4 Å represents the conventional limit for passage of water or K^+ ions. Note that both V496R and V496W constrict the tunnel. SF and CBS indicate locations of the selectivity filter and canonical binding site, respectively. (D) ATPase and transport assays of the Val496 mutants in 150 mM K^+ showing that introduction of positive charge (V496R) abolishes activity.

both transport and ATPase activity by about half, perhaps due to steric hindrance for the former and partial protonation for the latter.

Canonical ion binding site in KdpB

KdpB belongs to the P-type ATPase superfamily where ion binding occurs in a membrane embedded chamber next to a conserved proline along the M4 helix (Pro264 in KdpB). This proline interrupts the M4 helix and allows free carbonyls to form a binding site along its axis. For the well-studied P-type ATPases Sarcoendoplasmic Reticulum Ca^{2+} -ATPase (SERCA) and Sodium-Potassium ATPase (Na,K-ATPase), this canonical site is referred to as Ca2 and Na2, respectively and additional ions are accommodated in the cavity between M4, M5 and M6 (Ca1 site for SERCA and Na1 plus Na3 sites for Na,K-ATPase) (Kanai et al., 2013 [\[1\]](#); Toyoshima et al., 2000 [\[2\]](#)) (**Fig. 5 – figure suppl. 1** [\[3\]](#)). In the original X-ray structure of an inhibited conformation of KdpFABC, a spherical density was present at the site corresponding to Ca2/Na2 and was modeled as water due to the lack of anomalous K signal at that location (Huang et al., 2017 [\[4\]](#)). In subsequent E1-ATP structures of KdpFABC stabilized with AMPPCP, densities were seen in the cavity harboring the Na1 site and were modeled either as water (Sweet et al., 2021 [\[5\]](#)) or K^+ (Silberberg et al., 2021 [\[6\]](#)). A recent structure also captured under turnover conditions and ascribed to the E1~P conformation (7ZRK) showed a weak, elongated density leading toward the Ca2/Na2 site, which was modeled as K^+ , but was not fully resolved from the end of Lys586 which resides in this region (Silberberg et al., 2022 [\[7\]](#)). A second E1~P structure (7ZRM) from the same study did not resolve any densities in this area. Our new structure unambiguously resolves a discrete spherical density precisely at the canonical Ca2/Na2 site that is well coordinated by main chain carbonyl atoms from M4 (Val260, Cys261, Ile263) as well as the side chains of Thr265 and Asn624 with an average distance of $2.8 \pm 0.2 \text{ \AA}$ (**Fig. 5a** [\[8\]](#), **Fig. 5 – figure suppl. 2a** [\[9\]](#)). Given that this is the strongest density in KdpB, measuring 5.6x higher than the map densities shown in **Fig. 5** [\[10\]](#) (**Fig. 5 – figure suppl. 2b** [\[11\]](#)), we have modeled it as K^+ . Additional densities within this chamber have similar magnitude to the surrounding protein residues and have been modeled as water, including two that coordinate the central K^+ at distances of 2.9 and 3.4 $\text{\AA}$. These observations strongly support the canonical Ca2/Na2 site as the primary K^+ binding site for KdpB in the high energy E1~P-ADP conformation.

Asp583 and Lys586 are two conserved residues on M5 that have previously been shown to be essential for transport (Bramkamp and Altendorf, 2005 [\[12\]](#)). We made a series of substitutions to further probe their roles (**Fig. 5b** [\[13\]](#)). Most of these mutants show little to no activity, consistent with their crucial roles in shepherding K^+ through this binding site. This includes neutralization (D583N, K586Q) and reversal (D583K, K586E) of the charge. Swapping of the charge with a double mutant (D583K and K586D) also led to inactivation, showing that the geometry of the site is important. Interestingly, D583A mutant displayed substantial ATPase activity (30% of WT) but no transport, indicating that this particular mutation interfered with energy coupling.

K^+ release site in KdpB

In other P-type ATPases, conformational changes convert the high affinity canonical binding site to a low affinity binding site with access to the opposite side of the membrane, thus allowing ions to be released directly from these sites (Dyla et al., 2020 [\[14\]](#)). For KdpFABC, however, analogous conformational changes have not been observed and the low-affinity release site for K^+ ions is not well established. We previously proposed a novel mechanism (Sweet et al., 2021 [\[15\]](#)) in which Lys586 swings into the canonical Ca2/Na2 binding site (previously called the Bx site) in the transition to an E2 state, thus pushing K^+ into an alternate chamber (previously called the B2 site) between M1, M2 and M4 (**Fig. 6 – figure suppl. 1a** [\[16\]](#)). This alternate chamber has access to the cytoplasm via a water filled cavity and the relatively weak coordination of ions within this chamber is consistent with a low affinity release site. Although our new structure does not represent the state relevant for release, a non-protein density is seen in this chamber next to Thr75, which was previously postulated as the only potential ligand for ions or water in this site (**Fig. 6a** [\[17\]](#)).

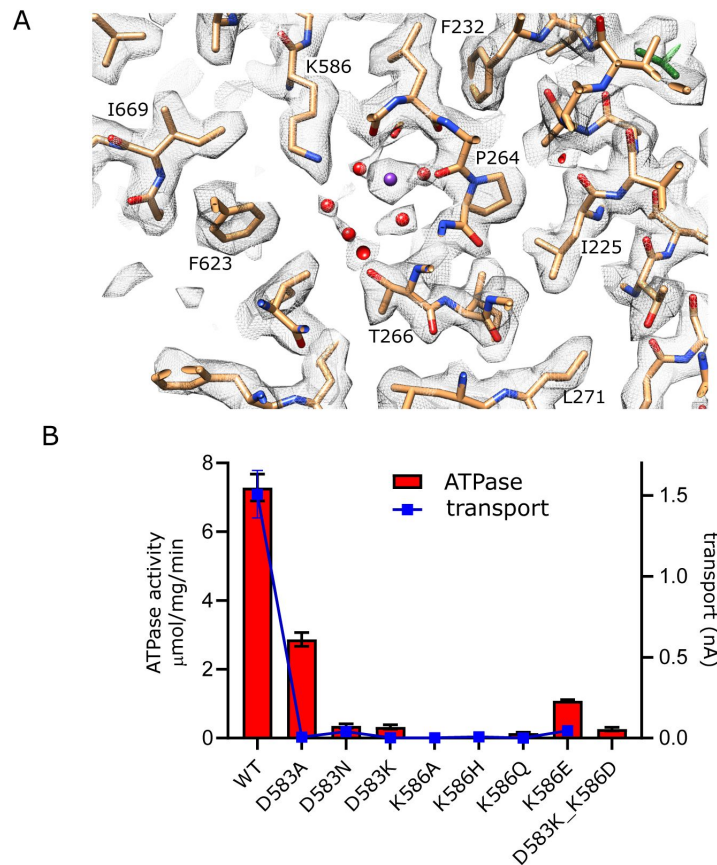


Figure 5.

Canonical binding site in KdpB.

(A) Several non-protein densities are visible at the canonical binding site, the most prominent of which occupies the conserved Na₂/Ca₂ site from Na,K-ATPase and SERCA (figure suppl. 1). This site has the highest density in KdpB (figure suppl. 2b) and thus has been modeled as K⁺ (purple sphere), whereas the others are modeled as water (red spheres). Grey mesh corresponds to the sharpened map at 5.8 σ. (B) Asp583 and Lys586 are the only charged residues in the transmembrane domain of KdpB. ATPase and transport assays in 150 mM K⁺ show that mutation of these residues is generally not tolerated, except for the D583A mutation which generates an uncoupled phenotype (ATPase activity without transport).

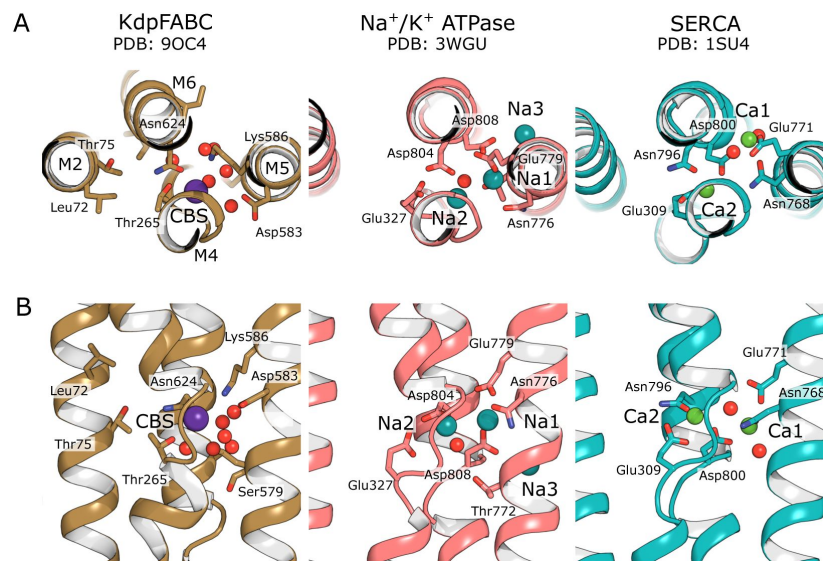


Figure 5 - figure supplement 1.

