Cloning, Expression, and Characterization of the Squid Na\textsuperscript{+}–Ca\textsuperscript{2+} Exchanger (NCX-SQ1)

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ABSTRACT We have cloned the squid neuronal Na\textsuperscript{+}–Ca\textsuperscript{2+} exchanger, NCX-SQ1, expressed it in Xenopus oocytes, and characterized its regulatory and ion transport properties in giant excised membrane patches. The squid exchanger shows 58% identity with the canine Na\textsuperscript{+}–Ca\textsuperscript{2+} exchanger (NCX1.1). Regions determined to be of functional importance in NCX1 are well conserved. Unique among exchanger sequences to date, NCX-SQ1 has a potential protein kinase C phosphorylation site (threonine 184) between transmembrane segments 3 and 4 and a tyrosine kinase site in the Ca\textsuperscript{2+} binding region (tyrosine 462). There is a deletion of 47 amino acids in the large intracellular loop of NCX-SQ1 in comparison with NCX1. Similar to NCX1, expression of NCX-SQ1 in Xenopus oocytes induced cytoplasmic Na\textsuperscript{+}-dependent Ca\textsuperscript{2+} uptake; the uptake was inhibited by injection of Ca\textsuperscript{2+} chelators. In giant excised membrane patches, the NCX-SQ1 outward exchange current showed Na\textsuperscript{+}-dependent inactivation, secondary activation by cytoplasmic Ca\textsuperscript{2+}, and activation by chymotrypsin. The NCX-SQ1 exchange current was strongly stimulated by both ATP and the ATP-thioester, ATP\textsubscript{γ}S, in the presence of F\textsuperscript{−} (0.2 mM) and vanadate (50 μM), and both effects reversed on application of a phosphatidylinositol-4,5-bisphosphate antibody. NCX1 current was stimulated by ATP, but not by ATP\textsubscript{γ}S. Like NCX1 current, NCX-SQ1 current was strongly stimulated by phosphatidylinositol-4,5-bisphosphate liposomes. In contrast to results in squid axon, NCX-SQ1 was not stimulated by phosphoarginine (5–10 mM). After chymotrypsin treatment, both the outward and inward NCX-SQ1 exchange currents were more strongly voltage dependent than NCX1 currents. Ion concentration jump experiments were performed to estimate the relative electrogenicity of Na\textsuperscript{+} and Ca\textsuperscript{2+} transport reactions. Outward current transients associated with Na\textsuperscript{+} extrusion were much smaller for NCX-SQ1 than NCX1, and inward current transients associated with Ca\textsuperscript{2+} extrusion were much larger. For NCX-SQ1, charge movements of Ca\textsuperscript{2+} transport could be defined in voltage jump experiments with a low cytoplasmic Ca\textsuperscript{2+} (2 μM) in the presence of high extracellular Ca\textsuperscript{2+} (4 mM). The rates of charge movements showed “U”-shaped dependence on voltage, and the slopes of both charge–voltage and rate–voltage relations (1,600 s\textsuperscript{−1} at 0 mV) indicated an apparent valency of ~0.6 charges for the underlying reaction. Evidently, more negative charge moves into the membrane field in NCX-SQ1 than in NCX1 when ions are occluded into binding sites.

KEY WORDS: sodium–calcium exchange • charge movements • Xenopus oocytes • patch clamp • phosphatidylinositol

INTRODUCTION

Na\textsuperscript{+}–Ca\textsuperscript{2+} exchangers mediate large Ca\textsuperscript{2+} fluxes across the plasma membranes of many cell types and thereby modulate diverse cell functions (for overview, see Hilgemann et al., 1996). Their major physiological role is to extrude Ca\textsuperscript{2+} in exchange for extracellular Na\textsuperscript{+}, which is subsequently extruded by the ATP-dependent Na\textsuperscript{+}–K\textsuperscript{+} pump.

Historically, two experimental models were used to characterize Na\textsuperscript{+}–Ca\textsuperscript{2+} exchange function: cardiac sarcolemmal vesicles (for review, see Philipson and Reeves, 1989) and perfused squid giant axons (for review, see DiPolo, 1989; DiPolo and Beaugé, 1991). Molecular studies of the cardiac Na\textsuperscript{+}–Ca\textsuperscript{2+} exchanger (NCX1) were made possible by its cloning (Nicoll et al., 1990). This was followed by the cloning and functional characterization of two other mammalian exchangers (NCX2 and NCX3; Li et al., 1994; Nicoll et al., 1996a, 1996b) and an NCX-type exchanger from Drosophila (Ruknudin et al., 1997; Schwarz and Benzer, 1997). Mutational studies of NCX1 have identified functional domains involved in both regulation and transport (Matsuoka et al., 1993, 1995, 1997; Levitsky et al., 1994; Nicoll et al., 1996a). Given the extensive nature of Na\textsuperscript{+}–Ca\textsuperscript{2+} exchange studies in squid giant axons, cloning and mo-
lecular characterization of the squid exchanger could provide many unique insights.

Basic properties of the cardiac and squid exchangers are clearly similar. These include a 3 Na\(^+\) to 1 Ca\(^{2+}\) stoichiometry, regulatory activation of exchanger-mediated Ca\(^{2+}\) influx by cytoplasmic Ca\(^{2+}\) (DiPolo, 1979; Kimura et al., 1986), and stimulation by ATP-dependent mechanisms (DiPolo, 1974; Baker and McNaughton, 1976; Hilgemann, 1990). Nevertheless, recent results suggest that there are important differences in the function and regulation of the different exchangers. (a) The ATP dependence of the squid exchanger appears to reflect its phosphorylation by a protein kinase (DiPolo and Beaugé, 1994; DiPolo et al., 1997), while the ATP-dependent activation of the cardiac exchanger appears to reflect the generation of phosphatidylinositol 4,5-bisphosphate (PIP\(_2\)) from phosphatidylinositol (Hilgemann and Ball, 1997). ATPPyS activates the squid exchanger but not the cardiac exchanger. Cat-ionic agents that bind anionic lipids inhibit the cardiac exchanger (Hilgemann and Collins, 1992), but agents such as pentalisine do not inhibit the squid exchanger (R. DiPolo and L. Beaugé, 1993). (b) The squid exchanger is regulated by a phosphoarginine-dependent process that may involve protein kinases unique to invertebrates (DiPolo and Beaugé, 1995); phosphoarginine is without effect on the cardiac exchanger (D.W. Hilgemann, unpublished observations). (c) The Ca\(^{2+}\)–Ca\(^{2+}\) exchange operation of the squid exchanger (DiPolo et al., 1985; DiPolo and Beaugé, 1990) and of barnacle muscle (Rasgado-Flores et al., 1996) appears to be strongly voltage dependent, while Na\(^+\)–Na\(^+\) exchange shows almost no voltage dependence in isotope flux studies. For the cardiac exchanger, on the other hand, Na\(^+\) transport has been shown to be strongly electrogenic (Hilgemann et al., 1991; Matsuoka and Hilgemann, 1992; Powell et al., 1993), while Ca\(^{2+}\)–Ca\(^{2+}\) exchange is only weakly voltage dependent (Hilgemann et al., 1991; Matsuoka and Hilgemann, 1992; Powell et al., 1993; Niggli and Lederer, 1991; Kappl and Hartung, 1996).

