Allosteric Activation of Sodium–Calcium Exchange Activity by Calcium: Persistence at Low Calcium Concentrations

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ABSTRACT The activity of the cardiac Na\(^+\)/Ca\(^{2+}\) exchanger is stimulated allosterically by Ca\(^{2+}\), but estimates of the half-maximal activating concentration have varied over a wide range. In Chinese hamster ovary cells expressing the cardiac Na\(^+\)/Ca\(^{2+}\) exchanger, the time course of exchange-mediated Ca\(^{2+}\) influx showed a pronounced lag period followed by an acceleration of Ca\(^{2+}\) uptake. Lag periods were absent in cells expressing an exchanger mutant that was not dependent on regulatory Ca\(^{2+}\) activation. We assumed that the rate of Ca\(^{2+}\) uptake during the acceleration phase reflected the degree of allosteric activation of the exchanger and determined the value of cytosolic Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_c\)) at which the rate of Ca\(^{2+}\) influx was half-maximal ([K\(_a\)]. After correcting for the effects of mitochondrial Ca\(^{2+}\) uptake and fura-2 buffering, K\(_a\) values of \(~\sim\)300 nM were obtained. After an increase in [Ca\(^{2+}\)]\(_i\), the activated state of the exchanger persisted following a subsequent reduction in [Ca\(^{2+}\)]\(_i\) to values \(<\)100 nM. Thus, within 30 s after termination of a transient increase in [Ca\(^{2+}\)]\(_i\), exchange-mediated Ca\(^{2+}\) entry began without a lag period and displayed a linear rate of Ca\(^{2+}\) uptake in most cells; a sigmoidal time course of Ca\(^{2+}\) uptake returned 60–90 s after the transient increase in [Ca\(^{2+}\)]\(_i\) was terminated. Relaxation of the activated state was accelerated by the activity of the endoplasmic reticulum Ca\(^{2+}\) pump, suggesting that local Ca\(^{2+}\) gradients contribute to maintaining exchange activation after the return of global [Ca\(^{2+}\)]\(_i\) to low values.

KEY WORDS: Chinese hamster ovary cells • fura-2 • digital imaging • mitochondria • SERCA

INTRODUCTION

The Na\(^+\)/Ca\(^{2+}\) exchange system is an important Ca\(^{2+}\) transporter in the plasma membrane of cardiac myocytes and many other kinds of cells (for reviews see Blaustein and Lederer, 1999; Shigekawa and Iwamoto, 2001). The exchanger couples the movement of Ca\(^{2+}\) across the membrane to the reciprocal movement of 3 Na\(^+\) (Reeves and Hale, 1984), or perhaps more (Fujioka et al., 2000b; Dong et al., 2002), Na\(^+\) ions. Under physiological conditions, it functions principally as a Ca\(^{2+}\) efflux mechanism, although it may also bring Ca\(^{2+}\) into the cell upon strong membrane depolarization or when cytosolic Na\(^+\) concentrations increase above normal. The exchanger itself is a protein of 938 amino acids containing 9 transmembrane segments, with a cytosolically disposed hydrophilic domain of 544 amino acids located between transmembrane segments 5 and 6. Removal of the central hydrophilic domain by proteolysis or mutation abrogates regulatory modulation of exchange activity (Matsuoka et al., 1993).

A major mechanism for the regulation of exchange activity involves the interaction of Ca\(^{2+}\) with two well-defined binding sites within the exchanger’s central hydrophilic domain (Levitsky et al., 1994). Ca\(^{2+}\)-dependent regulation, also called allosteric Ca\(^{2+}\) activation, was first observed in early experiments with squid giant axons (DiPolo, 1979) and has by now been documented by many investigators under a host of different experimental conditions (Blaustein and Lederer, 1999). Most studies indicate that the regulatory sites must be filled with Ca\(^{2+}\) for the exchanger to work in any of its known modes of operation—Na\(^+\)/Ca\(^{2+}\), Na\(^+\)/Na\(^+\), or Ca\(^{2+}\)/Ca\(^{2+}\) exchange. Allosteric Ca\(^{2+}\) activation provides a teleologically satisfying means of turning exchange activity on or off as needed. Importantly, shutting down the exchanger at a low cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_c\)) could in principle prevent exchange activity from reducing resting [Ca\(^{2+}\)]\(_i\), to the equilibrium value dictated by the transmembrane Na\(^+\) gradient under physiological conditions (\(~\sim\)20 nM for a 3:1 Na\(^+\)/Ca\(^{2+}\) exchange ratio at a cytosolic Na\(^+\) concentration of 10 mM and a membrane potential of \(~\sim\)80 mv)...

Abbreviations used in this paper: CHO, Chinese hamster ovary; Cl-CCP, carbonyl cyanide m-chlorophenylhydrazone; PSS, physiological salts solution; SERCA, sarco(endo)plasmic reticulum Ca\(^{2+}\)-ATPase; Tg, thapsigargin.

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activity is 200–600 nM (Collins et al., 1992; Hilgemann et al., 1992a; Matsuoka et al., 1995, 1997; Fujioka et al., 2000a). In contrast, measurements of exchange activity in intact cardiac myocytes yielded values ranging from 22 to 125 nM (Miura and Kimura, 1989; Noda et al., 1988; Weber et al., 2001). Previous work from our laboratory measured the Ca\(^{2+}\) dependence of exchange-mediated Ba\(^{2+}\) uptake in transfected Chinese hamster ovary (CHO) cells expressing NCX1.1 (Fang et al., 1998). Cytosolic Ca\(^{2+}\) activated Ba\(^{2+}\) influx cooperatively (Hill coefficient 1.6) with a K\(_h\) of 44 nM. It has been difficult to envision how such a high-affinity system could be an effective regulator of exchange activity, since the exchanger would be nearly fully activated at resting cytosolic Ca\(^{2+}\) concentrations of \(\sim 100 \text{ nM}\) (Reeves, 1998).

Here, we present a novel approach to measuring the K\(_h\) for the allosteric activation of exchange activity by Ca\(^{2+}\). We studied exchange activity operating in its “reverse mode”, when extracellular Na\(^{+}\) was removed so that Ca\(^{2+}\) influx occurred. We found that when reverse exchange activity was initiated, Ca\(^{2+}\) uptake did not begin immediately but showed a pronounced lag phase followed by a progressive increase in the rate of Ca\(^{2+}\) uptake. We assumed that the acceleration of Ca\(^{2+}\) uptake reflected increasing allosteric activation of the exchanger by Ca\(^{2+}\), and defined the K\(_h\) for activation as the value of [Ca\(^{2+}\)] when the rate of Ca\(^{2+}\) uptake was half-maximal. This approach yielded K\(_h\) values substantially higher (\(\sim 300 \text{ nM}\)) than we measured in our previous work on Ba\(^{2+}\) uptake. Remarkably, we also found that after a transient exposure to high [Ca\(^{2+}\)], the activated state of the exchanger persisted for tens of seconds after global [Ca\(^{2+}\)] had declined to values below the K\(_h\) for exchange activation. The persistence of exchange activation explains why our previous investigations yielded such a low apparent K\(_h\) for regulatory Ca\(^{2+}\) activation. Our results indicate that relaxation of the activated state is promoted by Ca\(^{2+}\) uptake into the ER, suggesting that local Ca\(^{2+}\) gradients may be involved in persistent activation.

