Effects of SEA0400 on Mutant NCX1.1 Na\(^+\)-Ca\(^{2+}\) Exchangers with Altered Ionic Regulation

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Effects of SEA0400 on Mutant NCX1.1 Na⁺-Ca²⁺ Exchangers with Altered Ionic Regulation

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A Thesis/Practicum submitted to the Faculty of Graduate Studies of The University of Manitoba in partial fulfillment of the requirements of the degree of

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LIST OF ABBREVIATIONS

\( \lambda \), rate of current decay

ARF, acute renal failure

BSA, bovine serum albumin

C, Cysteine

\( \text{Ca(NO}_3\text{)}_2 \), calcium nitrate

\( \text{CaCl}_2 \), calcium chloride

\( \text{CaCO}_3 \), calcium carbonate

CALX, \textit{Drosophila melanogaster} \( \text{Na}^+ \text{-Ca}^{2+} \) exchanger

CCCP, carbonyl cyanide m-chrolophenylhydrazone

CHX, \( \text{Ca}^{2+} \text{-H}^+ \) exchanger

CICR, \( \text{Ca}^{2+} \)-induced-\( \text{Ca}^{2+} \)-release

cRNA, complementary ribonucleic acid

cDNA, complementary deoxyribonucleic acid

CsOH, cesium hydroxide

DAD(s), delayed afterdepolarization(s)

DCB, (3',4'-dichlorobenzamil)

DPC, days post coitus

DMSO, dimethyl sulfoxide

E, Glutamic acid

EAD(s), early afterdepolarization(s)

EGTA, ethylene glycol-bis(\( \beta \)-aminoethyl ether)-N,N,N',N'-tetra-acetic acid
F, Phenylalanine
FMRF-amide, molluscan cardioexcitatory peptide
FRCRCF-amide, analogue of molluscan cardioexcitatory peptide
FSS, fraction of steady state current remaining
G, Glycine
HEPES, (N-[2-hydroxyethyl]piperazine-N’-[2-ethanesulfonic acid])
I1, Na+–dependent inactivation
I2, Ca2+–dependent regulation
IK1, inward rectifying potassium current
Itr, transient inward current
I-V, current-voltage relationship
K, Lysine
K2HPO4, potassium phosphate
KB-R7943, (2-[2-[4-(4-nitrobenzyloxy)-phenyl]ethyl]isothiourea methesulfonate)
KCl, potassium chloride
KOH, potassium hydroxide
L, Leucine
LiOH, lithium hydroxide
MES, (2-[N-morpholino]ethanesulfonic acid
Mg(OH)2, magnesium hydroxide
MgSO4, magnesium sulfate
MOPS, (3-[N-morpholino]propanesulfonic acid)
myristyl-FRCRCF-amide, analogue of molluscan cardioexcitatory peptide
N, Asparagine
NaCl, sodium chloride
NaHCO₃, sodium bicarbonate
NCKX, Na⁺-Ca²⁺/K⁺ exchanger
NCX, Na⁺-Ca²⁺ exchanger
NMG-MES, N-methyl-D-glucamine-(2-[N-morpholino]ethanesulfonic acid
PI, phosphatidylinositol
PIP₂, phosphatidylinositol 4,5-bisphosphate
PKA, protein kinase A
PKC, protein kinase C
PMA, phorbol 12-myristate 13-acetate
Q, Glutamine
R, Arginine
RT, room temperature
SEA0400, (2-4L-1Q,5-difluorophenyl)methoxylphenoxyl-5-ethoxyaniline
SL, Sarcolemma
SR, Sarcoplasmic Reticulum
T, Threonine
TEA-OH, tetraethylammonium hydroxide
TMS, transmembrane segment(s)
V, Valine
XIP, exchange inhibitory peptide
Y, Tyrosine
I. ABSTRACT

The plasmalemmal Na\(^+\)-Ca\(^{2+}\) exchanger protein plays an integral role in cardiac Ca\(^{2+}\) homeostasis. This unique protein allows for transsarcolemmal transport of Ca\(^{2+}\) across the cell membrane, in exchange for Na\(^+\). Removal of virtually all Ca\(^{2+}\) entering through L-type Ca\(^{2+}\) channels occurs on a beat-to-beat basis, allowing for the maintenance of a proper resting Ca\(^{2+}\) level. Calcium efflux occurs by removing Ca\(^{2+}\) in exchange for Na\(^+\) and is known as forward mode exchange. As the direction of exchange transport is based on both electrical and chemical gradients, there exists the potential for drawing Ca\(^{2+}\) into the cell. Calcium entry via the exchanger is labelled reverse mode exchange and is implicated in both cardiac injury and disease. Both exchanger modes are the subjects of intense investigation, as their precise physiological roles are controversial.

Although the physiological role of Na\(^+\)-Ca\(^{2+}\) exchange in cardiac muscle is unclear, the current body of knowledge suggests it contributes to cellular relaxation (i.e., forward mode) and cardiac pathophysiology (i.e., reverse mode). Under various pathological settings, reverse mode exchange (i.e., an elevated Na\(^+\), promoting Ca\(^{2+}\) entry) may be favoured. Thus, the exchanger may participate in cellular “Ca\(^{2+}\) overload”, creating a toxic intracellular environment. Toxicity via Ca\(^{2+}\) can eventually kill the cell through a variety of mechanisms (i.e., protease activation). Having a selective, reverse mode exchange inhibitor (leaving forward mode (i.e., Ca\(^{2+}\) removal) exchange unaffected) could prove most useful in the therapeutic setting. Hindering a detailed understanding of Na\(^+\)-Ca\(^{2+}\) exchange is the general lack of
selective and potent pharmacological agents. A few functional inhibitors, which were by-in-large weak, non-selective and potentially toxic (millimolar requirements for desired action) were described in the early 1990's. In 1996, a potent and reverse mode selective Na\(^+\)-Ca\(^{2+}\) exchange inhibitor, KB-R7943 (formerly No.7943) was described. However, as the results of KB-R7943 were presented, discrepancies over mode selectivity, site of action, and potency became apparent. In 2000, a new Na\(^+\)-Ca\(^{2+}\) exchange inhibitor was developed. SEA0400 was released as a the most potent and mode selective (preferential for reverse mode) inhibitor of Na\(^+\)-Ca\(^{2+}\) exchange ever produced. SEA0400 significantly inhibited reverse mode exchange and was without effect on forward mode exchange. This effect led to the search for its potential mechanism(s) of action. Previous work from our lab suggested that SEA0400 stabilizes or promotes the entry of Na\(^+\)-Ca\(^{2+}\) exchangers into a Na\(^+\)-dependent inactive state (I\(_1\)). Upon deregulation of the Na\(^+\)-Ca\(^{2+}\) exchanger with α-chymotrypsin, SEA0400's action was substantially reduced, suggesting an interaction/requirement of intact regulatory mechanisms for proper function. The two main regulatory mechanisms proposed to control the exchanger, are Na\(^+\)-dependent inactivation (I\(_1\)) and Ca\(^{2+}\)-dependent regulation (I\(_2\)). When deregulated (i.e., α-chymotrypsin), the exchanger no longer exhibited I\(_1\) and I\(_2\) regulation. To differentiate between the regulatory mechanisms that guide SEA0400's inhibition, previously characterized Na\(^+\)-Ca\(^{2+}\) exchanger mutants with defined alterations in their ionic regulatory properties were implemented. Mutants were categorized into three groups; those with accelerated I\(_1\) (F223E, K225Q), lacking I\(_1\) (K229Q, Δ229-232), and lacking I\(_2\) (Δ680-685). We then applied the giant, excised patch clamp technique
on *Xenopus laevis* oocytes expressing the cloned canine cardiac exchanger NCX1.1, and measured resulting inward and outward currents. To these currents, we applied SEA0400 and measured its effects. Using F223E, a mutant displaying strongly accelerated \( I_1 \) (Na\(^{+}\)-dependent inactivation), inhibition with SEA0400 was substantially increased. However, the second Group 1 mutant, K225Q, showed \( I_1 \) similar to that of wild-type. While, inhibition of K225Q with SEA0400 was comparable to that of wild-type. The first Group 2 mutant, K229Q, initially lacked \( I_1 \), yet upon SEA0400 application an \( I_1 \)-like inactive state appeared. Conversely, with the larger A229-232 mutant, which lacked \( I_1 \), SEA0400 was virtually without effect. Lastly, \( \Delta 680-685 \) (the sole Group 3 mutant), lacked \( I_2 \) (Ca\(^{2+}\)-dependent regulation) and inhibition with SEA0400 paralleled that of wild-type (WT). These results suggest that \( I_1 \) plays an integral, if not essential role in SEA0400’s inhibitory nature. As such, when \( I_1 \) is accelerated, SEA0400’s effects are increased. Conversely, when \( I_1 \) is absent, SEA0400 appears ineffective. These results may provide an explanation for SEA0400’s mode selectivity.
II. LITERATURE REVIEW

1. Role of Na\(^+\)-Ca\(^{2+}\) Exchanger

1.1. Excitation-Contraction Coupling

With every contraction and relaxation of the heart, intracellular Ca\(^{2+}\) concentration changes dramatically yet, for the most part, proper cardiac function is maintained. Through a series of pumps, channels, exchangers, and various signalling proteins, this order is upheld. Many factors that are so vital to internal cellular processes can ironically lead to organelle failure and eventually, cell death. For instance, in cardiac myocytes, Ca\(^{2+}\) is essential for cardiac contraction, yet at high levels it can become toxic to the cell’s internal environment, thereby interrupting cellular processes and/or killing the cell. Roughly, per beat, the internal Ca\(^{2+}\) concentration increases ~ 100 fold (0.1-10 μM). Calcium must be removed in order to preserve proper cellular function and allow for complete myocyte relaxation. The cell has several well-documented methods of maintaining Ca\(^{2+}\) homeostasis.

A primary mechanism for removing Ca\(^{2+}\) from cardiac tissue is the Na\(^+\)-Ca\(^{2+}\) exchanger. In 1968, this ion transporter was initially identified in cardiac muscle (275). The exchanger’s role is prominent in Calcium-Induced-Calcium-Release (CICR), a term used to describe the cellular process allowing for cardiac contraction (91; 92). During contraction initiation, Ca\(^{2+}\) travels down its favoured concentration gradient (into the cytoplasm), signals a further Ca\(^{2+}\) release, and thereby increases
Ca$^{2+}$ levels high enough to activate myofilament interaction (29; 43; 55; 221). From here, various cellular processes allow for: i) resequestering of Ca$^{2+}$ and ii) transsarcolemmal Ca$^{2+}$ removal. In doing so, this allows for proper cellular relaxation and diastolic filling, in preparation for the next consecutive beat. Although this process may seem initially simple, it is quite intricate.
Figure 1. Excitation-Contraction Coupling of the Na\(^+\)-Ca\(^{2+}\) exchanger. This figure represents the interplay among Ca\(^{2+}\) transporters involved in regulating cardiac contraction and Calcium-Induced-Calcium-Release (CICR).
For every cardiac beat to occur, a wave of depolarization is deployed, stimulating a cascade of events. First, the depolarizing wave travels along the sarcolemma reaching voltage-gated L-type Ca\(^{2+}\) channels and in turn activates them. These channels open and Ca\(^{2+}\) flows into the cell. This quantity of Ca\(^{2+}\) is appropriately termed "trigger Ca\(^{2+}\)", for it binds to the nearby sarcoplasmic reticular (SR) ryanodine receptors, and "triggers" the SR to release its internal Ca\(^{2+}\) stores (29; 43; 55; 221). This large intracellular Ca\(^{2+}\) increase initiates contraction through cardiac myofilament interaction (29; 43; 55; 92). The location of various cellular proteins and structures (L-type Ca\(^{2+}\) channels, SR ryanodine receptors, and Na\(^{+}\)-Ca\(^{2+}\) exchangers) are biologically organized to allow maximum efficiency. For example, a close association between L-type Ca\(^{2+}\) channels (on T-tubule invaginations) and ryanodine receptors has been shown (50; 51; 279; 306). However, diseases such as hypertrophy and heart failure may occur when these receptors and channels are no longer in close proximity, hinting at the importance of cellular location (103). As cardiac contraction requires specific machinery, so does relaxation.

After contraction, myocyte internal Ca\(^{2+}\) is high (~10 \(\mu\)M) and must be restored to resting diastolic levels (100 nM) (18). This decrease in Ca\(^{2+}\) is crucial for maintaining healthy cells, allowing them to relax, and promote diastolic filling. There are three primary components involved in cytosolic Ca\(^{2+}\) removal: the SR Ca\(^{2+}\)-ATPase, Na\(^{+}\)-Ca\(^{2+}\) exchanger and the Sarcolemmal (SL) Ca\(^{2+}\)-ATPase. These mechanisms of Ca\(^{2+}\) extrusion have all been thoroughly reviewed (11; 19; 21; 36; 130; 237; 250). To start, the SR Ca\(^{2+}\)-ATPase is responsible for resequestering Ca\(^{2+}\) (activator Ca\(^{2+}\)) initially released from the SR upon stimulation. It is the primary
mechanism of intracellular Ca\textsuperscript{2+} removal, with reports on rabbit myocytes (133) and rabbit ventricular muscle (22), suggesting that it contributes to \(~75\) % of the relaxation process. The ATPase utilizes energy derived from ATP hydrolysis, in order to pump Ca\textsuperscript{2+} back into the SR, against a large concentration gradient (135). In addition, results of transgenic mice over expressing SR Ca\textsuperscript{2+}-ATPase show an alteration in myocardial contractility with an increase in relaxation (6). Conversely, from experiments where SR Ca\textsuperscript{2+}-ATPase was functionally removed (245), a defect in relaxation was shown. Both over expression and functional SR Ca\textsuperscript{2+}-ATPase removal show this protein’s significance in contributing to proper cardiac relaxation.

The second ATPase involved in Ca\textsuperscript{2+} removal is the sarcolemmal (SL) Ca\textsuperscript{2+}-ATPase. Although a transmembrane Ca\textsuperscript{2+} removal mechanism, its role in cardiac relaxation remains quite ambiguous. It is thought to contribute minimally to Ca\textsuperscript{2+} removal (18; 330). For example, experiments having SL Ca\textsuperscript{2+}-ATPase as the single Ca\textsuperscript{2+} extrusion mechanism, demonstrated a massive decrease in relaxation rate (~1200%) compared to cells with intact mechanisms (22). Yet, there are increasing reports suggesting otherwise (13; 14; 54). Studies implementing carboxeosin, a SL Ca\textsuperscript{2+}-ATPase inhibitor, show the Ca\textsuperscript{2+}-ATPase to contribute 24%-45% of total membrane Ca\textsuperscript{2+} efflux (54; 237; 320). Furthermore, in 1999, a report demonstrated in rat ventricular myocytes lacking SR Ca\textsuperscript{2+}-ATPase, that Ca\textsuperscript{2+} efflux via the SL Ca\textsuperscript{2+}-ATPase was higher than previously reported (54). However, even a two-fold overexpression of SL Ca\textsuperscript{2+}-ATPase in transgenic rats, showed no substantial effect on cardiac function (106). Lastly, findings with the plasmalemmal Ca\textsuperscript{2+} pump absent, reported no change in cytoplasmic Ca\textsuperscript{2+} levels, further suggesting this pumps effects
are negligible (46). Functional importance of the Ca\textsuperscript{2+} pump has been suggested as a species dependent characteristic, wherein ferret ventricular muscle, the SL Ca\textsuperscript{2+}-ATPase was shown to have comparative level of importance to the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger (14; 19). Although a specific physiological role of SL Ca\textsuperscript{2+}-ATPase is not yet established, it is generally agreed upon that its contribution during relaxation is minimal and may only serve to perform minute alterations of resting Ca\textsuperscript{2+} levels. Even if the SL Ca\textsuperscript{2+}-ATPase does contribute to Ca\textsuperscript{2+} efflux to a greater degree than currently reported, its transport rate is much too slow to be effective (i.e., 15-30 times slower than Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange) (21). This leaves Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange as the major pathway for plasmalemmal Ca\textsuperscript{2+} extrusion, contributing to cardiac relaxation.

1.2. Forward Mode Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchange

Effective Ca\textsuperscript{2+} homeostasis is paramount for cardiac function. In short, any Ca\textsuperscript{2+} that enters a cell must be removed. If this delicate balance is not met over time, numerous cardiac dysfunctions will result. As stated previously, during contraction, Ca\textsuperscript{2+} enters through L-type Ca\textsuperscript{2+} channels and further stimulates Ca\textsuperscript{2+} release from the SR (CICR). SR Ca\textsuperscript{2+} is then resequestered via the SR Ca\textsuperscript{2+}-ATPase, leaving the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger to deal with the remaining Ca\textsuperscript{2+} (initially brought in through the L-type Ca\textsuperscript{2+} channels). Thus, Ca\textsuperscript{2+} removed by the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger must precisely balance L-type Ca\textsuperscript{2+} channel Ca\textsuperscript{2+} brought in. Removing Ca\textsuperscript{2+} in exchange for Na\textsuperscript{+} is known as forward mode exchange (i.e., Ca\textsuperscript{2+} efflux) (11; 19; 21; 36; 44; 130; 237; 250). It is essential that Ca\textsuperscript{2+} influx and efflux be matched. While the SR and the
mitochondria (sites of Ca\textsuperscript{2+} storage) can uptake Ca\textsuperscript{2+}, they only have limited capacities for storage, leaving the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger as the last mechanism to withdraw any residual Ca\textsuperscript{2+}. Several studies have shown the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger to play a powerful role in maintaining this balance between Ca\textsuperscript{2+} influx and efflux. An example of such was the 1975 study by Bridge et al., that measured Ca\textsuperscript{2+} efflux via the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger matching that of the inward L-type Ca\textsuperscript{2+} current (assuming a 3:1 stoichiometry) (41). Further studies by Niggli and Lederer, using flash photolysis and “caged Ca\textsuperscript{2+}”, found a significant parallel between current amplitudes from Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange and L-type Ca\textsuperscript{2+} channels (233). Lastly, caffeine-induced and rapid cooling contractions were used to quantify SR Ca\textsuperscript{2+} and showed a significant role for Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange in Ca\textsuperscript{2+} removal (23; 40; 133). Ca\textsuperscript{2+} entry can be quite varied physiologically (i.e., by changes in frequency, Ca\textsuperscript{2+} levels, and adrenergic stimulation) and pharmacologically (i.e., dyhydropyridines (negative), and adrenergic compounds (positive)) (20; 26; 246). This is a crucial function of the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger, for cardiac myocytes have a low tolerance to Ca\textsuperscript{2+} imbalance. However, as with other cellular proteins, different species and stages of development may vary exchanger function (36).

Compared to adult rat and rabbit hearts, newborn rat and fetal rabbit hearts appeared to have a six to eight-fold greater expression in Na\textsuperscript{+}-Ca\textsuperscript{2+} exchangers (38). It was proposed that Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange constitutes the majority of Ca\textsuperscript{2+} removal in these underdeveloped hearts. These hearts lack plasmalemmal invaginations, known as T-tubules, and a fully formed SR. Although results are derived from an underdeveloped organism, they strengthen the significance of Na\textsuperscript{+}-
Ca\(^{2+}\) exchange, whether in fetal development or in mature adults. Since the exchanger’s transport mode depends on electrical and chemical gradients, it has the potential to reverse directions. Thus, in theory, the Na\(^+-\)Ca\(^{2+}\) exchanger could contribute to systole as it does in diastole (11; 44; 237). This topic is currently contentious and has serious implications regarding the physiological and pathophysiological role(s) of the Na\(^+-\)Ca\(^{2+}\) exchanger.

1.3. Reverse Mode Na\(^+-\)Ca\(^{2+}\) Exchange

Understanding a physiological role for Ca\(^{2+}\) entry (i.e., reverse mode) via the Na\(^+-\)Ca\(^{2+}\) exchanger has been an ongoing process. It is suggested that Ca\(^{2+}\) entry through this protein can parallel the effects of “trigger Ca\(^{2+}\)” from the L-type Ca\(^{2+}\) channels, and initiate Calcium-Induced-Calcium-Release from the SR (24; 128) but there are also those that see no such result (39; 90). During the action potential upstroke, submembrane Na\(^+\) levels could initiate reverse mode Na\(^+-\)Ca\(^{2+}\) exchange but Ca\(^{2+}\) from the L-type Ca\(^{2+}\) channels increase cytoplasmic Ca\(^{2+}\) and reduces the drive for Ca\(^{2+}\) entry (11; 36). Not only is the drive for reverse mode exchange (Ca\(^{2+}\) influx) reduced, but the amount of Ca\(^{2+}\) entering through the Na\(^+-\)Ca\(^{2+}\) exchanger is marginal compared to that of L-type Ca\(^{2+}\) channels (292; 299). Nonetheless, reverse mode Na\(^+-\)Ca\(^{2+}\) exchange has been used to explain the positive inotropic effects of cardiac glycosides (34; 35; 187).

The positive inotropic effects of cardiac glycosides (digoxin, digitoxin, and ouabain) are well known. They block the Na\(^+-\)K\(^+\)-ATPase (enzymatic portion of the Na\(^+\) pump), allowing for an elevation of intracellular Na\(^+\) levels (~ 1-2 mM) (268).
This intracellular Na\(^+\) increase promotes reverse mode Na\(^+\)-Ca\(^{2+}\) exchange, bringing Ca\(^{2+}\) into the cell, thereby stimulating SR Ca\(^{2+}\) release channels. The elevated levels of cytoplasmic Ca\(^{2+}\), leads to greater SR Ca\(^{2+}\) loading, Ca\(^{2+}\) release and contraction. Yet, unlike the rapid increase of intracellular Ca\(^{2+}\) due to L-type Ca\(^{2+}\) channels, influx through reverse mode Na\(^+\)-Ca\(^{2+}\) exchange is slow (200). On a beat-to-beat basis, reverse mode Ca\(^{2+}\) influx, activating Ca\(^{2+}\)-induced-Ca\(^{2+}\)-release (CICR), is thought by some to be unlikely (39; 45).

It has also been proposed that the influx of Na\(^+\) upon depolarization may be enough to accumulate near the membrane and stimulate reverse mode exchange. This would bring Ca\(^{2+}\) into the cell and stimulate Ca\(^{2+}\)-induced-Ca\(^{2+}\)-release (CICR) (181). There are several reasons why this idea may be problematic. First, an intracellular Na\(^+\) concentration required to initiate reverse mode exchange does not occur under physiological conditions (but does in the experimental setting) (130). Secondly, this would imply that the SR was over-sensitive and spontaneously releases Ca\(^{2+}\) every instance Ca\(^{2+}\) concentration fluxed (39; 90; 221; 292; 293; 298; 299). However, certain adrenergic interactions with the voltage-dependent Na\(^+\) channels may result in a phenomenon termed “Slip Mode conductance”. Here, Ca\(^{2+}\) and Na\(^+\) enter via voltage-gated Na\(^+\) channels upon depolarization. Ca\(^{2+}\) entry here is sufficient to initiate Ca\(^{2+}\)-induced-Ca\(^{2+}\)-release (CICR).

If we consider the location of membrane proteins once more, we continue to find discrepancies. Ideally, for CICR, Na\(^+\)-Ca\(^{2+}\) exchangers should be in close proximity to the SR, such that they can act on ryanodine receptors efficiently. Certain reports localize the exchanger along T-tubules (96), while others describe exchanger
positioning as being uniformly placed along the sarcolemma (48; 160). If a uniform
distribution of the exchanger exists, then the probability of reverse mode Na⁺-Ca²⁺
exchange having an effect on the SR would be low. Reverse mode Na⁺-Ca²⁺
exchange has been proposed to modify the efficacy of L-type Ca²⁺ channels with
respect to CICR (12; 17). This mechanism, if possible, implies the Na⁺-Ca²⁺
exchanger could increase SR Ca²⁺ release and thereby, increase SR Ca²⁺ content
(load) (198; 335). Most recently, a report by Thomas et al., using a novel Na⁺-Ca²⁺
exchange antibody, combined with confocal microscopy (CLSM), found that
immunoreactivity is strongly expressed throughout the SL surface and intercalated
disk regions with sporadic labelling of the vertical transverse T-tubules but not the
longitudinal T-tubules (313). Using normal and detubulated rat cardiomyocytes they
suggested that the exchanger has multiple functions depending upon membrane
location such as; CICR modulation, SR Ca²⁺ load, and resting Ca²⁺ levels (313).