Canonical binding sites of P-type ATPases.

The primary K⁺ binding site in KdpB is structurally conserved with ion binding sites for Na⁺ and Ca²⁺ in Na,K-ATPase and SERCA. Whereas multiple ions are transported in each cycle by these related P-type ATPases, only a single ion appears to bind to KdpB at a position equivalent to the Na2 and Ca2 sites. Like those sites, the K⁺ in KdpB is coordinated by carbonyl oxygens exposed by the unwinding of the M4 helix near a conserved proline (figure suppl. 2). Additional ions bind in a pocket between M4, M5 and M6 in Na,K-ATPase and SERCA, which is occupied by weaker densities in KdpB that are modeled as water.

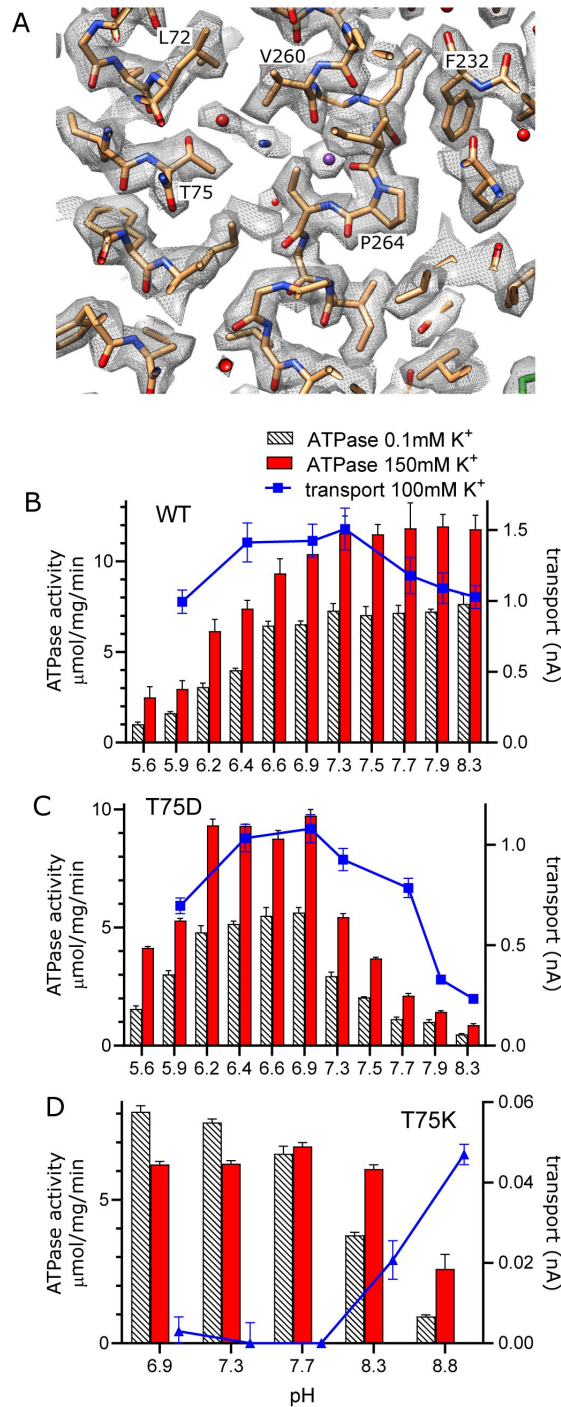


Figure 6.

Exit site from KdpB.

(A) The map shows a water filled cavity next to Thr75 that leads to the cytoplasm (sharpened map at 6.5 Å). (B-C) pH dependence of the T75D mutant is shifted relative to WT, consistent with the idea that high pH generates negative charge in this cavity and inhibits K⁺ release. (D) The T75K substitution not only interferes with transport, but also produces an uncoupled phenotype in which ATPase activity is not dependent on K⁺. At high pH there is a small recovery of transport activity as well as K⁺-dependence of ATPase activity, suggesting that these effects result from the positive charge from the lysine substitution.

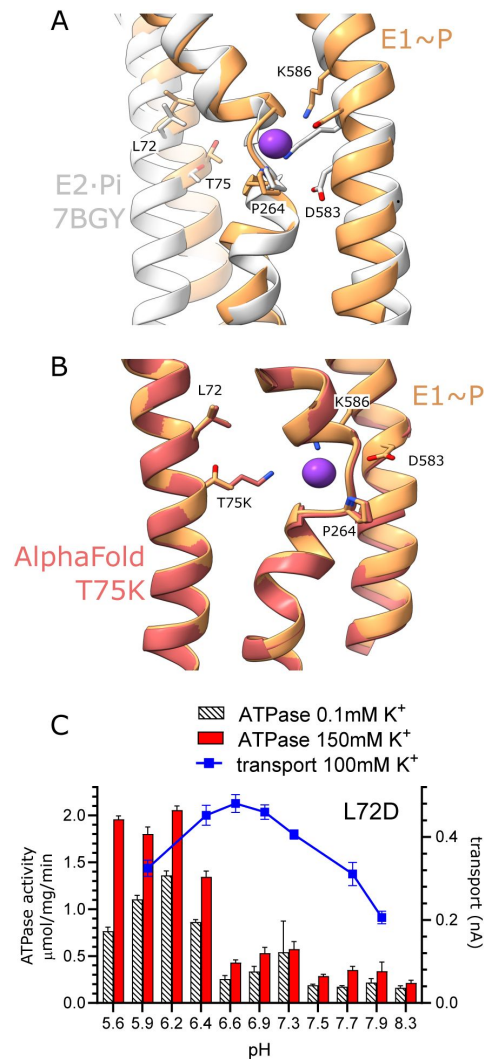


Figure 6 - figure supplement 1.

Exit site for K⁺ from KdpB.

(A) Comparison of the canonical binding site from E1~P (this study) and E2~Pi determined previously by Sweet et al. (Sweet et al., 2021) (PDB code 7BGY). In particular, movements of M5 cause Lys586 to swing into the canonical binding site in E2~Pi, thus displacing the K⁺. (B) Comparison of the exit site from wild-type E1~P model with a predicted model for the T75K mutant. In this predicted structure, the lysine side chain swings into the canonical binding site, thus potentially acting as a built in ion to stimulate ATPase activity in the absence of K⁺. (C) pH dependence of both ATPase activity and transport from the KdpB-L72D mutant. Like the T75D mutant, the pH dependence of activity is shifted towards low pH. Although the rates are lower than WT and T75D, the K⁺ dependence of ATPase activity indicates that energy coupling is intact.

To experimentally test this proposed release site, we substituted Thr75 with either Asp or Lys and measured effects on both ATPase and transport activity (**Fig. 6b-d**). For these measurements, we varied the pH to alter the charge at this site and looked for corresponding effects on the activity. At neutral pH, T75D produced K^+ -dependent ATPase and transport activities comparable to WT, indicating that the Asp substitution does not interfere with turnover or energy coupling. However, maximal activity of T75D is shifted toward more acidic pHs and, unlike WT, fall off dramatically at basic pH. Assuming that the pKa of T75D is shifted by this intramembrane environment, these results are consistent with the idea that negative charge generated by deprotonation of the Asp residue interferes with K^+ release. Indeed, pKa shifts have previously been studied both experimentally (Gayen et al., 2016; Isom et al., 2010; Morrison et al., 2015) and computationally (Henderson et al., 2020; Panahi and Brooks, 2015), showing pKa values up to 9 for acidic residues. In the case of T75D, we used PROPKA (Li et al., 2005) and a predicted model for this mutant, which suggested a substantial shift in pKa toward neutral pH (pKa of 6.8).