**MATERIALS AND METHODS**

**Cells**

CHO K1 cells or CHO T cells (CHO K1 cells expressing the human insulin receptor (Langille et al., 1999), provided by M. Czech (University of Massachusetts Medical Center), were transfected with the mammalian expression vector pcDNA3 containing the cDNA sequence for the bovine cardiac Na\(^{+}\)/Ca\(^{2+}\) exchanger (NCX1.1: accession number LO6438). Cells expressing exchange activity were selected using the ionomycin treatment of Iwamoto et al. (1998). No qualitative differences in the behavior of transfected CHO K1 or T1 cells were observed, but the T1 transfectants tended to exhibit higher levels of NCX expression and were used for most experiments shown here. Cells expressing the Δ(241–680) deletion mutant were prepared similarly. In this mutant, a large part of the exchanger’s central hydrophilic domain has been deleted; the Δ(241–680) mutant shows high activity in the absence of allosteric Ca\(^{2+}\) activation (Matsuoka et al., 1993). An additional mutant was used in one study (see Appendix), in which the positively charged residues within the so-called XIP region (251–270) were changed to alanines; this mutant is resistant to Na\(^{+}\)-dependent inactivation, although it displays normal allosteric Ca\(^{2+}\) activation (Condrescu and Reeves, 1999). Cells were grown in a 5% CO\(_2\) atmosphere in F12 medium supplemented with 10% fetal calf serum, 2 mM L-glutamine, 100 U/ml penicillin, 100 μg/ml streptomycin, and 20 μg/ml gentamicin. The cell-culture medium, including the fetal calf serum, was obtained from Life Technologies.

**Solutions**

Na-PSS contained 140 mM NaCl, 5 mM KCl, 1 mM MgCl\(_2\), 10 mM glucose, and 20 mM Mops, buffered to pH 7.4 with Tris. Li-PSS had an identical composition except that LiCl was substituted for NaCl. K-PSS contained 140 mM KCl, 1 mM MgCl\(_2\), 10 mM glucose, and 20 mM Mops, buffered to pH 7.4 with Tris. Unless otherwise indicated, experiments were performed at room temperature; for experiments at 37°C, the pH of the solutions was adjusted to pH 7.4 at 37°C. Biochemicals were purchased from Sigma-Aldrich.

**Fura-2 Imaging**

Cells grown on 25-mm circular coverslips were loaded with fura-2 by incubating them for 30 or 40 min at room temperature in Na-PSS containing 1 mM CaCl\(_2\), 1% fatty acid free bovine serum albumin, 0.25 mM sulfipyrazole (to retard fura-2 transport from the cell), and 3 μM fura-2 AM (Molecular Probes). The coverslips were then washed in Na-PSS + 1 mM CaCl\(_2\), placed in a stainless steel holder (bath volume \(~0.8 \text{ ml}\); Molecular Probes), and viewed in a ZEISS Axiovert 100 microscope coupled to an Astrofluor digital imaging system. Alternating excitation at 334 and 380 nm was obtained through the use of appropriate filters, and fluorescence was monitored with a long pass filter set for 510 nm. In most cases, 40–60 cells were individually monitored from each coverslip. Data points were taken at \(~1.7\) s intervals.

**Experimental Protocol**

To deplete internal Ca\(^{2+}\) stores and prevent Ca\(^{2+}\) uptake into the ER, cells were washed in Na-PSS + 0.3 mM EGTA and then treated with 200 μM ATP (a purinergic agonist) + 2 μM thapsigargin (Tg), an inhibitor of the sarco(endo)plasmic reticulum Ca\(^{2+}\) ATPase (SERCA) (Lyton et al., 1991). Experiments were begun \(~10\) min after the application of ATP + Tg. In experiments where ATP-induced Ca\(^{2+}\) release was monitored, fura-2–loaded cells were preincubated for 5–10 min in Na-PSS + 1 mM CaCl\(_2\); the cells were then washed with 4 ml of Na-PSS + 0.3 mM EGTA immediately before beginning the experiment. Changes in the bath medium were performed by manually applying \(~4\) ml of the desired medium with a cannulated syringe over a period of 15 s. Reverse exchange activity was initiated by applying K-PSS + 0.1 mM CaCl\(_2\).

**Calibration of [Ca\(^{2+}\)]**

Each cell was calibrated individually using the procedure of Grynkiewicz et al. (1985). During the course of this work, we examined various methods of determining R\(_{500}\), the 334/380 ratio at [Ca\(^{2+}\)]\(~=\)0. The procedure finally adopted involved applying Na-PSS + 0.3 mM EGTA to the cells after terminating the experiment, and then adding 10 μM ionomycin + 10 mM NH\(_4\)Cl in the same medium to release any residual Ca\(^{2+}\) from internal stores;
the NH₄Cl was added to facilitate ionomycin-induced Ca²⁺ release from acidic stores. After the resulting [Ca²⁺], transient decayed, 1 ml of 50 μM EGTA-AM in Na-PSS + 0.3 mM EGTA was applied and the 334/380 ratio was measured at 5-min intervals until no further decline was noted (usually >30 min). This ratio (average of 10 data points) was taken as Rmin after correcting for background fluorescence (see below). A 1:4 mixture of 100 mM CaCl₂ and K-PSS was then applied (along with an additional 10 μM ionomycin) and, after the fura signal stabilized, the average ratio over 10 data points, after background correction, was taken as Rcorr. Finally, 10 mM MnCl₂ in Na-PSS (1 ml) was applied to the cells to quench fura-2 fluorescence and the final fluorescence intensities at 334 nm and 380 nm excitation, averaged over 10 data points, were taken as background fluorescence. For each cell, individual background fluorescence values were subtracted from the experimental fluorescence values at 334 and 380 nm excitation, and Rmin, Rmax, and Sf/Sb values were computed; the latter value is the ratio of corrected fluorescence intensities at 380 nm under conditions for determination of Rmin (St) and Rmax (Sb). Experimental values for [Ca²⁺], were then calculated from the ratio of background-corrected fluorescence at 334/380 nm excitation (Rcorr), using the formula of Grynkiewicz et al. (1985), i.e., [Ca²⁺] = K_D,fura(Sf/Sb)(Rcorr - Rmin)/(Rmax - Rcorr). For experiments at room temperature, K_D,fura was assumed to be 274 nM (Grynkiewicz et al., 1985). The Kh for Allosteric Ca²⁺ Activation

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**Statistical Analysis**

Pooled results from multiple cells on a single coverslip are expressed as mean values ± SD. Pooled results from several coverslips are expressed as the mean value ± SEM. Significance testing was done using Student’s two-tailed t test for paired or unpaired samples, as indicated. **RESULTS**

**The Kh for Allosteric Ca²⁺ Activation**

Fig. 1 displays the time course of Ca²⁺ uptake after the initiation of the reverse (Ca²⁺ influx) mode of Na⁺/Ca²⁺ exchange in CHO cells expressing the wild-type Na⁺/Ca²⁺ exchanger (NCX1.1). In these experiments, the cells were pretreated with ATP + thapsigargin (Tg), an irreversible inhibitor of the SERCA pump, to release Ca²⁺ from internal stores and prevent subsequent Ca²⁺ uptake by the ER. The cells were then incubated in Ca²⁺-free Na-PSS for 10 min before beginning the experiment. Under these conditions, the initial cytosolic Ca²⁺ concentration was usually <25 nM. The trace labeled “I” in Fig. 1 (filled circles) demonstrates that when the reverse exchange activity was initiated by applying a Na⁺-free solution (K-PSS) containing 0.1 mM CaCl₂ (arrow labeled “K + Ca”), Ca²⁺ uptake was

**Figure 1.** Time course of exchange-mediated Ca²⁺ uptake in transfected CHO cells expressing NCX1.1. (A) Cells were treated with ATP + Tg in Na-PSS + 0.3 mM EGTA 10 min before beginning the experiment (see MATERIALS AND METHODS). At t = 50 s, Na-PSS containing either 0.1 mM EGTA (trace 1, filled circles) or 0.3 mM CaCl₂ (trace 2, open circles) was applied to the cells, and at t = 60 s, the medium was changed to K-PSS + 0.1 mM CaCl₂. Results are the average [Ca²⁺], values for 51 cells for trace 1 and 60 cells for trace 2. See MATERIALS AND METHODS for details of the calibration method. (B) Data in A on an expanded time scaled. The dotted lines are extrapolations of the linear phase of Ca²⁺ uptake. Similar results were obtained with four additional pairs of coverslips.
very slow for ~15 s but subsequently accelerated and attained a linear rate after an additional 10–20 s (see expanded scale in Fig. 1 B).

The other trace in Fig. 1 (labeled “2”, open circles) shows the effects of transiently increasing \([\text{Ca}^{2+}]_i\), before initiating reverse exchange activity. This was accomplished by applying Na-PSS containing 0.3 mM CaCl\(_2\) for 30 s before switching to K-PSS + 0.1 mM CaCl\(_2\). \([\text{Ca}^{2+}]_i\) increased from an initial value of 12 nM to 95 nM, and then declined to 80 nM just before the solution switch to K-PSS + 0.1 mM CaCl\(_2\). The increase in \([\text{Ca}^{2+}]_i\) in Na-PSS was due to \text{Ca}^{2+} entry through store-operated channels that had been activated by the prior release of \text{Ca}^{2+} from internal stores. Under these conditions, the initial delay in \text{Ca}^{2+} uptake was markedly reduced, and the cells attained a linear rate of uptake within ~10 s of the switch to K-PSS + 0.1 mM CaCl\(_2\) (Fig. 1 B). The reduced lag period is clearly evident from the dashed lines in Fig. 1 B, which represent the extrapolations of the linear phase of \text{Ca}^{2+} uptake back to its intersection with the time axis.

We interpret these results in the following way: For trace 1, exchange activity was initially low because \([\text{Ca}^{2+}]_i\) (~12 nM) was below the \(K_h\) for regulatory \text{Ca}^{2+} activation. As \([\text{Ca}^{2+}]_i\) slowly increased during the lag period, progressive activation of the exchanger by \text{Ca}^{2+} occurred, leading to an acceleration of \text{Ca}^{2+} uptake through positive feedback. For trace 2, the exchanger was partially activated by the \text{Ca}^{2+} entering through store-operated channels, leading to a marked reduction in the lag period and an increased initial rate of \text{Ca}^{2+} uptake. There was still a lag period of ~10 s for trace 2, suggesting that the exchanger was not fully activated by the level of \([\text{Ca}^{2+}]_i\) attained (~80 nM). Note that the store-operated channels did not contribute significantly to \text{Ca}^{2+} uptake in K-PSS because (a) the external \text{Ca}^{2+} concentration (0.1 mM) was too low to support extensive \text{Ca}^{2+} entry through store-operated channels and (b) channel activity was reduced in this depolarizing medium. In untransfected CHO cells, which lack \text{Na}^+/\text{Ca}^{2+} exchange, \([\text{Ca}^{2+}]_i\), increased by only ~30% in K-PSS + 0.1 mM CaCl\(_2\) (unpublished data).

Support for this interpretation is provided by the results of experiments with cells expressing a deletion mutant of NCX1.1, \(\Delta (241-680)\). This mutant is missing a large portion of the exchanger’s central hydrophilic domain and does not require \text{Ca}^{2+} for the activation of exchange activity. Trace 2 in Fig. 2 A (open circles) represents the average \([\text{Ca}^{2+}]_i\) for 59 \(\Delta (241-680)\) cells and shows that \text{Ca}^{2+} uptake began immediately upon initiating reverse exchange activity, i.e., without a lag beyond that required for mixing (~2–4 s). The trace with the filled circles is the same as trace 1 in Fig. 1, superimposed so that the time at which K-PSS + 0.1 mM CaCl\(_2\) was applied is the same for both traces. Fig. 2 B displays the rates of \text{Ca}^{2+} uptake for nine individual \(\Delta (241-680)\) cells. Although there was variation in the rates of \text{Ca}^{2+} uptake, no lag period was observed in any of these cells, or in the others not shown.

A very different pattern emerged when individual cells expressing the wild-type exchanger were examined. Fig. 3 A shows the fura-2 signals for 7 out of the 51 cells monitored for trace 1 in Fig. 1. As shown, the individual cells each displayed a lag period, but the duration of the lag period varied greatly. The trace labeled “1” in Fig. 3 A is for the cell that responded most rapidly in this experiment. The trace labeled “7” is for one
of the seven cells that failed to show Ca\(^{2+}\) uptake at all.
As shown in Fig. 3 B, there was a strong inverse correlation between the length of the lag period and the maximal rates of Ca\(^{2+}\) uptake (V\(_{\text{max}}\)) among the 44 responding cells in the population (P < \(10^{-3}\)). Because the cells showed such a high dispersion in their lag periods, further analysis of the results was done on a cell-by-cell basis.

Fig. 4 presents the rates of Ca\(^{2+}\) uptake plotted against the value of [Ca\(^{2+}\)]\(_{i}\) for each of the six responding cells shown in Fig. 3 A; the cells are numbered in order of increasing duration of the lag period. The rate of Ca\(^{2+}\) uptake at each data point was computed as the slope of the trace over an interval that included two data points on either side of the specified point (see materials and methods). A set of time points was chosen for analysis that ranged from the sigmoidal “foot” of the Ca\(^{2+}\) influx for seven individual cells. (A) Cells were selected from the coverslip used for trace 1 in Fig. 1. Upper arrow for each trace: data point corresponding to V\(_{\text{max}}\) (see materials and methods) Bottom arrow for each trace: data point corresponding to 0.5 V\(_{\text{max}}\). V\(_{\text{max}}\) is decorated with two arrows. The upper arrow corresponds to the data point at which the rate of Ca\(^{2+}\) uptake is maximal (V\(_{\text{max}}\)). The lower arrow (in bold) corresponds to the data point at which the slope is approximately half of the maximal slope. The [Ca\(^{2+}\)]\(_{i}\) value at this point provides an estimate of the K\(_{h}\) for allosteric Ca\(^{2+}\) activation. (B) Correlation between V\(_{\text{max}}\) and time-to-V\(_{\text{max}}\) using all the responding cells (V\(_{\text{max}}\) \(\geq\) 5 nM/s) from the coverslip for trace 1 in Fig. 1. r, correlation coefficient.

The above approach used only a small number of cells compared with the total monitored in each experiment. To estimate the value of K\(_{h}\) for the total population of responding cells, we used the spreadsheet matching procedure described in materials and methods; this approach is illustrated by the data for the six cells presented in Fig. 3 A. Each trace in Fig. 3 A is decorated with two arrows. The upper arrow corresponds to the data point where the rate of Ca\(^{2+}\) uptake is maximal (V\(_{\text{max}}\)). The lower arrow (in bold) corresponds to the data point at which the slope is approximately half of the maximal slope. The [Ca\(^{2+}\)]\(_{i}\) value at this point provides an estimate of the K\(_{h}\) for allosteric activation of the exchanger by Ca\(^{2+}\). The average value for the six cells analyzed in Fig. 3 A was 193 nM; this value is slightly less than that obtained by the curve-fitting procedure for the same cells (248 nM; paired t test P = 0.02) because the matching procedure finds data points where the rate may be less than precisely half of the maximal rate (see materials and methods). Excluding the seven cells that showed no Ca\(^{2+}\) uptake (V\(_{\text{max}}\) \(<\) 5 nM/s), the average K\(_{h}\) (±SD) for all of the cells on the coverslip was 237 ± 74 nM (n = 44). This analysis was performed for the six coverslips mentioned in the previous paragraph, three each for experiments performed at room temperature and 37°C. The values obtained at 37°C did not differ significantly from the data at room temperature and so the results for all 21 cells were pooled to yield the following mean values (±SEM): V\(_{\text{max}}\) = 153 ± 22 nM/s, n = 2.8 ± 0.15, K\(_{h}\) = 232 ± 18 nM.

We assume that these relations reflect the concentration dependence for regulatory activation of the exchanger by cytosolic Ca\(^{2+}\). V\(_{\text{max}}\) values varied over a fivefold range, from 35 nM/s (cell 6) to 182 nM/s (cell 2); the average value (± SD) for the Hill coefficient (n) was 2.6 ± 0.46. The values for K\(_{h}\) are presented in the panels for the individual cells. The average K\(_{h}\) (±SD) for these six cells was 248 ± 89 nM. This analysis was repeated for three additional cells from each of two additional coverslips; an additional nine cells were selected from three coverslips for experiments performed at 37°C. The values obtained at 37°C did not differ significantly from the data at room temperature and so the results for all 21 cells were pooled to yield the following mean values (±SEM): V\(_{\text{max}}\) = 153 ± 22 nM/s, n = 2.8 ± 0.15, K\(_{h}\) = 232 ± 18 nM.

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by curve-fitting for the 21 cells discussed in the previous paragraph. The values obtained by curve fitting were 33 ± 4.9% (±SEM) higher than the values obtained by the matching procedure. Correcting for this difference increased the average $K_h$ for the six coverslips to $\sim$300 nM.

**Persistence of Ca$^{2+}$ Activation**

In a previous publication (Vazquez et al., 1997), we demonstrated that exchange activity increased transiently after ATP-induced Ca$^{2+}$ release from internal stores. Stimulation was observed over a narrow range of low [Ca$^{2+}$] values and the results seemed compatible with our subsequent report of a high affinity for Ca$^{2+}$ activation. The following experiments were done to reevaluate this issue using the analytical approach described above. For the experiment in Fig. 5 A, the application of ATP (200 μM) and Tg (2 μM) in Na-PSS + 0.3 mM EGTA elicited a large Ca$^{2+}$ transient that decayed rapidly back toward baseline levels. 40 s after adding ATP/Tg, reverse exchange activity was initiated by applying K-PSS + 0.1 mM CaCl$_2$. [Ca$^{2+}$]$_i$ increased rapidly, without the lag period observed in the experiments described previously. The data are displayed on an expanded time scale in Fig. 5 B (filled circles), where at time $t = 0$ corresponds to the time at which K-PSS + 0.1 mM Ca$^{2+}$ was applied. The average [Ca$^{2+}$]$_i$ at the point where reverse exchange was initiated was 87 nM, well below the $K_h$ for activation of reverse exchange activity ($\sim$300 nM), and yet no lag period was observed. The traces with the open circles in Fig. 5 A and B, show the response of the same coverslip to a second application of K-PSS + 0.1 mM CaCl$_2$ after the cells had rested for a 10 min interval in Na-PSS + 0.1 mM EGTA. A long lag period was observed after the second application of K-PSS + 0.1 mM CaCl$_2$ (open circles), similar in duration to that observed for trace 1 in Fig. 1. We conclude that the exchanger remained activated for a while (see below) after Ca$^{2+}$ release from internal stores, even though global [Ca$^{2+}$]$_i$ had declined to values well below the $K_h$ for activation of exchange activity. We refer to this phenomenon as persistent Ca$^{2+}$ activation.

Fig. 5 C presents the results of an identical experiment performed with cells expressing the Δ(241–680) mutant, in which activity is not dependent on regulatory Ca$^{2+}$. As shown, the rates of Ca$^{2+}$ uptake were nearly the same for the first and second application of K-PSS + 0.1 mM CaCl$_2$, i.e., application of ATP + Tg did not stimulate the activity of the mutant exchanger, and no lag period was observed after the second application of K-PSS + 0.1 mM CaCl$_2$ (open circles), similar in duration to that observed for trace 1 in Fig. 1. We conclude that the exchanger remained activated for a while (see below) after Ca$^{2+}$ release from internal stores, even though global [Ca$^{2+}$]$_i$ had declined to values well below the $K_h$ for activation of exchange activity. We refer to this phenomenon as persistent Ca$^{2+}$ activation.

Analysis of the data for individual wild-type cells provides additional support for this interpretation. Fig. 6
displays the data for six cells selected from the population of 42 cells monitored in the experiment shown in Fig. 5 A. The traces following the first and second applications of K-PSS + 0.1 mM Ca$^{2+}$ are superimposed in Fig. 6 (filled and open circles, respectively), where $t = 0$ corresponds to the time at which exchange activity was initiated. In each case, [Ca$^{2+}$]$_i$ increased without a significant lag period when exchange activity was initiated 40 s after ATP + Tg (filled circles). The rates of Ca$^{2+}$ influx for cells 1 and 3 showed a tendency to accelerate after 20–30 s of uptake, and the $K_h$ values determined by the matching process described above were 295 nM and 231 nM, respectively. For the remaining cells, the matching analysis returned values that were essentially identical to the value of [Ca$^{2+}$]$_i$ at $t = 0$ ($K_h$ values given in figure).

For the second application of K-PSS + 0.1 mM CaCl$_2$ (open circles), lengthy lag periods were observed, followed by a strong acceleration of Ca$^{2+}$ uptake after periods ranging from 30 to >100 s, i.e., the cells behaved essentially identically to those shown in Fig. 3. The matching procedure returned $K_h$ values between 241 and 315 nM for five of the cells shown after the second application of K-PSS + 0.1 mM CaCl$_2$. The rate of Ca$^{2+}$ influx for cell 6 was <5 nM/s after the second application of K-PSS + 0.1 mM CaCl$_2$, and so it was not analyzed; note that this cell showed a much higher rate of Ca$^{2+}$ uptake when K-PSS + 0.1 mM CaCl$_2$ was applied after ATP addition (filled circles).

The following analyses pertain to the 31 cells on this coverslip that responded to both applications of K-PSS + 0.1 mM CaCl$_2$ ($V_{max}$ $\approx$ 5 nM/s). When exchange activity was initiated 40 s after ATP/Tg addition, the time required to attain a maximal rate of Ca$^{2+}$ uptake ($V_{max}$) after the initiation of exchange activity was 22 ± 12 s, as compared with 141 ± 57 s when activity was initiated 10 min later (± SD; P $<$ 10$^{-6}$, paired t test). The average $K_h$ values (±SD) measured 40 s and 10 min after ATP/Tg addition were 132 ± 133 nM and 269 ± 153 nM, respectively (P $<$ 2 × 10$^{-5}$, paired t test). It is notable that of the 11 cells that did not show significant Ca$^{2+}$ uptake after the second addition of K-PSS + 0.1 mM CaCl$_2$, 10 showed measurable Ca uptake when assayed 40 s after ATP addition, as exemplified by cell 6 in Fig. 6. Similar results were obtained with two additional coverslips. The behavior of the individual cells strongly supports our conclusion that exchanger remains in an activated state following the return of [Ca$^{2+}$]$_i$ to values below $K_h$.

The effect of the exposure to high Ca$^{2+}$ dissipates within about a minute after adding ATP. Fig. 7 shows three experiments where exchange activity was initiated 35, 45, or 60 s following the addition of ATP (65,
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75 and 90 s in the figure; see arrows at top of panel). Fig. 7 A shows the superimposed traces for the average [Ca²⁺], values for each coverslip and Fig. 7, B–D, show several individual cells for each experiment. For the experiment where exchange activity was initiated 35 s after ATP addition (Fig. 7 B), rapid Ca²⁺ influx was observed very shortly after the solution change to K-PSS/0.1 mM CaCl₂ (indicated by the arrow labeled “Ca” in the figure). The average time (±SD) required to attain V_max was 24 ± 15 s after the solution change (n = 64 cells). When exchange activity was initiated 45 s after ATP addition (Fig. 7 C), significant lag periods were observed for some cells; the average time required to attain V_max for this experiment was 44 ± 20 s after the solution change (n = 62). When exchange activity was initiated 60 s after ATP addition (Fig. 7 D), a well-developed pattern of variable lag periods was observed, with an average time to V_max of 74 ± 30 s (n = 68). The differences between the various times were highly significant (P << 10⁻⁶, unpaired t test). We conclude that the activated state of the exchanger dissipates over a time course of tens of seconds.

In addition to inducing an increase in [Ca²⁺], the activation of purinergic receptors initiates other signaling processes, such as activation of protein kinase C through diacylglycerol production, that could potentially contribute to persistent Ca²⁺ activation. To assess whether a rise in [Ca²⁺], was sufficient by itself for persistent exchanger activation, we incubated the cells in Na-PSS + 0.3 mM EGTA for 10 min after treating them with ATP + Tg, and then measured exchange activity after a transient application of external Ca²⁺. The results of one such experiment are shown in Fig. 8 A. The trace with the filled circles shows a typical response of the cells to the application of K-PSS + 0.1 mM CaCl₂ at t = 30 s; note the lengthy lag period. Na-PSS + 0.5 mM EGTA was then applied and at 300 s, 1 mM Ca²⁺ was applied in Na-PSS (open circles). Ca²⁺ entered the cells through store-operated channels and [Ca²⁺] increased to a plateau value of ~150 nM. It should be noted that this value of [Ca²⁺] was much lower than that observed for parental CHO cells under the same conditions, where [Ca²⁺] values of ~800 nM were attained (unpublished data). After 1 min, the external Ca²⁺ was removed with Na-PSS + 0.3 mM EGTA, and 20 s later, K-PSS + 0.1 mM CaCl₂ was applied to initiate a second round of reverse Na⁺/Ca²⁺ exchange. As shown in Fig. 8 A, [Ca²⁺], increased with only a short lag period. The data for the two traces are superimposed on an expanded time scale in Fig. 8 B where t = 0 corresponds to the time of application of K-PSS + 0.1 mM CaCl₂. Note that [Ca²⁺], just before the initiation of reverse exchange activity was the same (~25 nM) for the two traces.

Figure 6. Data for six cells from the coverslip used for the experiment shown in Fig. 5. Symbols as in Fig. 5 (see legend to Fig. 5 and text for further details). Apparent K_h values determined by the matching procedure are given next to each trace. ND, not determined.
Fig. 8, C–E, show how three individual cells behaved in this experiment. In each case, \([\text{Ca}^{2+}]_i\) increased more rapidly after the transient elevation of \([\text{Ca}^{2+}]_i\), although the \([\text{Ca}^{2+}]_i\) values at the time of initiation of exchange activity were identical in the two circumstances. The time course of \([\text{Ca}^{2+}]_i\) uptake was sigmoidal in each case, with similar apparent \(K_h\) values, determined by the matching procedure described above. The average \(K_h\) (\(\pm\) SD) required to attain \(V_{\text{max}}\) was 42 \(\pm\) 22 s when exchange activity was initiated following the transient increase in \([\text{Ca}^{2+}]_i\), whereas a value of 77 \(\pm\) 29 s was obtained for the cells analyzed in the absence of a prior \(\text{Ca}^{2+}\) transient (\(P < 10^{-5}\), paired \(t\) test; \(n = 54\)). For the 29 cells that responded to both applications of K-PSS + 0.1 mM \(\text{Ca}^{2+}\) (\(V_{\text{max}} \geq 5\) nM/s), the average \(K_h\) values (\(\pm\) SD) were 194 \(\pm\) 69 nM and 167 \(\pm\) 53 nM with or without a prior \(\text{Ca}^{2+}\) transient, respectively; the difference was not statistically significant (\(P = 0.13\), paired \(t\) test). (In this experiment, 17 cells showed \(V_{\text{max}} > 5\) nM/s after the \(\text{Ca}^{2+}\) transient although they did not respond in the absence of a prior \(\text{Ca}^{2+}\) transient; when these cells were included in the analysis, the \(K_h\) following the \(\text{Ca}^{2+}\) transient became 177 \(\pm\) 81 nM.) Thus, although the cells responded more rapidly after \(\text{Ca}^{2+}\) application in Na-PSS, their \(K_h\) for allosteric \(\text{Ca}^{2+}\) activation was identical to the same cells without prior \(\text{Ca}^{2+}\) application.

Fig. 9 shows the results of a similar experiment, except that in this case 1 mM \(\text{CaCl}_2\) was applied in Li-PSS instead of Na-PSS, \(\text{Ca}^{2+}\) entry under these conditions was due to both store-dependent \(\text{Ca}^{2+}\) channels and reverse \(\text{Na}^{+}/\text{Ca}^{2+}\) exchange, and \([\text{Ca}^{2+}]_i\) attained levels comparable to that seen after ATP-induced \(\text{Ca}^{2+}\) release. Two superimposed traces are shown in Fig. 9 A, representing average \([\text{Ca}^{2+}]_i\) values for all the cells monitored and corresponding to two pulses of Li-PSS +
1 mM Ca²⁺ applied to the same coverslip, separated by an interval of 5 min. 50 s after terminating the first pulse with Na-PSS + 0.3 mM EGTA, reverse exchange activity was initiated by applying K-PSS + 0.1 mM CaCl₂ (trace 1, filled circles). The cells were returned to Na-PSS + 0.3 mM EGTA and, after 5 min, a second 1-min pulse of Li-PSS + 1 mM CaCl₂ was applied; in this case, exchange activity was initiated 60 s after terminating the pulse (trace 2, open circles). The data are shown on an expanded time scale in Fig. 9 B but in this case, t = 0 corresponds to the time of initiation of reverse exchange activity with K-PSS + 0.1 mM CaCl₂ in each case. Symbols as in A. 

The results for four individual cells are shown in Fig. 10, C–F; the cells all show Ca²⁺ uptake without a lag period for trace 1, with variable lag periods for trace 2. Fig. 10 E shows a cell with high exchange activity that did not show a lag period for either trace 1 or trace 2. For trace 1, the average time (±SD) to reach V_max after the application of K + 0.1 mM CaCl₂ was 20 ± 9.6 s, while the corresponding value for trace 2 was 50 ± 33 s (P << 10⁻⁶, paired t test; n = 62).

The data in Figs. 8 and 9 indicate that the persistence of the activated state was induced by an increase in [Ca²⁺]ᵢ itself, and that other signaling processes initiated by purinergic receptor activation were not involved. The degree of activation appeared to be less for the experiment from Fig. 8, where the Ca²⁺ pulse was applied in Na-PSS and [Ca²⁺]ᵢ values of only ~150 nM were attained; the subsequent traces for reverse exchange activity showed some sigmoidal character, and the average time required to attain V_max was 42 s. For the experiment in Fig. 9, where [Ca²⁺]ᵢ levels were comparable to those attained with ATP, most cells responded immediately when exchange activity was initiated 30 s after termination of the Ca²⁺ transient, with an average time-to-V_max of 20 s (compare with 22 and 24 s for the experiments with ATP in Figs. 5 and 7 B).
We conclude that the degree of persistent activation of the exchanger increases as \([\text{Ca}^{2+}]_i\) increases.

**Local \(\text{Ca}^{2+}\) Gradients and Persistent \(\text{Ca}^{2+}\) Activation**

The persistence of regulatory \(\text{Ca}^{2+}\) activation in these experiments is compatible with either of three general mechanisms: (a) the exchanger retains the molecular configuration of the activated state after a reduction in \([\text{Ca}^{2+}]_i\), (b) the local concentration of \(\text{Ca}^{2+}\) in the vicinity of the exchanger remains elevated despite the fall of global \([\text{Ca}^{2+}]_i\), or (c) the elevated \([\text{Ca}^{2+}]_i\) initiates a signaling pathway, e.g., calmodulin-dependent protein kinase, which maintains the activated state. A full exploration of these possibilities is beyond the scope of the present report. Electrophysiological measurements of exchange currents in excised patches provide support for the first mechanism (see discussion).

In Fig. 10, we present data suggesting that local \(\text{Ca}^{2+}\) gradients are also involved. In this experiment, we used \(\text{La}^{3+}\) as a substitute for \(\text{Ca}^{2+}\) for measuring exchange activity. We recently reported that the exchanger transports \(\text{La}^{3+}\) at a low rate, and that \(\text{La}^{3+}\) interacts with the regulatory \(\text{Ca}^{2+}\) binding sites at low concentrations to activate the exchanger (Reeves and Condrescu, 2003). The advantage of using \(\text{La}^{3+}\) for the present purpose is that \(\text{La}^{3+}\) does not enter the cell through store-operated channels and does not appear to be transported into the ER by SERCA activity.

Two superimposed traces are shown in Fig. 10 A, corresponding to the addition of ATP, with or without 2 \(\mu\)M Tg, at \(t = 30\) s. At \(t = 50\) s, the effects of ATP were terminated by the addition of 0.5 mg/ml of hexokinase. With 10 mM glucose in the medium, the hexokinase rapidly converts ATP to ADP and terminates the ATP-induced \(\text{Ca}^{2+}\) release process since ADP is inactive with respect to the purinergic receptors in CHO cells.
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Figure 10. Local Ca²⁺ gradients and persistent Ca²⁺ activation. (A) Protocol: At t = 30 s, ATP (50 μM) in Na-PSS + 0.1 mM EGTA, with (filled circles) or without (open circles) 2 μM Tg, was applied. At t = 50 s, 1 ml of Na-PSS + 0.1 mM EGTA containing 0.5 mg/ml hexokinase (hex) was added, and at t = 65 s, K-PSS + 0.1 mM LaCl₃ was applied. (B) Effect of hexokinase in nontransfected CHO-T1 cells. ATP (50 μM) was applied at t = 30 s, 0.5 mg/ml hexokinase at t = 50 s, and K-PSS + 0.1 mM LaCl₃ was applied. For the control trace (open circles), 50 μM ATP as applied at 30 s and no other additions were made. (C) Filled circles: cells were treated with ATP + Tg 10 min before beginning of experiment and K-PSS + 0.1 mM LaCl₃ was applied at 65 s. Open circles: cells were not treated with ATP or Tg, but were washed with Na-PSS + 0.1 mM EGTA immediately before beginning the experiment and K-PSS + 0.1 mM LaCl₃ was applied at t = 65 s. For all traces, Na-PSS + 0.3 mM EGTA was applied at 4 min. (D) Rates of exchange-mediated La³⁺ uptake between 100–120 s for the conditions indicated. †, P < 0.002 versus +ATP, +Tg; ‡, P < 0.005 versus +ATP, +Tg, P < 0.01 vs. +ATP, −Tg (unpaired t test, n = 4).

(see below). K-PSS + 0.1 mM LaCl₃ was applied at t = 65 s to initiate exchange-mediated La³⁺ influx. For the trace obtained in the presence of Tg (filled circles), La³⁺ uptake began immediately upon the initiation of exchange activity. For the trace with the open circles, Tg was omitted, leaving SERCA activity intact; the rate of La³⁺ uptake was ∼1/3 the rate seen in the presence of Tg. The pooled results of four such experiments are presented in Fig. 10 D; the rate of La³⁺ uptake in the presence of Tg was significantly greater than in the absence of Tg (P < 0.002, unpaired t test). We conclude that the SERCA pump accelerates the relaxation of the activated state of the exchanger, presumably by transferring residual cytosolic Ca²⁺ into the ER. The presence of La³⁺ interfered with the calibration procedure in these studies but, by analogy to the results in Figs. 5–7, [Ca²⁺], had undoubtedly returned to values well below the Kₘ for exchange activation at the time La³⁺ uptake was initiated. The simplest interpretation of the results is that SERCA acts to reduce a local elevation of [Ca²⁺] that maintains exchanger activation, either by directly activating the exchanger or by maintaining the activity of a Ca²⁺-dependent signaling process.

The trace with the filled circles in Fig. 10 B documents the effectiveness of hexokinase in terminating the effects of ATP. Nontransfected CHO cells were used in this experiment; 50 μM ATP was added at 30 s, and hexokinase was added at 50 s. Note that the nontransfected cells show a much longer Ca²⁺ transient than the NCX-expressing cells; the difference between the two cell types undoubtedly reflects the efficiency of the exchanger in clearing Ca²⁺ from the cytosol. Upon the addition of hexokinase, [Ca²⁺], rapidly fell to low values (filled circles) compared with a control trace in which hexokinase was not added (open circles). For the hexokinase-treated cells, K-PSS + 0.1 mM LaCl₃ was applied at 70 s; as expected, no La³⁺ uptake was observed in the absence of NCX expression (Reeves and Condrescu, 2003).

Fig. 10 C displays La³⁺ uptake under two control conditions. For trace 1, the cells were treated with ATP and Tg 10 min before beginning the experiment. For trace 2, internal Ca²⁺ stores were left intact and the cells...
were simply washed with Na-PSS + 0.1 mM EGTA immediately before beginning the experiment. K-PSS + 0.1 mM LaCl₃ was applied at t = 65 s for both traces. The rates of La³⁺ uptake were similar in both instances and were significantly lower than seen after ATP addition either in the presence of Tg (P < 0.005, unpaired t test) or in the absence of Tg (P < 0.01, unpaired t test) (Fig. 10 D).

**DISCUSSION**

Much of our present understanding of allosteric Ca²⁺ activation stems from measurements of outward exchange currents in excised patches from cardiac myocytes or from *Xenopus* oocytes expressing NCX1.1. Kₘ values of 200–600 nM were obtained for the Ca²⁺-dependence of peak outward currents, with Hill coefficients close to 1 (Collins et al., 1992; Hilgemann et al., 1992a; Matsuoka et al., 1995, 1997; Fujioka et al., 2000a). Activation of exchange currents by Ca²⁺ is a rapid process, occurring in <200 ms (Hilgemann et al., 1992a; Weber et al., 2001), although one study reported a time constant for activation of ~600 ms (Kappl and Hartung, 1996). Only a few studies have examined the effects of ATP on Ca²⁺ activation. ATP modulates exchanger regulation by effecting the synthesis of phosphatidylinositol-4,5-bisphosphate in the patch membrane (Hilgemann and Ball, 1996). In some, but not all patches, ATP reduced the Kₘ for Ca²⁺ activation and increased the Hill coefficient (Hilgemann et al., 1992a). In experiments with patches from mouse myocytes, outward exchange currents became independent of cytosolic Ca²⁺ after application of ATP (Hilgemann and Ball, 1996; see Weber et al., 2001 for additional information on mouse myocytes).

In excised patches, the removal of cytosolic Ca²⁺ in the continued presence of cytosolic Na⁺ led to a time-dependent decay of exchange currents, reflecting relaxation of the Ca²⁺-activated state. The time constants for this decay were ~10 s in most experiments (Hilgemann et al., 1992a; Matsuoka et al., 1995); however, a much more rapid decay (~0.5 s) was observed in one study using a “macro-patch” from cardiac myocytes (Fujioka et al., 2000a). ATP was shown to slow the decay of the activated state following Ca²⁺ removal (see Fig. 12 in Hilgemann et al., 1992a). Since [Ca²⁺] was well controlled through the use of buffering agents in these experiments, these results (with one exception) demonstrate that the Ca²⁺-activated state persists for several seconds after a reduction of [Ca²⁺], to very low values. There is no comparable information available on the relaxation of the Ca²⁺-activated state of the exchanger in intact cells. Since intact cells have high ATP levels and their plasma membranes are presumably well stocked with phosphatidylinositol-4,5-bisphosphate, the regulatory behavior of the exchanger may be different in cells than in excised patches.

For the experiments reported here, the findings fall into four categories: (1) the unusual time course for exchange-mediated Ca²⁺ uptake, (2) the concentration dependence for allosteric Ca²⁺ activation of exchange activity, (3) the persistence of Ca²⁺ activation after the return of [Ca²⁺], to low values, and (4) the involvement of local ion gradients in persistent Ca²⁺ activation. A final category is the possible relevance of these findings to the physiological regulation of exchange activity in functioning cardiac myocytes.

**Time Course of Ca²⁺ Uptake**

The sigmoidal time course of Ca²⁺ uptake during reverse exchange activity (Fig. 1) can be divided into a lag phase, during which [Ca²⁺], increased only slowly, an acceleration phase, during which the rate of Ca²⁺ uptake increased rapidly toward Vₘₐₓ, a plateau of relatively high [Ca²⁺],, and a subsequent decline in [Ca²⁺],. The highly variable lag periods displayed by individual cells (Fig. 5) initially seemed quite puzzling. When we found that the lag periods could be reduced by elevating [Ca²⁺], (Fig. 1), and that cells expressing the Δ(241–680) deletion mutant of NCX1.1 did not display a sigmoidal time course (Fig. 2), it became evident that regulatory Ca²⁺ activation was involved. This interpretation was supported by our recent results with La³⁺, which activated exchange activity at very low concentrations (10–20 pM) and itself displayed a sigmoidal time-course of uptake. Indeed, preloading the cells with a small quantity of La³⁺ (~15 pM) abrogated the lag period during subsequent exchange-mediated Ca²⁺ uptake and stimulated the initial rate of Ca²⁺ uptake 20-fold (Reeves and Condrescu, 2003).

Ca²⁺ influx during the lag period seems to be mediated almost entirely by a low level of reverse exchange activity, although a small contribution by store-dependent Ca²⁺ channels cannot be completely excluded. Blockade of these channels with 50 μM SK&F 96365 did not obviously alter the pattern of lag periods displayed by the cells (unpublished data). The inverse correlation between the duration of the lag period and the Vₘₐₓ for Ca²⁺ uptake (Fig. 3 B) suggests that variability in exchange activity among the individual cells is an important factor in the high variability of the lag periods. Consistent with this interpretation, lag periods were shortened when exchange activity was increased by using K-PSS containing 1 mM Ca²⁺ instead of 0.1 mM Ca²⁺, or when cytosolic [Na⁺] was stabilized at 20 mM using gamicidin (unpublished data). An additional factor that must be considered is Ca²⁺ efflux by the plasma membrane Ca²⁺ ATPase, which would reduce net Ca²⁺ influx, particularly when exchange activity is low, and prolong the lag period (see below). In prin-
ciple, Ca\textsuperscript{2+} accumulation by internal compartments might also contribute to the duration of the lag periods. However, the Tg treatment blocked Ca\textsuperscript{2+} uptake into the ER, and mitochondria did not accumulate significant amounts of Ca\textsuperscript{2+} at the low [Ca\textsuperscript{2+}], values present during the lag phase (unpublished data). Although nonmitochondrial, Tg-resistant compartments are present in CHO cells (Condrescu et al., 1997), we could find no evidence that these compartments accumulated Ca\textsuperscript{2+} during the lag phase.

The acceleration phase reflected progressive activation of the exchanger as Ca\textsuperscript{2+} accumulated, leading to a rapid increase in Ca\textsuperscript{2+} uptake as [Ca\textsuperscript{2+}], approached the $K_h$ for the allosteric Ca\textsuperscript{2+} binding sites. Mitochondrial Ca\textsuperscript{2+} accumulation occurred during the acceleration phase (Opuni and Reeves, 2000; unpublished data). As discussed in the appendix, mitochondrial Ca\textsuperscript{2+} uptake produced a significant underestimate (44%) of the $K_h$ for allosteric Ca\textsuperscript{2+} activation of exchange activity. The plateau value appears to reflect a balance between continuing Ca\textsuperscript{2+} influx via the exchanger and mitochondrial Ca\textsuperscript{2+} accumulation (see appendix). An important factor in the decline in [Ca\textsuperscript{2+}], after the plateau phase is likely to be the loss of Na\textsuperscript{+} from the cell under the conditions of these experiments. Indeed, the use of ionophores such as gramicidin to stabilize [Na\textsuperscript{+}], at values ranging from 5 to 20 mM reduced the rate of decline, although it did not completely eliminate it. Importantly, $K_h$ values obtained with gramicidin-treated cells were similar to the values reported here throughout the range of [Na\textsuperscript{+}] studied (unpublished data), suggesting that the presence of the declining phase did not distort our analyses of allosteric Ca\textsuperscript{2+} activation.

**[Ca\textsuperscript{2+}] Dependence of Exchanger Activation**

Potential sources of error in our estimates of $K_h$ are considered in the appendix. In addition to the tendency of the matching procedure to underestimate $K_h$ by 33% compared with the curve-fitting procedure (see results), mitochondrial Ca\textsuperscript{2+} accumulation produces an additional underestimate of 44%, while the effects of fura-2 buffering on the measured velocities of Ca\textsuperscript{2+} uptake lead to an overestimate of $K_h$ in the range of 25–40% (see appendix). Using the overall value of $K_h$ obtained by the matching procedure from six coverslips ($231 \pm 14$ nM; see results), and applying the corrections indicated above, we conclude that the $K_h$ for allosteric Ca\textsuperscript{2+} activation is $\sim300$ nM.

This value is compatible with the electrophysiological measurements obtained in excised patches but does not agree with other measurements, including our own, that suggested a much higher affinity for the Ca\textsuperscript{2+} regulatory sites. Earlier electrophysiological measurements in cardiac myocytes (Noda et al., 1988; Miura and Kimura, 1989) may have suffered from inadequate control of [Ca\textsuperscript{2+}], despite the use of EGTA in the pipette solutions. The Ba\textsuperscript{2+} influx measurements that we conducted in transfected CHO cells (Fang et al., 1998) involved pretreating the cells with ionomycin and various external concentrations of Ca\textsuperscript{2+}, and then initiating exchange-mediated Ba\textsuperscript{2+} influx during the return of [Ca\textsuperscript{2+}], toward low values. This experimental design was similar in many respects to the experiments presented in Figs. 9 and 10, and would have induced persistent Ca\textsuperscript{2+} activation in the exchanger, accounting for the high rates of Ba\textsuperscript{2+} influx that we observed at low [Ca\textsuperscript{2+}], levels (Fang et al., 1998). Thus, the $K_h$ value of 44 nM we obtained in the previous studies reflected the decay of the Ca\textsuperscript{2+} activated states that were induced by a range of much higher [Ca\textsuperscript{2+}], values attained during the ionomycin/Ca\textsuperscript{2+} pretreatment.

The electrophysiological experiments of Weber et al. (2001) yielded a $K_h$ value of $\sim125$ nM for allosteric activation of exchange activity in ferret myocytes. Their experimental protocol involved estimating the $K_h$ from the [Ca\textsuperscript{2+}], dependence of the envelope of the outward branch of alternating inward and outward exchange currents. Their approach assumed that the allosteric activation machinery was in instantaneous equilibrium with the bulk [Ca\textsuperscript{2+}], an assumption that should be re-examined in view of the possible persistence of the Ca\textsuperscript{2+}-activated state in these cells.

The fits of the rate data to the Hill equation (Fig. 4) suggest that Ca\textsuperscript{2+} activation shows a high degree of cooperativity (Hill coefficient 2.8). This conclusion must be viewed cautiously, however, because the net rates of Ca\textsuperscript{2+} influx reflect not only exchange activity but the Ca\textsuperscript{2+} efflux activity of the plasma membrane Ca\textsuperscript{2+}-ATPase as well. The fits to the Hill equation are highly dependent on the net rates of Ca\textsuperscript{2+} uptake at the sigmoidal “foot” of the plots shown in Fig. 4, i.e., during the transition from the lag phase to the acceleration phase when exchange activity is low. The relative influence of Ca\textsuperscript{2+} efflux by the plasma membrane Ca\textsuperscript{2+}-ATPase on net Ca\textsuperscript{2+} influx will be profound when exchange activity is high, but less important when exchange activity is low and would therefore tend to increase the sigmoidal shape of the curve, perhaps yielding artificially high values for the Hill coefficient. Thus, the degree of cooperativity of allosteric Ca\textsuperscript{2+} activation in these experiments must remain uncertain.

**Persistence of the Ca\textsuperscript{2+}-activated State**

A transient elevation in [Ca\textsuperscript{2+}], to levels above the $K_h$ for Ca\textsuperscript{2+} activation led to a persistent state of activation with a lifetime of several tens of seconds (Figs. 7 and 9). This was observed whether the Ca\textsuperscript{2+} transient was induced by either ATP-mediated Ca\textsuperscript{2+} release (Figs. 5–7) or Ca\textsuperscript{2+} influx through a store-operated Ca\textsuperscript{2+} channels...
and reverse exchange activity (Figs. 8 and 9). In many cases, the initial time course of Ca\(^{2+}\) influx was linear under conditions where bulk cytosolic [Ca\(^{2+}\)] was well below the \(K_{h}\) for Ca\(^{2+}\) activation (Figs. 6 and 9). A linear time course of Ca\(^{2+}\) influx indicates that the exchanger brought about Ca\(^{2+}\) entry at a rate equal to \(V_{\text{max}}\), and was therefore fully activated despite the low [Ca\(^{2+}\)].

Persistent Ca\(^{2+}\) activation explains certain aspects of our earlier report showing that exchange activity was stimulated by ATP-induced Ca\(^{2+}\) release (Vazquez et al., 1997). A transient acceleration of Ca\(^{2+}\) or Ba\(^{2+}\) influx was observed even after the Ca\(^{2+}\) transient had decayed and [Ca\(^{2+}\)], had returned to values only slightly higher than control cells that had not been exposed to ATP. At the time, the results seemed consistent with a high affinity of the exchanger for regulatory Ca\(^{2+}\), but are now more readily explained by persistent Ca\(^{2+}\) activation.

Transient elevation of [Ca\(^{2+}\)], to levels less than the \(K_{h}\) for Ca\(^{2+}\) activation (e.g., \(\sim 150\) nM for the experiment in Fig. 8) led to partial activation of the exchanger, as indicated by the reduced duration of the lag period (Figs. 8, B–E). The cells exhibited a sigmoidal time course of Ca\(^{2+}\) uptake in the partially activated state and displayed \(K_{h}\) values that were not significantly different from those observed in the absence of a prior Ca\(^{2+}\) transient (see discussion of Fig. 8 in RESULTS). Indeed, the behavior of the cells under these conditions was essentially identical to that observed in trace 2 in Fig. 1, where a modest increase in [Ca\(^{2+}\)], shortened the duration of the lag period but did not eliminate the sigmoidity of the time course of Ca\(^{2+}\) uptake. In the experiment shown in Fig. 1 (trace 2), exchange activity was initiated while [Ca\(^{2+}\)], was still elevated, whereas in Fig. 8, [Ca\(^{2+}\)], was returned to low values for a short interval before initiating reverse exchange activity. It appears that the partially activated state of the exchanger persists to a degree that is comparable to the fully activated state induced by much higher levels of [Ca\(^{2+}\)].

**Local Ca\(^{2+}\) Gradients and Persistent Ca\(^{2+}\) