Visualizing the mapped ion pathway through the Na,K-ATPase pump

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The Na+,K+-ATPase pump achieves thermodynamically uphill exchange of cytoplasmic Na+ ions for extracellular K+ ions by using ATP-mediated phosphorylation, followed by autodephosphorylation, to power conformational changes that allow ion access to the pump's binding sites from only one side of the membrane at a time. Formally, the pump behaves like an ion channel with two tightly coupled gates that are constrained to open and close alternately. The marine agent palytoxin disrupts this coupling, allowing both gates to sometimes be open, so temporarily transforming a pump into an ion channel. We made a cysteine scan of Na+,K-ATPase transmembrane (TM) segments TM1 to TM6, and used recordings of Na+ current flow through palytoxin-bound pump-channels to monitor accessibility of introduced cysteine residues via their reaction with hydrophilic methanethiosulfonate (MTS) reagents. To visualize the open-channel pathway, the reactive positions were mapped onto a homology model of Na+,K-ATPase based on the structure of the related sarcoplasmic- and endoplasmic-reticulum (SERCA) Ca2+-ATPase in a BeF3--trapped state,1,2 in which the extra-cytoplasmic gate is wide open (although the cytoplasmic access pathway is firmly shut). The results revealed a single unbroken chain of reactive positions that traverses the pump from the extracellular surface to the cytoplasm, comprises residues from TM1, TM2, TM4 and TM6, and passes through the equivalent of cation binding site II in SERCA, but not through site I. Cavity search analysis of the homology model validated its use for mapping the data by yielding a calculated extra-cytoplasmic pathway surrounded by MTS-reactive residues. As predicted by previous experimental results, that calculated extra-cytoplasmic pathway abruptly broadens above residue T806, at the outermost end of TM6 that forms the floor of the extracellular-facing vestibule. These findings provide a structural basis for further understanding cation translocation by the Na+,K-ATPase and by other P-type pumps like the Ca2+- and H+,K-ATPases.

The Na+,K+ pump, a P-type ATPase (named for the phosphorylated intermediate), generates the steep transmembrane concentration gradients of Na+ and K+ that are essential for cell life. Each pump exchanges three intracellular Na+ ions for two external K+ ions per ATP molecule hydrolyzed, up to one hundred times a second. This cation exchange is accomplished by conformational changes linked to phosphorylation and dephosphorylation of the pump3,4 that permit strictly alternating access to its ion binding sites: from the extracellular side in the phosphorylated state, called E2P, but from the cytoplasm in the dephosphorylated state, E1. The marine toxin palytoxin somehow disrupts this strict coupling between the Na+,K+ pump's two gates, so allowing both to sometimes be open, whereupon the pump is temporarily transformed into an ion channel.5-7 Palytoxin thus provides an unprecedented opportunity to use direct recordings of ion flow through the Na+,K+ pump to monitor accessibility to small water-soluble probes of target cysteine residues introduced along the ion pathway,8 and thereby characterize the routes to and from the ion binding sites.9-13

Mapping the resulting outline of the pathway onto a 3-dimensional structure is not straightforward, however, as the

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transmembrane segments TM4 and TM6 (compare with refs. 10 and 11) contribute to an ion pathway characterized by a wide vestibule that opens to the extracellular surface\textsuperscript{12} and penetrates deep into the Na\textsuperscript{+},K\textsuperscript{+}-ATPase where it narrows and leads to a cation-selectivity filter\textsuperscript{13} comprising conserved acidic residues (E336 and D813) that help coordinate cations in the Na\textsuperscript{+},K\textsuperscript{+} pump equivalent of cation binding site II also, partly, site I; ref. 15) in SERCA.

Very recently, to obtain a snapshot of the entire ion pathway, we\textsuperscript{17} used positively charged MTSET\textsuperscript{+} (2-trimethylammonium-ethyl-methanethiosulfonate) to extensively scan TM1, the TM1-2 linker, TM2, TM5 and the TM5-6 linker, as well as strategic positions in TM3. We combined these results with those from our previous representative scans of TM4 and TM6, and we displayed them (Fig. 1A) on a homology model of the Na\textsuperscript{+},K\textsuperscript{+} pump based on the SERCA E2-BeF\textsubscript{3}\textsuperscript{−} structure. Reactive positions (where the ~6 Å x 8 Å MTSET\textsuperscript{+} adduct altered current ≥10%), with atoms shown as red spheres (Fig. 1A), enclose a continuous pathway through the Na\textsuperscript{+},K\textsuperscript{+} pump from one side of the membrane to the other, comprising residues from TM1, TM2, TM4 and TM6, and passing through cation binding site II, but were not found in TM5 or in binding site I. Non-responsive positions, shown as yellow sticks in Figure 1, surround the reactive positions, confirming that the scan was complete. And near abolition of Na\textsuperscript{+} current by modification of a single cysteine at several positions along the pathway argues that there is no other route for Na\textsuperscript{+} ion flow through palytoxin-bound Na\textsuperscript{+},K\textsuperscript{+} pump-channels\textsuperscript{17} (reviewed in ref. 18).

To further evaluate the scanning results, we used MOLE (reviewed in ref. 19; http://troll.chemik.muni.cz/whitezone/development/mole/online/moleonline1.3/) to examine the tunnels that connect binding site II, but were not found in TM5 or in binding site I. Non-responsive positions, shown as yellow sticks in Figure 1, surround the reactive positions, confirming that the scan was complete. And near abolition of Na\textsuperscript{+} current by modification of a single cysteine at several positions along the pathway argues that there is no other route for Na\textsuperscript{+} ion flow through palytoxin-bound Na\textsuperscript{+},K\textsuperscript{+} pump-channels\textsuperscript{17} (reviewed in ref. 18).

Figure 1. Snapshot of the ion pathway through the Na\textsuperscript{+},K\textsuperscript{+} pump. (A) Cysteine-scanning results mapped onto a homology model of the Na\textsuperscript{+},K\textsuperscript{+} pump transmembrane domain, based on the SERCA E2-BeF\textsubscript{3}\textsuperscript{−} structure, viewed from side (upper) and extracellular surface (bottom). Helices are colored grey except TM1 (pale blue), TM2 (magenta), TM4 (blue), TM5 (purple) and TM6 (green). Residues at MTSET\textsuperscript{+}-responsive and non-responsive positions are shown as red balls and yellow sticks, respectively. (B) MOLE analysis of the homology model shown in the same orientations as in (A). Gold: tunnels starting from S784, N785 and E788, equivalent to cation-binding site I in SERCA. Purple: tunnels starting from E336 and D813, equivalent to cation-binding site II. Red and yellow sticks mark MTSET\textsuperscript{+}-responsive and non-responsive positions. Scanning data were previously published.\textsuperscript{17}
are initially narrow and almost identical to each other (Fig. 2). Close to their origins, the three near identical site I tunnels are distinct from the three near identical site II tunnels (Fig. 2A), but they merge with the site II tunnels a few Å above site II (Fig. 2A and B). All tunnels then run together until, ~20 Å from binding site II, they relatively abruptly broaden and diverge somewhat, to form the irregular wide vestibule that opens to the exterior (Figs. 1B and 2). Once they broaden, each of the three site I tunnels is essentially identical to one of the three site II tunnels (Figs. 1B and 2). Notably, the site II tunnels and post-merger site I tunnels are surrounded by MTSET+-reactive positions (Figs. 1B and 2A), and a narrow extension of the site II tunnel reaches towards the deep reactive TM4 position, G337. Even MTSET+-responsive position N799 in the TM5-TM6 external loop, distant from most reactive positions in the homology model (Fig. 1A), contacts a pair of tunnels, supporting applicability of the model.

The tunnel branch leading to site I is somewhat narrower than the pathway to site II (Fig. 2B), in accord with the overall conclusion that the main pathway passes through site II but is linked to cation-binding site I via a narrower connection. In addition, the relatively abrupt change in radius of all the tunnels (~20 Å from site II; Fig. 2B) in the vicinity of residue T806, which is at the outermost end of TM6 within the floor of the extracellular-facing vestibule (Fig. 1A), corroborates our earlier electrophysiological results. In those findings, reaction of mutant G805C with variously charged MTS reagents yielded correspondingly varied electrostatic effects on Na⁺ current through palytoxin-bound Na⁺-K⁺-ATPase. In contrast, those same reagents all, regardless of charge, diminished Na⁺ current upon reaction with a cysteine introduced at the adjacent (deeper) position, in mutant T806C, presumably because the adducts all sterically impeded Na⁺ ion flow. This suggested that the wide extracellular pathway suddenly narrows at position T806.

Despite this reasonable correspondence between scanning results and model based on the SERCA BeF₃⁻ structure, the tunnel analysis implies that although a 1.9-Å diameter Na⁺ ion could reach site II (though not site I) the at least two times wider MTS reagents should not. Moreover, as the cytoplasmic-side pathway in the SERCA E2-BeF₃⁻ structure is closed by a 15–20 Å barrier, we attempted tunnel analysis using a homology model of Na⁺-K⁺-ATPase based on the E1-2Ca²⁺ structure (PDB code 1SU4) in which the cytoplasmic pathway appears open. However, whether starting from site I or site II, most of the tunnels calculated for this model exited the transmembrane domain at the level of the lipid bilayer, with several emerging between TM1 and TM2. Only one tunnel led to the cytoplasm, and it started in site II, passed between TM1,
TM2 and TM4, and contacted the deepest MTSET+-responsive positions, L339 (TM4), T145 (TM2) and G100 (TM1). Unfortunately, the significance of that tunnel remains unclear because two comparable tunnels, following almost the same route and also contacting L339, T145 and G100, were identified in a model based on a different E1·2Ca²⁺ structure (PDB code 1T5S), with bound AMPPCP, in which the cytoplasmic pathway appears closed.²¹

We conclude that the homology model of Na⁺,K⁺-ATPase built on the SERCA E2-BeF₃⁻ structure provides a reliable representation of the extracellular access pathway to the ion binding sites, but that a structural model of the Na⁺,K⁺ pump’s cytoplasmic access pathway for Na⁺ and K⁺ ions remains elusive. The incomplete nature of the atomic scaffold notwithstanding, the outline of the ion pathway through palytoxin-bound Na⁺,K⁺ pump channels obtained by combining electrophysiological data with homology modeling and cavity search analysis affords a structural framework for further understanding ion translocation by the Na⁺,K⁺-ATPase and by related P-type pumps, such as the Ca⁺- and H⁺,K⁺-ATPases.

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