As expected, the T75K mutation behaves quite differently. At pH<8, T75K has robust ATPase activity that does not depend on the presence of K^+ and does not generate any transport current (**Fig. 6d**). This behavior is consistent with a loss of energy coupling. At pH>8, K^+ -dependence of ATPase activity is restored and a small transport current is measured, suggesting partial recovery as charge on the Lys is reduced. Although the pKa of the lysine side chain in solution is 10.8, it would be expected to shift toward neutral pH in the membrane environment, again supported by a PROPKA calculation based on the predicted model for T75K (pKa of 6.3). Inspection of this predicted model reveals that the side chain of the lysine is inserted into the canonical binding site, analogous to the behavior of Lys586 in the E2 conformation (**Fig. 6 – figure suppl. 1b**). This suggests that T75K acts as a 'built-in cation' (Pedersen et al., 2007) that is detected at the canonical binding site to constitutively induce ATPase activity by KdpB, even in the absence of K^+ . Leu72 is another residue deeper in this release pocket and the L72D substitution was also tested. Although overall activity is diminished, it displays a pH dependence similar to T75D (**Fig. 6 – figure suppl. 1c**).

Discussion

We have used structure determination, mutagenesis and activity measurements to study the transport pathway for K^+ ions through KdpFABC. For cryo-EM analysis, we purified WT protein free from inhibitory effects of post-translational serine phosphorylation, reconstituted it into lipid nanodiscs, and imaged this sample under turnover conditions, thus providing the most native conditions visualized by structural biology to date. The largest population of particles adopted the E1~P-ADP conformation, consistent with it preceding the rate limiting step in the transport cycle. The resolution of the corresponding structure (2.1 Å) is much improved over previous structural studies, revealing new features relevant to this pathway.

The intersubunit tunnel is arguably one of the most intriguing elements of the KdpFABC complex. Although it has been postulated to conduct K^+ , experimental evidence has been lacking. However, our mutagenesis studies now firmly support this mechanism and illustrate key properties of this K^+ conduit. To start, the external surface of the selectivity filter carries negative charge, thus serving to attract ions. Mutations that reduce the net charge lower the apparent affinity, thus explaining the well-known effects of the Q116R mutant (Buurman et al., 1995).

Once in the selectivity filter, the ion appears to have a strong preference for the S3 site. This conclusion is supported by X-ray anomalous scattering (Huang et al., 2017; Madapally et al., 2025), the consistent presence of density at this site in previous lower-resolution structures (Silberberg et al., 2022, 2021; Sweet et al., 2021), and the exceedingly strong density in current map (**Fig. 2 – figure suppl. 1c**). Furthermore, coordination geometries in the selectivity filter are consistent with preferential binding at S3 and suggest that high apparent affinity and

selectivity towards K^+ comes from this site, and not from the canonical binding site in KdpB. Unlike K^+ channels, the relatively slow turnover of KdpFABC ($10\text{--}100\text{ sec}^{-1}$) allows for equilibrium binding at the selectivity filter, which may explain its ability to distinguish K^+ from its congener Rb^+ (Huang et al., 2017 [DOI](#); Liu and Lockless, 2013 [DOI](#)). As in previous work (Schrader et al., 2000 [DOI](#); van der Laan et al., 2002 [DOI](#)), the G232D mutation resulted in loss of selectivity and lowered affinity.

Given that Gly232 resides in the middle of the selectivity filter, these results are readily understood by its disruptive effects on the S3 site (Silberberg et al., 2021 [DOI](#)). Our demonstration of coupled transport of NH_4^+ and Rb^+ G232D not only confirms that the selectivity filter governs ion selection, but that the pump subunit, KdpB, is relatively promiscuous.

Like K^+ channels, the selectivity filter opens into a water-filled vestibule. Unlike channels, the vestibule in KdpFABC does not lead to the cytoplasm, but instead forms the entrance to the intersubunit tunnel. Nevertheless, mutations to Glu370 and Arg493 indicate that, like channels, this region controls the conductance of ions (Naranjo et al., 2016 [DOI](#)). These are the only charged residues in the membrane core of KdpA and, although they do not directly interact and do not project directly into the vestibule, they likely create an electrostatic field in the low dielectric environment of the membrane. Arg493, which is more highly conserved, is at the back of the vestibule, whereas Glu370 is along the direction of ion travel. Neutralization of either residue or reversal of their charge increases K_m , though V_{max} is unaffected. Indeed, alignment of diverse KdpA sequences shows that Glu370 is neutralized (Asn or Gln) in $\sim 50\%$ of the organisms (Huang et al., 2017 [DOI](#)). Interestingly, reversal of charge on the fully conserved Arg493 (R493E) has relatively modest effects, but reversal of Glu370 (E370K) essentially abolishes transport, perhaps indicating a role for an electrostatic field in moving K^+ through this region.

Non-protein spherical densities have been seen within the vestibule and tunnel in previous maps, but the current map reveals many more, likely due to its higher resolution. The identity of these densities has been contentious, sometimes modeled as a string of 7–9 K^+ ions (Silberberg et al., 2022 [DOI](#), 2021 [DOI](#)) whereas our previous structural models feature water molecules in similar positions (Sweet et al., 2021 [DOI](#)). Despite difficulty in distinguishing atomic species by cryo-EM, we continue to favor water molecules within this confined space for several reasons: First, a structure of the G232D mutant with K^+ substituted with Rb^+ did not change the normalized scattering density within the tunnel, as was seen at the S3 and canonical binding sites (Silberberg et al., 2021 [DOI](#)). Second, unconstrained all-atom MD simulations concluded that the tunnel accommodates at most one K^+ ion at a time, presumably due to electrostatic repulsion (Madapally et al., 2025 [DOI](#)). These simulations, which documented spontaneous entry of water into the vestibule and tunnel during the equilibration phase, resulted in a distribution of waters within the tunnel that is broadly consistent with our new high resolution map. Third, anomalous X-ray signal for K^+ has been observed at the S3 site but not in this region (Huang et al., 2017 [DOI](#); Madapally et al., 2025 [DOI](#)). Finally, map densities within the tunnel were either of comparable strength or weaker than surrounding side chain atoms, unlike at S3 and canonical binding sites. We therefore conclude that individual K^+ ions occupy the tunnel transiently as they transit between the selectivity filter and the canonical binding site. However, the energetics of this process remain obscure as the tunnel becomes narrow and dewetted as it approaches the KdpA/KdpB subunit interface. Furthermore, calculations of free energy indicate a massive energy barrier ($\sim 22\text{ kcal/mol}$) at this interface, suggesting that we are missing a key conformation that would facilitate passage (Madapally et al., 2025 [DOI](#)).

Both previous (Silberberg et al., 2022 [DOI](#)) and current cryoEM studies indicate that the E1~P state is the prevalent conformational state under turnover conditions. This observation suggests that the ensuing step, E1~P to E2-P, is rate limiting as also seen in other P-type ion pumps (Kuhlbrandt, 2004 [DOI](#)). Our higher resolution map shows an exceedingly strong, well-resolved density at the canonical binding site along the axis of the unwound M4 helix, providing compelling evidence that, like other P-type pumps, the transport ion is bound at this canonical site in this state. Several

additional, weaker densities are present in the adjacent pocket between M4, M5 and M6, coordinated either by this bound K^+ or by the surrounding residues. Their relative strength is consistent with water molecules, indicating that the stoichiometry of transport is likely to be 1 K^+ per ATP. The presence of these waters in the relatively large pocket together with the innate flexibility of M4 are consistent with promiscuity of this site. Indeed, when the selectivity filter is disrupted by the G232D mutation, the complex appears to be perfectly capable of transporting a broader range of cations.

Similar to the Glu370/Arg493 charge pair in KdpA, Asp583 and Lys586 are the only charged residues in the membrane core of KdpB. Although they have not been seen to interact directly, they coordinate accessory waters and likely produce an electrostatic field that guides the cation into the canonical binding site. Earlier structures have shown movement of the corresponding helix (M5) in the E2·Pi state that pushes the side chain of Lys586 into this binding site (Sweet et al., 2021 [DOI](#)), thus acting as a built-in counter ion as in related P-type ATPases ZntA (Wang et al., 2014 [DOI](#)) and AHA2 (Pedersen et al., 2007 [DOI](#)). As shown previously (Bramkamp and Altendorf, 2005 [DOI](#)), mutations of either residue generate an inactive complex, though current work reveals one exception: D583A resulted in uncoupled, K^+ -independent ATPase activity. One explanation for this behavior is that Asp583 exerts allosteric control over Lys586 and ensures that its movement into the binding site is coordinated with the transition from E1~P to E2·Pi. Accordingly, we speculate that truncation of the Asp583 side chain and elimination of its oxygen ligands eliminates this allosteric control and allows Lys586 to constitutively stimulate ATPase activity.