To better compare the function and structure of the exchangers, we have now cloned the squid neuronal

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1Abbreviations used in this paper: MES, 2-(morpholino)ethanesulfonic acid; NMG, N-methyl-D-glucamine; nts, nucleotides; PIP\(_2\), phosphatidylinositol-4,5-bisphosphate.
A oocyte was variable and depended on the oocyte batch. Na

The inhibition of an oocyte patch with symmetrical Na

The extracellular solution contained 4 mM Ca-sulfamic acid, 1 mM Mg-sulfamic acid, 40 mM Na-2-(morpholino)ethanesulfonic acid (MES), 20 mM Cs-MES, 20 mM tetraethyl ammonium (TEA)-MES, 40 mM N-methyl-d-glucamine (NGM)-MES, and 20 mM HEPES, adjusted to pH 7.0 with NMG. The cytoplasmic solution contained (mM): 10 EGTA, 6 Ca-sulfamic acid, 0.5 Mg-sulfamic acid, 60 Cs-MES, 20 TEA-MES, and either 40 additional Cs-MES or 40 Na-MES to activate outward exchange current, at pH 7.0 with NMG (pCa 6.5). Gigaohm seals were made in a solution containing (mM): 80 K-aspartate, 40 KCl, 4 MgCl₂, 5 EGTA, and 10 HEPES, at pH 7.0 with NMG. Experiments were performed at 32°C. Concentrated stock solutions of nucleotides were prepared as Mg²⁺/TRIS salts, with the Mg²⁺ concentration adjusted to 75% of the total nucleotide concentration. In this way, the free Mg²⁺ concentration (0.3 mM) is not changed on addition of nucleotide.

Endogenous conductances of the oocyte membrane were found to be activated by ATP and anionic lipids, so more extensive control experiments were needed to test for the influence of contaminating currents. Most importantly, we established conditions such that current changes in patches from uninjected (or water-injected) oocytes amounted to at most a few picoamperes using the same conditions and protocols employed in results (i.e., at 0 mV).

Since our efforts to understand and minimize endogenous conductances are relevant to many expression studies with Xenopus oocytes, we describe here the two major oocyte currents activated by ATP and PIP₂: Ca²⁺-activated Cl⁻ and voltage-activated Na⁺ currents. Activation of Ca²⁺-activated Cl⁻ current by ATP (Hilgemann, 1995) has been described previously. As shown in Fig. 1, Ca²⁺-activated Cl⁻ current is stimulated by many polyvalent anions (0.1–2 mM) and by F⁻ when cytoplasmic free Ca²⁺ is sub-maximal. Fig. 1 A shows results for phosphate. Activation by polyvalent anions typically takes ~1 min, but the effect decays on removal of anions in only a few seconds. Possibly, these effects reflect chelation of trace polyvalent cations in solutions (or from the pipette) by these anions (Hilgemann, 1997). Since the pipette tip can become contaminated with Cl⁻ during seal formation, there is a danger that outward Cl⁻ current can occur during exchange current measurements. Inclusion of Cl⁻ current blockers in the pipette solution (0.3 mM niflumic acid + 0.3 mM flurometric acid) effectively blocked the residual Cl⁻ currents, while results with PIP₂ and nucleotides in exchanger-expressing patches were not changed.

As shown in Fig. 2 A, a Na⁺ current can be activated in oocyte patches by depolarization for a few seconds beyond 0 mV. As described in previous two-electrode voltage clamp studies (Baud and Kado, 1984), this current activates and deactivates rapidly in response to voltage steps after it has first been primed by a long depolarization. Fig. 2 B shows the steady state current-voltage relation of an oocyte patch with symmetrical Na⁺-containing solutions (40 mM). We stress that the Na⁺ conductance was always negligible at potentials more negative than ~20 mV. In the absence of ATP and PIP₂, the magnitude of the Na⁺ conductance was variable and depended on the oocyte batch. Na⁺ conductance could be induced in the majority of patches by ATP or an-

**Preparation of NCX-SQ1 Fusion Protein and Antibody Production**

An expression construct containing the cDNA coding for the large intracellular loop of NCX-SQ1 was constructed by PCR. The 5' end of the forward primer (5'-AACGATGGGTGGATGCTGCATGTG-3'; nts 1677–1786) included an introduced SpH restriction site and the reverse primer (5'-TCTCCAGGATGGCTTACATAAAGCT-3'; nts 3097–2989) contained an introduced PstI restriction site at the 3' end. pBluescript SK² vector containing the NCX-SQ1 full-length cDNA (10 ng) was amplified using the TA Cloning Kit, and the identity of the clone was confirmed by sequencing. The PCR construct was ligated into the expression vector pQE (QIAGEN Inc., Chatsworth, CA). The loop fusion protein was expressed and purified as described by He et al. (1997) and used as antigen for generation of polyclonal antisera in rabbits (HRP Inc.; Denver, PA).

**Western Blot**

Samples from oocytes expressing NCX-SQ1 protein were prepared as follows: 10 oocytes were sonicated in 100 µl of homogenization buffer (0.1 M NaCl, 1% Triton X-100, 20 mM Tris-HCl, pH 7.6) and centrifuged at 12,000 rpm for 10 min. The supernatant was filtered twice through Spin-X filters to remove lipids and protease inhibitors buffer (0.1 M NaCl, 1% Triton X-100, 20 mM Tris-HCl, pH 7.4, 0.05% Tween 20). The membrane was incubated with the antiserum overnight (i.e., at 0 mV).
at the reversal potential of the Na$^+$-Ca$^{2+}$ exchanger. Potassium phosphate (with 3 mM Mg$^{2+}$, pH 7.0) enhances current by approximately fourfold over 1 min. The stimulatory effect reverses in just a few seconds on removal of phosphate. (B) Normalized stimulatory effects of other anions in the same protocol. “Basal” current corresponds to the current activated with 1 mM free Ca$^{2+}$. From left to right, results are shown for 2 mM Mg-ATP, 2 mM Mg-AMPPNP, 2 mM ATP to the absence of Mg$^{2+}$ from all solutions, 2 mM Mg-GTP, 1 mM Mg-GTPyS, 0.5 mM pyrophosphate (PP) in the absence of Mg$^{2+}$ from all solutions, 4 mM p-nitrophenylphosphate (PnPP), 4 mM Mg-phosphoarginine (P-Arg), 4 mM F$^-$ in the absence of Mg$^{2+}$ from all solutions, and 2 mM EDTA in the absence of Mg$^{2+}$ from all solutions.

