The cardiac Na⁺-Ca²⁺ exchanger has two cytoplasmic ion permeation pathways

Scott A. John, Jun Liao, Youxing Jiang, and Michela Ottolia

*Department of Medicine (Cardiology) and Cardiovascular Research Laboratory, David Geffen School of Medicine at the University of California, Los Angeles, CA 90095; ‡Department of Physiology and §Howard Hughes Medical Institute, University of Texas Southwestern Medical Center, Dallas, TX 75390; and ¶Cedars-Sinai Heart Institute, Cedars Sinai, Los Angeles, CA 90048

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The Na⁺-Ca²⁺ exchanger (NCX) is a ubiquitously expressed plasma membrane protein. It plays a fundamental role in Ca²⁺ homeostasis by moving Ca²⁺ out of the cell using the electrochemical gradient of Na⁺ as the driving force. Recent structural studies of a homologous archaeabacterial exchanger, NCX_Mj, revealed its outward configuration with two potential ion permeation pathways exposed to the extracellular environment. Based on the symmetry of NCX_Mj structure, an atomic model of an inward-facing conformation was generated showing similar pathways but directed to the cytoplasm. The presence of these water-filled cavities has yet to be confirmed experimentally, and it is unknown if the mammalian exchanger adopts the same structure. In this study, we mutated multiple residues within transmembrane segments 2 and 7 of NCX1.1 (cardiac isoform) to cysteines, allowing us to investigate their sensitivity to membrane-impermeable sulfhydryl reagents as exchanger current block. By trapping NCX1.1 in the inward-facing configuration, we have mapped two differently sized cytoplasmic aqueous cavities, the access of which is modified during exchange.

This data reveals movements of the protein associated with ion transport. Electrophysiological characterization shows that the conserved residues within transmembrane segments 2 and 7, coordinating Na⁺ and Ca²⁺ ions in NCX_Mj, play a fundamental role in NCX1.1. Our results suggest a similar architecture between the mammalian and archaeabacterial exchangers.

The Na⁺-Ca²⁺ exchanger (NCX) is a widely expressed electrogenic antiporter that exchanges one Ca²⁺ ion for three Na⁺ ions across the plasma membrane (1), thereby playing a fundamental role in Ca²⁺ homeostasis (1–3). Despite intensive investigation, how Na⁺ and Ca²⁺ access their binding sites is unknown, and there is limited knowledge about the residues involved in ion transport (4–6). The recent high-resolution crystal structure of an archaeabacterial NCX homolog (NCX_Mj) and its modeled inward configuration (7) have shown two potential pathways for ions to gain access to the core of the exchanger where one identified Ca²⁺ binding site and three inferred Na⁺ binding sites reside. Of twelve residues forming these sites, four residues are in transmembrane segment (TMS) 2, four residues are in TMS7, and two residues each are in TMS3 and -8. These TMSs, referred to as α-repeats (8), are conserved within the extended exchanger family (3) and important for ion translocation in both the mammalian NCX (4–6, 9) and the Na⁺/Ca²⁺-K⁺ exchanger (10).

The structure of NCX_Mj represents a milestone in the study of exchangers. However, for the mammalian homolog, functional studies supporting this structure are absent, and it is unknown if it adopts a similar architecture. Moreover, there is no information on the protein conformational changes that may occur during ion transport. The goals of this work were to identify the ion translocation pathways of the eukaryotic NCX (cardiac isoform NCX1.1) (11), their potential structural rearrangements during ion transport, and the residues involved in ion binding. Such studies allow determination of whether NCX1.1 functionally parallels the structural evidence from NCX_Mj and validate it under more physiological conditions. To these ends, we applied cysteine accessibility analysis to TMS2 and -7 of NCX1.1. Our results show that several residues lining TMS2 and -7 are exposed to the cytoplasm when the ion binding sites of NCX1.1 face the intracellular side of the cell (inward-facing configuration) but are inaccessible during transport. Results indicate that TMS2 and -7 contribute to the cytoplasmic ion permeation pathway and that they undergo conformational changes during ion translocation. Comparison between our accessibility data and the structural model of the inward-facing NCX_Mj model supports the presence of two cytoplasmic-facing aqueous cavities in NCX1.1. By using thiol-reactive compounds of different bulkiness, we also determined that these aqueous openings are of different sizes. Finally, electrophysiological analyses of the cysteine mutants reveal that those residues inferred to coordinate Na⁺ in NCX_Mj play an important role in the transport properties of NCX1.1. This data provides strong functional evidence for the identity of the residues forming the Na⁺ binding sites in the exchanger family. Similarly, mutations of residues analogous to those positions found to coordinate Ca²⁺ in NCX_Mj drastically alter NCX1.1 activity, further underlying similarity between the two proteins. The data indicate that the α-repeats of NCX play a fundamental role in transport function and that their architecture is maintained among distant family members.

Results

Effect of Cysteine Mutations: Role of TMS2 and -7 in Ion Translocation.

Twenty-three residues in TMS2 and -7 of NCX1.1 were individually replaced with cysteine with the final goal to investigate their reactivity with membrane-impermeable Methanethiosulfonate (MTS) reagents. First we analyzed the functional effects of these mutations, positions that are underlined in Fig. 1.A. By analogy to NCX_Mj structure (7), the residues of TMS2 and -7 coordinating ion binding are Ala⁶⁰, Ser¹⁰⁹, Ser¹¹⁰, Glu¹¹⁴ and Ala²⁰⁷, Thr¹⁰⁸, Ser¹¹¹, and Asp¹¹⁴ respectively (Fig. 1B). Some mutations produced inactive exchangers, namely E113C of TMS2 and T810C, S811C, and D814C of TMS7. Only mutant T810C could be rescued by the more conservative serine replacement. Functional data are consistent with the role of these residues in ion binding or transport. Indeed, as predicted by the crystal structure of NCX_Mj (7), Thr⁶¹⁰ and Ser⁷¹⁰ coordinate Na⁺ nearest to the external side, whereas the two negatively charged residues, E113 and D814, participate in multiple binding sites and play a central role in transport function.

Mutagenesis at the remaining ion coordinating sites generated active exchangers, although with somewhat reduced ionic currents.

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To whom correspondence should be addressed. E-mail: michela.ottolia@chs.org.

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as compared to WT. Exchangers A106C, S109C, and S110C of TMS2 and A807C and T810S of TMS7 were analyzed for their response to application of cytoplasmic Na\(^+\) and Ca\(^{2+}\). Fig. 2 displays an example showing the altered transport properties of the S109C mutant, namely a decreased affinity for cytoplasmic Na\(^+\) (K\(_{1/2}\), millimolar, values are S109C: 57 ± 5, n = 4; WT: 18 ± 1, n = 9) and increased affinity for Ca\(^{2+}\) (K\(_{1/2}\), millimolar, values are S109C: 0.64 ± 0.08, n = 4; WT: 2.1 ± 0.1, n = 10). The corresponding residue in NCX_Mj, Thr\(^{109}\), was inferred to coordinate Na\(^+\) at a site nearest to the cytoplasm, and our functional data support this role for Ser\(^{109}\).

The remaining mutants were examined, and Fig. 3 summarizes the concentrations of Na\(^+\) and Ca\(^{2+}\) needed to activate 50% of exchanger current (Hill coefficient values are reported in Fig. S1). Exchangers S109C, S110C, and A807C showed the most significant decrease in apparent affinity to intracellular Na\(^+\). Side chain size and polarity were also investigated by replacing Ser\(^{109}\) and Ser\(^{110}\) with the apolar residue alanine (S109A and S110A). Such replacement further decreased Na\(^+\) binding and reciprocally enhanced Ca\(^{2+}\) binding affinity. The result is consistent with Na\(^+\) and Ca\(^{2+}\) antagonizing each other when bound to their respective binding sites within the core of NCX1.1 (7, 12).

The amino acids within TMS2 and -7 found to coordinate Na\(^+\) and Ca\(^{2+}\) ions in NCX_Mj contribute to two highly conserved stretches of residues consisting of G/V/L/GSSAPE\(^{113}\) and GTSVPD\(^{114}\) in TMS2 and -7, respectively (Fig. L4). Within these sequences, the glycines and prolines do not directly participate in ion coordination (7). Nevertheless, these residues are highly conserved, suggesting an important role in NCX1.1 function. Although mutation of Gly\(^{108}\) to cysteine (G108C) abolished transport activity, insertion of an alanine (G108A) yielded an active exchanger with an unaffected apparent affinity for cytoplasmic Na\(^+\) (K\(_{1/2}\), millimolar, values are G108A: 15 ± 5, n = 5; WT: 18 ± 1, n = 9) but a significantly increased Ca\(^{2+}\) sensitivity (K\(_{1/2}\), micromolar, values are G108A: 0.9 ± 0.2, n = 9; WT: 2.1 ± 0.1, n = 10). Similarly, the equivalent mutation in TMS7, G809C, increased the cytoplasmic Ca\(^{2+}\) sensitivity (K\(_{1/2}\), micromolar, values are G809C: 1.39 ± 0.21, n = 6; WT: 2.1 ± 0.1, n = 10). These results indicate that a larger side chain at these two sites alters transport properties, presumably by altering the architecture of the adjoining ion binding sites. Introduction of cysteines at prolines 112 and 813 mainly decreased the sensitivity of NCX1.1 to Na\(^+\) (Fig. 3) without, however, preventing its ability to transport, indicating that these prolines are not essential for function.
cycling and inward-facing state conditions as summarized in Fig. 5A and B. The mutant M105C was blocked independently of NCX1.1 state, whereas L107C and S110C currents were inhibited only in the inward-facing state. Positions 105 and 109, which face the same surface of TMS2, seem to have greater exposure to the cytoplasm than residues 106, 107, and 110, allowing MTSET+ to also react with their side chains during cycling. Consistent with this hypothesis is the high rate of reactivity of M105C and S109C exchangers to MTSET+ compared to mutants A106C, L107C, and S110C. These results are summarized in Fig. 5 C and D, which compare the time course of the MTSET+ block of several NCX1.1 mutants when cycling and held in the inward-facing state. As shown, reactivity of cysteines at positions 106 and 110 is ~10-fold slower than the reactivity of M105C and S109C, suggesting limited accessibility of these residues.

The structural model of inward-facing NCX_Mj (7) predicts that TMS7 contributes to the cytoplasmic permeation pathway. We, therefore, introduced cysteine at eight strategic positions, resulting in the following active mutants: A807C, L808C, G809C, V812C, P813C, F816C, A817C, and V820C (summarized in Fig. 5). Mutants A807C, L808C, F816C, and V820C were not significantly affected by MTSET+, regardless of whether NCX1.1 was cycling or in its inward-facing conformation. The results indicate that these residues either are not functionally important or inaccessible from the cytoplasm. The facts that they are located on the same side of the TMS7 helix and that the homologous positions (Ala206, Phe207, Met215, and Leu219) in NCX_Mj appear to be immersed within the lipid bilayer are consistent with their inaccessibility.

Four mutants, G809C, V812C, F816C, and A817C were inhibited by cytoplasmic MTSET+ in both the absence and presence of Na+ (Fig. 5A and B). Because the NCX_Mj homologous residues to Gly809, Val812, and Pro813 are found in its core, the results indicate that NCX1.1 possesses a deep cavity that penetrates into the center of the protein. As shown in Fig. 5C, the time course of inactivation of these exchangers by MTSET+ was much slower when cycling, suggesting that this region of NCX1.1 undergoes conformational changes during transport.

To define further the TMSs forming a passage for Na+ and Ca2+, we also investigated the MTSET+ sensitivity of G833C of TMS8. As predicted by the crystal structure of NCX_Mj (7), this residue is expected to face the same aqueous cavity as residues Ala206 and Leu217 and therefore, should be exposed to cytoplasmic MTSET+. As shown in Fig. 5A, MTSET+ strongly inhibited the current of G833C only when in the inward-facing state. The result indicates that this segment of TMS8, near to the cytoplasmic surface, is solvent-accessible and contributes to the formation of an aqueous cavity that undergoes rearrangement during ion translocation.