In P-type ATPases such as SERCA and Na,K-ATPase, ions are bound and released from the canonical binding pocket next to the proline of M4. The ATPase cycle is associated with changing affinity and accessibility of this site to alternate sides of the membrane. Although KdpB belongs to this superfamily, it operates in a substantially different way with ions entering the canonical binding site from the intersubunit tunnel rather than directly from the cytosol or periplasm. Stock et al. initially proposed that release occurred directly from the canonical site via a pathway between M4, M5 and M6 (Stock et al., 2018 [DOI](#)). However, this proposed pathway is very narrow and does not depend on the enzymatic state of the protein. We previously proposed an alternative exit site for K^+ between M1, M2 and M4 that aligns with pathways seen in other P-type ATPases (Dyla et al., 2020 [DOI](#)). In KdpB, this alternative pathway appears to have low affinity and is constitutively open to the cytosol. We proposed that the key step in transport was moving the ion from the canonical site into this exit site, from which it would spontaneously exit (Sweet et al., 2021 [DOI](#)). Current results from Thr75 mutants provide the first experimental evidence supporting this hypothesis. In particular, the pH dependence of T75D indicates that introduction of negative charge into this exit site interferes with transport, most likely by preventing release of the ion from this site. This conclusion relies on the assumption that the pKa of the Asp is shifted toward neutral pH due to the membrane environment, which is supported by theoretical estimation of pKa. On the other hand, T75K generated an uncoupled phenotype with WT levels of ATPase activity even in the absence of K^+ . Structure prediction for this mutant shows that the substituted lysine inserts into the canonical binding site, similar to the insertion of Lys586 seen in the E2·Pi conformation (**Fig. 6 – figure suppl. 1** [DOI](#)). This hypothesis extends the idea of lysines acting as built-in ions capable of stimulating futile ATPase cycles.

More generally, this behavior is consistent with KdpB acting as a rather non-specific cation-stimulated ATPase capable of recognizing its natural ligand, K^+ , as well as Rb^+ , NH^+ or the positive charge from nearby lysine residues.

A final note of interest stems from the N-terminus of KdpB which was not resolved in previous maps. Our new structure shows Arg3 in an interaction network with Glu161 of the A-domain together with Asp552 and Lys557 from the P-domain. Glu161 is crucial for dephosphorylation of the catalytic Asp307 during the E2-P to E2 transition (Dyla et al., 2020 [DOI](#)), thus raising the possibility that this interaction could influence the kinetics of this step, similar to the

proposed role for the N-terminus of Na,K-ATPase in modulating the E2 to E1 transition and K⁺ deocclusion (Lev et al., 2023 [DOI](#)). More generally, there is extensive precedent for regulation of P-type pumps by their N- or C-terminal tails: e.g., autoinhibition by N-termini of plasma membrane Ca-ATPase (Cali et al., 2017 [DOI](#)) and ATP7a/b Cu-ATPases (Bitter et al., 2022 [DOI](#)), as well as C-termini of lipid flippases (Timcenko et al., 2019 [DOI](#)) and fungal H-ATPase (Zhao et al., 2021 [DOI](#)). In the case of KdpB, Glu161 resides in a conserved loop (TGES) that includes Ser162, the target of inhibitory phosphorylation by the dual sensor KdpD (Silberberg et al., 2024 [DOI](#)). Given our lack of insight into the structural mechanism of this posttranslational regulation, further study of the N-terminus is warranted.

Summary

Fig. 7 [DOI](#) summarizes features of the transport pathway elucidated by the current work.

Effects of mutants indicate that a negative surface charge attracts K⁺ into the selectivity filter of KdpA where it binds at the S3 site, which is the primary determinant of ion specificity. Like K⁺ channels, the selectivity filter opens into a vestibule that is filled with water, where ions are attracted into the intersubunit tunnel by Glu370. The proximal end of this tunnel is filled with water but becomes hydrophobic as it approaches the interface with KdpB, where ions are dewetted and encounter a large energy barrier. The current structure represents E1~P-ADP in which the K⁺ has arrived at the canonical binding site in KdpB, next to the conserved Pro264. In the ensuing, rate-limiting step to E2-Pi, the conserved Lys586 on M5 moves into the canonical binding site, thus displacing the ion into a novel exit site next to Thr75. This scenario differs from other P-type pumps which release ions directly from their initial binding site. Nevertheless, T75 as well as the entire cytoplasmic portion of the M2 helix are well conserved in Kdp systems (Huang et al., 2017 [DOI](#)), consistent with a crucial role in the transport pathway.

Methods

Expression and purification of KdpFABC mutants

Expression and purification of KdpFABC has been described previously (Hussein et al., 2024 [DOI](#); Sweet et al., 2021 [DOI](#)). Briefly, we used *E. coli* strain TK2281 (thi, rha, lacZ, nagA, trkA405, trkD1, Δ(kdpFABCDE)81) to express WT and the various mutant proteins from a pBAD plasmid. This strain lacks KdpD, which has recently been shown to phosphorylate Ser162 on KdpB and thus inhibit its activity under conditions used for overexpression of mutant proteins (Silberberg et al., 2024 [DOI](#); Sweet et al., 2020 [DOI](#)). For expression, cells were cultured in Luria Broth (LB) media to OD₆₀₀ of 0.6 and then expression of KdpFABC was induced with 1.3 mM arabinose for 3 h. After cell lysis, the membrane fraction was solubilized by overnight incubation in 50 mM Tris pH 7.5, 600 mM NaCl, 10 mM MgCl₂, 10% glycerol, 1 mM 0.5 mM tris (2-carboxyethyl) phosphine (TCEP), and 1.2% n-decyl-β-maltoside (DM) at 4°C (20 ml of buffer for each gram of the membrane pellet). The sample was then loaded onto a Ni-NTA HiTrap affinity column (Cytiva, Marlborough, MA) equilibrated with 50 mM Tris pH 7.5, 600 mM NaCl, 10 mM MgCl₂, 10% glycerol, 1 mM TCEP, and 0.15% DM and eluted with an imidazole gradient. Peak fractions were pooled, concentrated and further purified with a size exclusion column (Superdex 200 increase 10/300 GL, GE Healthcare) using a running buffer containing 100 mM NaCl, 25 mM Tris pH 7.5, 10% glycerol, 1 mM TCEP, and 0.15% DM.

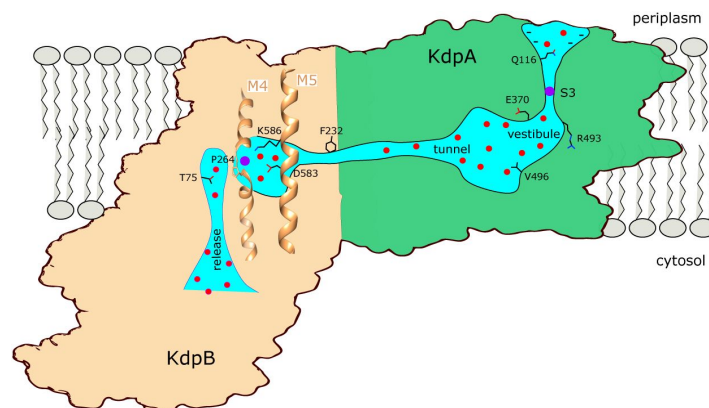


Figure 7.

Conduction pathway of KdpFABC.

Diagram illustrating features of the conduction pathway seen in the E1~P-ADP map together with residues that have been mutated in this study. Ions (purple spheres) are attracted from the periplasm to the mouth of the selectivity filter of KdpA by negative surface potential. Selectivity and affinity of the pump is governed by tight coordination of the ion at the S3 site. Ions become rehydrated upon release into a vestibule filled with water (red spheres) next to charged residues Glu370 and Arg493. Val496 resides in the widest part of the tunnel where introduction of positive charge abolishes transport. Phe232 resides at the subunit interface where there is a hydrophobic barrier that is essential to ion occlusion. K^+ binding to the canonical binding site next to Pro264 on M4 in KdpB triggers phosphorylation of Asp307 followed by allosteric movements of M5 that drive Lys586 to displace the ion into the exit site next to Thr75. A general lack of ligands in this exit site is consistent with low affinity, thus promoting K^+ release to the cytoplasm against the electrochemical gradient.