**R E S U L T S**

### Cloning the cDNA Coding for the Squid Na$^+-$Ca$^{2+}$ Exchanger

A fragment of the squid exchanger cDNA was obtained by PCR amplification of cDNA derived from a squid optic lobe library. Degenerate primers were designed based on conserved amino acid sequences of proposed transmembrane segments 6 and 9 of the mammalian exchangers NCX1 and NCX2. An appropriately sized PCR product (~320 bp) was subcloned. Sequence analysis indicated that the DNA coded for a protein homologous to NCX1. The PCR product was used to screen a squid optic lobe library and a partial clone of 1.5 kb was isolated with ~60% identity to NCX1 at the amino acid level. Longer clones were not found in the optic lobe library. A fragment of the partial 1.5-kb clone was then used to screen a squid stellate ganglion cDNA library. Two clones containing the complete coding sequence of the squid exchanger were isolated. These clones were identical at the 3’ end. The complete nucleotide and amino acid sequences of the longest clone, SG14, are shown in Fig. 3. SG14 is 4,906 bp long with an open reading frame of 2,676 nucleotides encoding for a protein of 892 amino acids, which we refer to as NCX-SQ1. The most 5’ ATG, in the proper reading frame, begins at nucleotide 994 with some features of a Kozak (1989) consensus initiation site. The 3’ end of the cDNA terminates with 16 adenosines preceded by multiple consensus polyadenylation sites.

The amino acid sequence of the squid stellate ganglion Na$^+$–Ca$^{2+}$ exchanger, aligned with the sequence of the canine cardiac Na$^+$–Ca$^{2+}$ exchanger (NCX1.1 in...
the terminology of Quednau et al., 1997) is shown in Fig. 4. Like NCX1, NCX-SQ1 has 12 hydrophobic segments that could form transmembrane segments. In NCX1, the first hydrophobic segment at the NH2 terminus is a cleaved leader peptide and is removed from the protein during biosynthesis in the endoplasmic reticulum (Durkin et al., 1991; Hryshko et al., 1993). NCX-SQ1 has a potential signal peptidase-recognition site
The Squid Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchanger (NCX-SQ1) (Von Heijne, 1983) after alanine 26, and we predict that a signal peptide is removed from this protein as well. Thus, the mature protein is modeled to have 11 transmembrane segments with a large intracellular loop separating transmembrane segments 5 and 6. The NH\textsubscript{2} terminus would be extracellular and the COOH terminus would be intracellular. NCX-SQ1 has nine potential sites for N-linked glycosylation. If the glycosylation pattern is similar to that of NCX1 (Hryshko et al., 1993), only the first site (asparagine 31) would be glycosylated. Overall, the mature NCX-SQ1 and NCX1 proteins are 58% identical, ignoring the gap in NCX-SQ1 in the intracellular loop region. More sequence analysis will be presented below.

The transcript size of NCX-SQ1 was determined by Northern blot analysis. Poly (A)\textsuperscript{+} RNA from squid optic lobe and stellate ganglion hybridized with an NCX-SQ1 probe at 9.6 kb (Fig. 5 A). The signal was substantially stronger in stellate ganglion than in optic lobe. The NCX-SQ1 mRNA is larger than that for NCX1 (7 kb), NCX2 (5 kb), or NCX3 (6 kb) (Nicoll et al., 1996). The NCX-SQ1 transcript apparently has a long 5' untranslated region since the clone ends with a poly (A)\textsuperscript{+} tail. Extensive untranslated regions also occur for the squid Na\textsuperscript{+} channel that has a transcript 12 kb (Rosenthal et al., 1993), which is larger than that of the mammalian Na\textsuperscript{+} channels.

A His-tagged fusion protein encompassing amino acids 259–665 of NCX-SQ1 was expressed in Escherichia coli. After affinity purification, the fusion protein was used for polyclonal antibody production in rabbit. Expression of NCX-SQ1 in membrane vesicles from squid optic lobe was detected by immunoblot (Fig. 5 B, lane 1). The antibody recognizes a major band at z120 kD and two bands at lower molecular weights. NCX1 produces a similar pattern on immunoblots (Nicoll et al., 1990). Preimmune serum from the same rabbit failed to produce any signal (not shown). Expression of exchanger protein in Xenopus oocytes injected with NCX-SQ1 cRNA was also examined. A protein band of z100 Figure 4. Amino acid comparison of the squid NCX-SQ1 and the canine NCX1 exchanger. Putative transmembrane segments, predicted by hydropathy analysis, are underlined and numbered. Highlighted in bold lettering are a potential signal peptide site (SigPase), potential N-linked glycosylation sites (NXS/T), and potential phosphorylation sites (RTIK, protein kinase C; TRKLT, cAMP-dependent kinase and Ca\textsuperscript{2+}/calmodulin-dependent kinase; DEHY and DDEEEY, tyrosine kinase). The two potential phosphorylation sites marked with an asterisk are unique to NCX-SQ1. The endogenous exchanger inhibitory peptide (XIP) region and Exon A are shaded, and the binding domain for regulatory Ca\textsuperscript{2+} is boxed. The triple aspartate motifs involved in Ca\textsuperscript{2+} binding are in bold. Dots in the NCX1 sequence indicate amino acids identical to those of NCX-SQ1.

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kD was detected in a membrane fraction from cRNA-injected oocytes, whereas no signal was detected in control water-injected oocytes (Fig. 5 B, lanes 2 and 3). The small difference in apparent molecular weight between the native squid exchanger and NCX-SQ1 expressed in oocytes is possibly due to a difference in glycosylation.

Functional Expression of NCX-SQ1 in Xenopus Oocytes

cRNA encoding NCX-SQ1 was synthesized from linearized plasmids and injected into Xenopus oocytes. Expression was optimized when the 5'- and 3'-untranslated regions of Xenopus β globin flanked the NCX-SQ1 coding region. Expression of Na⁺–Ca²⁺ exchange activity was assessed by measuring ⁴⁵Ca²⁺ fluxes into intact oocytes and by measuring exchanger currents using the giant excised patch technique. An example of Na⁺ gradient-dependent ⁴⁵Ca²⁺ uptake into Na⁺-loaded Xenopus oocytes is shown in Fig. 6. The first columns show that NCX-SQ1 RNA induces a substantial uptake of ⁴⁵Ca²⁺ in the presence of an outwardly directed Na⁺ gradient (K⁺o). This uptake is abolished in the absence of the Na⁺ gradient (Na⁺o). We have validated the use of this approach to measure Na⁺–Ca²⁺ exchange activity in previous studies (Longoni et al., 1988; Nicoll et al., 1990). No Ca²⁺ uptake was observed in control water-injected oocytes.

We used the ⁴⁵Ca²⁺ uptake assay to determine whether NCX-SQ1 exhibited secondary Ca²⁺ regulation by intracellular Ca²⁺. The presence of micromolar levels of [Ca²⁺], is required in squid giant axons to activate Na⁺-,dependent Ca²⁺ uptake (Baker and McNaughton, 1976; DiPolo and Beaugé, 1986). Oocytes expressing NCX1-SQ1 were injected with EGTA before the Ca²⁺ uptake assay. Chelation of internal Ca²⁺ blocked most Ca²⁺ uptake (Fig. 6), which is consistent with previous studies of squid axons and NCX1 (Baker and McNaughton, 1976; DiPolo and Beaugé, 1986; Kimura et al., 1986; Hilgemann et al., 1992a, 1992b).