2. Molecular Biology of the Na⁺-Ca²⁺ Exchanger

2.1. The Exchanger Superfamily

From the initial cloning of the canine cardiac Na⁺-Ca²⁺ exchanger NCX1.1
cDNA) in 1990 by the Philipson group, a large group of related proteins have emerged (227). This is now collectively known as the exchanger superfamily. It is a
group of Na⁺-Ca²⁺ exchangers and related proteins (227). Most of these proteins have
been identified through BLAST and PSI-BLAST searches of the GenBank database
(2). Many of these isolated proteins' function and significance are not known.
However, based on the sequence homology of other well-documented proteins, many of these homologous proteins are predicted to be membrane transporters. Based on hydrophobicity plots, every exchanger superfamily protein is found to have a conserved Calx-α motif and contain membrane-spanning regions (36; 130; 250). As reviewed by Philipson and Nicoll in 2000 (250), the exchanger superfamily has been divided into four categories: (1) NCKX family; (2) Bacterial family; (3) CHX family; (4) NCX family (Table 1).

2.1.1. The NCKX Family

The NCKX, has a similar homology to the Na⁺-Ca²⁺-K⁺ exchanger cloned from bovine rod photoreceptors. It is also however, found in neural tissue (316). The family consists of three mammalian NCKX exchangers, NCKX1 (273), NCKX2 (316), NCKX3 (174), six from Caenorhabditis elegans, one from Schizosaccharomyces pombe, and one from Arabidopsis thaliana. The cloned mammalian exchanger NCKX1 has a stoichiometry of 4 Na⁺: 1 Ca²⁺: 1 K⁺, with a requirement of K⁺ co-transport for all members (285; 286). This is the primary feature that separates them functionally from the NCX exchangers. Each exchanger has sequence similarity, a pair of Calx-α motifs, and a large intracellular loop.

2.1.2. The Bacterial Family

Although they constitute nine members of the superfamily, very little is actually known about this unique group. This exchanger family lacks the large intracellular loop and therefore, has a dramatically reduced exchanger size (~ 400
residues compared to those of NCX and NCKX; ~ 900 residues). Their primary point of homology is through the Calx-α motifs of each family member (263).

2.1.3. The CHX Family

This family consists of only a few members presently. CHX antiporters are believed to transport Ca$^{2+}$ in exchange for H$^+$. *Escherichia coli* (136), *Saccharomyces cerevisiae* (63), and *Arabidopsis thaliana* (126) all produce CHX exchangers.

2.1.4. The NCX Family

The most extensively studied and documented exchangers are found in this family. This is primarily due to cloning of the canine cardiac isoform NCX1.1 by Nicoll *et al.* (227). There are three mammalian exchangers: NCX1 (227), NCX2 (192), and NCX3 (230). The NCX1 gene is widely expressed in a variety of cells (*i.e.*, cardiac muscle, skeletal muscle, smooth muscle, neurons, astrocytes, kidney, lung and spleen (265). NCX2 and NCX3 are only found in brain and skeletal muscle (192; 265). These genes have similar hydropathy analysis results, suggesting a similar structure. They also appear to have functional similarity (36; 194). Genes sharing homology with NCX1 have been characterized by comparing to various mammalian species such as: humans (168; 170), rabbit (169), bovine tissue (1), guinea pig (317), mouse (163), and rat (100). In addition, there are exchangers isolated from non-mammalian species such as *Loligo opalescens*, squid exchanger NCX-SQ1 (114) and from the fruit fly, *Drosophila melanogaster*, CALX (131; 277;
These exchangers all have tight sequence similarity and analogous function. Other non-mammalian species include two proteins isolated from *Caenorhabditis elegans* (nematode) that share homology to various NCX proteins (331) and one from *Xenopus* species (146).

2.2. *Gene Structure*

NCX1, NCX2, and NCX3 are located on human chromosomes 2p21-23 (295), 19q13.2 (192), and 14q21-31 (230), respectively. The intron/exon structure of NCX1 has been characterized with respect to its organization on the gene (169; 173). The NCX1 gene runs more than 200 kb (169; 172; 173) and contains 12 exons, with the majority of the gene being coded in one exon of 1.8 kb. Also, each gene has a 5'-end alternatively spliced region prior to the open reading frame (184), which most likely contributes to regulating the expression of different isoforms. These regions are labelled as “tissue-specific” promoters. Recently, in 1999, the Philipson and Menick groups revealed that cardiac specific expression is regulated by a promoter region of roughly 200 bp (containing a GATA transcriptional element), a CarG element and by an E-box (49; 224). The gene is also regulated by β-adrenergic stimulation (102).
Table 1. The Exchanger Superfamily. The Exchanger Superfamily consists of four families: (1) the Na\(^+\)-Ca\(^{2+}\) exchanger (NCX) family, (2) the Na\(^+\)-Ca\(^{2+}\)/K\(^+\) exchanger (NCKX) family, (3) the Bacterial family, and (4) the Ca\(^{2+}\)/H\(^+\) exchanger (CHX) family. Listed below are known genes and the species from which the gene originated (Adapted from Philipson and Nicoll (10)).

<table>
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<tr>
<th>FAMILY</th>
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<th>GENE</th>
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2.3. **Topology of the Na\(^+\)-Ca\(^{2+}\) Exchanger NCX1.1**

Since the cloning of NCX1.1 from canine cardiac cDNA, there have been many advances in the Na\(^+\)-Ca\(^{2+}\) exchange field (227). Initially, studies from mammalian cardiac sarcolemmal vesicles, measuring Na\(^+\)-Ca\(^{2+}\) exchange transport revealed electrogenic movement of Na\(^+\) and Ca\(^{2+}\) with a 3:1 stoichiometry (27; 47; 257; 271; 272). These results were followed by protein purification work (249) that allowed for the initiation of NCX1.1 cloning (227). Nicoll *et al.* (227) reported polypeptides with molecular masses of 70, 120, and 160 kDa (249). Upon establishing polyclonal antibodies for these peptides, they were used to screen a cardiac expression library. Continuing, Nicoll *et al.* then cloned a sequence for a peptide that reacted with the polyclonal antibodies, and this was used to further identify the entire clone. From here, cRNA was produced and injected into the *Xenopus* oocyte expression system. Proper Na\(^+\)-Ca\(^{2+}\) exchanger expression was measured via giant, excised patch clamping and compared to a control of water-injected oocytes. Na\(^+\)-dependent Ca\(^{2+}\) influx could be measured in cRNA injected oocytes but not in the control oocytes (water-injected) (227). This demonstrated that the exchanger clone was complete, reliable, and displayed proper exchanger functions.

Initial molecular weight estimates for NCX1.1 ranged between 33 kD – 82 kD (105; 199). However, once the exchanger was cloned in 1990 (227), it revealed a full length NCX1.1 cDNA, with a reading frame of 2910 nucleotides. These nucleotides ultimately create a 970 amino acid protein weighing approximately 108 kD (227). Initially, the Philipson group modeled the exchanger as having twelve membrane
spanning segments with a large hydrophobic loop between segment 6 and 7 (labelled loop f) (227).

Essentially this cytoplasmic loop broke the exchanger into two segments. One segment contained TMS 1-6, and the other contained the remaining TMS 7-12. These results were based on hydropathy analysis (133). They also noted 6 potential N-linked glycosylation sites, and a potential phosphorylation site (227). However, in the native canine exchanger, only a single site at position 9 of the N-terminus (Asn-9) is glycosylated (81; 132; 229). Glycosylation of the exchanger appears to have no effect on the exchanger’s functionality (132). Identification of potential glycosylation sites helps determine the exchanger’s orientation, because amino acid glycosylation implies an extracellular location for that particular residue (36; 130). Furthermore, a would-be cleavable NH₂-terminal signal (171) and an amino acid segment resembling a calmodulin-binding domain (later known as the XIP region (193)) were reported (227). This potential cleavage site (between amino acid 32 and 33) was of interest for several reasons. Firstly, it appeared to be the start of the protein sequence and secondly, most membrane exchangers lack these sites (81). However, removing the cleavage site or deleting the signal sequence did not prevent proper membrane insertion (99; 278). A similar site was shown in both brain (192) and rod photoreceptors (273). Upon discovery of the signal sequence and through glycosylation studies, the exchanger was labelled to have 11 TMS: TMS 1-5 on the N-terminus and TMS 6-11 on the C-terminus. Later, more refined protocols, such as cysteine substitution, mutagenesis, and epitope tagging (61; 139; 228) narrowed down the TMS number to 9 (139; 228). After refining the number of TMS’s, further changes were made in the exchanger’s topology. For instance,
the large hydrophobic intracellular loop (loop f) was now situated between TMS 5 and 6 (228). In addition, short amino acid sequences known as Calx-α motifs (α-1 and α-2 repeats) were found to exist between TMS 2-3 (α-1) and TMS 7-8 (α-2) (139; 228). Moreover, they appeared on opposite sides of the membrane (228). TMS 6 was moved into the cytoplasmic loop f, while TMS 9 (part of α-2) was found to form a “P-loop”, or re-entrant loop that terminated in the cytoplasm.

The Na⁺-Ca²⁺ exchanger works as a monomer and can function properly when in a heterologous expression system (i.e., Xenopus) (115; 117; 122). Conversely, it has been suggested that the exchanger can function in a shortened form (lacking six C-terminal TMS) (101; 191). This gave rise to the idea that the exchanger becomes functional through dimerization. Another truncated exchanger, lacking 30 amino acids from the C-terminus, was isolated from the BALB/c mouse heart (294). The deletion is thought to occur through an early termination site. However, Ottolia et al. demonstrated compelling data on evenly divided exchangers (“split exchanger”) expressed individually or together. They found that co-expression of both domains is required for membrane targeting and functional exchange activity (243).

Although attempts are being made to crystallize the Na⁺-Ca²⁺ exchanger, an exact 3-D model is not yet available. Using cysteine susceptibility analysis and epitope tagging, Iwamoto et al. attempted to further characterize exchanger topology (139). This technique revealed that TMS 3 and 7 are in close proximity intracellularly, while TMS 2 and 8 are associated on the extracellular surface. These results suggest a role for TMS’s and Calx-α motifs in ion translocation.
2.3.1. The Calx-α Motifs (α-1 and α-2)

In 1997, the fruit fly (Drosophila melanogaster) Na\(^+\)-Ca\(^{2+}\) exchanger was cloned and termed CALX (277; 290). This gene shares a >50% structural homology with the NCX family (NCX1, NCX2 and NCX3) and has a similar transport function (82; 131; 240). The CALX protein contains two motifs comprised of ~40 amino acids each. They are termed Calx-α motifs and are conserved in every member of the NCX family (290) and in the mammalian and Drosophila NCKX exchangers (110; 316). These motifs, also known as α-1 and α-2 repeats, are suggested to reside in close proximity but with opposite orientations.

When site-specific mutations were made to residues in the Calx-α repeats, the result was a significant alteration in exchanger transport properties (226). The degree of change ranged from a subtle variation to complete transport alleviation (226). For example, a study that substituted carboxyl/hydroxyl containing amino acids in TMS 2, 3, and 7 (part of the Calx-α motifs), found a decrease or elimination of \(^{45}\)Ca\(^{2+}\) uptake in a Xenopus expression system (226). Mutations of conserved glycine residues; Gly138 (between TMS 2-3) and Gly837 (between TMS 7-8), shifted the exchanger I-V relationship. Further experiments were conducted in the hopes of tying Calx-α motifs to ion translocation. The Philipson group showed that in TMS 2, mutation of Asp101 to cysteine (N101C) removed the exchanger’s cytoplasmic Na\(^+\) regulation, while mutating threonine (T103C or T103V), increased the exchanger’s affinity for cytoplasmic Na\(^+\) (also allowing Li\(^+\) transport) (78). Iwamoto et al. further strengthened the role of Calx-α motifs through the use of chimeric exchangers between NCX1 and NCX3 (143). Using a
Ni\textsuperscript{2+} sensitive property of wild-type Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange, they were able to mimic and compare to native exchanger function using chimeras containing amino acid mutations (substitutions) in the \( \alpha \)-1 and \( \alpha \)-2 repeats (143). This suggested the importance of these regions in exchange properties. The same group, a year later, examined the properties of a proposed selective Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange inhibitor, KB-R7943 (discussed later on). Their findings suggested that drug selectivity between the exchanger NCX family may be a result of several residues in the \( \alpha \)-2 repeat (138). Specifically, it was shown through cysteine-scanning mutagenesis of the NCX1 \( \alpha \)-2 repeat, a single residue, Gly833, resulted in a \( \geq \) 30-fold reduction in KB-R7943 sensitivity (143). These results offer solid evidence indicating the motifs play a primary role in the ion translocation pathway and may be involved in drug inhibition (\( \alpha \)-2 repeat) (138).
Figure 3. *Putative 3-Dimensional Arrangement of the Na⁺-Ca²⁺ Exchanger*. The 3-dimensional topology of the transmembrane segments (TMS) comprising the Calx α-motifs currently are modeled to be oppositely oriented, possibly forming the ion translocation pathway (Adapted from Qui *et al.* (264)).
2.3.2. The Intracellular Loop

The large hydrophilic loop (initially termed loop f) is localized in the cytoplasm (227). It lies between TMS 5 and 6, and contains approximately 550 amino acids, most of which are not required for transport function (211). For instance, a mutant exchanger that lacked nearly the entire loop (Δ240-679) displayed normal transport/exchange activity (211). However, the intracellular loop is thought to be involved in exchanger regulation (210; 211) through intracellular Na⁺, Ca²⁺, H⁺, and various kinases (76; 140; 211). This suggests that either the entire loop or a portion of it is necessary for regulation. Furthermore, exchangers subjected to proteolytic enzyme treatment (α-chymotrypsin), displayed similar results to that of the mutant (Δ240-679) exchanger lacking an intracellular loop. As with the previous regulatory experiments, the exchanger maintains proper exchange function but lacks regulation (211). The long intracellular loop comprises more than 50% of the exchanger’s mass and is therefore, most likely to contribute to exchanger function and maintenance. There are to date, several key regions within the intracellular loop, and these are discussed below.

2.3.2.1. The Exchange Inhibitory Peptide Region (XIP)

A small 20 amino acid sequence of dispersed basic (positively charged) and hydrophobic residues (XIP; RRLLFYKYYKRYRAGKQRG) occurring just off TMS 5 in the intracellular loop is called the XIP region (193; 227). It is also known as the endogenous XIP region. This unique sequence of amino acids was initially identified for
having homology to a calmodulin-binding domain (227). Li et al. later characterized this segment as the XIP peptide (193). It is postulated that the XIP region contributes to exchanger regulation (autoregulation or autoinhibition) (193; 227). Several groups tested the hypothesis through the use of site-directed mutagenesis (deletion and substitution) and revealed that a form of Na⁺-dependent regulation could be removed (Δ229-232) or accelerated (F223E) upon residue alteration (209). Another group found that by mutating this region (XIP-4YW), they could remove Na⁺-dependent regulation (244). More so, mutants lacking Na⁺-dependent regulation were also insensitive to the effects of known regulators such as ATP, PIP₂, and PIP₂ antibody (112). A study by Pan et al. showed that these XIP mutants were insensitive to the removal of ATP (removal normally inhibits the exchanger) or to activation via phorbol esters (244). Although these exchangers maintained transport function, they could not be regulated. Just as the endogenous XIP region can exert regulatory effects, so can an exogenous form of XIP, otherwise known as the XIP peptide (193). Philipson and collaborators used a fabricated peptide with the exact 20 amino acid sequence of the original endogenous XIP region. When applied cytoplasmically to giant, excised patches from *Xenopus* oocytes, the XIP peptide completely inhibited reverse mode Na⁺-Ca²⁺ exchange (IC₅₀ ~ 0.1 μM) (193). The Philipson group also measured the effects of modified XIP peptides on ⁴⁵Ca²⁺ uptake in Na⁺ loaded cardiac sarcolemmal vesicles. Ultimately, it was found that a fully intact XIP peptide was required for maximum potency. Furthermore, this study revealed that the major inhibitory region of XIP occurs between amino acids 5 and 16, with basic and aromatic residues constituting the most important inhibitory components (113). Changes in these amino acids drastically alter the inhibitory effects of XIP peptide (113). Current
results, using a novel Na⁺-Ca²⁺ exchange inhibitor (SEA0400), are presented in the following pages and stress the importance of this small but vital region.

2.3.2.2. Regulatory Ca²⁺ Binding Region

The large intracellular loop is associated with regulation (ionic regulation via the XIP region) but other regions have been probed in the hopes of isolating further regulatory segments (193; 209). A technique known as ⁴⁵Ca²⁺ overlay was implemented on canine NCX1.1. Using fusion proteins representing various portions of the intracellular loop, they measured binding of radioactive ⁴⁵Ca²⁺. Ultimately, a 138 amino acid stretch (371-508) was isolated and characterized as the Ca²⁺ binding domain (188). In this region, there are two highly conserved portions, each containing three consecutive aspartic residues (188). Normal co-operative binding of Ca²⁺ with a range of 0.3 to 3 μM, was significantly diminished when these aspartic acid residues were mutated (188). Applying a giant, excised patch clamp identified four specific aspartic acid residues (447, 448, 498, and 500) that are responsible for a decrease in the domain's affinity for regulatory Ca²⁺. Interestingly, this high affinity Ca²⁺ binding region is highly conserved among Na⁺-Ca²⁺ exchangers, and functional Ca²⁺ regulation has been shown for NCX1, NCX2, and NCX3 (194). However, in the Drosophila exchanger, CALX, Ca²⁺ regulation is opposite to that of NCX1. In NCX1, Ca²⁺ stimulates exchange, whereas in CALX, Ca²⁺ inhibits exchange. The internal characteristics of these two exchangers prove to be interesting, for their Ca²⁺ regulatory regions appear to have an analogous
function (179). Further investigation into this regulatory mechanism could provide a better understanding of this important component.

2.3.2.3. Calx-β Motifs

On the intracellular loop is a pair of Calx-β motifs that were first identified on the Drosophila exchanger, like the Calx-α motifs (290). These highly conserved segments span ~ 70 amino acids. The first Calx-β motif is closely associated with the regulatory Ca\(^{2+}\) binding site, near the loop’s N-terminus. The second Calx-β motif, located near the C-terminus of the Ca\(^{2+}\) regulatory site, is thought to fold and interact with the regulatory Ca\(^{2+}\) region. Levitsky et al. ran an SDS-PAGE gel and found that Ca\(^{2+}\) binding to fusion proteins stimulated a large mobility shift (188). A potential folding and protecting of the regulatory Ca\(^{2+}\) binding domain could be a possible explanation for this result.

2.3.2.4. The Alternative Splice Site

The Na\(^{+}\)-Ca\(^{2+}\) exchanger is located in various tissues, but is particularly prominent in heart, nerve and kidney cells (96; 160; 202). However, as shown by Omelchenko et al., Ca\(^{2+}\) signaling and handling are different in each (83; 241). Their primary differences occur because they are tissue-specific isoforms of NCX1. These isoforms arise from an alternative splice site situated near the C-terminus of the large intracellular loop. In NCX1, this region is encoded by six cassette exons; A-F, of which
exon A and B never appear together in the same isoform. The last 4 exons can appear in different combinations for remaining splice variants. Variants containing exon A are usually located in excitable tissues, such as brain, skeletal and cardiac muscle (265). Variants containing the B exon are found in the remaining tissues (e.g., kidney, lung, and smooth muscle. There are 32 potential splice variants, however only 12 have been identified (162; 169; 184; 265; 324). The existence of these tissue specific variants may imply specific roles in varying cellular systems. For example, NCX1.4 (AD exon; Brain) is regulated by protein kinase A (PKA) whereas NCX1.3 (AB exon; Kidney) is not (111; 276). As mentioned previously, Ca$^{2+}$ signaling and ionic regulation is varied with different isoforms, especially in cardiac (NCX1.1), kidney (NCX1.3), and brain (NCX1.4) when tested via giant, excised patch clamping (83). Through these experiments, it appears that the presence of either exon A or B contributes to these differences. The *Drosophila* exchanger, CALX, contains two splice variants (CALX1.1 and CALX1.2), in a region consistent with the mammalian NCX (240; 277). A deeper investigation of the properties that govern regulation of Na$^+$-Ca$^{2+}$ exchanger splice variants must be conducted.
**Figure 3. Topology of the Na\(^+\)-Ca\(^{2+}\) Exchanger.** The Na\(^+\)-Ca\(^{2+}\) exchanger is currently modeled to span the membrane 9 times, with a large intracellular loop between transmembrane segments (TMS) 5 and 6 that houses the XIP region. There are 2 re-entrant loops, called Calx-α motifs, located between TMS 2 and 3, and TMS 7 and 8. There are two Calx-β motifs located in the central part of the intracellular loop (Adapted from Philipson and Nicoll (250)).
3. Transport Properties of the Na\(^+\)-Ca\(^{2+}\) Exchanger

3.1. Electrogenicity and Stoichiometry

Since the late 1960’s, Na\(^+\)-Ca\(^{2+}\) exchange stoichiometry has been a challenging topic and therefore, subject to debate. Initial evaluations of cardiac tissue and squid axons (7) suggested that Na\(^+\)-Ca\(^{2+}\) exchange was electrogenic. This would imply a stoichiometry > 2 Na\(^+\): 1 Ca\(^{2+}\), for electroneutral exchange would not produce current (i.e., 2:1). Many initial studies only measured unidirectional ionic movement and employed indirect methods for stoichiometric estimation (7). Currently, the Na\(^+\)-Ca\(^{2+}\) exchanger is considered an electrogenic ion countertransporter, with a well-documented stoichiometry of 3 Na\(^+\): 1 Ca\(^{2+}\) (40; 42; 62; 144; 164; 165; 179; 205; 207). However, new progress may suggest otherwise.

Initial studies following identification of the squid Na\(^+\)-Ca\(^{2+}\) exchanger (7), measured unidirectional ion movement (Na\(^+\)-dependent-Ca\(^{2+}\) efflux and Ca\(^{2+}\)-dependent Na\(^+\)-influx), and suggested exchange was coupled at a 3 Na\(^+\): 1 Ca\(^{2+}\) ratio (the initial stoichiometry proposed for Na\(^+\)-Ca\(^{2+}\) exchange) (7). Measuring Na\(^+\)-Ca\(^{2+}\) exchange activity, on any level, can be difficult. To date, it is impossible to isolate a single Na\(^+\)-Ca\(^{2+}\) exchanger and measure the resulting current. Even so, if a single exchanger were indeed isolated, the resulting current would be too small for accurate detection (unlike ion channels that allow the movement of large amount of ions, producing large elementary currents). It is understandable, therefore, that there are so many discrepancies among Na\(^+\)-Ca\(^{2+}\) exchange results, especially in the case of stoichiometry. Earlier investigations
measured voltage-dependencies, thermodynamics, and ionic competition as stoichiometric indicators. Through these results investigators concluded exchanger electrogenericity (87; 182; 183; 257; 269; 275). These results were based on the assumption that electrogenic systems must be voltage-sensitive, however the reciprocal statement does not always hold true. This was noted, and a new experiment was performed with results concluding that both electrogenic and electroneutral Na⁺-Ca²⁺ exchange are altered by membrane potential (mV) (87). In 1987, Kimura et al. (164) documented the first measurement of Na⁺-Ca²⁺ exchange current in whole cell patches, solidifying the exchanger’s electrogenericity. In 1984, Reeves and Hale re-evaluated the exchanger’s stoichiometry. Using a novel method involving inside-out bovine sarcolemmal vesicles, they measured ⁴⁵Ca²⁺ flux (269). By balancing ionic concentrations they essentially removed the net driving force of both Ca²⁺ and Na⁺. This allowed them to manipulate the exchanger and measure stoichiometry based on a response to changes in membrane potential (mV). Membrane potential was altered using an ionophore (valinomycin: K⁺ channel former), which could generate positive or negative potentials. The result was a stoichiometry of 2.97 Na⁺ to 1 Ca²⁺. The Reeves and Hale experiment (269) complimented a less convincing study completed five years prior (257). It is imperative that a correct stoichiometry be determined for Na⁺-Ca²⁺ coupled exchange, as many bodies of work have based their results on an assumption of 3 Na⁺: 1 Ca²⁺ coupled exchange. Measurements of Ca²⁺ entry, efflux, transport rates (number of ions transported per unit time) and kinetics are all affected by stoichiometric designation. Once electrophysiological techniques became readily available, more attempts at identifying a correct stoichiometry were made. Certain studies chose to
measure the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger reversal (\(E_{\text{rev}}\)) potential using formulaic predictions made at various stoichiometric ratios. Using whole-cell voltage clamping and measuring reversal potentials, several groups reported a consistent stoichiometry of 3:1, which has remained the popular convention (30; 86).

Recently, there have been several reports that challenge the classic 3:1 stoichiometry. Reports suggesting a 4:1 ratio have appeared over the last few years (15; 79). This ratio was initially proposed in 1977 by Mullins (219), and then later in 1983 by Ledovra and Hegyvary (182). In 2000, Fujioka et al, re-introduced the notion of a 4:1 Na\textsuperscript{+}-Ca\textsuperscript{2+} stoichiometry, using inside-out “macro” patches excised from guinea-pig ventricular myocytes (98). They concluded that the stoichiometry was 4:1 but had the ability to shift with changing ionic conditions (98). Measuring Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange stoichiometry accurately in native cell systems is difficult, primarily due to confounding variables such as: 1) ionic channels and 2) various pumps. Both can affect Na\textsuperscript{+} and Ca\textsuperscript{2+} movement and concentration. Therefore, Dong et al. suggested that the cell systems and techniques used in previous stoichiometric studies could have had confounding variables thereby, altering final results (79). They chose to express NCX1 in a system devoid of potential contaminating ionic transport pathways. Using a heterologous cell system (allowing high levels of recombinant exchange) of HEK-293 cells, they expressed NCX1.1 from rat hearts. Applying electrophysiology they measured the thermodynamic equilibrium for Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange under varied ionic conditions. As was mentioned, only one other group, Szerencsei et al., reported using this system to measure stoichiometry (309). In addition to minimizing confounding variables, they noted that HEK cells were electrically quiet, with small amounts of resting membrane current (79).
By applying various voltage steps (ramps), they set out to establish a $\text{Na}^+\text{-Ca}^{2+}$ exchange reversal potential ($E_{\text{rev}}$). Through this, they found a charge ratio of 2 to 1, making the stoichiometry (if $\text{Na}^+$ has one charge and $\text{Ca}^{2+}$ has two charges) 4:1.

A report from the Matsuoka group also suggested a 4:1 stoichiometry. Using inside-out macropatches excised from guinea-pig sarcolemma, XIP-sensitive currents revealed that stoichiometry was primarily 4:1. However, this ratio fluxed to that of 3:1 upon lowering cytoplasmic $\text{Na}^+$ concentrations (206). Kimura (125) contested these recent 4:1 findings. In their experiment, guinea pig ventricular myocytes were used to assess stoichiometry. Both the $\text{Na}^+\text{-Ca}^{2+}$ exchanger's reversal potential and intracellular $\text{Ca}^{2+}$ concentration (via confocal microscopy) were measured. Placing the cell at a holding potential (HP) that coincided with the theoretical reversal potentials (for different stoichiometries), they found that the $E_{\text{NCX}}$ measured coincided with the $E_{\text{holding}}$ (HP) and the theoretical $E_{\text{rev}}$ (same as HP in this case). Also, when measuring at a HP equal to the predicted $E_{\text{rev}}$ for a 3:1 ratio, the membrane potential remained constant. At a holding potential (HP) equal to a 4:1 stoichiometry, however, the membrane potential initially began as a 4:1 value, but soon shifted to a 3:1 value. Based on these results, a conclusion of 3:1 was given (125).

Establishing an accurate stoichiometry is crucial to further $\text{Na}^+\text{-Ca}^{2+}$ exchange study. Future studies require an accurate determination of stoichiometry to assess other various exchange functions. It has been suggested that if a 4:1 ratio existed, then a serious re-evaluation of previous literature would be required (130). Properties such as transport and turnover rates (98; 118; 124; 150), probability of mode occurrence under physiological conditions (i.e., forward vs. reverse), resting ionic concentrations, and even
the exchanger's level of physiological importance would change. Although not globally accepted, the idea of a 4:1 stoichiometry is not entirely illogical. It is reported that a related exchanger, the NCKX (Na\(^+\)-Ca\(^{2+}\)/K\(^+\) exchanger), has a similar 4:1 stoichiometry (261; 316). The NCKX family are found in photoreceptor cells (273) and neural tissue (261; 316), and are functionally related but hold little structural similarity to the NCX family. However, both families are ion counter-transporters that move Ca\(^{2+}\) across the membrane. With the NCKX family, a Ca\(^{2+}\)-dependent regulatory mechanism was reported, preventing a sub-nanomolar Ca\(^{2+}\) range from occurring (287; 288). This said, the potential of a 4:1 stoichiometry existing, may be more realistic then initially considered.

3.2. *Turnover Rates and Exchanger Density*

A transport/turnover rate refers to the number of ions transported by a single exchanger/channel per unit time. However, an ion channel allows a large number of ions to pass, making it easier to measure elementary current (the magnitude of current passing through a single channel) and to detect turnover rates. Considering the difficulty of isolating single exchangers and that resulting current is very small, direct measurements cannot be made. Therefore, most Na\(^+\)-Ca\(^{2+}\) exchanger turnover rates have been made through indirect approaches. As in the case of stoichiometry, there remains significant disparity regarding correct turnover rates. Reports estimating values from <75 s\(^{-1}\) (262), to 5000 s\(^{-1}\) (118; 124) have appeared in the literature. Such a diverse range of turnover rates largely occurs because of indirect and different methods applied by investigators. In
1996, Hilgemann *et al.* implemented a new, more direct technique equating current noise associated with exchanger inactivation to transport rate measurements (124). Maximum unitary exchange currents were estimated at 0.6 - 1.3 fA (10-15 A), with a charge movement rate of 5000 s\(^{-1}\) (124).

Lacking experimental tools (*i.e.*, radioligands) and unable to accurately assess elementary current (no way to determine an exchanger count), investigators turned to measuring exchanger densities. Combined results of exchanger densities and turnover rates could provide insight into the protein's physiological and pathophysiological importance. Nevertheless, exchanger densities proved equally ambiguous. Density values range from 250 (232), 300-400 (in giant, excised patches) (118; 124), to 1235 (262) exchangers per \(\mu\)m\(^2\). Equally indefinite to exchanger density is exchanger location. Understanding the exchanger density in various cellular regions (*i.e.*, T-tubules or other membrane invaginations) would be valuable in characterizing exchanger function. However, most recent evidence shows the Na\(^+-\)Ca\(^{2+}\) exchange to be located on the surface of the SL, intercalated disc region and vertical T-tubule region. Furthermore, using the rat model, Thomas *et al.* implemented double labelling studies and found that the Na\(^+-\)Ca\(^{2+}\) exchanger is as close to the SR Ca\(^{2+}\) release channels as the L-type Ca\(^{2+}\) channels are. This may provide additional support for the Na\(^+-\)Ca\(^{2+}\) exchanger having more than a Ca\(^{2+}\) removal role in the cell. Such that, under certain circumstances (increased [Na\(^+\)], and diseased states) reverse mode exchange could play an important role in E-C coupling.
3.3. Transport Mechanism

The Na\(^+\)-Ca\(^{2+}\) exchanger has two proposed transport mechanisms. The first mechanism, dubbed “Consecutive” or “Ping-Pong” transport, has ion binding sites that shift between intracellular or extracellular positions (124; 151; 155; 190; 242). Essentially, the process can be broken down into two-steps. First, Na\(^+\) binds and is translocated across the membrane. Next, Ca\(^{2+}\) binds and is carried through the membrane. This process is considered analogous to that of the Na\(^+\) pump (Na\(^+\)-K\(^+\)-ATPase) (36; 214). During the translocation of these ions, they pass through several occluded states. The process is reversible and occurs at a very rapid rate. The second mechanism is known as “simultaneous” transport. Basically, ionic transport only occurs if both external and internal sites become occupied (i.e., Na\(^+\) binds extracellularly and Ca\(^{2+}\) binds intracellularly), and are transported at the same time. There is, however, a third mechanism termed “sequential” transport, initially proposed by Milanick and Frame (214), stating that an exchanger requires both ions to be bound on the same side, in order for an ion to dissociate (214). Consecutive transport is largely agreed upon. In 1977, Blaustein produced results consistent with the simultaneous model (37). This analysis was based on two results. First, there was no increase in Ca\(^{2+}\) efflux upon an increase of intracellular Ca\(^{2+}\). Secondly, the exchanger mediated both Na\(^+\)-Ca\(^{2+}\) exchange and Ca\(^{2+}\)-Ca\(^{2+}\) exchange (alkali metal ion-activated), implying that both exchanger sides must be occupied for transport to occur (36; 37). If this were not the case, they would have noticed unidirectional or uncoupled transport occurring, but they did not. Perhaps the assumption that uncoupled exchangers are still functional and that recordings were not
performed under “zero-trans conditions” (i.e., concentration of counter ion on the opposite side should be zero) lead to these conclusions (36). However, in 1990 and 1991, Khananshvili and Kimura respectively, repeated the Blaustein experiment under “zero-trans conditions” and generated results that were in agreement with the consecutive transport mechanism (155; 189).
Figure 4. The Consecutive Eight-State Model for Na\(^+\)-Ca\(^{2+}\) Exchange. Eight kinetic states and one inactive state (I\(_1\)) are represented in this transport model. E\(_{oi}\) and E\(_{oo}\) represent empty states, E\(_{n}\) and E\(_{e}\) are Na\(^+\) and Ca\(^{2+}\) occluded states, E\(_{no}\) and E\(_{co}\) correspond to extracellular Na\(^+\) and Ca\(^{2+}\) bound states, while E\(_{ni}\) and E\(_{ci}\) stand for intracellular Na\(^+\) and Ca\(^{2+}\) bound states. The representation is shown with both slow (front) and fast (back) transport configurations. Clockwise cycling represents reverse mode (i.e., Ca\(^{2+}\) influx) exchange and counter-clockwise cycling represents forward mode (i.e., Ca\(^{2+}\) efflux) exchange.
In 1991, Hilgemann et al. applied an electrophysiological approach to the measurement of transport (124). Using the giant, excised patch technique, they measured $\text{Na}^+$ and $\text{Ca}^{2+}$ partial transport reactions and suggested a consecutive mechanism of transport. Low extracellular (pipette) concentrations of either $\text{Na}^+$ (5 mM) or $\text{Ca}^{2+}$ (10 $\mu$M) were applied initially in the absence of intracellular $\text{Na}^+$ or $\text{Ca}^{2+}$. With only one type of ion in the pipette at a time (either $\text{Na}^+$ or $\text{Ca}^{2+}$), the exchanger would bind the ion and translocate it. At this point, the exchanger-binding site is facing the cytoplasmic side, devoid of any ions. In a consecutive model, all the ion-binding sites orient themselves facing the cytoplasmic side, and therefore, all the exchangers were facing the cytoplasm prior to experiment initiation. A positive charge movement was measured with cytoplasmic $\text{Na}^+$ application, but not with $\text{Ca}^{2+}$ (150). Thus, $\text{Na}^+$ translocation and/or unbinding were considered to be the electrogenic step(s) (150).

Using flash photolysis (releasing “caged $\text{Ca}^{2+}$” upon stimulation), Niggli and Lederer reported unidirectional charge movements termed “conformational currents”, in voltage-clamped myocytes. These currents were reportedly different compared to $\text{Na}^+-$ $\text{Ca}^{2+}$ exchange current (232). Also using photoreleased “caged $\text{Ca}^{2+}$”, Kappl and Hartung found that a negative charge movement was observed during $\text{Ca}^{2+}$ translocation (under specific pipette ion concentrations) (151). Partial reaction charge movements examined in the squid $\text{Na}^+-$ $\text{Ca}^{2+}$ exchanger (NCX-SQ1), were found using either transport substrate ($\text{Na}^+$ or $\text{Ca}^{2+}$), suggesting that ion translocation is electrogenic (114). Overall, the most recent evidence suggests that the exchanger operates via a consecutive transport mechanism.
Initial transport cycle analysis described the exchanger as having eight kinetic states (124) (Figure 4). The $E_{oo}$ and $E_{oi}$ states are exchangers when their binding sites are empty and ready to bind ions. The $E_{no}$ and $E_{co}$ are extracellular states when the ions (either $Na^+$ or $Ca^{2+}$) are bound, while the $E_{ni}$ and $E_{ci}$ are the occluded ion bound states. The $E_{ni}$ and $E_{ci}$ are the bound states on the intracellular side of the exchanger (124). There is also an inactive state known as the $I_1$, which is entered through the $E_{ni}$ state, also known as $E_{3ni}$ (for it binds 3 $Na^+$ ions) (123). Not only does the exchanger transport $Na^+$ and $Ca^{2+}$ ions, it is regulated by them (121; 123). Specifically, $Na^+$ regulates the exchanger by activating it and then brings it into an inactive state known as $I_1$. Initially described by Hilgemann in 1990 (116), this process is known as $Na^+$-dependent inactivation ($I_1$) and occurs only during reverse mode (clockwise cycling) exchange (where cytoplasmic $Na^+$ is present). It is termed $Na^+$-dependent inactivation ($I_1$), because as $Na^+$ concentration increases, there is a corresponding increase in current decay to a steady state level (123). In addition, the eight state model presented includes the hypothetical “fast” and “slow” cycles. It was proposed by Omelchenko et al. that the exchanger could transition between “slow” and “fast” cycles depending on the presence or absence of regulatory $Ca^{2+}$ (242). It is believed that recovery from the $I_1$ inactive state is accelerated by increasing cytoplasmic $Ca^{2+}$ and this promotes the exchanger to transition into a “fast” cycle, in which the likelihood of entry into $I_1$ is decreased, while exit from $I_1$ is increased. This may explain why less $I_1$ is observed upon increasing regulatory $Ca^{2+}$ concentrations.
Figure 5. Na⁺-Dependent Inactivation (Iᵢ). The above figure shows a typical outward (i.e., reverse mode) Na⁺-Ca²⁺ exchange current trace obtained using the giant, excised patch technique. Current decay (i.e., inactivation) increases as the cytoplasmic Na⁺ concentration is raised.
4. Regulation of Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchange

4.1. Na\textsuperscript{+}-Dependent Regulation (I\textsubscript{1})

Currently, it is unclear whether Na\textsuperscript{+}-dependent regulation (I\textsubscript{1}) plays an important role in the physiological setting. Since we do not know for certain when and where reverse mode exchange occurs, it is difficult to determine this mechanism’s importance. In a whole cell system of intact cardiac guinea-pig myocytes, Na\textsuperscript{+}-dependent inactivation was reported (208). However, like many other experimental procedures, this study was conducted under atypical physiological conditions. Specifically, intracellular Na\textsuperscript{+} concentrations used were > 30 mM, nearly three times that of physiological levels (~ 10 mM) (208). Most experimental settings measuring Na\textsuperscript{+}-dependent inactivation use concentrations of 25-100 mM Na\textsuperscript{+} to achieve significant exchange current and inactivation (current decay; I\textsubscript{1}). Otherwise, such results are unlikely under normal physiological Na\textsuperscript{+} concentrations. Na\textsuperscript{+} pooling (during an action potential) near the sarcolemma is suggested to be sufficient for the initiation of reverse mode exchange (5; 185; 328; 329). Like many other cellular phenomenon, this may be physiologically possible but highly unlikely. At very high (positive) membrane potentials, the reverse mode exchange is favoured. This is due to the positive repulsion of intracellular Na\textsuperscript{+}, which would favour exiting the cell, thereby stimulating outward Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange (reverse mode). Weber et al. demonstrated exchange activity at membrane potentials well above physiological levels (+50 mV) (329). This study showed that a very high potential must be reached to achieve reverse mode exchange. However, there are times
when normal physiological conditions are exceeded and could produce the “unlikely” reverse mode exchange. Specifically, in the case of a positive inotropic increase, where frequency of stimulation is increased, augmenting local Na\(^+\) levels with every depolarization. These Na\(^+\) levels would rapidly accumulate and contribute to the positive inotropic response seen under such circumstances (i.e., increasing CICR and resting Ca\(^{2+}\) levels).

Understanding Na\(^+\)-Ca\(^{2+}\) exchange function pathophysiologically is equally important as understanding it physiologically. Insight into the regulatory role of Na\(^+\)-dependent inactivation remains a key research interest, as I\(_1\) regulation is reverse mode specific, which in turn, is primarily associated with pathophysiology. An explanation into the I\(_1\) mechanism and its implications in cardiac pathophysiology could lead to novel therapeutic approaches (ideally pharmaceutical). When \(\alpha\)-chymotrypsin-induced proteolysis of the Na\(^+\)-Ca\(^{2+}\) exchanger is analysed, the I\(_1\) mechanism is apparently removed, while the resulting exchanger appears deregulated (116). It is suggested that cleavage occurs along the large intracellular loop, because a loop-less exchanger lacks I\(_1\) (211). The XIP region has been implicated in the development of I\(_1\), for mutagenesis experiments reveal significant alterations in Na\(^+\)-dependent inactivation (113; 209; 244). Usually mutants targeting Na\(^+\)-dependent inactivation also affected Ca\(^{2+}\) regulation (209). Moreover, it has been shown that both regulatory Ca\(^{2+}\) and mutations in the regulatory Ca\(^{2+}\) binding domain, affect I\(_1\) suggesting an interaction between these two mechanisms (in the intact exchanger) (121; 209; 210). Recent studies using a novel reverse mode specific Na\(^+\)-Ca\(^{2+}\) exchange inhibitor, SEA0400, suggest a mechanism of action requiring I\(_1\) (data currently in review).
4.2. \( \text{Ca}^{2+}\)-Dependent Regulation (I$_2$)

Intracellular \( \text{Ca}^{2+}\) contributes to the \( \text{Na}^+\)-\( \text{Ca}^{2+}\) exchanger’s regulatory profile. Intracellular \( \text{Na}^+\) inactivates \( \text{Na}^+\)-\( \text{Ca}^{2+}\) exchange. Cytoplasmic \( \text{Ca}^{2+}\) stimulates it. This regulatory \( \text{Ca}^{2+}\) remains non-transported and is essential for \( \text{Ca}^{2+}\)-dependent regulation, otherwise known as I$_2$ (116; 123; 208). DiPolo, who found that in the absence of intracellular \( \text{Ca}^{2+}\) no \( \text{Na}^+\)-dependent \( \text{Ca}^{2+}\) influx occurs (67), initially reported \( \text{Ca}^{2+}\)-dependent regulation in the giant squid axon. Furthermore, \( \text{Ca}^{2+}\)-dependent regulation, or I$_2$, alleviates \( \text{Na}^+\)-dependent inhibition (121) and stimulates exchange activity thereby, increasing current. This has been shown in several different exchanger types (such as SQ1-NCX (114), NCX1, NCX2, and NCX3 (194)) and in varying experimental settings (52; 93; 210; 215). Comparing techniques, the electrophysiological approach of giant, excised patch clamping provided the most detailed characterization of \( \text{Ca}^{2+}\)-dependent regulation (I$_2$) (116; 121; 210). These data suggest regulatory \( \text{Ca}^{2+}\) is not transported and is primarily an activator of \( \text{Na}^+\)-\( \text{Ca}^{2+}\) exchange. The \( \text{Ca}^{2+}\) affinity (\( K_m \)) for the regulatory \( \text{Ca}^{2+}\) binding domain ranges from nanomolar to micromolar values. Various experimental settings from guinea-pig myocytes to Xenopus giant, excised patches have reported \( K_m \) values ranging from 22 nM – 0.4 \( \mu \)M (97; 121; 209; 210; 215). With such a broad spectrum of \( K_m \) values, it is difficult to assess the proper physiological role of \( \text{Ca}^{2+}\)-dependent activation. For example, if a low \( K_m \) value is correct (\textit{i.e.}, nanomolar affinity), then even at resting diastolic \( \text{Ca}^{2+}\) values, this regulatory site would be saturated, implying that it has no modulatory role. However, if the \( K_m \) value were in the low micromolar range, then this would be more responsive to the physiological environment (activating at systolic \( \text{Ca}^{2+}\) values and inactivating at lower diastolic \( \text{Ca}^{2+}\) values). Bers et
al. gave a projected physiological Ca\textsuperscript{2+} range (during systole and diastole) of ~0.1 - 1.0 μM (19). Part of this K\textsubscript{m} discrepancy may be a result of different experimental conditions. In the case of whole cell patches versus giant, excised patches, the former has limited capabilities for intracellular Ca\textsuperscript{2+} change. Cells cannot tolerate high Ca\textsuperscript{2+} as they become toxic. This toxicity could either produce incorrect results and/or kill the cell. Therefore, results at higher Ca\textsuperscript{2+} concentrations may be overlooked due to these concentration borders. In contrast, giant, excised patches can be subjected to a wide range of cytoplasmic Ca\textsuperscript{2+} concentrations (10-30 μM range), and allows for a broader analysis (130).
Figure 6. **Ca\(^{2+}\)-Dependent Regulation (I\(_2\)).** Illustrated above is a typical outward (i.e., reverse mode) Na\(^{+}\)-Ca\(^{2+}\) exchange current trace obtained from giant, excised patch clamp experiments. Increasing cytoplasmic Ca\(^{2+}\), or regulatory Ca\(^{2+}\), stimulates Na\(^{+}\)-Ca\(^{2+}\) exchange current and alleviates I\(_1\) inactivation. At concentrations > 10 µM regulatory Ca\(^{2+}\), Na\(^{+}\)-Ca\(^{2+}\) exchange activity decreases presumably due to competition between transport Ca\(^{2+}\) and Na\(^{+}\).
As mentioned earlier, there appears to be substantial interaction between these two regulatory mechanisms. It has been demonstrated by Hilgemann et al., that upon increasing regulatory Ca\textsuperscript{2+}, the exchanger is stimulated while Na\textsuperscript{+}-dependent inactivation is attenuated (121). If the regulatory Ca\textsuperscript{2+} levels are high enough, I\textsubscript{1} can appear abolished. Furthermore, mutations in the regulatory Ca\textsuperscript{2+} binding region performed by Matsuoka et al., decreased relative cytoplasmic Ca\textsuperscript{2+} affinities and altered Na\textsuperscript{+}-dependent inactivation (210). This report established Ca\textsuperscript{2+} regulation for both forward (Ca\textsuperscript{2+} efflux) and reverse mode (Ca\textsuperscript{2+} influx) exchange (210). Conversely, mutations specific for Na\textsuperscript{+}-dependent inactivation altered Ca\textsuperscript{2+} regulation (209). Continuing, α-chymotrypsin treatments eliminated Ca\textsuperscript{2+}-dependent regulation as well as Na\textsuperscript{+}-dependent inactivation (116; 151; 211). There is a significant body of literature suggesting these two mechanisms interact with one another and play important roles in reciprocal regulation. Investigations of both I\textsubscript{1} and I\textsubscript{2} may provide further insight as to how the exchanger regulates itself as a global and/or single unit.

4.3. Regulation of Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchange by pH

When the cell is exposed to varying pH levels, a corresponding change in Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange activity occurs. It has been shown that the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger is highly sensitive to slight variances in resting cellular pH (~ 7.2) (75-77; 159; 247). In general, it appears that an alkalization of the environment increases exchange activity, while an acidification inhibits. The cytoplasmic surface of the exchanger looks to be affected to a greater extent than the extracellular surface. Take for example, a study by DiPolo et al.,
which displayed a ~ four-fold increase in exchanger activity upon *intracellular* alkalinization to pH 8.8, yet no effect was seen with an *extracellular* alkalinization up to pH 9.0 (68). Upon acidification, exchanger inhibition by way of proton block is suggested to occur through two separate components (76). The first inhibitory component is slow forming and dependent on the presence of cytoplasmic Na\(^+\). The second component was noted as rapid and Na\(^+\)-independent. In giant, excised patches exposed to α-chymotrypsin, the proteolyzed exchanger appears unaffected by acidification, which would normally result in inactivation. Considering that the proteolyzed exchanger lacks ionic regulation and that a component of proton block required the presence of Na\(^+\) (cytoplasmic), it is possible that the effects of pH on the exchanger involve I\(_1\), I\(_2\), and/or the intracellular loop (77).