Expression and purification of MSP protein

A circularized membrane scaffold protein designated spNW25 was used for nanodisc formation (Zhang et al., 2021 [DOI](#)). For expression, *E. coli*, BL21 STAR (DE3) cells carrying spNW25 on a pET vector were cultured in LB media at 37°C, 200 rpm to an OD₆₀₀ of 0.7 followed by addition of 0.25 mM Isopropyl β-D-1-thiogalactopyranoside for 16 hours at 16°C. Cells were collected by centrifugation at 3500 g for 20 min at 4°C and then resuspended in 50 mM Tris-HCl pH 8, 100 mM NaCl, 5% glycerol, and 2 mM β-mercaptoethanol supplemented with protease inhibitor cocktail tablet. Cells were lysed by three passages through an Emulsiflex C3 high-pressure homogenizer (Avestin, Ottawa Canada). Lysed cells were centrifuged at 12,000 g for 45 min and the supernatant was loaded onto a 5 ml Ni-NTA column equilibrated with 50 mM Tris-HCl (pH 8), 400 mM NaCl, 5% glycerol, 20 mM imidazole. spNW25 was eluted after washing the column with buffers containing 25 mM and 50 mM imidazole followed by a 50-500 mM imidazole gradient. Fractions containing purified protein were pooled, dialyzed against 50 mM Tris-HCl pH 8, 100 mM NaCl, 5% glycerol, 2 mM β-mercaptoethanol, then aliquoted, frozen and stored at -80°C.

Preparation of lipid nanodisc

Stock solutions of lipid were prepared by mixing 1-palmitoyl-2-oleoyl-glycero-3-phosphocholine (POPC) and 1,2-dioleoyl-sn-glycero-3-phosphate (DOPA, Avanti Polar Lipids, Alabaster AL) in 20:1 weight ratio in chloroform. A thin film was produced by drying this mixture under Argon for 1-2 h and this film was then resuspended in 20 mM Tris pH 7.4, 100 mM KCl, 0.5 mM TCEP and 10 mg/ml Triton X-100 at a final lipid concentration of 5 mg/ml. After incubation at room temperature for 30 min, 0.18 mg of lipid were mixed with 0.48 mg of KdpFABC and 0.72 mg of spNW25 (total volume of 0.5 ml) and incubated for 60 min. Detergent was removed by sequential addition of BioBeads (BioRad Hercules, CA): 1 mg for 30 min, 1 mg for 30 min, and 40 mg overnight. This preparation was optimized by evaluating size distribution of nanodiscs by HPLC size exclusion chromatography and negative stain electron microscopy. In addition, the composition of nanodiscs was determined by SEC-MALS analysis (Fig. 1 – figure suppl. 1 [DOI](#)).

For preparation of cryo-EM samples, reconstituted nanodiscs were filtered with a 0.22 μm syringe filter and loaded onto a Superdex 200 size exclusion column that was equilibrated with 20 mM Tris pH 7.4, 100 mM KCl, and 0.5 mM TCEP. Peak fractions were concentrated to 4.25 mg/ml using a 50 kDa cutoff centrifugal concentrator (AMICON, Millipore Sigma, Burlington MA). The concentrated sample was aliquoted into PCR tubes and mixed with MgATP buffer to yield a final concentration of 2 mM MgCl₂, 1 mM ATP, and 3.4 mg/ml KdpFABC nanodisc. 3 μl of sample was added to glow-discharged EM grids (C-Flat 1.2/1.3-4Cu-50; Protochips, Inc) and blotted under 100% humidity at 4°C prior to plunging into liquid ethane using a Vitrobot (Thermo Fisher Scientific Inc, Bridgewater NJ).

Activity assays

ATPase rates were determined using the coupled enzyme assay (Warren et al., 1974 [DOI](#)). The standard buffer was composed of 75 mM TES-Tris pH 7.3, 7.5 mM MgCl₂, 0.5 mM phosphoenolpyruvate, 2.4 mM ATP, 0.8 mM NADH, 0.3% DM, 9.6 U/mL lactate dehydrogenase, and 9.6 U/mL pyruvate kinase. For potassium titration, 25 mM TES-Tris, 2.5 mM MgCl₂ and 1 mM ATP were used. All reagents were acquired from Sigma-Aldrich (St. Louis MO).

For ion transport, KdpFABC was reconstituted into proteoliposomes composed of a 4:1 weight ratio of POPC and DOPA as previously described (Hussein et al., 2024 [DOI](#)). Briefly, liposomes were prepared by resuspending a thin film of lipid in 50 mM HEPES-Tris pH 7.1, 5 mM MgSO₄, followed by three freeze-thaw cycles and extrusion using a 0.4 nm polycarbonate filter (Whatman-Cytiva).

Triton X-100 was added to liposomes at lipid to detergent weight ratio of 2.5:1 followed by addition of KdpFABC to yield a final lipid concentration of 5 mg/ml and a lipid-to-protein ratio of 5:1. The detergent was slowly removed by the sequential addition BioBeads.

Ion transport was measured using solid-supported membrane electrophysiology as implemented by the SURFE²RN1 (Nanion Technologies, Livingston NJ). Proteoliposomes were diluted 1:10 with inactivation buffer (50 mM HEPES-Tris, pH 7.1, 5 mM MgSO₄, 150 mM K₂SO₄), bath sonicated four times for 10 s each. Gold sensors were passivated with 1,2-diphytanoyl-sn-glycero-3-phosphocholine according to the manufacturer's instructions. After adding 50 µl of inactivation buffer to each sensor, 10 µl of diluted proteoliposomes were added and sensors were centrifuged at 2500 g for 30 min at 4°C. For measurement of transport, the sensor was washed with inactivation buffer for 1 s, followed by 1 s of activation buffer (inactivation buffer with 300 µM of ATP) and again with inactivation buffer for 1 s. For K⁺ titration, a rinsing step was added to introduce the new K⁺ concentration followed by 4-min incubation to allow equilibration across the membrane. For these titrations, 80 µM of nigericin was mixed with the diluted proteoliposomes prior to addition to sensors to facilitate the K⁺ concentration changes. For analysis of transport, peak current at 1.25 s was measured and plotted using GraphPad Prism10 software (GraphPad Software Inc, San Diego CA).

Potassium analysis of stock solutions was conducted using a simultaneous inductively coupled plasma-mass spectrometry (si-ICP-MS), known as Mattauch–Herzog-ICP-MS (MH-ICP-MS) instrument (SPECTRO MS, SPECTRO Analytical Instruments GmbH, Kleve, Germany) (Rabieh et al., 2020 [DOI](#)). Samples were acidified to 2% (vol/vol) with 65% Suprapur nitric acid (analytical-reagent grade, Merck, Darmstadt, Germany) and transferred via a Cetac autosampler (ASX-560, Teledyne Cetac Technologies, Omaha, NE, USA) to the nebulizer of MH-ICP-MS instrument. Each sample was analyzed in triplicate. A single-element potassium standard solution (Inorganic Ventures, Christiansburg, VA, USA) was used to prepare five K standard solutions ranging from blank to 2,500 parts per billion, which were used for constructing the calibration curve. This standard was also used to prepare control and spiked samples, which were analyzed at the beginning and end of the run. The average potassium recoveries for the control and spiked (pyruvate kinase enzyme) samples were 95.5% and 91.2%, respectively.

Cryo-EM structure determination

Samples of WT KdpFABC in lipid nanodiscs were imaged on a Krios G3i electron microscope equipped with Selectris imaging filter and Falcon4i detector (Thermo Fisher Scientific, Inc) at a pixel size of 0.93 Å and total dose of ~50 electrons/Å². A total of ~50,000 images were collected from 4 grids prepared from a single sample, which were processed with cryoSPARC v4.6 (Punjani et al., 2017 [DOI](#)). After removing images with crystalline ice, excessive contamination, or imaging artifacts, the remaining images were divided into five groups of 3000-5000 for preliminary processing (**Fig. 1 – figure suppl. 2** [DOI](#)). Particles were picked from each group using TOPAZ (Bepler et al., 2019 [DOI](#)), extracted with a box size of 335 Å with 3-fold binning (120 pixels) and subjected to two rounds of 2D classification to remove false positives. Selected particles were subjected to successive rounds of ab initio reconstruction with C1 symmetry and three classes, with the best two classes advancing to the next round. Particles selected from ab initio jobs were reextracted without binning (box size of 360 pixels) and used for successive rounds of heterogeneous refinement to separate the various conformations adopted under these active turnover conditions. Non-uniform refinement was ultimately used to produce structures from homogeneous classes. The largest class, corresponding to the E1~P-ADP conformation is being reported on here.