Outward Exchange Current of the Squid (NCX-SQ1)

Fig. 7 shows basic properties of outward Na⁺–Ca²⁺ exchange current in patches from oocytes expressing the NCX-SQ1 exchanger. In brief, no obvious property of the current is different from NCX1 exchange current. On application of cytoplasmic Na⁺, the current activates in the solution switch time and then shows partial

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**Figure 6.** Functional expression of NCX-SQ1 in Xenopus oocytes. Oocytes injected with cRNA for NCX-SQ1 (A) or control (C), water-injected oocytes were assayed for Na⁺–Ca²⁺ exchanger activity. ⁴⁵Ca²⁺ uptake into Na⁺ (90 mM)-loaded oocytes was measured in cells diluted into ⁴⁵Ca²⁺-containing medium in the presence (extracellular K⁺) or absence of an outwardly directed Na⁺ gradient (extracellular Na⁺). In the middle pair of columns (B), 46 nl of a 100 mM EGTA solution was injected into the oocytes before loading the cells with Na⁺ to deplete intracellular Ca²⁺.

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**Figure 5.** (left) Northern blot analysis of NCX-SQ1 RNA. mRNA (1 μg) from squid optical lobe (lane 1) and stellate ganglia (lane 2) was probed with a fragment of the NCX-SQ1 cDNA. (right) Western blot analysis of NCX-SQ1 protein. Protein from squid optical lobe vesicles (lane 1) and oocytes injected with water (lane 2) or cRNA for NCX-SQ1 (lane 3) was probed with an antibody raised against a histidine-tagged fusion protein fragment of NCX-SQ1.
inactivation over several seconds (see Fig. 7, first record, in the presence of 1 μM cytoplasmic free Ca^{2+}). When cytoplasmic Ca^{2+} is removed, the current magnitude decreases, and inactivation on application of Na^{+} is subsequently faster. Exchange current remains substantial (second record). With 5 μM free cytoplasmic Ca^{2+} (third record), or higher concentrations (not shown), the current did not increase further and ran down with time. Current run-down prevented us from determining Ca^{2+}–current relations in more detail. Application of α-chymotrypsin (1 mg/ml) activated the exchange current over 1 min. The final current magnitude was typically more than twice the peak current magnitude obtained on applying Na^{+}. After chymotrypsin, the exchange current was insensitive to changes of cytoplasmic free Ca^{2+} from 0 to 5 μM.

Stimulation of NCX1 Exchange Current by ATP and Reversal of Stimulation by Anti–PIP_2 Antibody

ATP strongly activates exchange current in excised cardiac membrane patches and vanadate is without effect on the stimulatory effect or its reversal after removal of ATP (Collins et al., 1992). However, in our initial studies of the cardiac exchanger (NCX1) in oocyte patches, no effects of ATP were observed (Matsuoka et al., 1993). We now report that ATP indeed can be effective when the phosphatase inhibitors, F− and vanadate, are included in the cytoplasmic solution. The concentrations of F− and vanadate are selected to maintain free Mg^{2+} in the range of 0.1–0.3 mM, and vanadate alone was usually effective. Fig. 8 describes typical results with 0.1 mM F− and 50 μM vanadate (0.5 mM total Mg^{2+} in nucleotide-free solution). The outward current activated by Na^{+} shows the usual inactivation when 40 mM cytoplasmic Na^{+} is applied. As shown in Fig. 8 A, 2 mM of the nonhydrolyzable ATP analogue, AMP-PNP, was without effect, while 2 mM ATP increased the current to a magnitude somewhat greater than the peak current on initial application of cytoplasmic Na^{+}. ATP strongly activates exchange current in excised cardiac membrane patches and vanadate is without effect, whereas 2 mM Mg-ATP stimulates the current to a magnitude somewhat greater than the initial peak current on application of cytoplasmic Na^{+}. The stimulatory effect reverses partially over 2 min. (B) Application of 2 mM Mg-ATPgS has only a small stimulatory effect compared with 2 mM Mg-ATP, applied subsequently. The stimulatory effect reverses by ∼20% over 2 min, and it reverses almost completely in 1 min on application of PIP_2 antibody (AB).

Stimulation of Squid (NCX-SQ1) Na^{+}–Ca^{2+} Exchange by ATP and Its Reversal by Anti–PIP_2 Antibody

Fig. 9 shows typical effects of ATP on the outward NCX-SQ1 exchange currents. In Fig. 9 A, the current was first activated by cytoplasmic Na^{+}, and it then acti-
Stimulation of the Squid (NCX-SQ1) Na\(^+\)–Ca\(^{2+}\) Exchange Current by ATP

The NCX-SQ1 Na\(^+\)–Ca\(^{2+}\) exchange current, in contrast to NCX1 current, was usually strongly stimulated by applying ATP\(\gamma\)S (four of six results). Typical results are shown in Fig. 10. In the presence of F\(^-\) and vanadate, application of a nonhydrolyzable ATP derivative, AMP-PNP (2 mM) was nearly without effect (Fig. 10 A); the small inhibition of exchange current may be due to a small increase of free Mg\(^2+\). Thereafter, application of 2 mM ATP\(\gamma\)S stimulates the exchange current roughly to the extent observed with ATP in NCX-SQ1-expressing patches. The stimulatory effect reverses within a few minutes on removal of the nucleotide. As shown in Fig. 10 B, PIP\(_2\) antibody can reverse the effect of ATP\(\gamma\)S over the course of 90 s, suggesting a possible involvement of PIP\(_2\) in the effects of ATP\(\gamma\)S as well as those of ATP.

Stimulation of the Squid (NCX-SQ1) Na\(^+\)–Ca\(^{2+}\) Exchange Current by PIP\(_2\) Antibody

Fig. 11 A shows the typical stimulatory effect of PIP\(_2\) liposomes on NCX-SQ1 exchange current. Exchange current was activated by application of cytoplasmic Na\(^+\), and it was allowed to run-down for 3 min to a very small magnitude. Then, 50 \(\mu\)M PIP\(_2\) was applied and current increased over several minutes, comparable to the effect of chymotrypsin (see Fig. 7). To check for con-
fllicting ionic currents, current–voltage relations were acquired at several times. The records 1, 2, and 3, indicated in Fig. 1 A, were subtracted so as to define the current activated by PIP₂ (±PIP₂; Fig. 11 B, 2–I) and the current switched off by removing Na⁺ at the end of the recording period (±Na⁺; Fig. 11 B, 2–3). The two current–voltage relations are very similar, indicating that PIP₂-activated outward current is also activated by cytoplasmic Na⁺, and the shapes of current–voltage relations are typical for outward exchange current. We note also that similar results were obtained in patches from HEK cells in which NCX-SQ1 was expressed by transient transfection, although the maximum exchange currents were only ~15 pA (not shown).

Fig. 11 C shows the inactivation time courses observed on application of cytoplasmic Na⁺, before and after applying PIP₂ in another NCX-SQ1-expressing patch. Single exponential functions were fitted to the inactivation phases, and they are plotted as dotted lines. The time constant of inactivation increases from 3.9 s in control records to 8.2 s after application of PIP₂. This is similar to the slowing of inactivation observed with cardiac exchange current when ATP is applied (Hilgemann et al., 1992b), indicating that similar molecular mechanisms might be involved.