4.4. *Phosphorylation*

4.4.1. *Protein Kinase C (PKC) and Protein Kinase A (PKA)*

Na\(^+\)-Ca\(^{2+}\) exchange phosphorylation has been studied for nearly 30 years, yet still remains a controversial subject. Initially characterized on the squid axon exchanger in 1977 by DiPolo et al., phosphorylation via ATP and various analogues (with hydrolysable terminal phosphates) was shown to stimulate Na\(^+\)-Ca\(^{2+}\) exchange (Na\(^+\)-dependent Ca\(^{2+}\) efflux) (36; 66; 70; 72). Exchange activity, stimulated through ATP application, displayed an increased affinity for intracellular Ca\(^{2+}\) and extracellular Na\(^+\), with a decreased Na\(^+\)-dependent inactivation (33). Recently, DiPolo et al. reported a 13
kDa peptide, isolated from the squid axon cytoplasm, which was required for MgATP-dependent exchanger activation (73). When exogenous alkaline phosphatases were injected directly into the axoplasm, the ability for ATP to activate the exchanger was reversed (73). Even in the previous experiment (73), Dipolo found that under Na\(^+\)-free conditions, ATP does not increase exchanger activity when measuring forward mode exchange. As a note, this mode of exchanger lacks Na\(^+\)-dependent inactivation (I1). Later, Hilgemann used giant, excised patches from cardiac myocytes, and revealed that the exchanger was regulated and inactivated by intracellular Na\(^+\) (123). He further examined this property (I1) and found that it could be alleviated through cytoplasmic MgATP application (123). Perhaps this explains the lack of stimulation by ATP on forward mode exchange (in the absence of Na\(^+\)) observed in the DiPolo study (73). The Na\(^+\)-dependent inactivation process is implicated in this effect. However, Hilgemann and Ball suggested that ATP-dependent activation was the result of phosphatidylinositol 4,5-bisphosphate (PIP\(_2\)) production by phosphorylation of phosphatidylinositol (PI) (119).

Mene \textit{et al.} proposed that vasoconstrictors (angiotensin II and/or endothelin-1) promote Na\(^+\)-Ca\(^{2+}\) exchange by a mechanism independent of Ca\(^{2+}\) and PKC, but potentially linked to Na\(^+\) influx (213). Other investigators found that phenylephrine, endothelin 1, growth factors, and angiotensin II all stimulated Na\(^+\)-Ca\(^{2+}\) exchange activity through a G protein (G\(_{\text{q}}\))-mediated mechanism (8; 9), whereas blocking PKC action prevented exchanger stimulation (9; 140; 141; 305). In mammalian exchangers, PKC mediated phosphorylation was first presented by Iwamoto on aortic smooth muscle (144), followed by its identification in cardiac muscle (141). These studies further indicated that brief exposure to phorbol esters (phorbol 12-myristate 13-acetate or PMA) allowed
PKC-dependent phosphorylation, activating the exchanger (129; 141). However, long-term exposure resulted in the removal of any excitatory response that occurred by growth factors in aortic smooth and cardiac muscle myocytes (141; 144). Another agent found to activate the exchanger through phosphorylation was the platelet-derived growth factor BB (PDGF-BB) (144). However, this effect did not occur when applied to exchangers previously subjected to PMA incubation (144). On the contrary, another study showed that 24-hour pre-treatment with PMA could prevent exchanger activation via angiotensin II (213). Prolonged exposure to both PKC (arteriole myocytes) and PMA is shown to decrease exchanger expression (302; 303). Using phosphopeptide analysis and site-specific mutagenesis, Iwamoto et al. identified multiple phosphorylation sites. There were three sites identified, all on serine residues (Ser-249, Ser-250, and Ser-357) (141; 144). The Shigekawa group, however, created a mutant exchanger where all identified phosphorylation residues were substituted with alanine. Following mutation, it was discovered that the exchangers retained responsiveness to PMA (phorbol ester), suggesting direct phosphorylation of the Na\(^+\)-Ca\(^{2+}\) exchanger was not required for PKC-dependent stimulation. Yet, in NCX1, PMA responsiveness was missing when the large intracellular loop was mutated (244). In this case, an NCX1 mutant, XIP-4YW was not activated by PMA and was resistant to inhibition by ATP removal. The results suggested that the XIP region is involved in exchanger regulation by protein kinase C and/or ATP (244).

Collins et al. used the giant, excised patch clamp technique to observe ATP-induced Na\(^+\)-Ca\(^{2+}\) exchange activation (57). Through the use of various kinase inhibitors, phosphatases, and phosphatase inhibitors they found no increase in exchange activity.
In a later report using bovine cardiac exchanger expressed in Chinese hamster ovary cells, Condescu et al. showed that both calyculin A and okadaic acid (phosphatase inhibitors) significantly inhibited reverse mode exchange (Ca\(^{2+}\) influx). They further reported that mutant exchangers lacking their intracellular loops (central part ~ 420 amino acids) were still responsive to inhibition by calyculin A and okadaic acid, suggesting that phosphorylation exerts its effects indirectly through other cellular proteins. Forward mode exchange was slightly inhibited by calyculin A and okadaic acid.

Another modulator of Na\(^{+}\)-Ca\(^{2+}\) exchange is protein kinase A, or PKA. In the case of PKA-dependent stimulation, NCX1 isoform-specific results have emerged. For instance, the PKA-dependent stimulation of NCX1.4 only displayed a 40% increase in exchange activity. Conversely, NCX1.3 was not stimulated by PKA (16). The use of different cell types, recording techniques, materials, protocols, exchanger species and isoforms all contribute to varying results. A variety of proteins and enzymes may be present and/or lacking among individual experimental methods, potentially contributing to these final results. Although a complete agreement regarding exchanger phosphorylation is lacking, results suggest an involvement in exchanger function.

4.4.2. Phosphatidylinositol 4,5-bisphosphate (PIP\(_2\))

For over 25 years, it has been shown that ATP stimulates the Na\(^{+}\)-Ca\(^{2+}\) exchanger. The giant, excised patch clamp technique has been instrumental in the recent explanation for this mechanism. In 1992, Hilgemann and Ball, using giant, excised
patches, investigated ATP-dependent stimulation on outward (Ca\(^{2+}\) influx) Na\(^+-\)Ca\(^{2+}\) exchange currents. They revealed that the conversion of phosphatidylinositol (PI) to phosphatidylinositol 4,5-bisphosphate (PIP\(_2\)) via phosphorylation was the primary mechanism (119). While cytoplasmic application of PIP\(_2\) mimicked stimulatory effects of ATP, a PI-specific phospholipase C eliminated them. However, the effect could be restored once again, by direct PI application. Also, reversing the stimulatory effect of ATP were PIP\(_2\)-specific phospholipase, anti-PIP \(_2\) antibody, and aluminium (binds PIP\(_2\) highly) (119). In addition, Hilgemann and Ball noted that high intracellular [Ca\(^{2+}\)]\(_i\) reversed the effect of ATP, possibly through the activation of a PIP\(_2\)-specific phospholipase C (119).

The stimulatory effect of ATP on the Na\(^+-\)Ca\(^{2+}\) exchanger is only observed during reverse mode exchange (Ca\(^{2+}\) influx) (73; 123). Na\(^+-\)-dependent inactivation (I\(_1\)), and its attenuation by ATP, implicated the regulatory XIP region. Therefore, the Philipson group examined potential interaction(s) between PIP\(_2\) and the XIP region (112). Exchangers containing mutated endogenous XIP regions were created. These mutants displayed altered Na\(^+-\)-dependent inhibition (I\(_1\)). Findings revealed that regardless of the resulting changes in I\(_1\), any functional response to PIP\(_2\) or PIP\(_2\) antibodies was removed (112). Furthermore, they used immobilized phospholipid vesicles (containing low concentrations of PIP\(_2\)) to which exogenous, iodinated XIP peptide was applied and found to bind. These results further accentuate the complexity and involvement of various phospholipids in Na\(^+-\)Ca\(^{2+}\) exchange regulation.
4.5. Other Modulators of Na\(^+\)-Ca\(^{2+}\) Exchange Activity

Several reports have shown that the Na\(^+\)-Ca\(^{2+}\) exchanger can be directly and indirectly modulated by its lipid environment (56; 120; 176; 177; 251-254; 321; 322). It is known that other ion transporters, such as the Ca\(^{2+}\)-ATPase and Na\(^+\)-K\(^+\)-ATPase, are sensitive to their lipid environment (322), perhaps even to specific lipids. Therefore, it was suggested that a unique lipid group might regulate Na\(^+\)-Ca\(^{2+}\) exchange. Na\(^+\)-Ca\(^{2+}\) exchange is sensitive to charged lipid components, specifically anionic amphiphiles (248). Philipson et al. demonstrated in cardiac sarcolemmal vesicles that anionic phospholipids (namely phosphatidylserine and phosphatidylincholine), when incorporated, could stimulate exchange activity (252; 253; 321). Using solubilized Na\(^+\)-Ca\(^{2+}\) exchangers reformed in proteoliposomes and cardiac sarcolemmal vesicles, phospholipase C and D (which increase negatively charged lipids in the membrane) were both shown to stimulate Na\(^+\)-Ca\(^{2+}\) exchange activity (248; 251). Phospholipase D in particular, displayed at times, a 400% increase in stimulation with a paralleling 9-fold increase in membrane phosphatidic acid (251). The exchanger was shown function optimally under a certain lipid ratio (PC-30% / PS-50% / Cholesterol-20 %) (321). This 20 % component of cholesterol proved important in a previous report by Kutryk et al., when the cholesterol content in ventricular dog sarcolemma was modified via incubation of phosphatidylincholine liposomes with cholesterol (in varied amounts) (177). Cholesterol-enriched vesicles displayed up to a 48% increase in exchange activity, while cholesterol-depleted vesicles decreased in activity by 15% (177). Furthermore, a study investigating the effects of fatty acids on Na\(^+\)-Ca\(^{2+}\) exchange, reported stimulation of
exchanger activity (252). In 1987, Collins et al. utilized giant, excised membrane patches (from cardiac sarcolemma) to study phospholipid involvement (56). By direct application of “vehicle-suspended” phospholipids onto patches, their results conferred a strong stimulatory effect of phosphatidylserine on outward exchange current and altered exchanger kinetics (56). Phosphatidylserine treated patches were deregulated by α-chymotrypsin (56) and showed no further stimulatory effect. This suggested phosphatidylserine exerts its effect on the “protease-sensitive regulatory domain”. Using deregulated exchangers (not already treated with phosphatidylserine) a stimulation with phosphatidylserine was maintained. This suggested an interaction with the exchange mechanism rather than regulatory mechanisms. In general, there is an agreement that the Na⁺-Ca²⁺ exchanger is affected by specific sterol/protein interactions in the cellular environment.

5. **Pharmacology of the Na⁺-Ca²⁺ Exchanger**

Pharmacology can reveal information on specific characteristics, mechanisms, and function(s) of various proteins. Conversely, understanding exchanger function can help understand drug mechanism(s). This results in the production of newer and more effective agents. Unfortunately, tools used to manipulate the exchanger have been significantly limited, especially in the “blockers” category.
5.1. Inorganic Cations: Di- and Trivalent Cations

For many years, studies have revealed that certain divalent and trivalent cations can serve as substrates and inhibitors of various Ca$^{2+}$-dependent mechanisms. The Lanthanides, a group of trivalent cations (e.g., La$^{3+}$, Nd$^{3+}$, Tm$^{3+}$, Y$^{3+}$, and Gd$^{3+}$) and their effects on Na$^+$-Ca$^{2+}$ exchange have been investigated (7; 164; 267; 315). Primarily, La$^{3+}$ is regarded as a blocker of Na$^+$-Ca$^{2+}$ exchange, with a low apparent affinity ($\geq 0.5$ mM) (296). Yet extracellularly applied La$^{3+}$ was shown to enter cells through Na$^+$-Ca$^{2+}$ exchange, substituting for Ca$^{2+}$ (296). It is suggested that if this occurs, inhibition may act through the intracellular surface. La$^{3+}$ also blocks voltage-gated L-type Ca$^{2+}$ channels and the plasmalemmal Ca$^{2+}$-ATPase (with a higher affinity than for Na$^+$-Ca$^{2+}$ exchange) (267; 280). This property of La$^{3+}$ is used to isolate Na$^+$-Ca$^{2+}$ exchange by inhibiting other forms of transport activity, when applied at a low concentration ($i.e., < 0.5$ mM) (100; 296). Although applied in whole cell systems (164), measuring La$^{3+}$ inhibition should be performed in a system where the exchanger is in isolation. Therefore, when high amounts of La$^{3+}$ are added (to induce significant inhibition), no other transporter/exchange system is affected. As mentioned previously, other lanthanides such as Tm$^{3+}$, Nd$^{3+}$, Y$^{3+}$, were tested by Philipson in cardiac sarcolemmal vesicles (315). They were relatively poor Na$^+$-Ca$^{2+}$ exchange inhibitors in comparison to La$^{3+}$. Gd$^{3+}$ had no inhibitory effect (315).

Like trivalent cations, divalent cations can adhere to Ca$^{2+}$ binding sites, either inhibiting or acting as transport substrates. However, Mg$^{2+}$, a weak inhibitor (cardiac tissue – IC$_{50}$ $\sim 12.5$ mM) (164; 301), does not show transport activity with the Na$^+$-Ca$^{2+}$
exchanger (315). There is, however, Na\(^+\)-coupled Mg\(^{2+}\) countertransport by Na\(^+\)-Mg\(^{2+}\) exchange (69). Sr\(^{2+}\) and Ba\(^{2+}\), two other divalent cations, have demonstrated Ca\(^{2+}\) substitution and can act as transport substrates (albeit at slower rates) (37) for the Na\(^+\)-Ca\(^{2+}\) exchanger (58; 164; 231; 314). These ions may function in a mode-dependent manner and have been shown to disrupt Ca\(^{2+}\) transport during Na\(^+\)-Ca\(^{2+}\) exchange (37; 315). For example, Trac et al. demonstrated in giant, excised patches virtually no inward current (forward model; Ca\(^{2+}\) extrusion), even upon application of up to 300 \(\mu\)M Ba\(^{2+}\) (314). Unlike Sr\(^{2+}\) and Ba\(^{2+}\), Ni\(^{2+}\) is used as a Na\(^+\)-Ca\(^{2+}\) exchange inhibitor, with an effective range of 2-5 mM (30; 164; 232). However, the inhibitory action of Ni\(^{2+}\) is not very specific and will inhibit other cellular transporters. Yet in many electrophysiological studies, Ni\(^{2+}\) is used to block Na\(^+\)-Ca\(^{2+}\) exchange current (known as "Ni\(^{2+}\) sensitive" current), therefore isolating it (164; 232; 233). Egger et al. demonstrated that Ni\(^{2+}\), like La\(^{3+}\), could inhibit Na\(^+\)-Ca\(^{2+}\) exchange and be simultaneously transported (in a neutral manner) across the membrane (84; 85). Overall, these ions are considered non-selective and must be applied with caution. The combination of high experimental concentrations mixed with low selectivity, makes for risky tools when measuring Na\(^+\)-Ca\(^{2+}\) exchange. However, if used properly, they can provide reliable results.
5.2. Peptides

5.2.1. The Exchange Inhibitory Peptide (XIP)

Upon cloning and sequencing the cardiac Na\(^+\)-Ca\(^{2+}\) exchanger (NCX1.1) (227), the Philipson group found a unique 20 amino acid segment residing in the large intracellular loop. This sequence is presently known as the XIP peptide (193). This small peptide trails off the fifth TMS and contains a mix of hydrophobic and basic residues resembling a calmodulin-binding site (193).

The endogenous XIP sequence (RRLLFYKVKYRYRAGKQRG) spans amino acids 219-238 of the Na\(^+\)-Ca\(^{2+}\) exchanger and is highly conserved among exchanger isoforms; NCX1, NCX2, and NCX3 (194). This peptide was exogenously applied to the cytoplasmic surface and displayed potent exchange inhibition (193). Through various preparations such as cardiac sarcolemmal membrane patches (132; 193), giant, excised membrane patches (Xenopus oocytes) (131; 192), and other various methods (53; 113), the XIP peptide has shown strong inhibitory properties, holding an IC\(_{50}\) range of 0.15-1.5 \(\mu\)M. It was speculated that the endogenous XIP region played a role in exchanger regulation, primarily through a form of autoinhibition. Through the functional comparison of NCX isoforms (NCX1, NCX2 and NCX3), it was shown that the corresponding exchanger XIP peptides could actively inhibit NCX1, however, with a slightly reduced potency (194). Carrying a high degree of inhibition and potency, the properties of the XIP peptide became of interest to investigators. Mutagenesis studies were performed to examine specific amino acids and their relative contribution to the XIP
peptide’s unique characteristics (113). Studies looking at the effects of mutant XIP peptides on the sarcolemmal Ca^{2+}-ATPase are also reported (332; 333). As mentioned previously, Philipson demonstrated that the entire intact XIP peptide is required for proper inhibition, especially amino acids 5 through 16 (270). He et al. uncovered that basic and aromatic residues are most important for the inhibitory function of XIP (113). Substitutions of arginine 12 and arginine 14 with alanine or glutamine (neutralizing them), significantly decreased XIP’s potency (113). In contrast, Xu et al. conferred the opposite, suggesting that arginine 12 and 14 need not be positively charged (but cannot be negative either) for proper inhibitory function (332). Furthermore, Xu et al. demonstrated through Sulfosuccinimidyl acetate (SNA)-modified XIP peptides (with neutralized lysine residues) that one lysine residue out of three (positions 7, 11, or 17) was essential for inhibition (332). In an alternate study, He et al. proposed that most aromatic residues participate in binding and do so through hydrophobic interactions (113). Results from mutagenesis suggested tyrosine-6 was required for maximal inhibition, while phenylalanine 5, tyrosine 8, tyrosine 10 and tyrosine 13 are potentially important players in XIP function (113). The XIP peptide, although a highly potent inhibitor, must be used with caution, for it can bind and interfere with endogenous calmodulin and calmodulin binding proteins (e.g., Ca^{2+}-ATPase) (89). Furthermore, XIP peptide can only act on the cytoplasmic surface and does not cross the membrane readily. Therefore, XIP peptide must be internally dialyzed for certain experimental techniques (i.e., whole cell preparations).
5.2.2. Other Peptides

Other than XIP peptide, there are several similar peptides that have shown to inhibit (some with significant potency) the Na\(^+-\)Ca\(^{2+}\) exchanger. In the cardiac Na\(^+-\)Ca\(^{2+}\) exchanger, there exists an opiate-like binding site (157). The molluscan cardioexcitatory peptide Phe-Met-Arg-Phe-amide (FMRF-amide) is known to bind opiate binding sequences (157). While studying the effects of opiate agonists and antagonists on Na\(^+-\)Ca\(^{2+}\) exchange, Khananshvili et al. found that the FMRF-amide inhibited Na\(^+-\)Ca\(^{2+}\) exchange in cardiac sarcolemmal vesicles (156; 157). The inhibitory action of the FRMF-amide has also been reported in giant squid axons (71) and pancreatic β-cells (319). Khananshvili applied a group of newly synthesized (positively charged) cyclic hexapeptides (156). These hexapeptides have different amino acid sequences, containing two arginines, two phenylalanine, and two cysteines (to form an S-S bond) (156). Altogether, seven cyclic hexapeptides inhibited \(^{45}\)Ca\(^{2+}\) uptake by Na\(^+-\)Ca\(^{2+}\) exchange with different potencies and IC\(_{50}\) values (200-300 μM) (158). When tested on rabbit ventricular myocytes, the study revealed that one of the seven cyclic hexapeptides, dialyzed Phe-Arg-Cys-Arg-Cys-Phe-amide (FRCRCF-amide), showed complete inhibition of Na\(^+-\)Ca\(^{2+}\) exchange at 1 μM (127; 158). However, a subsequent study by the Philipson group, comparing all three Na\(^+-\)Ca\(^{2+}\) exchange isoforms (NCX1, NCX2 and NCX3), demonstrated FRCRCF-amide to have a reduced potency, such that 50 μM resulted in a range of 32-41 % inhibition (194). Inhibition for the FRCRCF-amide peptide occurs on the cytoplasmic surface, just as in the case of XIP peptide. Therefore a cell-permeant analogue of FRCRCF-amide was created and termed Myristyl-(Myr)-
FRCRCF-amide. When applied exogenously on isolated rabbit ventricular myocytes in the whole cell patch configuration, Myrystyl-FRCRCF-amide moderately inhibited Na\(^+\)-Ca\(^{2+}\) exchange current but at a reduced potency (60).

5.3. *Organic Inhibitors*

5.3.1. *Amiloride Derivatives*

The acylguanidine diuretic termed amiloride is a weak inhibitor (IC\(_{50}\) ~ 1 mM) of the Na\(^+\)-Ca\(^{2+}\) exchanger, however, its derivatives are potent (148; 149). Two examples of stronger amiloride derivatives are benzamil and 3,4-dichlorobenzamil (DCB). The first analogue, benzamil, is considered a moderately potent Na\(^+\)-Ca\(^{2+}\) exchange inhibitor with IC\(_{50}\) values ranging from 17 \(\mu\)M in guinea pig vesicles and papillary muscles (297), 36 \(\mu\)M in bovine sarcolemmal vesicles, and 100 \(\mu\)M in rat anterior pituitary cells (147). The second potent analogue inhibitor, DCB, has a reported IC\(_{50}\) of ~ 17 and 30 \(\mu\)M in guinea-pig atrial myocytes (195; 327). These compounds can be difficult to work with due to their limited solubility in aqueous solutions and lack of specificity. Therefore, they can non-selectively block other membrane transport systems (Ca\(^{2+}\)-ATPase, voltage-gated Ca\(^{2+}\) channels, Na\(^+\)-H\(^+\) exchange, Na\(^+\)-K\(^+\)-ATPase and Na\(^+\) channels) (31; 149). These analogues can also bind to other transport systems at higher affinities than that for the Na\(^+\)-Ca\(^{2+}\) exchanger. However, if a system in which other transport mechanisms are either absent or already blocked (perhaps measuring current in whole cell voltage clamping), Na\(^+\)-Ca\(^{2+}\) exchange can be identified by its sensitivity to the chosen inhibitor.
Therefore, studies using Ni$^{2+}$ and amiloride derivatives (i.e., benzamil, DCB or benzobenzamil) are used to isolate and inhibit Na$^+\text{-Ca}^{2+}$ exchange (232). A report from Kaczorowski et al. suggested that amiloride analogues mimic Na$^+$ ions in Na$^+$-transport systems and can therefore, bind and inhibit the exchanger (149). Slaughter et al. found that DCB could inhibit both forward (Ca$^{2+}$ removal) and reverse (Ca$^{2+}$ extrusion) mode exchange (300). DCB was also shown to inhibit Na$^+\text{-Na}^+$ exchange but not Ca$^{2+}\text{-Ca}^{2+}$ exchange. At low concentrations, DCB preferentially binds to a specific Na$^+$-binding site, while at higher concentrations, the inhibitor interferes with Na$^+$, Ca$^{2+}$ and K$^+$ binding (300).