For structure refinement, we started with coordinates for KdpA-Q116R/KdpB-S162A mutant bound to AMPPCP (PDB 7LC3). After reversing the mutations, replacing AMPPCP with ADP + 2Mg²⁺ and adding phosphate to Asp307, the model was fitted as a rigid body to the map using CHIMERA (Pettersen et al., 2004 [DOI](#)). This model was adjusted to the experimental map using COOT (Emsley et al., 2010 [DOI](#)) and then subjected to rounds of real space refinement in PHENIX (Adams et al.,

2010 [\[link\]](#)). The A- and N-domain of KdpB displayed somewhat lower resolution (**Fig. 1 – figure suppl. 2** [\[link\]](#)), presumably due to their flexibility. As a result, these domains were refined against the unsharpened map, which displayed better connectivity in the peripheral structural elements. These two cytoplasmic domains were then joined to the membrane domains in COOT to produce the final model. Model and map depictions were done in CHIMERA (Pettersen et al., 2004 [\[link\]](#)), CHIMERAX (Meng et al., 2023 [\[link\]](#)) and PyMOL Molecular Graphics System (Schrödinger, LLC).

For analysis of the water tunnel, we used Caver Analyst 2.0 Beta (Jurcik et al., 2018 [\[link\]](#)). Tunnel profiles were derived from a starting point near KdpB-Ile628 using the following settings: Approximation 12, Minimum probe radius 0.5, Clustering threshold 3.5, Shell depth 2.5, Shell radius 4.5, Residues included AA and 9Y0. To evaluate structural effects of Val496 mutants, we used AlphaFold 3 (Jumper et al., 2021 [\[link\]](#)) to predict structures and then analyzed the tunnel profile with the above parameters. To estimate the pKa values of the exit site mutants, we generated structures of the mutants using AlphaFold 3, which were then used as input for online version of PropKa (Li et al., 2005 [\[link\]](#)).

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References

1. Adams PD, Afonine PV, Bunkoczi G, Chen VB, Davis IW, Echols N, Headd JJ, Hung LW, Kapral GJ, Grosse-Kunstleve RW, McCoy AJ, Moriarty NW, Oeffner R, Read RJ, Richardson DC, Richardson JS, Terwilliger TC, Zwart PH (2010) **PHENIX: a comprehensive Python-based system for macromolecular structure solution** *Acta Crystallogr D Biol Crystallogr* **66**:213–21 <https://doi.org/10.1107/S0907444909052925> | [Google Scholar](#)
2. Albers RW (1967) **Biochemical aspects of active transport** *Ann Rev Biochem* **36**:727–756 [Google Scholar](#)
3. Bepler T, Morin A, Rapp M, Brasch J, Shapiro L, Noble AJ, Berger B (2019) **Positive-unlabeled convolutional neural networks for particle picking in cryo-electron micrographs** *Nat Methods* **16**:1153–1160 <https://doi.org/10.1038/s41592-019-0575-8> | [Google Scholar](#)
4. Bitter RM, Oh S, Deng Z, Rahman S, Hite RK, Yuan P (2022) **Structure of the Wilson disease copper transporter ATP7B** *Sci Adv* **8**:eabl5508 <https://doi.org/10.1126/sciadv.abl5508> | [Google Scholar](#)
5. Bramkamp M, Altendorf K (2005) **Single amino acid substitution in the putative transmembrane helix V in KdpB of the KdpFABC complex of Escherichia coli uncouples ATPase activity and ion transport** *Biochemistry* **44**:8260–6 [Google Scholar](#)
6. Buurman ET, Kim KT, Epstein W (1995) **Genetic evidence for two sequentially occupied K⁺ binding sites in the Kdp transport ATPase** *J Biol Chem* **270**:6678–85 <https://doi.org/10.1074/jbc.270.12.6678> | [Google Scholar](#)
7. Cali T, Brini M, Carafoli E (2017) **Regulation of Cell Calcium and Role of Plasma Membrane Calcium ATPases** *Int Rev Cell Mol Biol* **332**:259–296 <https://doi.org/10.1016/bs.ircmb.2017.01.002> | [Google Scholar](#)
8. Doyle DA, Cabral JM, Pfuetzner RA, Kuo A, Gulbis JM, Cohen SL, Chait BT, MacKinnon R (1998) **The structure of the potassium channel: molecular basis of K⁺ conduction and selectivity** *Science* **280**:69–77 [Google Scholar](#)
9. Durell SR, Bakker EP, Guy HR (2000) **Does the KdpA subunit from the high affinity K⁺-translocating P-type KDP-ATPase have a structure similar to that of K(+) channels?** *Biophys J* **78**:188–99 [https://doi.org/10.1016/S0006-3495\(00\)76584-2](https://doi.org/10.1016/S0006-3495(00)76584-2) | [Google Scholar](#)
10. Dyla M, Kjærgaard M, Poulsen H, Nissen P (2020) **Structure and Mechanism of P-Type ATPase Ion Pumps** *Annu Rev Biochem* **89**:583–603 <https://doi.org/10.1146/annurev-biochem-010611-112801> | [Google Scholar](#)
11. Emsley P, Lohkamp B, Scott W, Cowtan K (2010) **Features and development of Coot** *Acta Crystallogr Sect - Biol Crystallogr* **66**:486–501 [Google Scholar](#)
12. Epstein W, Davies M (1970) **Potassium-dependant mutants of Escherichia coli K-12** *Bacterial* **101**:836–843 [Google Scholar](#)

13. Epstein W, Whitelaw V, Hesse J (1978) **A K⁺ transport ATPase in Escherichia coli** *J Biol Chem* **253**:6666–8 [Google Scholar](#)
14. Gassel M, Mollenkamp T, Puppe W, Altendorf K (1999) **The KdpF subunit is part of the K⁺-translocating Kdp complex of Escherichia coli and is responsible for stabilization of the complex in vitro** *J Biol Chem* **274**:37901–37907 [Google Scholar](#)
15. Gayen A, Leninger M, Traaseth NJ (2016) **Protonation of a glutamate residue modulates the dynamics of the drug transporter EmrE** *Nat Chem Biol* **12**:141–5 <https://doi.org/10.1038/nchembio.1999> | [Google Scholar](#)
16. Henderson JA, Huang Y, Beckstein O, Shen J (2020) **Alternative proton-binding site and long-distance coupling in Escherichia coli sodium-proton antiporter NhaA** *Proc Natl Acad Sci U A* **117**:25517–25522 <https://doi.org/10.1073/pnas.2005467117> | [Google Scholar](#)
17. Hesse JE, Wieczorek L, Altendorf K, Reicin AS, Dorus E, Epstein W (1984) **Sequence homology between two membrane transport ATPases, the Kdp-ATPase of Escherichia coli and the Ca²⁺-ATPase of sarcoplasmic reticulum** *Proc Natl Acad Sci USA* **81**:4746–4750 [Google Scholar](#)
18. Huang CS, Pedersen BP, Stokes DL (2017) **Crystal structure of the potassium-importing KdpFABC membrane complex** *Nature* **546**:681–685 <https://doi.org/10.1038/nature22970> | [Google Scholar](#)
19. Hussein A, Zhang X, Stokes DL (2024) **Numerical model for electrogenic transport by the ATP-dependent potassium pump KdpFABC** *Biophys Rep N* **4**:100169 <https://doi.org/10.1016/j.bpr.2024.100169> | [Google Scholar](#)
20. Isom DG, Castaneda CA, Cannon BR, Velu PD, Garcia-Moreno EB (2010) **Charges in the hydrophobic interior of proteins** *Proc Natl Acad Sci U A* **107**:16096–100 <https://doi.org/10.1073/pnas.1004213107> | [Google Scholar](#)
21. Jumper J, Evans R, Pritzel A, Green T, Figurnov M, Ronneberger O, Tunyasuvunakool K, Bates R, Zidek A, Potapenko A, Bridgland A, Meyer C, Kohl SAA, Ballard AJ, Cowie A, Romera-Paredes B, Nikolov S, Jain R, Adler J, Back T, Petersen S, Reiman D, Clancy E, Zielinski M, Steinegger M, Pacholska M, Berghammer T, Bodenstein S, Silver D, Vinyals O, Senior AW, Kavukcuoglu K, Kohli P, Hassabis D (2021) **Highly accurate protein structure prediction with AlphaFold** *Nature* **596**:583–589 <https://doi.org/10.1038/s41586-021-03819-2> | [Google Scholar](#)
22. Jurcik A, Bednar D, Byska J, Marques SM, Furmanova K, Daniel L, Kokkonen P, Brezovsky J, Strnad O, Stourac J, Pavelka A, Manak M, Damborsky J, Kozlikova B (2018) **CAVER Analyst 2.0: analysis and visualization of channels and tunnels in protein structures and molecular dynamics trajectories** *Bioinformatics* **34**:3586–3588 <https://doi.org/10.1093/bioinformatics/bty386> | [Google Scholar](#)
23. Kanai R, Ogawa H, Vilsen B, Cornelius F, Toyoshima C (2013) **Crystal structure of a Na⁺-bound Na⁺,K⁺-ATPase preceding the E1P state** *Nature* **502**:201–6 <https://doi.org/10.1038/nature12578> | [Google Scholar](#)
24. Kuhlbrandt W (2004) **Biology, structure and mechanism of P-type ATPases** *Nat Rev Mol Cell Biol* **5**:282–295 [Google Scholar](#)
25. Laimins LA, Rhoads DB, Altendorf K, Epstein W (1978) **Identification of the structural proteins of an ATP-driven potassium transport system in Escherichia coli** *Proc Natl Acad Sci* **75**:3216–9 [Google Scholar](#)

26. Lev B, Chennath M, Cranfield CG, Cornelius F, Allen TW, Clarke RJ (2023) **Involvement of the alpha-subunit N-terminus in the mechanism of the Na⁺,K⁺-ATPase** *Biochim Biophys Acta BBA - Mol Cell Res* **1870**:119539 <https://doi.org/10.1016/j.bbamcr.2023.119539> | [Google Scholar](#)
27. Li H, Robertson AD, Jensen JH (2005) **Very fast empirical prediction and rationalization of protein pKa values** *Proteins Struct Funct Bioinforma* **61**:704–721 <https://doi.org/10.1002/prot.20660> | [Google Scholar](#)
28. Liu S, Lockless SW (2013) **Equilibrium selectivity alone does not create K⁺-selective ion conduction in K⁺ channels** *Nat Commun* **4**:2746 <https://doi.org/10.1038/ncomms3746> | [Google Scholar](#)
29. Madapally HV, Hussein A, Eriksen MW, Pedersen BP, Stokes DL, Khandelia H (2025) **On the mechanism of K⁺ transport through the inter-subunit tunnel of KdpFABC** *bioRxiv* <https://doi.org/10.1101/2025.03.04.641403> | [Google Scholar](#)
30. Meng EC, Goddard TD, Pettersen EF, Couch GS, Pearson ZJ, Morris JH, Ferrin TE (2023) **UCSF ChimeraX: Tools for structure building and analysis** *Protein Sci* **32**:e4792 <https://doi.org/10.1002/pro.4792> | [Google Scholar](#)
31. Miller C (2000) **An overview of the potassium channel family** *Genome Biol* **1**:reviews0004.1 <https://doi.org/10.1186/gb-2000-1-4-reviews0004> | [Google Scholar](#)
32. Morais-Cabral JH, Zhou Y, MacKinnon R (2001) **Energetic optimization of ion conduction rate by the K⁺ selectivity filter** *Nature* **414**:37–42 <https://doi.org/10.1038/35102000> | [Google Scholar](#)
33. Morrison EA, Robinson AE, Liu Y, Henzler-Wildman KA (2015) **Asymmetric protonation of EmrE** *J Gen Physiol* **146**:445–61 <https://doi.org/10.1085/jgp.201511404> | [Google Scholar](#)
34. Naranjo D, Moldenhauer H, Pincuntureo M, Diaz-Franulic I (2016) **Pore size matters for potassium channel conductance** *J Gen Physiol* **148**:277–91 <https://doi.org/10.1085/jgp.201611625> | [Google Scholar](#)
35. Panahi A, Brooks CL (2015) **Membrane environment modulates the pKa values of transmembrane helices** *J Phys Chem B* **119**:4601–7 <https://doi.org/10.1021/acs.jpcc.5b00289> | [Google Scholar](#)
36. Pedersen BP, Buch-Pedersen MJ, Morth JP, Palmgren MG, Nissen P (2007) **Crystal structure of the plasma membrane proton pump** *Nature* **450**:1111–4 [Google Scholar](#)
37. Pedersen BP, Stokes DL, Apell HJ (2019) **The KdpFABC complex - K⁺ transport against all odds** *Mol Membr Biol* **35**:21–38 <https://doi.org/10.1080/09687688.2019.1638977> | [Google Scholar](#)
38. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, Ferrin TE (2004) **UCSF Chimera--a visualization system for exploratory research and analysis** *J Comput Chem* **25**:1605–1612 [Google Scholar](#)
39. Post RL, Kume S, Tobin T, Orcutt B, Sen AK (1969) **Flexibility of an active center in sodium-plus-potassium adenosine triphosphatase** *J Gen Physiol* **54**:306–26 [Google Scholar](#)
40. Punjani A, Rubinstein JL, Fleet DJ, Brubaker MA (2017) **cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination** *Nat Methods* **14**:290–296 <https://doi.org/10>

.1038/nmeth.4169 | [Google Scholar](#)

41. Puppe W, Siebers A, Altendorf K (1992) **The phosphorylation site of the Kdp-ATPase of *Escherichia coli*: site-directed mutagenesis of the aspartic acid residues 300 and 307 of the KdpB subunit** *Mol Microbiol* **6**:3511–20 [Google Scholar](#)
42. Rabieh S, Bayaraa O, Romeo E, Amosa P, Calnek K, Idaghdour Y, Ochsenkühn MA, Amin SA, Goldstein G, Bromage TG (2020) **MH-ICP-MS Analysis of the Freshwater and Saltwater Environmental Resources of Upolu Island, Samoa** *Molecules* **25**:4871 <https://doi.org/10.3390/molecules25214871> | [Google Scholar](#)
43. Rhoads DB, Waters FB, Epstein W (1976) **Cation transport in *Escherichia coli*. VIII. Potassium transport mutants** *J Gen Physiol* **67**:325–41 [Google Scholar](#)
44. Schrader M, Fendler K, Bamberg E, Gassel M, Epstein W, Altendorf K, Drose S (2000) **Replacement of glycine 232 by aspartic acid in the KdpA subunit broadens the ion specificity of the K⁺-translocating KdpFABC complex** *Biophys J* **79**:802–813 [Google Scholar](#)
45. Silberberg JM, Corey RA, Hielkema L, Stock C, Stansfeld PJ, Paulino C, Hanelt I (2021) **Deciphering ion transport and ATPase coupling in the intersubunit tunnel of KdpFABC** *Nat Commun* **12**:5098 <https://doi.org/10.1038/s41467-021-25242-x> | [Google Scholar](#)
46. Silberberg JM, Ketter S, Bohm PJN, Jordan K, Wittenberg M, Grass J, Hanelt I (2024) **KdpD is a tandem serine histidine kinase that controls K(+) pump KdpFABC transcriptionally and post-translationally** *Nat Commun* **15**:3223 <https://doi.org/10.1038/s41467-024-47526-8> | [Google Scholar](#)
47. Silberberg JM, Stock C, Hielkema L, Corey RA, Rheinberger J, Wunnicke D, Dubach VRA, Stansfeld PJ, Hanelt I, Paulino C (2022) **Inhibited KdpFABC transitions into an E1 off-cycle state** *eLife* **11** <https://doi.org/10.7554/eLife.80988> | [Google Scholar](#)
48. Stock C, Hielkema L, Tascon I, Wunnicke D, Oostergetel GT, Azkargorta M, Paulino C, Hanelt I (2018) **Cryo-EM structures of KdpFABC suggest a K(+) transport mechanism via two intersubunit half-channels** *Nat Commun* **9**:4971 <https://doi.org/10.1038/s41467-018-07319-2> | [Google Scholar](#)
49. Stumpe S, Schlosser A, Schleyer M, Bakker EP (1996) **K⁺ circulation across the procaryotic cell membrane: K⁺-uptake systems** In: Konings WN, Kaback HR, Lolkema JS, editors. *Handbook of Biological Physics* Amsterdam: Elsevier Science BV pp. 473–499 [Google Scholar](#)
50. Sweet ME, Larsen C, Zhang X, Schlame M, Pedersen BP, Stokes DL (2021) **Structural basis for potassium transport in prokaryotes by KdpFABC** *Proc Natl Acad Sci U S A* **118** <https://doi.org/10.1073/pnas.2105195118> | [Google Scholar](#)
51. Sweet ME, Zhang X, Erdjument-Bromage H, Dubey V, Khandelwal H, Neubert TA, Pedersen BP, Stokes DL (2020) **Serine phosphorylation regulates the P-type potassium pump KdpFABC** *eLife* **9** <https://doi.org/10.7554/eLife.55480> | [Google Scholar](#)
52. Timcenko M, Lyons JA, Janulienė D, Ulstrup JJ, Dieudonné T, Montigny C, Ash M-R, Karlsen JL, Boesen T, Kühlbrandt W, Lenoir G, Moeller A, Nissen P (2019) **Structure and autoregulation of a P4-ATPase lipid flippase** *Nature* **571**:366–370 <https://doi.org/10.1038/s41586-019-1344-7> | [Google Scholar](#)