Lack of Effect of Phosphoarginine on Outward Squid (NCX-SQ1) Na⁺–Ca²⁺ Exchange Current in Oocyte Patches

As noted in the introduction, the high-energy compound phosphoarginine stimulates the Na⁺–Ca²⁺ exchange process in squid axons, probably by a mechanism that involves phosphorylation (DiPolo and Beaugé, 1995). As shown in Fig. 12, phosphoarginine (5 mM) was without effect on the outward exchange current in oocyte patches (four observations, applying phosphoarginine with Mg²⁺). In the same patch, 50 μM PIP₂ was highly effective. As with all nucleotides, other phosphates, and citrate, large stimulatory effects were observed in patches when phosphoarginine was applied without added Mg²⁺ (not shown). The probable explanation for our results in patches is that all of these anions chelate Mg²⁺ and thereby relieve an inhibition of exchange current by cytoplasmic Mg²⁺.

Voltage Dependence of Chymotrypsin-deregulated Outward NCX-SQ1 Exchange Currents

To compare the voltage dependencies of NCX-SQ1 exchange current with those of cardiac exchange current, we examined both the outward and inward exchange currents under “zero trans” conditions (i.e., with Na⁺ and no Ca²⁺ on one membrane side, and Ca²⁺ but no Na⁺ on the other side). This was carried out in chymotrypsin-treated patches, so that regulatory mechanisms were absent. We allowed chloride and sodium currents to “run down” for a few minutes in Mg²⁺-containing solution before application of α-chymotrypsin (1 mg/ml for 30 s). Current changes were negligible in patches from uninjected oocytes, using the same conditions and protocols.

Fig. 13 A shows current–voltage relations for outward NCX-SQ1 exchange current with 4 mM extracellular Ca²⁺ and no extracellular Na⁺. With the Ca²⁺-free/10 mM EGTA cytoplasmic solution employed in these experiments, Ca²⁺-activated Cl⁻ conductance is zero, and

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**Figure 11.** Stimulation of outward NCX-SQ1 Na⁺–Ca²⁺ exchange current by PIP₂. (A) Current was activated by cytoplasmic Na⁺ and was then allowed to run down to <5 pA. PIP₂ was applied and the outward current increased over 5 min to a magnitude more than twofold greater than the peak current on application of Na⁺. Current–voltage relations were taken just after application of PIP₂ (I), after the maximum stimulatory effect was obtained (2), and after exchange current was turned off by removing cytoplasmic Na⁺ (3). (B) Current–voltage relation of the exchange current. The exchange current is defined by subtracting records before PIP₂ application from those with PIP₂ (2–I, ○) and those after removing Na⁺ from those with Na⁺ plus PIP₂ (2–3, ○). (C) Comparison of current transients obtained on activating exchange current before (control) and after (+PIP₂) applying PIP₂. The time constant (τ) of inactivation increases from 3.9 to 8.2 s. Results are from a different patch.
the same solutions (with 20 mM Cl\textsuperscript{−}) could be used as in previous measurements with cardiac membrane patches (Hilgemann et al., 1992a). Cytoplasmic Na\textsuperscript{+} was varied from 5 to 90 mM, substituting it for Cs\textsuperscript{+}, and baseline current–voltage relations in the absence of Na\textsuperscript{+} were subtracted. The current–voltage relations are similar in shape and they can be scaled well to each other (not illustrated). Fig. 13 B shows the Na\textsuperscript{+} dependence of outward exchange current at −60 and +60 mV; the K\textsubscript{50} (half-maximal concentration) for Na\textsuperscript{+} is 27 mM at +60 mV and 24 mM at −60 mV, and the Hill slopes are 1.2 and 1.7, respectively. For the cardiac exchanger, by contrast, current–voltage relations become less steep with high cytoplasmic Na\textsuperscript{+}, and the K\textsubscript{50} for Na\textsuperscript{+} decreases somewhat at positive potentials (Matsuoka and Hilgemann, 1992).

**Fig. 13.** Outward NCX-SQ1 Na\textsuperscript{+}–Ca\textsuperscript{2+} exchange current in a chymotrypsin-treated patch. The pipette solution contains 4 mM Ca\textsuperscript{2+} and no Na\textsuperscript{+}; the cytoplasmic solution contains 10 mM EGTA and no Ca\textsuperscript{2+}. (A) Current–voltage relations at the given Na\textsuperscript{+} concentrations from 5 to 90 mM. Data points for descending and ascending voltage steps show no hysteresis. (B) Cytoplasmic Na\textsuperscript{+} dependence of outward exchange current at +60 and −60 mV. The data points are fit to a Hill equation; the slope is 1.2 at +60 and 1.7 at −60 mV; the K\textsubscript{50} is 27 mM at +60 mV and 24 mM at −60 mV.

**Fig. 12.** Lack of effect of phosphoarginine (p-ARG) on NCX-SQ1 exchange current in an excised oocyte patch. After activation of the exchange current by applying Na\textsuperscript{+}-containing solution, 5 mM phosphoarginine was applied with 3 mM Mg\textsuperscript{2+} (pH 7.0). There is no stimulatory effect, whereas application of 50 μM PIP\textsubscript{2} strongly stimulates the exchange current to a magnitude more than twofold greater than the peak obtained on applying Na\textsuperscript{+} initially.

As mentioned in the introduction, Ca\textsuperscript{2+} flux studies in squid axons suggest that Ca\textsuperscript{2+} translocation is substantially more voltage dependent than Na\textsuperscript{+} translocation (DiPolo et al., 1989; DiPolo and Beaugé, 1990) and similar results are described for the barnacle Na\textsuperscript{+}–Ca\textsuperscript{2+} exchanger (Rasgado-Flores et al., 1996). To test directly whether Ca\textsuperscript{2+} translocation is electrogenic in NCX-SQ1, ion concentration jump experiments were

Fig. 14 compares inward exchange current–voltage relations for NCX-SQ1 and NCX1, expressed in the same batch of oocytes and using the same Cl\textsuperscript{−}-free solutions. The cytoplasmic solution was the same as described in materials and methods without Na\textsuperscript{+}. The pipette solution contained 120 mM Na\textsuperscript{+} (see Fig. 14, legend, for complete composition). Fig. 14, A and B, shows results for NCX1 and NCX-SQ1, respectively, with 0.2, 2, 10, and 300 μM free cytoplasmic Ca\textsuperscript{2+}. The current–voltage relations of the NCX-SQ1 exchanger are substantially steeper than those for the NCX1 exchanger and have a more “exponential” form. Those for NCX1 are nearly linear over the entire voltage range. For NCX-SQ1, current doubles in ∼30 mV in the steepest region of the current–voltage relations. This is close to the slope expected for a single charge movement across the entire electrical field. Fig. 14, C and D, shows the Ca\textsuperscript{2+} dependencies of the inward current at −150 and −30 mV. Both data sets are well-described by Hill equations with slopes of 1, and there are small shifts of the K\textsubscript{50}'s to higher concentrations at more negative potentials; 2.5–4.2 μM for NCX1, and 3.5–7.2 μM for NCX-SQ1.
The Squid Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchanger (NCX-SQ1)