5.3.2. *Isothiourea Derivative: KB-R7943 (No.7943)*

Very few pharmacological agents are available for Na$^+\text{-Ca}^{2+}$ exchange research. Furthermore, those agents that already exist are used exhaustively and remain poor in terms of selectivity, potency and administration. An exception to this is the XIP peptide, however, it is not membrane permeable. However, in 1996 KB-R7943 (\textit{2-[2-[4-(4-nitrobenzyloxy)phenyl]ethyl]}isothiourea methanesulfonate), a selective Na$^+\text{-Ca}^{2+}$ exchange inhibitor was introduced. KB-R7943, a sulphonylurea derivative, is an amphipilic molecule containing a positively charged isothiourea group (at neutral pH) (32). To date, there are no conclusive or unanimous agreements towards the action and potency of KB-R7943. However, most investigators agree that specific pharmacological tools used to manipulate the exchanger must be derived so that we may further our understanding of this important ion countertransporter. As it stands, KB-R7943 was a
step the right direction, for no other compound resembling KB-R7943 ever existed over
the past 30 years. Combined with the use of various (available) experimental tools (e.g.
mutagenesis, chimeric analysis, whole cell and giant, excised patch clamping), a specific
Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange inhibitor could provide detailed analysis and insight into exchange
mechanisms, kinetic properties, ion specificity, and physiological and pathophysiological
functions. As mentioned above, discrepancies still persist in regards to KB-R7943’s
properties (\textit{i.e.}, inhibitory mechanism, mode specificity, isoform specificity and potency).
Figure 7. Chemical Structure of KB-R7943 (No. 7943),
(2-[2-[4-(4-nitrobenzyloxy)phenyl]ethy1]isothiourea methanesulphonate).
The KB-R7943 inhibitor, initially described by two groups (145; 327), was evaluated as being reverse mode (Ca\(^{2+}\) influx) selective in various experimental settings such as guinea-pig ventricular myocytes (327), smooth muscle cells, and NCX1-transfected fibroblasts (145). Reported IC\(_{50}\) values ranged from 0.3-2.4 \(\mu\)M (reverse mode exchange) and 17-30 \(\mu\)M (forward mode exchange) (145; 327). However, in a report by Kimura et al., Na\(^+\)-Ca\(^{2+}\) exchange currents from guinea pig myocytes were measured using whole cell patch clamping and reported that under bi-directional conditions, KB-R7943 inhibition was mode-independent (inhibiting both modes equally with an IC\(_{50}\) \(\approx\)1 \(\mu\)M) (166). Kimura et al. (167), however, potentially explained this effect in a later report. From their previous studies (327), KB-R7943 was suggested to be competitive with respect to external Ca\(^{2+}\). Therefore, they suggested that KB-R7943's direction-dependent block was related to external Ca\(^{2+}\) concentration. They found that if Ca\(^{2+}\) was applied after KB-R7943, inhibition was \(\approx\) 90\% (IC\(_{50}\) \(\sim\) 0.3 \(\mu\)M) and when KB-R7943 was applied following Ca\(^{2+}\), inhibition was reduced to only \(\approx\) 50\% (IC\(_{50}\) \(\sim\) 1.5 \(\mu\)M) (167). With Na\(^+\) affinity for the exchanger being low and Ca\(^{2+}\) affinity being high (and that both compete with each other), when they are both present under bi-directional conditions, Ca\(^{2+}\) binds more readily. Therefore, this would explain why inhibition was equal for both modes of exchange under bi-directional conditions (167).

As mentioned above, competition between KB-R7943 and transported ions (Na\(^+\) and Ca\(^{2+}\)) was suggested. The preliminary reports differed in their views regarding drug-ion competition. Iwamoto et al. proposed that KB-R7943 was non-competitive with respect to Na\(^+\) and Ca\(^{2+}\) in cardiac myocytes, vascular smooth muscle cells and sarcolemmal vesicles (145). Watano et al., using guinea pig myocytes, stated that KB-
R7943 was a competitive antagonist with external Ca$^{2+}$, when measuring outward Na$^+$-Ca$^{2+}$ exchange (Ca$^{2+}$ influx) (327). Furthermore, Iwamoto et al. (who initially stated that KB-R7943 was non-competitive), found while comparing NCX isoforms, that NCX1 and NCX2 showed a mixed type of competition (both competitive and non-competitive with extracellular Ca$^{2+}$), whereas NCX3 remained non-competitive (142). A competitive inhibition with Ca$^{2+}$ was supported by a following report from Watano and Kimura, who showed that after-treatment of KB-R7943 inhibited outward Na$^+$-Ca$^{2+}$ exchange current with an IC$_{50}$ of 3 µM, a value 10 times higher than that of KB-R7943 pre-treatment (326). Clearly, this component of KB-R7943 requires clarification.

When tested on different isoforms, KB-R7943 demonstrated a three fold higher level of inhibition for Na$^+$-dependent $^{45}$Ca$^{2+}$ uptake in CCL-39 fibroblast cells expressing NCX3, compared to those expressing NCX1 and NCX2 (IC$_{50}$ range 1.5 – 4.9 µM) (142). However, another study by Philipson, expressing the three isoforms in BHK cells, found no differential selectivity between isoforms and only demonstrated a 35 – 45 % range of inhibition at 10 µM KB-R7943 (194). It is possible that KB-R7943 binds to regions specific to only certain isoforms, resulting in varied levels of inhibition. Currently, KB-R7943’s mechanism(s) of action is still unknown. However, Iwamoto et al. (138) have proposed a potential binding site or region for KB-R7943. Considering Iwamoto et al. had shown NCX3 to be three fold more sensitive to KB-R7943 than for either NCX1 or NCX2, they chose to further examine this result (326). Using chimeras of NCX1 and NCX3, with site-directed mutagenesis, they demonstrated that the highly conserved re-entrant membrane loop, Calx-α (α-2) repeat, is responsible for differences in KB-R7943’s response among isoforms (138). It was reported that the single residue
substitutions V820G, Q826V, A809V and A890I were important for drug sensitivity. Further testing through cysteine-scanning mutagenesis of the Calx-a (α-2) repeat revealed that a single residue Gly833 caused a ≥30 fold reduction in drug sensitivity. However, substituting Gly833 to Threonine produced an even larger difference (138). This region was proposed to form part of the receptor for KB-R7943. In addition, KB-R7943 was suggested to exert its effects from the extracellular surface, as 0.8 μM KB-R7943 significantly inhibited whole-cell outward exchange, but 30 μM applied intracellularly had minimal effect (138). Although the α-2 repeat (implicated in KB-R7943 inhibition) is accessible from the intracellular side, while inhibition is induced upon extracellular application, a conformational change of the exchanger could account for this result (138; 166). Results obtained using inside-out giant, excised patches from Xenopus laevis oocytes expressing NCX1.1, have demonstrated the opposite; strong inhibition with intracellular application (88). Furthermore, the initial report by Iwamoto, suggested that KB-R7943 works primarily on external sites but can inhibit the exchanger from both sides when applied intracellularly in cardiac sarcolemmal vesicles, smooth muscle cells, and CCL-39 fibroblasts (145). This was not the case in a following Iwamoto report using CCL-39 fibroblasts (138).

The capacity of KB-R7943 to inhibit Na⁺-Ca²⁺ exchange in several pathophysiological models was tested. Experiments performed on both cellular and whole organ models. Firstly, several studies have shown KB-R7943 to reduce or suppress various models of cardiac arrhythmia. Mukai et al. found that 10 μM KB-R7943 significantly decreased the incidence of reoxygenation-induced arrhythmias and shortened the duration of arrhythmias in guinea-pig papillary muscles (218). In the case
of ouabain-induced arrhythmias, Watano et al. reported KB-R7943 to suppress arrhythmias in isolated guinea pig whole atria and induce changes in the measured electrocardiogram (ECG) (325). Elias et al. implemented re-oxygenation-induced arrhythmias on intact Langendorff-perfused whole rabbit hearts and demonstrated a suppressive effect of 3 μM KB-R7943. The result was a marked attenuation of ventricular arrhythmia (both tachycardia and fibrillation) (88). In a study presenting negative effects of KB-R7943, Lu et al. employed anaesthetized rats (subjected to 5 min of coronary artery occlusion followed by 10 min of reperfusion) and found that pre-treatment with KB-R7943 did not decrease ischemia/reperfusion-induced ventricular arrhythmias (201). With the dog model, mixed results have been shown. While Miyata et al. found KB-R7943 to prevent atrial fibrillation-induced cell shortening (217), Miyamoto et al. demonstrated no suppression of induced arrhythmias (216). In general, positive results have been associated with KB-R7943 treatment in various arrhythmia-inducing models.

Reperfusion injury models (i.e., ischemia / reperfusion) have also been applied in the study of KB-R7943 and Na+–Ca2+ exchange. In 1998, Nakamura et al. studied the effects of KB-R7943 in a reperfusion model using isolated rat hearts (222). In normal hearts, KB-R7943 (1 and 10 μM) did not reduce contractile function. After ischemia-reperfusion, administration of KB-R7943 resulted in a significant recovery of left ventricular developed and end-diastolic pressure, either by pre-ischemic or post-ischemic treatment (222). Later, Ladilov et al. measured rat cardiomyocytes with simulated ischemia. They found 20 μM KB-R7943 completely inhibited reverse mode exchange, while treatment with the same concentration during anoxia did not influence the onset of
rigor contracture but significantly reduced the cytosolic accumulation of Ca\textsuperscript{2+} and Na\textsuperscript{+}. However, upon re-oxygenation, myocytes developed hypercontracture that was significantly reduced by KB-R7943 administration (178). With the use of perfused rat hearts they found that 20 μM KB-R7943 (during anoxia) significantly attenuated reoxygenation-induced enzyme release (lactate dehydrogenase and creatine kinase) (178). Continuing, Satoh et al. used 5 μM KB-R7943 on rat cardiomyocytes, to selectively block Ca\textsuperscript{2+} entry through the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger (~90%) (281). KB-R7943 did not affect steady state twitches, Ca\textsuperscript{2+} transients, SR Ca\textsuperscript{2+} load or post-rest potentiation. It did, however, prolong the action potential plateau. They concluded that Ca\textsuperscript{2+} entry through the exchanger is not important for normal excitation-contraction coupling but can contribute to cardiac arrhythmogenesis (281). Lastly, a recent study using KB-R7943 in whole rat hearts showed considerably suppressed intracellular Ca\textsuperscript{2+} levels during an ischemic period. Following reperfusion, diastolic Ca\textsuperscript{2+} was normalized more rapidly in KB-R7943-treated hearts (291).

Nishida et al. chose a model for Ca\textsuperscript{2+} overload, produced by metabolic inhibition (by carbonyl cyanide m-chrolophenylhydrazone (CCCP)), in isolated guinea pig myocytes (234). When applied, 10 μM KB-R7943 reduced Ca\textsuperscript{2+} overload, whereas other ion transport inhibitors did not (234). Using guinea pig myocytes for a Ca\textsuperscript{2+} paradox model (a massive Ca\textsuperscript{2+} influx during a Ca\textsuperscript{2+} repletion phase following a period of time in the absence of Ca\textsuperscript{2+}), Iwamoto et al. reported no hypercontracture on Ca\textsuperscript{2+} repletion, using 10 μM KB-R7943 (145). Hydroxyl-induced Ca\textsuperscript{2+} overload through the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger was examined by Zeitz et al. (336). Their results supported a protective effect by KB-R7943, showing a preservation of diastolic function in both rabbit and rat.
myocardium (336). Also, in failing human myocytes, Ca\textsuperscript{2+} transients and contractions exhibit an SR-related, phasic component and a slow, reverse mode Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange-related tonic component (slow decay of the Ca\textsuperscript{2+} transient) (65). Dipla et al. demonstrated that KB-R7943 removed the action potential's tonic component and reduced the phasic component, in failing human ventricular myocytes (65).

In other non-cardiac related models, KB-R7943 has proven to be quite useful. For example, in CA1-neurons from rat hippocampal slices, KB-R7943 protected against hypoxic-hypoglycemic injury (289). In cultured primary osteoblasts, KB-R7943 prevented extracellular mineral deposition (304). Studies in the rat have used KB-R7943 in hepatocytes (134), forebrain neurons (28), cerebral cortex (255), and with acute renal failure (ARF) in kidney cells (175; 334).

The specificity of KB-R7943 over other ion translocation mechanisms and cellular proteins is important. Non-selective inhibitors (i.e., Ni\textsuperscript{2+}, amiloride, and DCB) must be applied with caution, as results can be ambiguous when applied in a natural setting. KB-R7943’s specificity has been assessed on several occasions. Iwamoto et al., in their original report stated that KB-R7943 inhibits voltage-sensitive Na\textsuperscript{+} channels, L-type Ca\textsuperscript{2+} channels, and inward rectifying K\textsuperscript{+} channels with concomitant IC\textsubscript{50} values of 14, 8, and 7 \textmu M (145). Iwamoto et al. also reported that KB-R7943 was without effect on the Na\textsuperscript{+}-Ca\textsuperscript{2+}-K\textsuperscript{+} exchanger, Na\textsuperscript{+}-H\textsuperscript{+} exchanger, SR Ca\textsuperscript{2+}-ATPase, SL Ca\textsuperscript{2+}-ATPase, and the Na\textsuperscript{+}-K\textsuperscript{+}-ATPase at concentrations as high as 30 \textmu M (145). KB-R7843 can block nicotinic acetylcholine receptors (256). Most recently, KB-R7943 was proposed to directly modulate Na\textsuperscript{+}-Mg\textsuperscript{2+} exchange in a Ca\textsuperscript{2+}-dependent manner. Using pig carotid artery smooth muscle, the result was an increase in Mg\textsuperscript{2+} (318).
Results presented on KB-R7943 are quite varied. This greasy compound has been reported as competitive, non-competitive and mixed-competitive. Potency (exchanger and mode selectivity) ranged from substantially low to very high, never quite acting as an ideal inhibitor. However varied these results may be, KB-R7943 has still proven to be a useful inhibitor and has given promising insight into the future of Na$^+$-Ca$^{2+}$ exchange pharmacology.

5.3.3. SEA0400

Recently, a new Na$^+$-Ca$^{2+}$ exchange inhibitor, SEA0400 (2-[4-[(2,5-difluorophenyl)methoxy]phenoxy]-5-ethoxyaniline) was introduced. Currently, there are only seven reported publications involving SEA0400. SEA0400 may be the most selective and potent Na$^+$-Ca$^{2+}$ exchanger inhibitor to date (137; 203; 204; 239; 310; 311). This inhibitor was isolated through the screening of a compound library, measuring Na$^+$-dependent Ca$^{2+}$ uptake in cardiac sarcolemmal vesicles and astrocytes (204). The ethoxyaniline derivative was initially tested and compared to KB-R7943 using cerebral tissue and cerebral pathology models (204). In the initial SEA0400 study by Matsuda et al., inhibition was measured in a cerebral ischemia-reperfusion injury (both in vitro and in vivo) model. SEA0400 showed significantly greater potency than KB-R7943 (IC$_{50}$ ~ 2-3 μM), inhibiting Na$^+$-dependent Ca$^{2+}$ uptake in neurons (IC$_{50}$ ~ 33 nM), astrocytes (IC$_{50}$ ~ 5.0 nM), and microglia (IC$_{50}$ ~ 8.3 nM) (204). Conversely, KB-R7943 produced IC$_{50}$ values of 3.8, 2.0, and 3.1 μM respectively. SEA0400 demonstrated only minor affinities for Ca$^{2+}$ channels, Na$^+$ channels, K$^+$ channels, norepinephrine transporter, and
14 other receptors (204). In addition, SEA0400 did not alter the activities of Na⁺-H⁺ exchange, Na⁺-K⁺-ATPase, Ca²⁺-ATPase and five enzymes (phospholipase A₂, phospholipase C, 5-lipoxygenase, iNOS and cNOS). As previously reported (4), KB-R7943 inhibited store-operated Ca²⁺ entry (SOCE) in cultured astrocytes yet SEA0400 did not until at very high concentrations (10 μM - where solubility may be questionable). SEA0400 dose-dependently attenuated; paradoxical Ca²⁺ induced reactive oxygen species (ROS) production, DNA ladder formation, and nuclear condensation in neonatal astrocytes (204). In vivo studies reported that SEA0400 reduced infarct volumes upon transient middle cerebral artery occlusion in rat cerebral cortex and striatum (204). As a result, Matsuda et al. concluded that SEA0400 was a specific and potent Na⁺-Ca²⁺ exchange inhibitor that has protective properties for cerebral ischemia (204). SEA0400 was then employed in the cardiac setting, where Tanaka et al. isolated ventricular myocytes from guinea pig hearts and measured the inhibition of Na⁺-Ca²⁺ exchange current (311). SEA0400 at 1 μM inhibited myocardial exchange current (~ 80%) to the same degree as 10 μM KB-R7943. At the same concentrations, KB-R7943 resulted in >50 % inhibition of Na⁺ channel currents, L-type Ca²⁺ channel currents, delayed and inwardly rectifying K⁺ channel currents, while SEA0400 (1 μM) had no significant effect (311). These results support SEA0400 as a potent and highly selective inhibitor of Na⁺-Ca²⁺ exchange.
Figure 8. Chemical structure of SEA0400, (2-[4-[2,5-difluorophenyl]methoxy]phenoxy]-5-ethoxyaniline (SEA0400)).
Exchange activity is difficult to measure accurately and therefore, must be analysed from a variety of angles. Various research groups apply unique methods and models of analysis, in the hopes of improving our understanding of Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange. One model recently published by the Philipson group implemented Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange knockout mice (NCX\textsuperscript{-/-}) to test the specificity of KB-R7943 and SEA0400 (274). Using heart tubes from fetal mouse embryos at 9.5 days post coitus (dpc) and fura-2 loading, they were able to measure the resulting Ca\textsuperscript{2+} transients. The study revealed that KB-R7943 (5 µM) and SEA0400 (0.1 µM) decrease Ca\textsuperscript{2+} transients in both control cells and cells from knockout mice (274). Another laboratory, the Shigekawa group, used heterozygous NCX1 knockout and wild-type mice to test various inhibitors. They concluded that their model was useful, providing low NCX1 expressing mice (137).

The effects of SEA0400 were also tested in an ischemia-reperfusion setting for the rat, rabbit and dog (in vivo and in vitro) (203; 310). The first study examined isolated Langendorff-perfused rabbit hearts exposed to regional ischemia and reperfusion (203). It was found that both KB-R7943 (1 µM) and SEA0400 (0.1 µM) reduced infarct size independent of the time of administration (before or after ischemia). However, SEA0400 was more effective, reducing the infarct by ~75 %, whereas, KB-R7943 was only 40 % effective. At concentrations ≥ 1 µM, KB-R7943 impaired cardiac function. However, SEA0400 showed a continuous recovery of function. These data stress that SEA0400 is significantly better at cardioprotection, even at a ~10 fold lower concentration. Another ischemia-reperfusion study in canine cardiac sarcolemmal vesicles and rat cardiomyocytes prolonged the trend for SEA0400’s positive results (310). Measuring Na\textsuperscript{+}-dependent \textsuperscript{45}Ca\textsuperscript{2+} uptake, they reported an SEA0400 IC\textsubscript{50} value of 90 nM.
(sarcolemmal vesicles) and 92 nM (rat cardiomyocytes). In vivo rat coronary ligation and reperfusion experiments showed a decrease in ventricular fibrillation and mortality with SEA0400. Furthermore, Takahashi et al. (310) were able to reduce Ca\(^{2+}\)-paradox-induced cell death in their study. The latest evidence supporting SEA0400 as having protective effects was reported by Ogata et al., using renal ischemia-reperfusion and hypoxia-reoxygenation injury in the rat model (239). It was demonstrated that SEA0400 could dose-dependently reduce ischemia-reperfusion induced renal dysfunction and tubular necrosis. Furthermore, in isolated cultured porcine tubular cells, application of SEA0400 before or after hypoxia-reoxygenation was found to protect against cellular damage (239).

It is evident that SEA0400 is an effective and useful compound. The majority of results confer that SEA0400 has great potential in the study of Na\(^{+}\)-Ca\(^{2+}\) exchange. A potent and mode selective inhibitor is precisely what is required in a field lacking pharmacological tools. This could prove useful in the pathophysiological studies of Na\(^{+}\)-Ca\(^{2+}\) exchange. Understanding its mechanism(s) of action, therefore, would be valuable information. This could lead to a better knowledge of Na\(^{+}\)-Ca\(^{2+}\) exchange function, but may also give rise to more potent and specific Na\(^{+}\)-Ca\(^{2+}\) exchange inhibitors. Therefore, presented in this thesis are results suggesting a potential mechanism of action for SEA0400.
6. **Na\(^+\)-Ca\(^{2+}\) Exchange Pathophysiology**

The physiological role of the cardiac Na\(^+\)-Ca\(^{2+}\) exchanger, NCX1.1, is not entirely clear. However, it is generally agreed upon that the exchanger's primarily role is transarcolemmal Ca\(^{2+}\) removal. Theoretically, Na\(^+\)-Ca\(^{2+}\) exchange should remove the exact amount of Ca\(^{2+}\) initially brought in through L-type Ca\(^{2+}\) channels thereby, balancing Ca\(^{2+}\) influx with efflux. Accurate Ca\(^{2+}\) balance is crucial for cellular and whole heart relaxation (19; 21; 36; 44; 237; 250). This process must occur on a beat-to-beat basis, otherwise Ca\(^{2+}\) overload or depletion will result. Cardiomyocytes are very sensitive to changes in intracellular Ca\(^{2+}\) and depend on the maintenance of proper Ca\(^{2+}\) homeostasis for survival. Calcium sensitivity paired with a reversible Na\(^+\)-Ca\(^{2+}\) exchanger can be dangerous for the heart and is, therefore, implicated in cardiac pathophysiology. Under a variety of conditions, the exchanger could contribute to specific types of cardiac disease and injury. Whether or not these conditions actually contribute to pathophysiology is not entirely evident. However, there is rapidly amassing evidence suggesting this possibility.

Whether Na\(^+\)-Ca\(^{2+}\) exchange function is normal, reversed, increased or decreased, there are a multitude of cellular properties that can be altered (both positive and negative). For example, cardiac action potentials (5; 61; 67; 186; 329), SR Ca\(^{2+}\) loading (153), Ca\(^{2+}\) transients and diastolic Ca\(^{2+}\) levels (30; 65; 161) are all affected by alterations in Na\(^+\)-Ca\(^{2+}\) exchange. Therefore, an understanding of Na\(^+\)-Ca\(^{2+}\) exchange activity and regulation, in both cardiac disease and injury, is essential if we wish to apply this to a clinical setting (i.e., new pharmaceuticals or surgical techniques). In heart failure, it is suggested that Na\(^+\)-Ca\(^{2+}\) exchanger protein, transcript, and activity levels are altered.
Certain reports find an increase in all three (protein, transcript, and activity) (95; 107; 108; 307). Other reports demonstrate minimal change (170) or even a decrease (64; 74) in exchanger levels. In certain cases where exchanger levels are upregulated, this is considered as a mechanism compensating for a decrease in other Ca\(^{2+}\) handling proteins, such as the SR Ca\(^{2+}\)-ATPase (197; 223; 238; 238). Recently, O’Rourke et al., inhibited SR function (Ca\(^{2+}\) uptake) with cyclopiazonic acid and found a greater requirement of Na\(^{+}\)-Ca\(^{2+}\) exchange for cytosolic Ca\(^{2+}\) removal. The Na\(^{+}\)-Ca\(^{2+}\) exchanger expression was increased by 104% (238). Meanwhile, Schillinger et al. demonstrated that poor diastolic function occurs in hearts with a decreased level of SR Ca\(^{2+}\)-ATPase but with unchanged Na\(^{+}\)-Ca\(^{2+}\) exchanger levels (109; 283). Conversely, some studies proposed that an increase in Na\(^{+}\)-Ca\(^{2+}\) exchange activity was deleterious to cardiac function (260; 266; 282).

In 2000, Schillinger et al. reported that overexpression of the Na\(^{+}\)-Ca\(^{2+}\) exchanger in rabbit cardiomyocytes (adenoviral gene transfer), resulted in a decrease in contractility (282). In summary, whether an increased, decreased or unchanged level of Na\(^{+}\)-Ca\(^{2+}\) exchangers actually occurs in heart failure remains to be clarified. Furthermore, understanding how the change in exchanger function affects contractile behaviour under such diseased conditions, would be important information to both the field of Na\(^{+}\)-Ca\(^{2+}\) exchange and cardiology.