Probing the Relative Size of the Two NCX1.1 Cavities. The crystal structure of NCX_Mj predicts the presence of two independent solvent-accessible pathways leading to Na+ and Ca2+ binding sites (7). Our accessibility data are consistent with the presence of these two water-filled cavities, as shown in Fig. 6 and Fig. S2, where the residues of NCX1.1 reactive with MTSET+ are indicated on the NCX_Mj inward state model. Residues of both TMS2 (Met105 and Ser109) and -7 (Gly809, Pro813, Val812, Ala817, and Val820) face the crevice inferred as the Ca2+ pathway in NCX_Mj, whereas cysteines at positions 106, 107, and 109 of TMS2 are exposed to the lumen suggested to be the Na+ passageway in NCX_Mj. To determine the relative sizes of these aqueous cavities, we tested cysteine modification by the membrane-impermeable MTS reagents 1-(Trimethylammonium)methyl Methanethiosulfonate Bromide (MTSMT+) and 2-(Triptylaminomethyl)ethyl Methanethiosulfonate Bromide (MTS-TPAE). MTSMT+ is slightly shorter than MTSET+ because of the presence of a methyl group instead of an ethyl chain, whereas MTS-TPAE is a larger reagent with pentyl chains (Fig. S3). Reactivity of these reagents with S110C and G809C was investigated. These positions are the deepest residues to react with MTSET+, and each lies in separate cavities, as predicted by the NCX_Mj inward-facing model. Results indicate modification of S110C currents only by the smaller reagent MTSMT+ (Fig. S3). Although the extent of block during cycling was minimal (28 ± 11%, n = 5), S110C current was extensively inhibited by MTSMT+ in the inward-facing state (82 ± 2%, n = 5) and more rapidly blocked than by MTSET+ (24 ± 2 s, n = 5 vs. 41 ± 3 s, n = 4, respectively) (Fig. S3E). In contrast, G809C currents were completely inhibited by both cytoplasmic MTSMT+ and MTS-TPAE independently of its state.

The data indicate that the two ion translocation pathways of NCX1.1 have different sizes. The one represented by G809C is sufficiently wide to accommodate the larger MTS-TPAE reagent, whereas the other cavity is too narrow to allow MTS-TPAE to react with S110C; however, it is reachable by the smaller MTSMT+ and MTSET+ reagents.

Discussion

X-ray crystallography of the archaeabacterial NCX_Mj in its outward-facing conformation (7) revealed that NCX is organized into two sets of five TMSs with similar structure but opposite...
of a steric hindrance destabilizing Na\(^+\) at this site and allowing Ca\(^{2+}\) to bind more efficiently. In contrast, Ser\(^{109}\) coordinates sites coordinating both Na\(^+\) and Ca\(^{2+}\) in NCX1.1 are two highly conserved regions: 108GSSAPE113 (TMS2) and 209GTSVPD214 (TMS7). As predicted by the NCX_Mj structure, the glycine and proline residues within these sequences are conserved among the exchangers, and both residues have essential structural roles in many membrane proteins. The homologous glycines 108 and 809 and prolines 112 and 813 did not drastically increase the percent of block while NCX1.1 was held in the inward-facing state, the glycine and proline residues within these sequences are located deep in the protein but are not directly involved in Na\(^+\) and Ca\(^{2+}\) coordination (7). Our data showed that replacement of both glycines 108 and 809 and prolines 112 and 813 did not drastically alter NCX1.1 activity, suggesting that they are not essential for function. This result is striking because these amino acids are highly conserved among the exchangers, and both residues have essential structural roles in many membrane proteins. The homologous prolines in NCX_Mj participate in forming part of TMS2 and -7 helices instead of creating a helix break or kink, which may explain why they are more resistant to mutation (7). In NCX_Mj, these conserved prolines are also part of a hydrophobic patch, Ala\(^{106}\) and Leu\(^{115}\) of TMS2 and Val\(^{812}\) and Phe\(^{816}\) of TMS7, with cysteine, we showed the most substantial shifts in both Na\(^+\) and Ca\(^{2+}\) binding sites, which were only inferred in the NCX_Mj X-ray structure (7).

Previously, Na\(^+\)-dependent 45Ca\(^{2+}\) uptake measurements indicated that mutations at sites 109, 110, and 810 lead to inactive transporters, despite their plasma membrane localization (4). Although the ionic currents generated by these mutants were low compared with WT, we could routinely measure their transport properties using the electrophysiological approach, enabling their initial characterization. Possibly, the large decrease in Na\(^+\) affinity for these mutants prevented measurements of a significant 45Ca\(^{2+}\) influx.
Ca$^{2+}$ apparent affinities for L115C (Fig. 4 and Fig. S1), substantiating its important role in ion transport.

TMS2 and -7 Are Part of the Exchanger Cytoplasmic Ion Translocation Pathway. By examining the effects of the membrane-impermeable thiol-reactive agent MTSET$^+$ on mutants with cysteine substitutions within TMS2 and -7, we have mapped the residues accessible from the cytoplasm. Within TMS2, cysteines at positions 105, 106, 107, 109, and 110 were effectively modified by MTSET$^+$ when the ion binding sites were trapped facing the cytoplasm (inward-facing state). As shown in Fig. 6 and Fig. S2, the model for the inward configuration of NCX_Mj shows a similar ionic-accessible surface of this portion of TMS2 with the exception of serine 51 (equivalent to Ser$^{110}$ of NCX1.1), which is modeled as buried within the protein. However, our data clearly indicate exposure of this residue (S110C) to the cytoplasm when NCX1.1 is in the inward-facing configuration, suggesting a pocket deeper than predicted by the structural model. This result may underlie potential differences between the mammalian and archaeabacterial isoforms. However, it is worth noting that the structural model represents a static picture. The protein during electrophysiological measurements is dynamic, and therefore, the cavity size likely fluctuates, making certain residues transiently accessible.

We investigated the accessibility of residues during transport initiated by the addition of Na$^+$ at the intracellular surface. Currents generated by mutant exchangers M105C and S109C were completely blocked by MTSET$^+$ in both the presence and absence of Na$^+$. Cytoplasmic Na$^+$, however, eliminated cysteine reactivity at positions 106, 107, and 110. Conformational changes occurring during transport may render these positions inaccessible, and/or a short-lived inward state may not provide sufficient time for cysteine modification. In contrast, mutants M105C and S109C seem to have greater exposure, allowing MTSET$^+$ to also react during cycling. Note that, in the absence of Na$^+$, MTSET$^+$ reacted faster with these cysteines than mutations at positions 106, 107, 109, and 110. Therefore, our data clearly indicate exposure of these residues to the cytoplasm when NCX1.1 is in the inward-facing configuration, suggesting a pocket deeper than predicted by the structural model.
and 110 (Fig. C and D), supporting the idea that residues 105 and 109 have maximal accessibility from the cytoplasm when NCX1.1 is in the inward-facing state. An alternative explanation is that positions 105 and 109 are always exposed to the cytoplasmic aqueous environment, independent of the transport cycle. We consider the latter to be the less probable scenario, because residue 109 is part of one of the Na⁺ binding sites, as revealed by the NCX_Mj structure, and likely to become occluded during the transport cycle.

To map further the aqueous cavity, we analyzed the cytoplasmic accessibility of residues within TMS7. Among the eight mutants (A807C, L808C, G809C, V812C, P813C, F816C, A817C, and V820C), only currents recorded from the exchangers G809C, V812C, P813C, and A817C were inhibited by cytoplasmic MTSET⁰. By analogy to NCX_Mj, residues Gly⁸⁰⁹, Val⁸¹², and Ala⁸¹⁷ are close to the binding site for Ca⁺⁺ near the center of the protein, and their accessibility indicates the presence of a deep cavity reaching the core of NCX1.1. As seen for positions 105 and 109, the rates at which MTSET⁰ blocked G809C, V812C, and P813C were extremely fast in the absence of transport, but the kinetics of reactivity markedly slowed during ion exchange.

The present accessibility data provide strong evidence that TMS2 and -7 are part of the ion translocation pathway of NCX1.1, and the changes in accessibility under different states of the transport cycle show that these regions rearrange during exchange.

The Eukaryotic Exchanger Seems to Have Two Cytoplasmic Ion Permeation Pathways of Different Sizes. The consistency between our accessibility results and the recently proposed structural model of NCX_Mj (Fig. 6 and Fig. S2) strongly supports the presence of two inferred deep aqueous cavities that may allow Na⁺ and Ca⁺⁺ to reach their ion-coordinating sites. The first aqueous cavity is delimited by both TMS2 and TMS7, whereas the second is defined by residues Gly⁸⁰⁹, Val⁸¹², and Ala⁸¹⁷, and it is quite wide to accommodate the bulky reagent MTS-TPAE. The presence of a large cavity at these positions is also supported by kinetic studies revealing the fast time course of the MTSET⁰ block for P813C and V812C, regardless of whether NCX1.1 is held in the inward state or cycles. The result suggests that cytoplasmic MTSET⁰ can rapidly reach these positions as NCX1.1 transitions through the inward state during transport. The second cavity seems to be narrower, as judged by the slow reactivity of MTSET⁰ with the facing side chains of cysteines at positions 106, 107, and 110 of TMS2 and 833 of TMS8, in the inward configuration and the absence of MTSET⁰ modification of the same residues during transport. Consistent with this hypothesis is the faster reactivity of position 110 with the smaller MTSMT⁰ reagent when NCX1.1 is held in the inward state and the inability of MTS-TPAE to significantly modify S110C currents.

It has been proposed that the two ion permeation pathways allow Na⁺ and Ca⁺⁺ to access their respective binding sites separately (7). Whether the kinetic data and size-dependent MTS reagents accessibility confirm the notion of the wider cavity being responsible for Ca⁺⁺ access and the narrower cavity being responsible for Na⁺ access will require additional investigation. Nevertheless, our data provide experimental evidence of two aqueous cavities of different sizes accessible from the cytoplasmic face of NCX1.1 that undergo conformational changes during transport. Comparison between the accessibility data presented here and the structural model of the inward-facing configuration of NCX_Mj suggests that these two proteins have similar architecture within the α-repeats. The data validates the structural information and suggests that the high-resolution structure of the archaeabacterial NCX provides a foundation for studying the eukaryotic exchanger.

Methods

Molecular Biology. Mutations were generated and sequenced in 400- to 1000-bp cassettes from the full-length exchanger using QuikChange Mutagenesis (Stratagene). RNA was synthesized from linearized DNA using T3 mMessage mMachine (Ambion) and injected into Xenopus laevis oocytes as described in ref. 6.

Electrophysiology. The apparent affinities for Na⁺ and Ca⁺⁺ and sensitivity to MTS reagents were assessed in giant inside-out patches (6, 16). Before recording, the excised patch is briefly exposed to chymotrypsin to remove NCX1.1 Na⁺ and Ca⁺⁺ secondary regulation (17). Experiments are performed at Vₒ = 0 mV and 35 °C. Intracellular solutions were rapidly changed using a computer controlled 20-channel solution switcher. Pipettes were filled with 100 mM N-methylglucamine, 20 mM Hepes, 20 mM Heps, 20 mM tetraethylammonium hydroxide, 8 mM Ca(OH)₂, 0.1 mM niflumic acid, and 0.15 mM ouabain, pH 7 (using methanesulfonic acid). Bath solution contained 100 mM CsOH or NaOH, 20 mM tetraethylammonium hydroxide, 20 mM Hepes, and 10 mM EGTA, pH 7 (using methanesulfonic acid). MTS reagents (Toronto Research Chemicals) (Fig. S3) solutions were prepared immediately before each experiment because of the instability of compounds. MTSET⁰ (5 mM), MTSMT⁰ (5 mM), and MTS-TPAE (100 μM) were applied cytoplasmically in presence of either cytoplasmic Ca⁺⁺ or Na⁺ and constant 8 mM external Ca⁺⁺ (pipette).

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