NCX1.1

NCX-SQ1

\begin{figure}
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\includegraphics[width=\textwidth]{figure14}
\caption{Comparison of inward NCX1 and NCX-SQ1 Na\textsuperscript{+}–Ca\textsuperscript{2+} exchange currents in chymotrypsin-treated patches. The pipette solution contains (mM): 120 Na\textsuperscript{+}, 10 EGTA, 20 Cs\textsuperscript{+}, 20 HEPES, 4 Mg\textsuperscript{2+}, and no Ca\textsuperscript{2+} (pH 7.0 with NMG); the cytoplasmic solution contains 10 mM EGTA and no Na\textsuperscript{+}. The inward current–voltage relations are defined by subtracting records with Ca\textsuperscript{2+} from records without Ca\textsuperscript{2+}. In descending order, the current–voltage relations are with 0.2, 2, 10, and 300 \textmu M Ca\textsuperscript{2+}. (A) Current-voltage relations for NCX1. (B) Current-voltage relations for NCX-SQ1. Same batch of oocytes as in A. (C) Cytoplasmic Ca\textsuperscript{2+} dependence of the inward NCX1 exchange current at \(-150\) and \(-30\) mV. The $K_{50}$ indicated by an arrow, is 2.5 \textmu M at \(-30\) mV and 4.2 \textmu M at \(-150\) mV. (D) Cytoplasmic Ca\textsuperscript{2+} dependence of the inward NCX-SQ1 exchange current at \(-150\) and \(-30\) mV. The $K_{50}$ is 3.5 \textmu M at \(-30\) mV and 7.2 \textmu M at \(-150\) mV.

\end{figure}

designed to isolate the possible charge movements of outward Ca\textsuperscript{2+} and Na\textsuperscript{+} translocation. To do so, the outward translocation of ions is initiated by moving the patch pipette tip through the interface between two solution streams in 1 ms, whereby one stream contains no substrate and the other contains a high substrate concentration. A computer-controlled Piezo-type manipulator is used to move the patch clamp head stage together with the patch pipette (Hilgemann and Lu, 1998). The speed of the substrate concentration change occurring at the membrane surface in these experiments (~150-ms time constant) is determined by diffusion from the pipette orifice to the membrane surface (20–70 \textmu m). We point out, however, that the speed of current activation, observed upon applying a substrate, can be much faster. That is because binding sites can be saturated quickly, compared with average diffusion times, if substrate concentrations are high with respect to binding site affinity. This is the case for experiments with Na\textsuperscript{+} jumps, but it appears not to be the case for Ca\textsuperscript{2+} jumps.

\begin{figure}
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\includegraphics[width=\textwidth]{figure15}
\caption{Identification of electrogenic reactions of NCX1 and NCX-SQ1 using concentration jumps. (A) Current transients recorded from NCX1-expressing patch when 40 mM cytoplasmic Na\textsuperscript{+} is applied and removed in the presence of 20 mM extracellular Na\textsuperscript{+}. (B) Typical lack of current transients recorded from NCX-SQ1-expressing patch when 40 mM cytoplasmic Na\textsuperscript{+} is applied and removed in the presence of 20 mM extracellular Na\textsuperscript{+}. (C) Inward NCX1 current activated when a solution with 5 \textmu M free Ca\textsuperscript{2+} is applied as in A. (D) Inward NCX-SQ1 current activated when a solution with 5 \textmu M free Ca\textsuperscript{2+} is applied as in B. (E) Typical lack of current transients for a Ca\textsuperscript{2+} jump to 5 \textmu M free Ca\textsuperscript{2+} in NCX1-expressing patch; 50 \textmu M extracellular Ca\textsuperscript{2+}. (F) Current transients recorded from NCX-SQ1-expressing patch when a solution with 5 \textmu M free Ca\textsuperscript{2+} is applied and removed in the presence of 50 \textmu M extracellular Ca\textsuperscript{2+}. (G) Outward current activated by applying 40 mM Na\textsuperscript{+} to an NCX1 patch with 50 \textmu M extracellular Ca\textsuperscript{2+}. (H) Outward current activated by applying 40 mM Na\textsuperscript{+} to an NCX1 patch with 50 \textmu M extracellular Ca\textsuperscript{2+}. See text for details.

Fig. 15 illustrates the major experimental results. All patches were chymotrypsin treated and it is noted that experiments were successful only in oocyte batches with high exchanger expression (approximately one batch out of six). The protocols, based on the predicted function of an alternating-access exchange model, were the same as used previously to monitor “half-cycles” of ion transport (Hilgemann et al., 1991): in the presence of substrate on the extracellular side and no substrate on the cytoplasmic side, the exchanger binding sites will orient to the cytoplasmic side and will be free of substrate. When a high concentration of substrate is applied to the cytoplasmic side, substrate will bind and
the binding sites will reorient and open to the extracellular side. Substrate will be released, and binding sites will remain in the extracellular orientation, on average, if the extracellular substrate concentration is relatively low. Thus, charge movement observed during this protocol should reflect the electrogenicity of ion transport for the substrate added.

Fig. 15, A and B, shows results for jumping cytoplasmic Na\(^+\) from 0 to 40 mM in the presence of 20 mM extracellular Na\(^+\) (no Ca\(^{2+}\) on either membrane side). Results for NCX1 are the same as described previously (Hilgemann et al., 1991). An outward current transient, \(\sim 150\) ms in duration with a peak of \(\sim 3\) pA, is observed on application of Na\(^+\), and a slower inward current transient with a peak of \(\sim 1\) pA is observed on removing Na\(^+\). The areas defined by the current transients correspond to \(\sim 300,000\) elementary charges, which in turn corresponds to \(\sim 300\) charges/\(\mu\)m\(^2\) with a 10-pF patch. Using the same protocols in patches expressing the squid exchanger, current transients were either absent or very small (five observations; Fig. 15 B). That these results reflect a real difference in exchanger function is supported by the observation that inward currents activated by applying cytoplasmic Ca\(^{2+}\) were of similar magnitude in the NCX1 and NCX-SQ1 patches (Fig. 15, C and D). We point out that the current activated by 5 \(\mu\)M Ca\(^{2+}\) corresponds to \(\sim 10\)% of the current activated in patches from the same oocyte batches when 150 mM Na\(^+\) was included in the pipette.

Fig. 15, E and F, shows the equivalent results for jumps of cytoplasmic Ca\(^{2+}\) from 0 to 5 \(\mu\)M in the presence of 50 \(\mu\)M extracellular Ca\(^{2+}\). As described previously (Hilgemann et al., 1991), charge movements are small or absent for this protocol with NCX1. For NCX-SQ1, however, inward current transients are obtained when 150 mM Na\(^+\) is included on the extracellular side. Without cytoplasmic Ca\(^{2+}\), all exchangers should orient to the cytoplasmic side with empty binding sites. Voltage pulses are applied first in the absence of cytoplasmic Ca\(^{2+}\), and then in the presence of 2 \(\mu\)M cytoplasmic Ca\(^{2+}\) to activate the ion occlusion reaction. Charge rather than current was recorded, and the holding potential was \(-40\) mV. Records presented in Fig. 16 A are a subtraction of records with cytoplasmic Ca\(^{2+}\) from records without Ca\(^{2+}\), whereby 16 records were acquired in alternating order with and without cytoplasmic Ca\(^{2+}\), and results were averaged. The charge movements show small fast components that appear as charge jumps on changing voltage, and they show slower components that saturate progressively as larger voltage pulses are applied to +160 and \(-200\) mV. Fig. 16 B shows the voltage dependence of the charge movements, fitted to a Boltzmann relation \(1/[1 + \exp^{(Em - 465)/26.5}]\), which gives an equivalent charge \(q\) of 0.46 underlying the charge movement. This is twice the value obtained for