Two major components of heart failure and cardiac disease are arrhythmias and reperfusion injury. Arrhythmias are an irregular heart rhythm, or an abnormality in heart beat pattern, potentially resulting in tachychardia and then fibrillation. The primary causes of arrhythmia are non-re-entrant mechanisms created by abnormal automaticity in
the heart (236; 258). This is also known as triggered arrhythmogenesis. It was shown by Blaustein et al. that an induced increase in intracellular Ca\(^{2+}\) could influence SR Ca\(^{2+}\) stores. A slight increase in cytosolic Ca\(^{2+}\) can lead to a substantial increase in SR uptake, allowing this Ca\(^{2+}\) to participate in the next release (35). In the heart, if the Ca\(^{2+}\) gain becomes very large, the SR can release Ca\(^{2+}\) spontaneously. This is known as SR Ca\(^{2+}\) overload (153). Ca\(^{2+}\) overload is suggested to be a result of an increase in Ca\(^{2+}\) due to reverse mode Na\(^+-\)Ca\(^{2+}\) exchange, and/or the reduced drive of forward mode Na\(^+-\)Ca\(^{2+}\) exchange (by elevated intracellular Na\(^{+}\) levels). This increase in Ca\(^{2+}\) is still accompanied by normal Ca\(^{2+}\) influx through L-type Ca\(^{2+}\) channels. Ultimately, this leads to a “trigger sensitive” SR, instigating spontaneous release of SR Ca\(^{2+}\) stores, creating a wave of Ca\(^{2+}\), that in turn produces a transient inward current (I\(\text{di}\)). Transient inward current occurs by two mechanisms. The first is through intracellular Ca\(^{2+}\) activated non-selective channels (152; 154). The second mechanism is through Na\(^+-\)Ca\(^{2+}\) exchange (94; 195; 196; 259; 284; 337). While the non-selective channels bring ions into the cell, the Na\(^+-\)Ca\(^{2+}\) exchanger (forward mode) removes Ca\(^{2+}\) concomitantly bringing Na\(^{+}\) into the cell, ultimately depolarizing the membrane. Depolarization, if strong enough, can become an all-or-none response and lead to a premature contraction. This depolarizing I\(\text{di}\) is directly related to triggered arrhythmias through events called early afterdepolarizations (EAD) and delayed afterdepolarizations (DAD) (152; 154; 235). EAD’s occur during a prolongation of the cardiac action potential plateau in which the resulting inward transient current is sufficient for depolarization, resulting in a contraction. A second depolarization prior to the completion of the original depolarization results in an increased Ca\(^{2+}\) load, which could produce DAD’s (308; 323).
The heart is a highly aerobic muscle, requiring a constant supply of oxygen that is fed through a distinct set of vasculature. Deoxygenated blood enters the lungs, reoxygenates and returns to the heart. Ultimately, the heart’s constant pumping is directly related to insuring its own oxygen supply. When blood flow is reduced or completely occluded, the heart is subjected to a period of ischemia (low or no blood flow) or anoxia (absence of oxygen). Upon reperfusion, the heart undergoes large, rapid ionic shifts. Ultimately, this has been attributed to an intracellular rise in Ca\(^{2+}\) (3; 80). As previously mentioned, injury also results from a sudden rise in Ca\(^{2+}\) when the cells are exposed to Ca\(^{2+}\) following a “Ca\(^{2+}\)-free” period. This is termed “Ca\(^{2+}\) paradox”. Reverse mode Na\(^{+}\)-Ca\(^{2+}\) exchange contributes to the “Ca\(^{2+}\) overload” associated with ischemia-reperfusion injury (104; 220; 312). As well, during ischemia, an internal acidification occurs as a result of H\(^{+}\) ion production as the cell switches to anaerobic metabolism (glycolysis) (3). ATP is degraded, resulting in the inhibition of the Na\(^{+}\) pump (Na\(^{+}\)-K\(^{+}\)-ATPase). This in turn, increases intracellular Na\(^{+}\) levels. Upon reperfusion, the Na\(^{+}\)-H\(^{+}\) exchanger attempts to restore intracellular pH levels by removing H\(^{+}\) ions, and therefore brings Na\(^{+}\) into the cell (180). With such a large rise in intracellular Na\(^{+}\), forward mode Na\(^{+}\)-Ca\(^{2+}\) exchange becomes less favourable. Conversely, the reverse mode becomes more favourable. This not only leads to a rise in Ca\(^{2+}\) but also a decrease in Ca\(^{2+}\) removal. Eventually, this leads to Ca\(^{2+}\) overload and may seriously damage the cell (104; 220; 312). With respect to roles for the exchanger in physiology and pathophysiology, there is sufficient disagreement to merit further investigation.
III. METHODS

1. Preparation of Xenopus laevis Oocytes

*Xenopus laevis* oocytes are large and robust single-celled biological hosts. Furthermore, they have a large membrane surface area, express the protein and localize exchangers to the membrane. As current produced by a single exchanger is small, a large exchanger population is required to create a measurable signal. Therefore, using this technique allows us to excise a large surface area and increase our experimental exchanger population. Giant, excised patch clamping also allows for rapid cytoplasmic access, which is difficult or absent in other techniques.

*Xenopus laevis* were anaesthetized in a solution of 250 mg/l ethyl p-aminobenzoate (Sigma) in deionized ice-water for 30 minutes prior to sacrificing. Oocytes were surgically removed and then washed in Solution A containing (in mM): 88 NaCl, 15 HEPES, 2.4 NaHCO₃, 1.0 KCl, 0.82 MgSO₄; pH 7.6 at room temperature (RT). Connective follicle(s) surrounding the oocytes was teased apart to assist digestion. Continuing, the oocytes were transferred into Solution A containing ≈ 16,000 units of collagenase (Type II; Worthington). Next, the same batch of oocytes was subjected to an incubation period accompanied by gentle agitation at RT for 1.5-2 hours. This rotating motion helps clean the egg by removing excess follicle. Following the incubation period, the oocytes were washed several times in Solution B containing (in mM): 88 NaCl, 15 HEPES, 2.4 NaHCO₃, 1.0 KCl, 0.82 MgSO₄, 0.41 CaCl₂, 0.3 Ca(NO₃)₂, 1 mg/ml BSA (Fraction V; Sigma); pH 7.6 at RT, and then transferred to a RT 100 mM K₂HPO₄ (pH
6.5) solution, containing 1 mg/ml BSA. After incubation for 12-20 minutes (with gentle agitation at RT), the oocytes were washed in Solution B. Defolliculated stage V-VI oocytes were selected and incubated in glass scintillation vials at 18 °C filled with Solution B (without BSA) until the cRNA is injected.

2. Synthesis of NCX1.1 cRNAs

Hind III (New England Biolabs) was used to linearize complementary DNA (cDNA encoding NCX1.1) residing in pBluescript II SK(+) (Stratagene Inc.). Complementary RNA (cRNA) was synthesized using either mCAP mRNA capping kit (Stratagene Inc.) or mMessage mMachine (Ambion) transcription kit. Oocytes were injected with ≈ 5 ng/50 nl of cRNA encoding NCX1.1, and stored in Solution B (without BSA) at 18 °C. Electrophysiological measurements of Na⁺-Ca²⁺ exchange activity were obtained 3-7 days post-injection.

3. Preparation of Mutant Na⁺-Ca²⁺ Exchangers

Mutations were generated using the Sculptor in vitro mutagenesis kit (Amersham Corp.) as previously described (82; 209; 225; 226). Both large and site-specific amino acid deletions were created for this study. The large deletions were made on the C-terminus end of the XIP region, while the site-specific amino acid mutations were created throughout. These mutations were used to determine the involvement of both I₁ and I₂ regulatory mechanisms in NCX1.1 current inhibition by SEA0400. A previous study by
He et al. looked at the effects of mutant, exogenous XIP peptides and their inhibitory consequences (113). They noted that a potent XIP peptide inhibitor must be intact in order to exert its full inhibitory effect. They also revealed that specific amino acids were important for proper inhibitory function. For example, they noted that basic and aromatic residues were the most important for XIP’s inhibitory action. Similar results for the XIP region were found in the Drosophila melanogaster exchanger CALX1.1 (82). Therefore, basic and aromatic residues were the primary experimental targets, due to results from corresponding experiments involving amino acid substitutions, indicating the importance of these residues in exchanger inhibition (113). Basic residues were switched to neutral hydrophilic glutamine residues, while aromatic tyrosines were changed to threonine in order to remove the aromatic characteristic, while maintaining the residue’s hydroxyl characteristic. In one case, an aromatic residue, F223, located in a strongly conserved amino NH2-terminal of the XIP region, was mutated to glutamate so as to eliminate the aromatic group and introduce an acidic residue into a primarily basic exchanger region (209). All deletion and site-specific XIP region mutations produced functional exchangers. The mutations were categorized into three groups based on their corresponding regulatory phenotypes. Group 1, contained two site-specific mutants (K225Q, F223E) and exhibited accelerated Na+-dependent inactivation (I1). Group 2 mutants were comprised of one site-specific (K229Q) and one deletion mutant (Δ229-232) that exhibited suppressed Na+-dependent inactivation (I1). Lastly, the single Group 3 deletion mutant (Δ680-685) exhibited suppressed I2 regulation. Although not a XIP region mutant, Δ680-685 was previously shown to have altered Ca2+-dependent regulation (I2), in both giant, excised patches from Xenopus laevis oocytes and whole cell
patches from mouse cardiomyocytes (212). These distinct mutant phenotypes are presented in Figure 18 and ultimately allow an assessment of SEA0400's inhibitory effect on exchanger's with altered ionic regulatory mechanisms (i.e., I₁ and I₂).

4. Measurement of Exchange Activity

Na⁺-Ca²⁺ exchange currents (outward and inward) were measured using the giant, excised patch clamp technique as previously described (116) (Figure 9). *Xenopus* oocytes were placed in a hypo-osmotic solution containing (in mM): 250 Sucrose, 67 KCl, 10 D-Glucose, 10 HEPES, 5 MgCl₂, 2.5 EGTA; pH 7.0 at RT (with KOH) for ≈ 15 minutes, to assist in removal of the vitellin layer surrounding the oocyte membrane. After vitellin layer dissection using blunt forceps, oocytes were placed in a solution containing: (values in mM): 100 KOH, 100 MES, 20 HEPES, 5 EGTA, 5 MgCl₂; pH 7.0 at RT (with MES). Borosilicate glass pipettes were pulled and polished to a final inner diameter of ≈ 20 - 30 µm and coated with a Parafilm™:Mineral oil mixture to enhance patch stability and reduce electrical noise. Pipettes were filled with the appropriate pipette solution for desired current, secured tightly inside the pipette holder (Axon Instruments), and positioned onto the animal pole surface (dark surface) of the oocyte by way of a micromanipulator (Narshige). Gigaohm seals were formed via gentle suction. Membrane patches (inside-out configuration) were excised through progressive horizontal and vertical pipette tip movements. Pipettes containing excised patches were then relocated to a carved divot in front of the perfusion tip. This pipette was connected
to a computer-controlled, 20-channel solution-switching machine, which allowed rapid 
(i.e., \(\approx 200 \text{ ms}\)) solution changes of the bath (i.e., intracellular) solution.
Inject cRNA

Form GΩ Seal

Excise Patch

**Figure 9. The Giant, Excised Patch Technique.** Complimentary RNA (cRNA) encoding cardiac Na\(^+\)-Ca\(^{2+}\) exchanger protein (NCX1.1) was injected into *Xenopus laevis* oocytes (*left panel*). After 3 – 7 days, allowing time for protein expression, gigaohm (GΩ) seals were formed by gentle suction of the pipette tip onto the egg surface (*middle panel*). Excision of the membrane patch (inside-out configuration, i.e., the cytoplasmic surface is exposed to the outside) was achieved by small mechanical manipulations of the pipette tip (*right panel*).
For outward (i.e., reverse mode) Na\textsuperscript+-Ca\textsuperscript{2+} exchange currents, pipette (extracellular) solution contained (in mM): 100 NMG-MES, 30 HEPES, 30 TEA-OH, 16 sulfamic acid, 8.0 CaCO\textsubscript{3}, 6 KOH, 0.25 ouabain, 0.1 niflumic acid, 0.1 flufenamic acid; pH 7.0 at RT with MES. Outward currents (Figure 10) were initiated by rapidly switching from a Li\textsuperscript{+} to a Na\textsuperscript{+}-based bath solution containing (in mM): 100 [Na\textsuperscript{+} + Li\textsuperscript{+}]-aspartate, 20 CsOH, 20 MOPS, 20 TEA-OH, 10 EGTA, 0 - 9.91 CaCO\textsubscript{3}, 1.0 - 1.5 Mg(OH)\textsubscript{2}; pH 7.0 at 30\textdegree C (with MES or LiOH).
Figure 10. *Initiation of a Typical Outward (i.e., Reverse Mode) Na⁺-Ca²⁺ Exchange Current.* For outward exchange, pipettes contain 8 mM Ca²⁺ (*left panel*). The pipette was then perfused with 100 mM Na⁺ and 1 μM Ca²⁺, initiating Na⁺-Ca²⁺ exchange. As exchange occurs, 3 Na⁺ ions enter the pipette in exchange for 1 Ca²⁺ ion. A net single positive charge associated with Na⁺ translocation, moves across the membrane into the pipette (analogous to leaving the cell) and generates an *outward* current. The *right panel* shows a typical outward (*i.e.*, reverse mode) Na⁺-Ca²⁺ exchange trace, recorded using the giant, excised patch technique.
For inward (Figure 11) Na\(^+\)-Ca\(^{2+}\) exchange current measurements, the pipette (extracellular) solution contained (in mM): 100 Na-MES, 20 CsOH, 20 TEA-OH, 10 EGTA, 10 HEPES, 8 sulfamic acid, 4 Mg(OH)\(_2\), 0.25 ouabain, 0.1 niflumic acid, 0.1 flufenamic acid; pH 7.0 at R.T. with MES. Inward currents were activated by switching between Ca\(^{2+}\)-free and Ca\(^{2+}\)-containing, Li\(^+\)-based bath solutions, as described above.

For combined inward-outward current measurements (on the same patch), pipettes contained (in mM): 100 Na-MES, 20 CsOH, 20 HEPES, 20 TEA-OH, 4 sulfamic acid, 2 CaCO\(_3\), 0.25 ouabain, 0.1 niflumic acid, 0.1 flufenamic acid; pH 7.0 at RT (with MES). Outward and inward currents were activated using the same solutions as those used for initiating pure outward and pure inward currents, as described above.
Figure 11. *Initiation of a Typical Inward (i.e. Forward Mode) Na⁺-Ca²⁺ Exchange Current.* Here, pipettes contain 100 mM Na⁺ (*left panel*). The pipettes were perfused in 100 mM Li⁺ and 10 μM Ca²⁺ initiating inward Na⁺-Ca²⁺ exchange. As exchange occurs 1 Ca²⁺ ion is brought into the pipette in exchange for 3 Na⁺ ions. Therefore, the single positive charge associated with Na⁺ translocation moves out of the pipette (analogous to entering the cell), consequently generating an *inward* current. The *right panel* shows a typical inward (*i.e.* forward mode) Na⁺-Ca²⁺ exchange trace, recorded using the giant, excised patch technique.
Axon Instruments hardware (Axopatch 200a) and software (Axotape) were used for data acquisition and analysis. Origin software was used for curve-fitting (e.g. IC$_{50}$ values) and statistical analyses. Pooled data are presented as mean ± SEM. Two-tailed Student’s t-tests were used for comparison of unpaired data, and $P < 0.05$ was considered significant. Free Mg$^{2+}$ and Ca$^{2+}$ concentrations were calculated using MAXC software (25). Unless indicated otherwise, a holding potential of 0 mV was implemented for current measurements. All experiments were conducted at 30 ± 1°C.

SEA0400 was prepared by dissolving in DMSO as a 20-40 mM stock and further diluted directly into bath solutions. With each drug concentration change, a time period of at least 32 seconds was allowed to lapse prior to re-examining current levels. The concentration of DMSO never exceeded 0.075% and was without effect on inward or outward Na$^+$-Ca$^{2+}$ exchange current characteristics. The solubility of SEA0400 in water is 0.03 μg/ml. In these studies however, SEA0400 was dissolved in pure DMSO, and further diluted into other experimental (buffered) solutions. The highest SEA0400 concentration used in any experimental setting was 3 μM. Taisho Pharmaceutical Co. Ltd. generously provided SEA0400.
IV. RESULTS

1. SEA0400 and Na⁺-Ca²⁺ Exchange Currents

The effects of SEA0400 on both outward (reverse mode) and inward (forward mode) exchange current were measured in giant, excised membrane patches from *Xenopus laevis* oocytes expressing the canine cardiac exchanger (NCX1.1). To determine mode selectivity of SEA0400, combined outward-inward exchange currents were recorded in the absence and presence of 0.1 μM SEA0400 (Figure 12A). Both modes (inward and outward) were recorded on the same membrane patch, containing the same population of exchangers. For outward-inward current recordings, pipettes contained 2 mM Ca²⁺ and 100 mM Na⁺. Outward or reverse mode exchange currents were initiated by applying 100 mM Na⁺ to the cytoplasmic patch surface with 1 μM regulatory Ca²⁺. Inward currents were activated by switching from a 100 mM Li⁺ and 1 μM Ca²⁺ solution to a 100 mM Li⁺ and 10 μM Ca²⁺ solution (in the absence of Na⁺). Control outward and inward Na⁺-Ca²⁺ exchange currents exhibited typical characteristics. That is, outward currents displayed their usual rise to a peak, followed by a progressive decay (a result of Na⁺-dependent inactivation (Iᵢ)) to a steady state level. Inward (forward mode) currents, however, showed no inactivation (Iᵢ) and appear as square waveforms. When 0.1 μM SEA0400 was applied, the resulting effect was a substantial inhibition of outward Na⁺-Ca²⁺ exchange current.
Figure 12. The Inhibitory Effects of SEA0400 on Combined Inward-Outward Exchange Activity. Figure A (top) shows a typical NCX1.1 mediated exchange current recording from an excised membrane patch measuring inward and outward exchange. The pipette contained 2 mM Ca\(^{2+}\) and 100 mM Na\(^{+}\). Outward currents were initiated by applying 100 mM Na\(^{+}\) to the cytoplasmic surface of the patch with 1 μM regulatory Ca\(^{2+}\). Inward currents were initiated by applying 100 mM Li\(^{+}\) and 10 μM Ca\(^{2+}\) to the cytoplasmic surface. Currents were recorded before and after the application of 0.1 μM SEA0400. Figure 2B shows the % inhibition of peak and steady state for both inward and outward currents, in the presence of 0.3 μM SEA0400.
Results demonstrated minimal inhibition of the inward Na\(^{+}\)-Ca\(^{2+}\) exchange currents, with 0.1 μM SEA0400. Shown in Figure 12B, are the pooled results of 0.3 μM SEA0400 on inhibition of peak and steady state currents for both outward and inwards currents. For outward currents, 0.3 μM SEA0400 reduced peak and steady state currents by 61 ± 1% and 87 ± 4% respectively, whereas inward currents were inhibited by 29 ± 4%. Therefore, these data show a preferential inhibition of outward (reverse mode) Na\(^{+}\)-Ca\(^{2+}\) exchange by SEA0400, under bi-directional conditions.

Once SEA0400’s reverse mode exchange selectivity had been established, an evaluation of inhibitory potency on Na\(^{+}\)-Ca\(^{2+}\) exchange was performed. In this case, measurements were conducted under unidirectional conditions, where only a single transport mode could occur. Figure 13A illustrates the inhibitory effects of SEA0400 on a representative outward Na\(^{+}\)-Ca\(^{2+}\) exchange current. Switching from a bath solution (cytoplasmic surface) containing 100 mM Li\(^{+}\) to 100 mM Na\(^{+}\) activated the outward Na\(^{+}\)-Ca\(^{2+}\) exchange currents. As required for exchanger activation, 1 μM Ca\(^{2+}\) (regulatory Ca\(^{2+}\)) was constantly present during recording, while pipette Ca\(^{2+}\) was maintained at 8 mM (transport Ca\(^{2+}\)). Application of 0.1 and 1 μM SEA0400 to the cytoplasmic surface of the patch resulted in a dose-dependent inhibition of both peak and steady state outward Na\(^{+}\)-Ca\(^{2+}\) exchange currents. Figure 13B represents the pooled data of SEA0400 over a range of concentrations, further illustrating SEA0400’s concentration dependent effect.

To establish this range of values, SEA0400 was applied in continuous increments and the resulting inhibition was recorded. These increments were used primarily because a satisfactory washout of SEA0400 could not be obtained. In both typical traces and pooled values, SEA0400 appeared to have a greater inhibitory potency (IC\(_{50}\)) for steady
state currents than for peak currents, with IC$_{50}$ values of 23 ± 4 nM and 78 ± 15 nM, respectively.
Figure 13. Concentration-Dependent Inhibition of Wild-Type NCX1.1 Outward Exchange Current with SEA0400. Panel A (top) shows typical outward Na⁺-Ca²⁺ exchange currents from an excised membrane patch in the absence and presence of different SEA0400 (0, 0.1, 1 μM) concentrations. Panel B (bottom) shows the SEA0400 concentration dependency for inhibition of outward Na⁺-Ca²⁺ exchange currents (peak and steady state). Pooled data show means ± S.E. obtained from 3-13 patches at each individual SEA0400 concentration.
2. **SEA0400 and Na\(^+\)-Ca\(^{2+}\) Exchange Regulatory Mechanisms (I\(_1\) and I\(_2\))**

To investigate SEA0400's inhibitory mechanism(s), experiments were conducted on outward Na\(^+\)-Ca\(^{2+}\) exchange currents in the absence of any primary-structure or protein based regulatory mechanisms (i.e., I\(_1\) and I\(_2\)). Membrane patches were deregulated by treatment with 1 mg/ml \(\alpha\)-chymotrypsin for ~1-2 minutes. This limited proteolysis resulted in a loss of both Na\(^+\)-dependent inactivation (I\(_1\)) and Ca\(^{2+}\)-dependent regulation (I\(_2\)), leaving the exchanger in a fully activated state (116). The exchanger displayed a square waveform and no longer required cytoplasmic Ca\(^{2+}\) (regulatory Ca\(^{2+}\)) for activation. Figure 14A displays a representative outward current following \(\alpha\)-chymotrypsin treatment, in the absence and presence of 0.1 and 1.0 \(\mu\)M SEA0400. Compared to intact Na\(^+\)-Ca\(^{2+}\) exchangers, the level of inhibition was notably reduced, even at a 10-fold increase in concentration. Figure 14B shows the pooled values for percent inhibition of SEA0400 following proteolytic treatment. A complete inhibitory profile of SEA0400 could not be obtained, because the compound began to precipitate out of solution at concentrations above 1-3 \(\mu\)M. However, an estimated IC\(_{50}\) value of 1.2 ± 0.6 \(\mu\)M is given. Regardless of limited solubility, these results suggested that SEA0400 requires intact regulatory mechanisms for a complete inhibitory effect.
Figure 14. Effects of SEA0400 on Outward Exchange Currents After Proteolytic Treatment. Panel A (top) shows typical outward Na\(^+\)-Ca\(^{2+}\) exchange currents from an excised membrane patch that was deregulated by the cytoplasmic application of 1 mg/ml \(\alpha\)-chymotrypsin for \(\sim\) 1 minute. Currents are shown in the absence and presence of 0, 0.1, and 1.0 \(\mu\)M SEA0400. Pooled results are shown in panel B (bottom), where each point represents the mean \(\pm\) S.E obtained from 3-12 patches.
3. **Effect of SEA0400 with Increasing Regulatory Ca²⁺**

Increases in regulatory Ca²⁺ can stimulate outward Na⁺-Ca²⁺ exchange activity (116; 121). Activation occurs only when Ca²⁺ is applied to the cytoplasmic surface of the patch. This mechanism is referred to as Ca²⁺-dependent regulation, or I₂ (121). Inward Na⁺-Ca²⁺ exchange currents do not exhibit this type of behaviour for the regulatory Ca²⁺ binding site is saturated (K_d ~ 0.3 μM) under inward current recording conditions (210). Therefore, it was suggested that during outward (reverse mode) Na⁺-Ca²⁺ exchange, SEA0400 might interact with varying regulatory Ca²⁺ levels. Results between SEA0400 and Ca²⁺ would be more pronounced during outward Na⁺-Ca²⁺ exchange, as regulatory Ca²⁺ levels are low (i.e., 1 μM) compared to those for inward current (i.e., 10 μM). Figure 15A shows the effect of regulatory Ca²⁺ concentrations on outward Na⁺-Ca²⁺ exchange currents, in the absence and presence of 1 μM SEA0400. Ca²⁺-dependent regulation (I₂) is visible during the control trace, producing a square waveform and essentially eliminates Na⁺-dependent inactivation (I₁) at 10 μM Ca²⁺. However, at the lower regulatory Ca²⁺ concentration of 1 μM, Na⁺-dependent inactivation (I₁) is present. Upon application of 1 μM SEA0400, Na⁺-Ca²⁺ exchange currents are inhibited at both 1 and 10 μM Ca²⁺. Compared to inhibition at 1 μM Ca²⁺, SEA0400’s effects are reduced at 10 μM Ca²⁺. This result is most prominent for peak currents. Furthermore, the elimination of Na⁺-dependent inactivation (I₁) at 10 μM Ca²⁺ is absent in the presence of SEA0400. The effects of SEA0400 and regulatory Ca²⁺ on F_ss values (ratio of steady state to peak currents) are shown in Figure 15B (pooled data). These data suggest that when in the presence of SEA0400, 10 μM Ca²⁺ is unable to increase F_ss values as it does
in control recordings. Moreover, these results suggest a potential interaction between SEA0400 and the Na$^+$-dependent inactivation (I$_I$) regulatory mechanism.
Figure 15. Effects of SEA0400 with Increasing Regulatory Ca\(^{2+}\) Concentrations.

Panel A (top) shows the effects of regulatory Ca\(^{2+}\) on typical outward Na\(^{+}\)-Ca\(^{2+}\) exchange currents before and after treatment with 1 μM SEA0400. Currents were initiated by applying 100 mM Na\(^{+}\) to the cytoplasmic surface of the patch with either 1 or 10 μM regulatory Ca\(^{2+}\) present. Panel B (bottom) shows pooled data (mean ± S.E., n = 3-43 patches) illustrating the effect of regulatory Ca\(^{2+}\) on F\(_{SS}\) (ratio of steady state to peak current) for control and SEA0400 treated patches.
4. **SEA0400 and Exchanger Recovery**

To examine the hypothesis that SEA0400 interacts with Na\(^+\)-dependent inactivation (I\(_I\)), paired-pulse experiments were performed to determine exchanger rate of recovery from I\(_I\). Outward currents were initiated through the cytoplasmic application of 100 mM Na\(^+\) with 1 \(\mu\)M Ca\(^{2+}\). Following exchanger activation, an inter-pulse was initiated 6.4 seconds into the recovery period, thereby stimulating outward exchange (Figure 16A). The paired-pulse experiments were conducted in the presence and absence of 0.1 \(\mu\)M SEA0400, over a range of time intervals and regulatory Ca\(^{2+}\) concentrations. Figure 16B demonstrates SEA0400’s ability to reduce the rate of recovery for peak outward currents at an increased regulatory Ca\(^{2+}\) level of 3 \(\mu\)M. In addition, recovery rates measured over a wide range of regulatory Ca\(^{2+}\) values supports the contention that SEA0400 can interfere with exchanger recovery from I\(_I\), as is shown in Figure 16C. In Na\(^+\)-Ca\(^{2+}\) exchangers with full regulatory function, increased regulatory Ca\(^{2+}\) significantly augmented the rate of recovery from I\(_I\). This recovery is strongly attenuated by the presence of SEA0400. These results suggest that SEA0400 essentially stabilizes or promotes the entry of the exchanger into its Na\(^+\)-dependent inactive state (I\(_I\)).
Figure 16. Effects of SEA0400 on Exchanger Recovery. Panel A (top) shows the effects of SEA0400 on representative paired-pulse recovery experiments where an interpulse interval of 6.4 sec is given. Outward Na⁺-Ca²⁺ exchange currents were initiated by applying 100 mM Na⁺ and 1 µM regulatory Ca²⁺ to the cytoplasmic surface of the patch. Pooled data are shown in panel B (bottom left) for results obtained from 6-7 patches (means ± S.E.) over a range of interpulse intervals (0.7-32 sec) to determine recovery rates. This data was used to generate panel C (bottom right), which shows recovery rates over a range of Ca²⁺ concentrations, in the presence and absence of 0.1 µM SEA0400. Points in panel C (bottom right) represents pooled data from 3-10 patches.
5. **Effect of SEA0400 on Inward Na\(^{+}\)-Ca\(^{2+}\) Exchange**

To further analyze SEA0400's mechanism of action, the compound was applied to intact and deregulated (1 mg/ml α-chymotrypsin) Na\(^{+}\)-Ca\(^{2+}\) exchangers and inward currents were measured (Figure 17A). Inward currents were initiated by applying 100 mM Li\(^{+}\) with 10 μM Ca\(^{2+}\) to the cytoplasmic membrane surface, in exchange for 100 mM Na\(^{+}\) (pipette). The inhibitory potency of SEA0400 on intact exchangers exhibiting forward mode exchange (inward current) is substantially less than that of reverse mode exchange (outward current). Consequently, the inhibitory efficacy of SEA0400 was further attenuated in deregulated exchangers (even at 3 μM SEA0400). As mentioned previously, SEA0400 demonstrated poor solubility at higher concentrations (> 1 μM); therefore it was difficult to assess an IC\(_{50}\) value for inward current. However an IC\(_{50}\) of ~8 μM was estimated for inward Na\(^{+}\)-Ca\(^{2+}\) current in intact patches.
Figure 17. *Effects of SEA0400 on Inward Exchange Currents After Proteolytic Treatment.* Panel A (top) shows the effects of SEA0400 for typical inward Na⁺-Ca²⁺ exchange currents before and after limited patch proteolysis with 1 mg/ml α-chymotrypsin, which deregulates the exchanger. In panel B (bottom), pooled data show mean values ± S.E. obtained from 3-15 patches.
6. **NCX1.1 Mutant Exchangers**

To test the hypothesis that SEA0400 stabilizes or facilitates entry of the exchanger into I₁, the effects of SEA0400 were tested on mutant exchangers exhibiting altered I₁ and I₂ regulatory characteristics. These three distinctive mutant exchanger groups, shown in Figures 18A, 18B and 18C, were previously characterized by the Philipson group in 1997 (209). Figure 18A illustrates Group 1 mutants (F223E and K225Q). Both of these mutants exhibited an accelerated rate of inactivation for outward current, which is measured as λ (discussed later) but differed in their extents of I₁. The extent of I₁ is measured by a ratio of steady state to peak current and is known as an $F_{ss}$ value. The F223E mutant displayed an increase in I₁, whereas K225Q demonstrated a decrease in the extent of I₁, as compared to wild-type NCX1.1. In the Group 2 mutants (K229Q and Δ229-232), the extent of I₁ inactivation is greatly diminished compared to wild-type NCX1.1. Figure 18B shows outward currents of both Δ229-232 and K229Q, which display square waveforms, suggesting a lack of Na⁺-dependent inactivation (I₁).

The last mutant group, Group 3, contains the Δ680-685 mutation shown in Figure 18C. This mutant appears to have a greatly reduced I₂ regulatory mechanism. Illustrated in Figure 18C, are the resulting outward currents at 1 and 10 μM regulatory Ca²⁺, that suggest a relative absence of both ionic regulatory mechanisms (I₁ and I₂). For the Δ680-685 mutant, there appears to be a decrease in current magnitude at 10 μM regulatory Ca²⁺. This is most likely due to competition between intracellular Na⁺ and Ca²⁺ for the exchanger transport sites as a result of the higher Ca²⁺ concentration.
Figure 18. *Electrophysiological Profiles of Wild-Type and NCX1.1 Mutants.* This figure illustrates the three distinct groups of mutant NCX1.1 exchangers used to investigate the inhibitory mechanism of SEA0400 at 1 and 10 μM Ca$^{2+}$. Panel A (*top*) displays two mutant exchangers with increased Na$^{+}$-dependent inactivation ($I_1$). Panel B (*middle*) displays two mutants that have suppressed Na$^{+}$-dependent inactivation ($I_1$). Lastly, Panel C (*bottom*) displays a single mutant with altered Ca$^{2+}$-dependent regulation ($I_2$). Currents were initiated by the application of 100 mM Na$^+$ with either 1 or 10 μM regulatory Ca$^{2+}$. 
7. Group 1 NCX1.1 Mutants (Outward Current)

From previous results, it was suggested that SEA0400 interacts with the Na$^+$-dependent inactivation mechanism, either by stabilizing I$_1$ in the presence of high Ca$^{2+}$, or by promoting the entry of the exchanger into an inactive state. Furthermore, SEA0400 only exerted its effects strongly when I$_1$ was present (i.e., reverse mode). However, in a deregulated setting (i.e., α-chymotrypsin), where I$_1$ is absent, SEA0400 shows greatly reduced effects. Furthermore, for inward currents (forward mode), I$_1$ is absent and so is the inhibitory effect of SEA0400. These results suggested an interaction between SEA0400 and the exchanger’s ionic regulatory mechanisms. Mutants displaying increased Na$^+$-dependent inactivation were tested in the presence of SEA0400. If a SEA0400-1$_1$ interaction existed, then perhaps SEA0400 could inhibit to a greater degree, when the extent of I$_1$ was increased. Figure 19A shows the effect of 0.1 μM SEA0400 on outward Na$^+$-Ca$^{2+}$ exchange currents for mutant K225Q, where the rate of inactivation was increased, but the extent of I$_1$ inactivation (F$_{ss}$) was decreased or comparable to wild-type values. In this mutant, the inhibitory effects of SEA0400 on peak and steady state outward currents were similar to that of wild-type at both 1 and 10 μM regulatory Ca$^{2+}$ and in the presence or absence of SEA0400 (Table 3). Demonstrated in Figure 19B, however, is the second Group 1 mutant, F223E. Prior to SEA0400 application, this mutant displays a clearly accelerated current decay, coupled with an increase in the extent of I$_1$. Table 2 shows that the extent of I$_1$ inactivation (F$_{ss}$) is greater at 1 μM Ca$^{2+}$ than at 10 μM Ca$^{2+}$. Upon SEA0400 application, there was a further increase in the extent of I$_1$, bringing steady state current levels near baseline. This however, made the calculation of
percent inhibition and F223E more difficult. In the presence of 1 and 10 μM Ca^{2+}, SEA0400 inhibited steady state currents by ~100% and 56 ± 4%, respectively. Peak currents for F223E were less affected by SEA0400’s presence. Compared to F223E, results of K225Q with or without SEA0400 may be a consequence of being a more subtle mutation than that of F223E.
Figure 19. Effects of SEA0400 on Outward Na\(^+-\)Ca\(^{2+}\) Exchange Currents of Mutant NCX1.1 Exchangers Exhibiting Enhanced I\(_{ij}\) Inactivation. Panels A (top) and B (bottom) illustrate the effects of 0.1 µM SEA0400 on exchange currents from K225Q (A) and F223E (B), in the presence of 1 and 10 µM regulatory Ca\(^{2+}\), respectively. Outward currents were initiated by applying 100 mM Na\(^{+}\) to the cytoplasmic surface of the patch with 1 or 10 µM Ca\(^{2+}\).
8. **Group 2 NCX1.1 Mutants (Outward Current)**

Experiments were conducted on mutants lacking Na\(^+\)-dependent inactivation to address the hypothesis that SEA0400 and I\(_1\) interact. If SEA0400 interacted as suggested, inhibition should be suppressed in these mutants (lacking Na\(^+\)-dependent inactivation (I\(_1\))). The Group 2 mutants, K229Q and \(\Delta 229-232\) both exhibited square current waveforms, indicating the absence of I\(_1\). However, Figure 20A shows prominent inhibition of both peak and steady state current (at 1 \(\mu\)M regulatory Ca\(^{2+}\)) upon application of 0.1 \(\mu\)M SEA0400 on K229Q outward exchange currents. Furthermore, as shown in the right panel (Figure 20A), the inhibitory effects of SEA0400 were attenuated upon switching to a higher regulatory Ca\(^{2+}\) value (i.e., 10 \(\mu\)M). As peak current inhibition at 1 and 10 \(\mu\)M Ca\(^{2+}\) remains relatively the same (26 ± 6 % and 22 ± 6 %, respectively), steady state current inhibition was significantly attenuated in the presence of 10 \(\mu\)M Ca\(^{2+}\). SEA0400 may induce I\(_1\)-like behaviour in K229Q, which can be alleviated by increasing regulatory Ca\(^{2+}\) concentrations. In addition, this indicates that the I\(_1\)-like inactivation induced by SEA0400 behaves in a similar manner to the native I\(_1\) inactivation process (with respect to regulatory Ca\(^{2+}\)), even though K229Q apparently lacks this form of regulation in SEA0400’s absence. The second Group 2 mutant, \(\Delta 229-232\) also shows a lack of I\(_1\) in the absence of SEA0400. However, unlike K229Q, outward exchange current from \(\Delta 229-232\) is minimally affected by 0.1 \(\mu\)M SEA0400 (Figure 20B). This slight inhibitory effect of SEA0400 was recorded at both high and low regulatory Ca\(^{2+}\) concentrations (i.e., 1 and 10 \(\mu\)M). These differences, may be a result of \(\Delta 229-232\) being a deletion mutant starting at the same point as K229Q. This
gross mutation (Δ229-232) may remove an entire portion of the XIP region involved in I1 formation. These results continue to support the hypothesis that SEA0400 interacts with I1; because when I1 is absent, SEA0400 is almost without effect.
Figure 20. **Effects of SEA0400 on Outward Na\(^+\)-Ca\(^{2+}\) Exchange Currents of Mutant NCX1.1 Exchangers Exhibiting Suppressed \(I_1\) Inactivation.** Panels A (top) and B (bottom) illustrate the effects of 0.1 μM SEA0400 on outward exchange currents mediated by Δ229-232 (A) and K229Q (B) in the presence of 1 μM (left) and 10 μM (right) regulatory Ca\(^{2+}\). Outward currents were initiated by rapidly applying 100 mM Na\(^+\) to the cytoplasmic surface of the patch with either 1 or 10 μM regulatory Ca\(^{2+}\).
9. **Group 3 NCX1.1 Mutants (Outward Current)**

The last category of NCX1.1 mutants consisted of only one member, Δ680-685. This non-XIP mutant (created in the COOH-terminal of the intracellular loop) demonstrated suppressed Ca$^{2+}$-dependent regulation ($I_2$) (212). Represented in Figure 21 are the effects of 0.1 μM SEA0400 on outward Δ680-685 exchange currents in the presence of 0, 1, and 10 μM regulatory Ca$^{2+}$. Ca$^{2+}$-dependent regulation ($I_2$) is lacking in Δ680-685, for this mutant exchanger is fully active in the absence of regulatory Ca$^{2+}$ (i.e., 0 μM). Consecutive increases in regulatory Ca$^{2+}$ for control currents show no alteration in effect, save for a small decrease in peak current at 10 μM Ca$^{2+}$. This can be explained as transport-ion competition. However, regardless that the exchanger lacks $I_2$, inhibition with SEA0400 mirrors that of wild-type. Table 1 shows that for Δ680-685, % peak inhibition was 32 ± 8 and 31 ± 6, while the wild-type was 57 ± 6 and 31 ± 8 at 1 and 10 μM regulatory Ca$^{2+}$, respectively. These results suggest that SEA0400 does not interact with and/or require Ca$^{2+}$-dependent regulation ($I_2$) for proper inhibition.
Figure 21. Effects of SEA0400 on Outward Na⁺-Ca²⁺ Exchange Currents of a Mutant NCX1.1 Exchanger Exhibiting Altered I₁ and I₂ Regulation. This figure demonstrates the effects of 0.1 μM SEA0400 on Δ680-685 exchange currents in the presence of 0 (left), 1 (middle) and 10 (right) μM regulatory Ca²⁺. Outward exchange currents were initiated by applying 100 mM Na⁺ to the cytoplasmic surface of the patch with 0, 1 or 10 μM regulatory Ca²⁺.
10. \( F_{ss} \) and \( \lambda \)

Tables 3 and 4 summarize the \( F_{ss} \) and \( \lambda \) results for NCX1.1 exchangers and all three mutant groups, in the absence and presence of 0.1 \( \mu \)M SEA0400. As stated by Hilgemann et al., \( F_{ss} \) values are the ratio of steady state to peak outward currents (121; 181). A high \( F_{ss} \) value represents a lower extent of Na\(^{+}\)-dependent (I\(_1\)) inactivation and vice versa. \( F_{ss} \) values also represent the fraction of active exchangers producing current at steady state (121). The parameter, \( \lambda \), is a measured rate constant for outward current inactivation. Furthermore, \( \lambda \) is indicative of I\(_1\) development represented by the formula:

\[
\lambda = \alpha E_{3ni} + \beta
\]

Here, \( \alpha \) is a rate constant for the transition of exchangers in the cytoplasmic \( E_{3ni} \) state into the I\(_1\) inactivation state. Basically, \( E_{3ni} \) represents the fraction of 3 Na\(^{+}\)-loaded exchangers with an intracellular orientation, while \( \beta \) is a rate constant for the recovery of exchangers from I\(_1\) to the \( E_{3ni} \) state. In short, \( \lambda \) is a measure of the rate of I\(_1\) inactivation, whereas \( F_{ss} \) measures the level or extent of I\(_1\) inactivation.

Interpreting \( F_{ss} \) and \( \lambda \) data in Tables 3 and 4 further strengthens the argument for an involvement of SEA0400 with the I\(_1\) process. Both K225Q and \( \Delta 680-685 \) had similar \( F_{ss} \) values to that of NCX1.1 wild-type at both 1 and 10 \( \mu \)M regulatory Ca\(^{2+}\), and in the absence and presence of SEA0400. This suggests that either the fine mutation or the lack of I\(_2\) does not affect the inhibitory function of SEA0400. Both \( \Delta 229-232 \) and K229Q, mutants lacking I\(_1\), show an increase in \( F_{ss} \) values in the absence or presence of SEA0400, and without regard for regulatory Ca\(^{2+}\) concentrations. These results parallel those of \( \alpha \)-chymotrypsin treated patches, lacking regulatory mechanisms I\(_1\) and I\(_2\). Here SEA0400 was without effect, leaving the deregulated exchanger with an \( F_{ss} \) value nearing \( \sim 1 \).
Conversely, the opposite occurs for the Group 1 mutant, F223E, which displays an accelerated and enhanced $I_1$. In this case, $F_{ss}$ was reduced and $\lambda$ was increased under control conditions, while an additional increase was observed following the addition of SEA0400. Results post-SEA0400 application on F223E (enhanced $I_1$) and Δ229-232 (suppressed $I_1$) mutants are in agreement with the belief that SEA0400 exerts its inhibitory effects through interactions involving the $I_1$ inactivation regulatory process.

11. **Group 1 NCX1.1 Mutants (Inward Exchange)**

To measure SEA0400’s inhibitory action in an exchanger mode lacking Na$^+$-dependent inactivation ($I_1$), all mutant inward exchange currents were tested in the absence and presence of SEA0400. $I_1$ inactivation is imperceptible for inward currents as Na$^+$ is lacking in the bath solution. All inward Na$^+$-Ca$^{2+}$ exchange currents (forward mode) were initiated by switching from a “Na$^+$ and Ca$^{2+}$ free” (but Li$^+$ containing) bath solution to one containing 3 or 10 μM regulatory Ca$^{2+}$. The pipette contained 100 mM Na$^+$ and was “Ca$^{2+}$-free”. Figure 22 illustrates a representative inward current of wild-type NCX1.1 in the absence and presence of 0.1 μM SEA0400. Here, SEA0400 is without effect on inward exchange currents at both 3 and 10 μM Ca$^{2+}$, as is expected.
Figure 22. Effects of SEA0400 on Inward Na\(^+\)-Ca\(^{2+}\) Exchange Currents of Wild-Type NCX1.1 Exchangers. This figure demonstrates a representative inward, NCX1.1-mediated exchange current, in the absence (Control) and presence of 0.1 \(\mu\)M SEA0400, are shown. Inward Na\(^+\)-Ca\(^{2+}\) exchange currents were initiated by rapidly applying either 3 or 10 \(\mu\)M Ca\(^{2+}\) to the cytoplasmic surface of the patch. The pipette solution contained 100 mM Na\(^+\).
Following the same order, Group 1 mutants (K225Q, F223E) were subjected to 0.1 μM SEA0400 during inward exchange current generation at both 3 and 10 μM regulatory Ca$^{2+}$. As is seen in Figure 23A, K225Q is somewhat inhibited by SEA0400 at both regulatory Ca$^{2+}$ concentrations (3 and 10 μM). Whereas, inward current of the second group 2 mutant F223E (exhibited in Figure 23B), is strongly inhibited by SEA0400 (% inhibition of steady state ~ 54 ± 6). These results suggest that SEA0400 maintains its inhibitory function in mutants displaying significantly increased $I_{1}$, even under ionic conditions where $I_{1}$ is absent. All percent inhibitions for inward exchange are summarized in Table 5.
Figure 23. Effects of SEA0400 on Inward $\text{Na}^+-$Ca$^{2+}$ Exchange Currents of Mutant NCX1.1 Exchangers Exhibiting Enhanced $I_I$ Inactivation. Panels A (top) and B (bottom), show the effects of 0.1 $\mu$M SEA0400 on inward exchange currents from K225Q (A) and F223E (B), respectively in the absence (left) and presence (right) of 0.1 $\mu$M SEA0400. Inward $\text{Na}^+-$Ca$^{2+}$ exchange currents were initiated by rapidly applying either 3 or 10 $\mu$M Ca$^{2+}$ to the cytoplasmic surface of the patch in exchange for 100 mM pipette Na$^+$. 
12. **Group 2 and 3 Mutant Inward Exchange**

When inward exchange currents for Group 2 mutants (K229Q, Δ229-232) were examined, it was observed in Figure 24, that both mutants were relatively unaffected by SEA0400, resulting in a % inhibitory range of 3 ± 1 to 13 ± 3. As expected, these I$_{1}$-suppressed mutants were similarly insensitive to SEA0400 on inward exchange currents, as they were on outward exchange currents. Lastly, inward exchange current for the sole Group 3 mutant, Δ680-685, was recorded. Shown in Figure 25, SEA0400 allows moderate inhibition of inward current at both regulatory Ca$^{2+}$ concentrations.
Figure 24. Effects of SEA0400 on Inward Na\textsuperscript{+}-Ca\textsuperscript{2+} Exchange Currents of Mutant NCX1.1 Exchangers Exhibiting Suppressed I\textsubscript{i} Inactivation. Panels A (top) and B (bottom) show representative tracings from K229Q (A), Δ229-232 (B) in the absence (left) and presence (right) of 0.1 μM SEA0400. Inward Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange currents were generated by rapidly applying either 3 or 10 μM Ca\textsuperscript{2+} to the cytoplasmic surface of the patch in exchange for 100 mM pipette Na\textsuperscript{+}. 
Figure 25. Effects of SEA0400 on Inward Na⁺-Ca²⁺ Exchange Current of a Mutant NCX1.1 Exchangers Exhibiting Altered I₁ and I₂ Regulation. This figure represents inward exchange current of Δ680-685 in the absence (left) and presence (right) of 0.1 µM SEA0400. Inward Na⁺-Ca²⁺ exchange currents were initiated by rapidly applying either 3 or 10 µM Ca²⁺ to the cytoplasmic surface of the patch in exchange for 100 mM pipette Na⁺.
Table 2. Effects of 0.1 μM SEA0400 on Peak and Steady State Outward Current in NCX1.1 and Mutant Na⁺-Ca²⁺ Exchangers.

<table>
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<th>Mutant</th>
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<th>n</th>
<th>I_{ss} (% Inhibition)</th>
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<td>1 μM Ca²⁺</td>
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<tr>
<td>WT</td>
<td>57 ± 6</td>
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</tr>
<tr>
<td>Δ229-232</td>
<td>9 ± 3*</td>
<td>8</td>
<td>21 ± 10*</td>
<td>8</td>
</tr>
<tr>
<td>K229Q</td>
<td>14 ± 5</td>
<td>13</td>
<td>36 ± 3*</td>
<td>13</td>
</tr>
<tr>
<td>Δ680-685</td>
<td>31 ± 6</td>
<td>14</td>
<td>65 ± 5</td>
<td>14</td>
</tr>
</tbody>
</table>

I_{peak}: peak outward current; I_{ss}: steady-state outward current; WT: wild-type NCX.1; *: P<0.05 compared to WT; ‡: block is to zero current level (see text for details).

Table 3. Effects of 0.1 μM SEA0400 on F_{ss} in NCX1.1 and Mutant Na⁺-Ca²⁺ Exchangers.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Control</th>
<th>SEA0400</th>
</tr>
</thead>
<tbody>
<tr>
<td>F_{ss}</td>
<td>n</td>
<td>F_{ss}</td>
</tr>
<tr>
<td>1 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.17 ± 0.01</td>
<td>43</td>
</tr>
<tr>
<td>F223E</td>
<td>0.07 ± 0.02*</td>
<td>18</td>
</tr>
<tr>
<td>K225Q</td>
<td>0.24 ± 0.02*</td>
<td>13</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>0.82 ± 0.05*</td>
<td>12</td>
</tr>
<tr>
<td>K229Q</td>
<td>0.90 ± 0.03*</td>
<td>28</td>
</tr>
<tr>
<td>Δ680-685</td>
<td>0.63 ± 0.05*</td>
<td>13</td>
</tr>
<tr>
<td>10 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.72 ± 0.05</td>
<td>13</td>
</tr>
<tr>
<td>F223E</td>
<td>0.34 ± 0.04*</td>
<td>17</td>
</tr>
<tr>
<td>K225Q</td>
<td>0.83 ± 0.03</td>
<td>11</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>0.84 ± 0.02*</td>
<td>17</td>
</tr>
<tr>
<td>K229Q</td>
<td>0.94 ± 0.02*</td>
<td>20</td>
</tr>
<tr>
<td>Δ680-685</td>
<td>0.62 ± 0.05</td>
<td>17</td>
</tr>
</tbody>
</table>

F_{ss}: I_{ss}/I_{peak}; WT: wild-type NCX.1; *: P<0.05 compared to WT.
### Table 4. Effects of 0.1 μM SEA0400 on λ in NCX1.1 and Mutant Na⁺-Ca²⁺ Exchangers.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Control</th>
<th>SEA0400</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>λ₀</td>
<td>n</td>
</tr>
<tr>
<td>1 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.20 ± 0.02</td>
<td>43</td>
</tr>
<tr>
<td>F223E</td>
<td>1.53 ± 0.06*</td>
<td>18</td>
</tr>
<tr>
<td>K225Q</td>
<td>0.33 ± 0.03*</td>
<td>13</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>1.07 ± 0.24*</td>
<td>10</td>
</tr>
<tr>
<td>K229Q</td>
<td>0.51 ± 0.08*</td>
<td>15</td>
</tr>
<tr>
<td>Δ680-685</td>
<td>0.34 ± 0.04*</td>
<td>13</td>
</tr>
<tr>
<td>10 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.86 ± 0.22</td>
<td>12</td>
</tr>
<tr>
<td>F223E</td>
<td>1.47 ± 0.15*</td>
<td>17</td>
</tr>
<tr>
<td>K225Q</td>
<td>1.13 ± 0.25</td>
<td>8</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>1.57 ± 0.33</td>
<td>11</td>
</tr>
<tr>
<td>K229Q</td>
<td>0.68 ± 0.32</td>
<td>9</td>
</tr>
<tr>
<td>Δ680-685</td>
<td>0.32 ± 0.04*</td>
<td>17</td>
</tr>
</tbody>
</table>

λ: rate of current inactivation s⁻¹; WT: wild-type NCX.1; *: P<0.05 compared to WT.

### Table 5. Effects of 0.1 μM SEA0400 on Steady State Inward Current in NCX1.1 and Mutant Na⁺-Ca²⁺ Exchangers.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Ims (%, Inhibition)</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>3 ± 1</td>
<td>6</td>
</tr>
<tr>
<td>F223E</td>
<td>52 ± 4*</td>
<td>6</td>
</tr>
<tr>
<td>K225Q</td>
<td>15 ± 5</td>
<td>10</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>13 ± 3*</td>
<td>4</td>
</tr>
<tr>
<td>K229Q</td>
<td>5 ± 2</td>
<td>12</td>
</tr>
<tr>
<td>10 μM Ca²⁺</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>3 ± 1</td>
<td>15</td>
</tr>
<tr>
<td>F223E</td>
<td>28 ± 5*</td>
<td>10</td>
</tr>
<tr>
<td>K225Q</td>
<td>10 ± 5</td>
<td>8</td>
</tr>
<tr>
<td>Δ229-232</td>
<td>2 ± 1</td>
<td>8</td>
</tr>
<tr>
<td>K229Q</td>
<td>7 ± 2</td>
<td>10</td>
</tr>
</tbody>
</table>

Ims: steady-state outward current; WT: wild-type NCX.1; *: P<0.05 compared to WT.
V. DISCUSSION

1. Transport Mode Selectivity

The primary aim was to examine the inhibitory effects of SEA0400, a novel Na\(^+\)-Ca\(^{2+}\) exchange inhibitor and provide insight into its inhibitory mechanism(s). Using NCX1.1 (canine cardiac Na\(^+\)-Ca\(^{2+}\) exchanger), expressed in *Xenopus laevis* oocytes, we applied the giant, excised patch clamp technique. We wished to investigate SEA0400’s mode selectivity on NCX1.1, if any. Previous results indicate that SEA0400 may exhibit reverse mode (i.e., outward) selectivity in both neural cells (204) and cardiomyocytes (310; 311) but in some instances it was not selective for the Na\(^+\)-Ca\(^{2+}\)exchanger itself (274). However, under bi-directional conditions, inward and outward currents were measured using the same population of exchangers (i.e., same patch) and indicated inhibition of outward exchange current by SEA0400 to a greater degree than for inward current. Under these conditions, 0.3 \(\mu\text{M}\) SEA0400 reduced outward current by ~ 90 % but only affected inward currents by ~ 30 %. With pooled data in Figure 12B, it is apparent that SEA0400 inhibits steady state outward currents to a greater extent than peak outward currents. SEA0400 was also tested under unidirectional conditions that measure outward and inward exchange currents separately. Even under unidirectional conditions, as Figures 13A illustrates, SEA0400 still exerts its potent effects on outward current, while Figure 17A shows SEA0400 to be virtually without effect on inward currents at concentrations up to 3 \(\mu\text{M}\) SEA0400. An \(I_C^{50}\) value for outward exchange inhibition was placed at ~ 23 nM (0.023 \(\mu\text{M}\)), whereas an \(I_C^{50}\) value for inward exchange currents could
not be obtained due to SEA0400’s poor solubility at higher concentrations. As mentioned previously, the effects of SEA0400 on inward current at higher concentrations (i.e., > 3 μM) could not be tested due to solubility issues. Although an accurate IC₅₀ could not be isolated, an estimated value of ~ 8 μM was suggested. Despite the tendency for SEA0400 to precipitate out of solution it is quite evident under both conditions, SEA0400 is reverse mode selective. This parallels results from previous studies (204; 310). However, there are reports of SEA0400 functioning with equal potency for both forward and reverse mode. In a study by Tanaka et al., both SEA0400 and KB-R7943 were applied to guinea pig ventricular myocytes, and resulted in an equipotent inhibition of both outward and inward Na⁺-Ca²⁺ exchange currents (IC₅₀ ~ 32 and 40 nM, respectively) (311). Results for KB-R7943 provided a similar trend, however with higher IC₅₀ values (311). It was suggested that mode-selective drug binding occurs through the preferential recognition of certain exchanger conformations. Thus, binding to these positions would render the exchanger inactive. Nevertheless, under certain ionic conditions (favouring forward or reverse mode), the exchangers position themselves according to their preferred mode of exchange. This adjusts the relative percentage of exchangers displaying certain conformational forms. This implies that there are a different percentage of exchanger conformations under unidirectional and bi-directional conditions. Therefore, under unidirectional conditions, exchangers will be in one of two conformations (facing cytoplasmically or extracellularly). As the drug binds, it chooses a favourable conformation and this renders the exchanger temporarily inactive. The drug will affect both modes differently, for they each contain different conformations. However, in the bi-directional setting, the possibility for either exchange mode to occur
exists and so does the potential for both conformational positions. Therefore, in certain circumstances, equipotent inhibition is observed (311). Conversely, Takahashi et al. reported that in canine sarcolemmal vesicles and rat myocytes, SEA0400 inhibits reverse mode exchange, with IC$_{50}$ values of 90 nM and 92 nM, respectively (310). Iwamoto et al. also reported mode selective inhibition for both SEA0400 and KB-R7943, while measuring Na$^+$-dependent Ca$^{2+}$ uptake (204). Placing these mode preference discrepancies aside, our results suggest that SEA0400 does exert reverse mode selectivity.

2. **SEA0400 and Ionic Regulatory Mechanisms**

In an attempt to reveal a mechanism(s) for SEA0400 inhibition, the exchanger was removed of its ionic regulatory processes. Using $\alpha$-chymotrypsin, both significant ionic regulatory mechanisms, $I_1$ and $I_2$, were eliminated, leaving the exchanger in a deregulated state. Both of these mechanisms have been suggested to play important roles in maintaining exchanger function, under various ionic conditions (116; 121; 123). As mentioned previously, $I_1$ is known as Na$^+$-dependent inactivation. As a result of cytoplasmic Na$^+$ application during outward Na$^+$-Ca$^{2+}$ exchange current, the exchanger is stimulated to a peak current level, upon which rapid current decay begins, levelling off to a steady state. However, $I_2$ or Ca$^{2+}$-dependent regulation refers to the requirement of cytoplasmic (regulatory) Ca$^{2+}$ for both outward and inward exchange current to occur. In the absence of cytoplasmic Ca$^{2+}$, no exchange activity is observed, whereas by gradually increasing regulatory Ca$^{2+}$, exchange current increases as the regulatory binding site
becomes saturated ($K_d \sim 0.3 \mu M$). This increase, however, results in the removal of the $I_1$ mechanism, leaving a square waveform. At higher regulatory $Ca^{2+}$ levels (> 10 $\mu M$ $Ca^{2+}$), current values begin to decrease as this probably reflects a competition between transported $Na^+$ and $Ca^{2+}$ ions. As a note, regardless of favourable ionic conditions, $Na^+$-$Ca^{2+}$ exchangers devoid of regulatory $Ca^{2+}$ will not exhibit exchange. As Figure 14A demonstrates, SEA0400 appears to have a decreased inhibitory effect on outward exchange current in deregulated exchangers (lacking $I_1$ and $I_2$). This result that when regulatory mechanisms are absent or disrupted, SEA0400 cannot fully exert its inhibitory action. This predicates a requirement or utilization of either or both $I_1$ and $I_2$. However, inward exchange currents were also measured after proteolysis, and SEA0400 was found to be without effect even at 3 $\mu M$ SEA0400. Nevertheless, $\alpha$-chymotrypsin’s site(s) of proteolysis is unknown, and may very well interact with certain exchanger regions specific for SEA0400 binding and inhibition. Therefore, although SEA0400 appears to require intact ionic regulatory mechanisms, the potential for an alternate binding site cannot be completely ruled out.

It is understood that both $I_1$ and $I_2$ interact (121; 123). It has been shown that for outward exchange, upon increasing regulatory $Ca^{2+}$ levels, there is an alleviation of $I_1$. Figure 15A clearly shows that switching from 1 to 10 $\mu M$ regulatory $Ca^{2+}$ strongly attenuates $Na^+$-dependent inactivation ($I_1$). Therefore, to further investigate the involvement of regulatory mechanisms in SEA0400’s mode of inhibition, experiments were performed at 1 and 10 $\mu M$ $Ca^{2+}$. SEA0400 was, however, able to prevent the alleviation of $I_1$ in the presence of high regulatory $Ca^{2+}$ concentrations (i.e., 10 $\mu M$). It would appears that SEA0400 is able to promote entry or maintain the exchanger’s $I_1$
mechanism, such that Na\(^{+}\)-dependent inactivation is present under conditions where it is normally absent. In addition, SEA0400 minimally inhibits inward exchange current (Figure 12), where Na\(^{+}\)-dependent inactivation is absent. This result implicated I\(_{1}\) involvement for inhibition through SEA0400.

When comparing the results of SEA0400 inhibition, current decay rate (\(\lambda\)) was greatly increased in comparison to control (decay) rates. Thus, the rate at which I\(_{1}\) entry occurred upon SEA0400 application is increased (Table 2). Further supporting the involvement of I\(_{1}\), were the results stemming from paired-pulse experiments. Data from measured recovery rate (\(\beta\)) in the presence and absence of 0.1 \(\mu\)M SEA0400 revealed that under the same conditions, \(\beta\) was for the most part unchanged (Figure 16C). With increasing regulatory Ca\(^{2+}\) concentrations, SEA0400 appears to prevent the alleviation of I\(_{1}\) normally associated with such increased Ca\(^{2+}\) levels. It appears that SEA0400 assists the entry into I\(_{1}\) during lower regulatory Ca\(^{2+}\) levels (\(i.e.,\ 1 \mu\)M) and does not affect the rate of recovery from I\(_{1}\). Upon increased regulatory Ca\(^{2+}\) levels, SEA0400 stabilizes I\(_{1}\) with minimal alleviation, even at high Ca\(^{2+}\) values (\(i.e.,\ 10 \mu\)M) (Figure 16C). These results strongly implicated the involvement of ionic regulatory mechanisms in the process of SEA0400 inhibition, with a tendency towards favouring I\(_{1}\). The next rational measure was to test the effects of SEA0400 on mutants displaying distinct altered ionic regulatory phenotypes.
3. **SEA0400 and Mutant Na\(^+\)-Ca\(^{2+}\) Exchangers**

Prior mutant exchanger experiments, we suggested that SEA0400 was reverse mode selective and required intact regulatory mechanisms for proper inhibition. Considering that forward mode exchange lacks \(I_1\) and that following exchanger deregulation, SEA0400’s potency was greatly reduced, it was suggested that the \(I_1\) regulatory mechanism was involved. Changing regulatory Ca\(^{2+}\) values showed SEA0400 to interact with \(I_1\) and maintain this inhibitory state, even at excessively high Ca\(^{2+}\) levels. The primary purpose for implementing mutated Na\(^+\)-Ca\(^{2+}\) exchangers with altered ionic regulatory properties was to further determine the requirement of Na\(^+\)-dependent inactivation (\(I_1\)) for SEA0400 inhibition. Subsequent to Na\(^+\)-Ca\(^{2+}\) exchanger cloning (227), the XIP region was identified and proposed to contribute to exchanger autoregulation (193). Therefore, the XIP region was a target for mutation for the exploration of exchange regulation. In addition, the small 20 amino acid sequence (219-238) XIP region, was shown to be involved in ionic regulation when Matsuoka et al. created both point and deletion mutants, resulting in exchangers displaying altered ionic regulation (209; 211). Therefore, we chose to study these defined mutants in conjunction with SEA0400 analysis. The mutants were grouped according to their regulatory phenotype. In short, there were mutants with increased \(I_1\), lacking \(I_1\) and lacking \(I_2\). Our results were most convincing with mutants with accelerated \(I_1\) and those lacking \(I_1\).
4. Mutant Na\(^+\)-Ca\(^2+\) Exchangers with Altered I\(_1\) (Outward Exchange)

We found that the K225Q mutant had similar I\(_1\) levels to that of wild-type, whereas F223E had significantly increased I\(_1\) levels. Inhibition by SEA0400 on K225Q was nearly identical to that of wild-type. However, in the F233E mutant, carrying a \(~2\) fold increase in the rate and level of I\(_1\), the degree of inhibition by SEA0400 was appreciably increased, bringing steady state current levels near baseline. Such inhibition made it difficult to quantify values for % inhibition (peak and steady state) and F\(_{ss}\) values. In the Group 2 mutants (K229Q, Δ229-232), both displayed suppressed I\(_1\), resulting in square waveforms. When the first mutant, K229Q, was tested, SEA0400 appeared to induce I\(_1\)-like inactivation. Furthermore, increasing regulatory Ca\(^2+\) alleviated this I\(_1\)-like inactive state, just as Ca\(^2+\) does with wild-type. This suggested that the observed result was in fact, genuine I\(_1\) being formed. The second Group 2 mutant, Δ229-232, displayed a more definitive effect, showing minimal inhibition with SEA0400. Therefore, both F223E and Δ229-232 complement one another; such that when I\(_1\) is augmented, so is SEA0400’s ability to inhibit and when I\(_1\) is absent, so is SEA0400’s ability to inhibit. However, the effects of SEA0400 on both K225Q and K229Q should not be dismissed. The resulting “lack of effect” of SEA0400 on K225Q may be due to its more subtle mutation compared to F223E. Furthermore, K225Q, in the absence of SEA0400, does not have significantly increased I\(_1\). Thus, following the hypothesis that SEA0400 requires I\(_1\), this may explain why SEA0400 inhibits to the same degree as wild-type. Conversely, K229Q initially appears to lack functional I\(_1\). However, SEA0400 application produces an I\(_1\)-like inactivation that is alleviated by increasing regulatory
Ca\textsuperscript{2+} levels. Perhaps this ability of SEA0400 to form I\textsubscript{i} may reside in the position and severity of the mutation. Both group 2 mutants begin at amino acid 229. While K229Q is only a point mutation, Δ229-232 is a large deletion of 4 amino acids. It may be that the K229Q point mutation only alters a portion of the region governing I\textsubscript{i} entry, whereas Δ229-232 removes the entire segment. This could allow K229Q to retain the ability to enter the I\textsubscript{i} inactive state from the cytoplasmic E\textsubscript{3ni} state. However, it may be that the potential of entry into I\textsubscript{i} exists; yet it may be less stable and therefore less favourable. We have previously suggested that SEA0400 may promote entry into I\textsubscript{i} or stabilize this inactive state. Therefore, in the case of K229Q, SEA0400 may stabilize the I\textsubscript{i} state and subsequently allow its accumulation. Lastly, this I\textsubscript{i}-like state is alleviated by increasing regulatory Ca\textsuperscript{2+} concentrations, which follows the same convention of wild-type I\textsubscript{i}. Due to the use of varied mutants, results with SEA0400 would most likely be mutant specific. Therefore, results found with SEA0400 on both K229Q and Δ229-232 do not detract from results indicating direct interaction with I\textsubscript{i}, but complement them. However, we cannot unequivocally state that these explanations comprise the actual mechanism(s) of action for SEA0400.

5. Mutant Na\textsuperscript{+}-Ca\textsuperscript{2+} exchangers with altered I\textsubscript{2} (Outward exchange)

The last mutant, Δ680-685, belongs to Group 3 and displays suppressed Ca\textsuperscript{2+}-dependent regulation (I\textsubscript{2}). As described previously, cytoplasmic Ca\textsuperscript{2+} regulates the exchanger, such that if it is not present, exchanger activation does not occur. We refer to this requirement of Ca\textsuperscript{2+} to activate the exchanger and to stimulate outward exchange
current with increasing levels, as Ca$^{2+}$-dependent regulation. However, at very low cytoplasmic Ca$^{2+}$ levels, exchangers remain inactive or exhibit decreased exchange. This inactive state is referred to as I$_2$ inactivation. By this convention, we can say that both I$_1$ and I$_2$ inactivation depend on Ca$^{2+}$. Since both of these regulatory mechanisms are Ca$^{2+}$-dependent, we decided to examine a mutant Na$^+$-Ca$^{2+}$ exchanger exhibiting altered I$_2$, previously described by Maxwell et al. (212). Our results in Figure 21, displaying outward exchange currents, demonstrate that in the presence of SEA0400, inhibition was equal to that of wild-type for peak and steady state currents (Table 3). However, this suggests that I$_2$ does not play an important role for SEA0400 inhibition, yet results have varied in the extent of I$_1$ production between Δ680-685 mutants. Therefore, I$_2$ cannot be wholly excluded as a potential mechanism of SEA0400 action.

6. Inward Mutant Na$^+$-Ca$^{2+}$ Exchange (Group 1-3)

We continued to test inward Na$^+$-Ca$^{2+}$ exchange currents, known to lack I$_1$, in hopes of further establishing I$_1$ as the dominant mechanism for SEA0400 inhibition. The I$_1$ inactive state arises from E$_{3ni}$, which is more likely to occur during outward exchange currents, where cytoplasmic [Na$^+$]$_i$ is high. Conversely, during inward exchange current, Na$^+$ translocates from the extracellular side to a virtually "Na$^+$-free" side, where unbinding is most favourable. Therefore, E$_{3ni}$ is less likely to accumulate here and enter the I$_1$ state. This is the primary reason for its absence during inward exchange. As we expected, most inward exchange currents were insensitive to SEA0400, except for that of F223E. The Group 1 mutant displayed moderate inhibition with SEA0400 exposure. A
potential explanation may be that F223E shows greatly increased I₁ for outward current but may form a small amount of I₁ during inward exchange currents, which is further acted upon by SEA0400 (which uses I₁ to exert its inhibitory effects).

7. Peak vs. Steady State Current Inhibition

As Table 2 indicates, during outward exchange current, SEA0400 inhibits steady state current to a greater extent than peak current. However, peak current is thought to be independent of Na⁺-dependent inactivation. Therefore, if we suggest that SEA0400 inhibits through I₁, and we see a resulting inhibition of peak current, SEA0400 may have an alternate site of interaction. It remains difficult to determine whether the resulting peak current is due to SEA0400, for peak current represents the net outcome between developing current and that of I₁ entry. At present, we lack the temporal resolution to distinguish between these two possibilities.

8. Pathophysiological Implications

As previously discussed, the primary physiological role of the Na⁺-Ca²⁺ exchanger is to maintain Ca²⁺ homeostasis by directly removing the proportional amount of Ca²⁺ that enters through L-type Ca²⁺ channels. If this coupling is not met, the cell can become either Ca²⁺ overloaded or depleted. In addition, the exchanger can function bi-directionally, as its mode of action depends on both electrical and concentration gradients. Therefore, under various pathophysiological conditions, reverse mode (Ca²⁺
influx) exchange can occur, leading to not only a reduction in the removal of Ca\textsuperscript{2+} but an increased Ca\textsuperscript{2+} influx. This can be severely detrimental to the cell and can quickly result in Ca\textsuperscript{2+} overload (with associated dysfunctions) and cell death. In cases such as ischemia-reperfusion injury (either surgically induced or naturally occurring) and arrhythmogenesis, where reverse mode Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange is considered to be a prime contributor, having a reverse mode selective inhibitor would be invaluable. A selective Na\textsuperscript{+}-Ca\textsuperscript{2+} inhibitor that preferentially blocks Ca\textsuperscript{2+} entry but leaves Ca\textsuperscript{2+} efflux unaltered could play a crucial role in the therapeutic treatment of cardiac ailments. Even more appealing would be an inhibitor that selectively targeted reverse mode exchange with a very high potency, thus requiring low doses for treatment. Until recently, KB-R7943 showed promise, as it appeared significantly more potent and reverse mode selective than any other available inhibitor (145; 327). However, results on KB-R7943’s selectivity (both mode and cellular protein) and potency were varied, and in some cases investigators discounted KB-R7943 as a potentially useful clinical inhibitor.

In summary, we have shown SEA0400 to be a potent, reverse mode selective inhibitor and suggest that it or its analogue derivatives could serve as potential therapeutic agents (cardiac disease/injury, renal disease/injury and cerebral disease/injury). Understanding SEA0400’s mechanism(s) of action could give insight into the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger’s properties and lead to the rational approach in future pharmacological design. The exchanger has been shown to have a crucial physiological role in maintaining proper cardiac function and, however, has been implicated in several cardiac injuries, namely ischemia-reperfusion injury. It is essential to fully understand the mechanism(s) of Na\textsuperscript{+}-Ca\textsuperscript{2+} exchange function in order to develop specific tools
directly targeting this membrane transporter. Therefore, we must examine SEA0400 and identify its inhibitory mechanism(s) and mode(s) of action. If fully understood, SEA0400 or putative derivatives could be applied to various diseases/injuries. For example, during periods of induced ischemia-reperfusion injury (i.e., cardiac surgery), preceding drug administration would be most desirable. Here, the majority of damage is a result of reperfusing the recently operated heart. With SEA0400, we could give the drug before inflicting unavoidable damage (reperfusion) and hopefully improve patient recovery. Furthermore, access to a viable reverse mode selective Na\(^+\)-Ca\(^{2+}\) exchange inhibitor would allow for prophylactic treatment. Theoretically, an ideal inhibitor would exert its effects only during reverse mode exchange, leaving forward mode exchange unaltered. The population who would benefit most, are those with a history of heart disease and/or are at risk for cardiac injury (via obesity, diabetes, high cholesterol, smoking and genetic predisposition). If the heart becomes ischemic, it is essential that blood flow be re-established. Although reperfusion is imperative (without it, the tissue will die), it is this very act that induces the most serious and irreversible damage. Therefore, having the inhibitor present prior to the insult, would provide the most effective protection against the necessary reperfusion. To conclude, SEA0400, in conjunction with effective reperfusion therapy could greatly increase a patient’s prospect for recovery following a cardiac insult.

As cardiac disease and injury are reaching epidemic proportion in North America alone, further study of cardiac Na\(^+\)-Ca\(^{2+}\) exchange and its pharmacology will play a vital role in both the treatment of pre-existing cardiac disease and its prophylactic utilization.
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