53. Toyoshima C, Nakasako M, Nomura H, Ogawa H (2000) **Crystal structure of the calcium pump of sarcoplasmic reticulum at 2.6 Å resolution** *Nature* **405**:647–655 [Google Scholar](#)
54. van der Laan M, Gassel M, Altendorf K (2002) **Characterization of amino acid substitutions in KdpA, the K⁺-binding and -translocating subunit of the KdpFABC complex of *Escherichia coli*** *J Bacteriol* **184**:5491–5494 [Google Scholar](#)
55. Walderhaug MO, Polarek JW, Voelkner P, Daniel JM, Hesse JE, Altendorf K, Epstein W (1992) **KdpD and KdpE, proteins that control expression of the kdpABC operon, are members of the two-component sensor-effector class of regulators** *J Bacteriol* **174**:2152–9 <https://doi.org/10.1128/jb.174.7.2152-2159.1992> | [Google Scholar](#)
56. Wang K, Sitsel O, Meloni G, Autzen HE, Andersson M, Klymchuk T, Nielsen AM, Rees DC, Nissen P, Gourdon P (2014) **Structure and mechanism of Zn²⁺-transporting P-type ATPases** *Nature* **514**:518–522 <https://doi.org/10.1038/nature13618> | [Google Scholar](#)
57. Warren GB, Toon PA, Birdsall NJM, Lee AG, Metcalfe JC (1974) **Reversible lipid titrations of the activity of pure adenosine triphosphatase-lipid Complexes** *Biochemistry* **13**:5501–5507 [Google Scholar](#)
58. Zhang S, Ren Q, Novick SJ, Strutzenberg TS, Griffin PR, Bao H (2021) **One-step construction of circularized nanodiscs using SpyCatcher-SpyTag** *Nat Commun* **12**:5451 <https://doi.org/10.1038/s41467-021-25737-7> | [Google Scholar](#)
59. Zhao P, Zhao C, Chen D, Yun C, Li H, Bai L (2021) **Structure and activation mechanism of the hexameric plasma membrane H⁺-ATPase** *Nat Commun* **12**:6439 <https://doi.org/10.1038/s41467-021-26782-y> | [Google Scholar](#)

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Reviewer #1 (Public review):

Summary:

This study on potassium ion transport by the protein complex KdpFABC from *E. coli* reveals a 2.1 Å cryo-EM structure of the nanodisc-embedded transporter under turnover conditions. The results confirm that K⁺ ions pass through a previously identified tunnel that connects the channel-like subunit with the P-type ATPase-type subunit.

Strengths:

The excellent resolution of the structure and the thorough analysis of mutants using ATPase and ion transport measurements help to strengthen new and previous interpretations. The evidence supporting the conclusions is solid, including biochemical assays and analysis of mutants. The work will be of interest to the membrane transporter and channel communities and to microbiologists interested in osmoregulation and potassium homeostasis.

Weaknesses:

There is insufficient credit and citation of previous work.

<https://doi.org/10.7554/eLife.107397.1.sa2>

Reviewer #2 (Public review):

Summary:

The paper describes the high-resolution structure of KdpFABC, a bacterial pump regulating intracellular potassium concentrations. The pump consists of a subunit with an overall structure similar to that of a canonical potassium channel and a subunit with a structure similar to a canonical ATP-driven ion pump. The ions enter through the channel subunit and then traverse the subunit interface via a long channel that lies parallel to the membrane to enter the pump, followed by their release into the cytoplasm.

Strengths:

The work builds on the previous structural and mechanistic studies from the authors' and other labs. While the overall architecture and mechanism have already been established, a detailed understanding was lacking. The study provides a 2.1 Å resolution structure of the E1-P state of the transport cycle, which precedes the transition to the E2 state, assumed to be the rate-limiting step. It clearly shows a single K⁺ ion in the selectivity filter of the channel and in the canonical ion binding site in the pump, resolving how ions bind to these key regions of the transporter. It also resolves the details of water molecules filling the tunnel that connects the subunits, suggesting that K⁺ ions move through the tunnel transiently without occupying well-defined binding sites. The authors further propose how the ions are released into the

cytoplasm in the E2 state. The authors support the structural findings through mutagenesis and measurements of ATPase activity and ion transport by surface-supported membrane (SSM) electrophysiology.

Weaknesses:

While the results are overall compelling, several aspects of the work raised questions. First, the authors determined the structure of the pump in nanodiscs under turnover conditions and observed several structural classes, including E1-P, which is detailed in the paper. Two other structural classes were identified, including one corresponding to E2. It is unclear why they are not described in the paper. Notably, the paper considers in some detail what might occur during the E1-P to E2 state transition, but does not describe the 3.1 Å resolution map for the E2 state that has already been obtained. Does the map support the proposed structural changes?

The paper relies on the quantitative activity comparisons between mutants measured using SSM electrophysiology. Such comparisons are notoriously tricky due to variability between SSM chips and reconstitution efficiencies. The authors should include raw traces for all experiments in the supplementary materials, explain how the replicates were performed, and describe the reproducibility of the results. Related to this point above, size exclusion chromatography profiles and reconstitution efficiencies for mutants should be shown to facilitate comparison between measured activities. For example, could it be that the inactive V496R mutant is misfolded and unstable?

Similarly, are the reduced activities of V496W and V496H (and many other mutants) due to changes in the tunnel or poor biochemical properties of these variants? Without these data, the validity of the ion transport measurements is difficult to assess.

The authors propose that the tunnel connecting the subunits is filled with water and lacks potassium ions. This is an important mechanistic point that has been debated in the field. It would be interesting to calculate the volume of the tunnel and estimate the number of ions that might be expected in it, given their concentration in bulk. It may also be helpful to provide additional discussion on whether some of the observed densities correspond to bound ions with low occupancy.

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Reviewer #3 (Public review):

Summary:

By expressing protein in a strain that is unable to phosphorylate KdpFABC, the authors achieve structures of the active wild-type protein, capturing a new intermediate state, in which the terminal phosphoryl group of ATP has been transferred to a nearby Asp, and ADP remains covalently bound. The manuscript examines the coupling of potassium transport and ATP hydrolysis by a comprehensive set of mutants. The most interesting proposal revolves around the proposed binding site for K⁺ as it exits the channel near T75. Nearby mutations to charged residues cause interesting phenotypes, such as constitutive uncoupled ATPase activity, leading to a model in which lysine residues can occupy/compete with K⁺ for binding sites along the transport pathway.

Strengths:

Although this structure is not so different from previous structures, its high resolution (2.1 Å) is impressive and allows the resolution of many new densities in the potassium transport pathway. The authors are judicious about assigning these as potassium ions or water

molecules, and explain their structural interpretations clearly. In addition to the nice structural work, the mechanistic work is thorough. A series of thoughtful experiments involving ATP hydrolysis/transport coupling under various pH and potassium concentrations bolsters the structural interpretations and lends convincing support to the mechanistic proposal.

Weaknesses:

The structures are supported by solid membrane electrophysiology. These data exhibit some weaknesses, including a lack of information to assess the rigor and reproducibility (i.e., the number of replicates, the number of sensors used, controls to assess proteoliposome reconstitution efficiency, and the stability of proteoliposome absorption to the sensor).

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