Voltage Jump-induced Charge Movements of Ca\(^{2+}\) Transport of the NCX-SQ1 Exchanger

For the cardiac exchanger (NCX1), charge movements of Ca\(^{2+}\) transport have been isolated in voltage-jump experiments (Hilgemann, 1996). The charge movements were of small magnitude and showed weak voltage dependence with rates of \(\sim 5,000\) s\(^{-1}\) at 0 mV. Results for NCX-SQ1, after chymotrypsin treatment, are shown in Fig. 16. The rationale of the experiment is that most of the voltage dependence of Ca\(^{2+}\) transport comes about during occlusion of Ca\(^{2+}\) from the cytoplasmic side. Therefore, a high concentration of Ca\(^{2+}\), 4 mM, is included on the extracellular side. Without cytoplasmic Ca\(^{2+}\), all exchangers should orient to the cytoplasmic side with empty binding sites. Voltage pulses are applied first in the absence of cytoplasmic Ca\(^{2+}\), and then in the presence of 2 \(\mu\)M cytoplasmic Ca\(^{2+}\) to activate the ion occlusion reaction. Charge rather than current was recorded, and the holding potential was \(-40\) mV. Records presented in Fig. 16 A are a subtraction of records with cytoplasmic Ca\(^{2+}\) from records without Ca\(^{2+}\), whereby 16 records were acquired in alternating order with and without cytoplasmic Ca\(^{2+}\), and results were averaged. The charge movements show small fast components that appear as charge jumps on changing voltage, and they show slower components that saturate progressively as larger voltage pulses are applied to +160 and \(-200\) mV. Fig. 16 B shows the voltage dependence of the charge movements, fitted to a Boltzmann relation \(1/[1 + \exp^{(Em - 465)/26.5}]\), which gives an equivalent charge \(q\) of 0.46 underlying the charge movement. This is twice the value obtained for
NCX1 (Hilgemann, 1996). Fig. 16 C shows the voltage dependence of the rates of the charge movements, obtained by fitting the slow components of Fig. 16 A to single exponential functions. The rates have a “U-shaped” dependence on voltage, as expected for a simple reaction with voltage dependence of both the forward and reverse rates. The rates can be well-described by the sum of two exponentials, \( K_a \cdot e^{-\frac{v}{55}} + K_b \cdot e^{-\frac{v}{55}} \), where \( K_a \) and \( K_b \) are the forward and backward rates at 0 mV. The fit gives an equivalent charge of 0.59. The overall rate at 0 mV is \( \sim 1,600 \text{s}^{-1} \), which is about three-fold lower than rates obtained in equivalent experiments with NCX1 (Hilgemann, 1996).

**Discussion**

The Squid Na\(^{+}\)-Ca\(^{2+}\) Exchanger, NCX-SQ1

We have cloned and expressed the squid neuronal Na\(^{+}\)-Ca\(^{2+}\) exchanger, NCX-SQ1. The squid exchanger is a member of the family of NCX-type exchangers, including three mammalian exchangers, NCX1 (Nicoll et al., 1990), NCX2 (Li et al., 1994), and NCX3 (Nicoll et al., 1996), as well as the *Drosophila* Na\(^{+}\)-Ca\(^{2+}\) exchanger (Schwarz and Benzer, 1997; Ruknudin et al., 1997). NCX-SQ1 is 58% identical at the amino acid level to the canine cardiac Na\(^{+}\)-Ca\(^{2+}\) exchanger, NCX1, and has similar identities (51–64%) to the other NCXs. There are several features of note. Regions determined to be of functional importance in previous studies of NCX1 are well conserved. For example, we have identified specific acidic residues within the binding site for regulatory Ca\(^{2+}\) (Fig. 4, boxed area) that are important for Ca\(^{2+}\) binding (Levitsky et al., 1994; Matsuoka et al., 1995). These triple aspartate (DDD) motifs (Fig. 4, bold) are perfectly conserved in all NCX exchangers. Likewise, we have described the endogenous XIP region of NCX1 and have proposed that the XIP region is involved in Na\(^{+}\)-dependent inactivation (Li et al., 1991; Matsuoka et al., 1997). A homologous region (Fig. 4, shaded area) is also present in NCX-SQ1 and is conserved among the exchangers.

The predicted topology of NCX-SQ1 is similar to that of the other exchangers with 11 transmembrane segments and a large intracellular loop. Sequence conservation among the exchangers is highest in the proposed transmembrane segments consistent with a catalytic role of the hydrophobic domains in ion translocation. We have described that exchanger function is especially sensitive to mutations within portions of transmembrane segments 2, 3, 8, and 9 (Nicoll et al., 1996). These regions are known as the \( \alpha \) repeats (\( \alpha \)-1 and \( \alpha \)-2). Significantly, the \( \alpha \) repeats are highly conserved in species as divergent as squid and dog, consistent with a proposed role in ion transport. Proposed transmembrane segment 11 is the least well conserved transmembrane domain among the NCX exchangers. Perhaps the COOH terminus of the protein has a lesser role in exchange function. As noted previously (Tsuruya et al., 1994), the NH\(_2\) terminus of the exchanger, which represents a signal peptide region, is poorly conserved among NCX proteins.

The squid Na\(^{+}\)-Ca\(^{2+}\) exchanger has previously been reported to be stimulated by phosphorylation reactions and by phosphoarginine (DiPolo and Beaugé, 1994; DiPolo et al., 1997) and we analyzed the NCX-SQ1 sequence for potential phosphorylation sites. Potential sites for phosphorylation by protein kinases A and C, Ca\(^{2+}\)/calmodulin-dependent kinase, and tyrosine kinases are shown in Fig. 4. The PKC site (threonine 184) between transmembrane segments 3 and 4 and the tyrosine kinase site in the Ca\(^{2+}\) binding region (tyrosine 462) are unique among the exchangers. Phosphorylation at either site might be expected to have functional effects.

As shown in Fig. 4, there is a deletion of 47 amino acids in the large intracellular loop of NCX-SQ1 in comparison with NCX1. This region of NCX1 displays extensive alternative splicing (Nakasaki et al., 1993; Kofuji et al., 1994; Lee et al., 1994; Quednau et al., 1997). Six small exons are used in different combinations in a tissue-specific manner. The first two of these exons (exons A and B) are mutually exclusive. The NCX1 splice variant in Fig. 4 is NCX1.1 (Quednau et al., 1997) using exons A and C–F. Of the six small exons, NCX-SQ1 apparently uses only exon A. The homologous exons A of NCX-SQ1 and NCX1 are shaded in Fig. 4. We performed reverse transcriptase–PCR using optic lobe and stellate ganglion RNA to detect other splice variants of NCX-SQ1. Nine clones were sequenced, but no other splicing isoforms were detected.

Regulation of Na\(^{+}\)-Ca\(^{2+}\) exchange by intracellular Ca\(^{2+}\) was first described in the squid axon (Baker and McNaughton, 1976; DiPolo, 1979). Micromolar levels of free intracellular Ca\(^{2+}\) are required to activate function. That is, in addition to transporting Ca\(^{2+}\), the exchanger is separately regulated by Ca\(^{2+}\). This secondary effect of intracellular Ca\(^{2+}\) has been analyzed at the molecular level for the cloned cardiac exchanger NCX1 (Levitsky et al., 1994; Matsuoka et al., 1995). Using \(^{45}\)Ca\(^{2+}\) fluxes, we find that Ca\(^{2+}\) regulation is apparently intrinsic to the NCX-SQ1 exchanger protein. Chelation of intracellular Ca\(^{2+}\) with EGTA prevents Na\(^{+}\)-dependent Ca\(^{2+}\) uptake (Fig. 6). This is not surprising since, as noted above, the regulatory Ca\(^{2+}\) binding site of NCX1 is conserved in the sequence of NCX-SQ1.

**Regulatory Mechanisms Acting on NCX-SQ1 in Oocyte Giant Patches**

Given the sequence similarities to NCX1, it is not surprising that many of the regulatory properties of NCX-
SQ1 are similar to those of NCX1. This includes the properties of Na⁺-dependent inactivation, secondary activation by cytoplasmic Ca²⁺, and deregulation by chymotrypsin (Fig. 7). Furthermore, similarities include stimulation by ATP and PIP₂, and reversal of the stimulatory effects of ATP by a PIP₂ antibody (Figs. 8 and 9). However, there are also evident differences between NCX-SQ1 and NCX1. In the absence of phosphatase inhibitors, NCX-SQ1 exchange current is usually stimulated by ATP, which is not the case with NCX1. Either the squid exchanger has higher affinity for PIP₂ or it is modulated by an additional ATP-dependent reaction. Consistent with this possibility, the thioester of ATP, ATPγS, stimulates NCX-SQ1 in oocyte patches but not NCX1 current (Figs. 8 and 10). An attractive speculation is that we are observing phosphorylation of NCX-SQ1 at one of its consensus phosphorylation sites. We cannot, however, completely eliminate a possibility that effects of the ATPγS preparations employed reflect contaminating ATP and an increased sensitivity of the squid to PIP₂.

The reversal of ATP effects on Na⁺–Ca²⁺ exchange current by a PIP₂ antibody is reported here for the first time. The fact that the PIP₂ antibody can reverse stimulatory effects of ATPγS on the squid exchanger indicates that PIP₂ is involved. Perhaps phosphorylation increases the affinity of NCX-SQ1 for PIP₂. Precedents for such a mechanism come from recent work with inward rectifier potassium channels. With GIRK-type channels, activation by G-protein βγ subunits is accompanied by an increase of affinity for PIP₂ (Huang et al., 1998), and, with ROMK-type channels, activation by cAMP-dependent protein kinase is accompanied by an increase of the apparent affinity for PIP₂ (Dr. C.L. Huang, personal communication). Since ATP increases the apparent affinity for Ca²⁺ at exchanger regulatory sites (DiPolo and Beaugé, 1987; Collins et al., 1992), a primary effect of phosphorylation on Ca²⁺ affinity is also a possibility. In contrast to our results, it is reported that the PIP₂ antibody is without effect on Na⁺–Ca²⁺ exchange in squid giant axons, either with or without ATP (DiPolo and Beaugé, 1998). Either the antibody does not reach the membrane in axons or the exchanger is regulated in a fundamentally different way in the squid axon environment. The fact that phosphoarginine does not stimulate the cloned squid exchanger presumably reflects the absence of appropriate phosphoarginine-dependent kinases in oocyte giant membrane patches.

**Voltage Dependence of NCX-SQ1 in Oocyte Giant Patches**

The ion dependencies of the NCX-SQ1 exchange currents, determined here for the cytoplasmic side (Figs. 13 and 14), are consistent with those determined in dialyzed squid axons (DiPolo, 1989) and are only slightly different from results for the NCX1 exchanger. As described in Figs. 13 and 14, the voltage dependencies of outward and inward exchange currents of NCX-SQ1 are stronger than for NCX1, and the isolation of charge movements of Ca²⁺ transport for the squid exchanger (Figs. 15 and 16) verifies that electrogenic reactions are indeed different in the squid exchanger. Na⁺ transport is relatively less electrogenic, while Ca²⁺ transport is more electrogenic (Figs. 15 and 16). An important observation, which allows interpretation of the current–voltage relations of the squid exchanger, is that Na⁺–Na⁺ exchange by the squid exchanger is substantially greater than Na⁺–Ca²⁺ or Ca²⁺–Ca²⁺ exchange (DiPolo et al., 1989; DiPolo and Beaugé, 1990; Dr. L. Beaugé, personal communication). Thus, Ca²⁺ transport may in general be rate limiting and the voltage dependence of Ca²⁺ transport will determine the overall voltage dependence of transport current. Consistent with this interpretation, the rates of Ca²⁺-dependent charge movements determined for the squid exchanger at 0 mV and 33°C are substantially less (1,700 s⁻¹) than those determined for the NCX1 exchanger (5,000 s⁻¹; Hilgemann et al., 1991; Hilgemann, 1996).

Our interpretation is tempered to some extent because an alternating access model, or "consecutive" exchange mechanism, has not been rigorously verified for the squid exchanger. In a consecutive mechanism, the apparent affinity of one transported ion should increase as the concentration of the countertransported ion is decreased. For the squid exchanger, it has been reported that the Na⁺ dependence of Ca²⁺ efflux does not change as the cytoplasmic Ca²⁺ concentration is reduced (DiPolo, 1989). Thus, further work is required on transport properties of the squid exchanger; the application of techniques to photo-release Ca²⁺ in giant patches within microseconds (Kappl and Hartung, 1996), rather than 100 ms, will allow much better resolution of these issues.

The differences in the charge-moving reactions of NCX1 and NCX-SQ1 may provide an important key to elucidating the physical basis of exchanger electrogenicity. Ion occlusion reactions may result from the movement of charged residues anywhere in the exchanger protein, but only residues that enter or leave the membrane electrical field together with ions will generate charge movements. Evidently, more than two negative charges must move into the membrane electrical field when Ca²⁺ is occluded by the squid exchanger from the cytoplasmic side. Studies of chimeras of the two exchangers, and ultimately point mutation studies, should be able to define the involvement of specific groups and provide an understanding of the conformational changes underlying ion transport.

In summary, the squid Na⁺–Ca²⁺ exchanger, NCX-SQ1, has been cloned and expressed in *Xenopus* oocytes, and its function has been characterized electrophysiologically in
giant membrane patches. The differences between the sequence and functional properties of NCX-SQ1 and the mammalian NCX1 now provide a new basis to elucidate both regulatory and transport properties of Na\(^{+}\)-Ca\(^{2+}\) exchange. NCX-SQ1 is strongly activated by the anionic phospholipid, PIP\(_2\), and the presence of phosphorylation sites, not present in NCX1, may correlate with stimulation of the squid exchanger by thioester derivatives of ATP. The molecular basis of differences in the voltage dependence of cardiac and squid exchangers can now be pursued by the combined methods of molecular biology and electrophysiology.

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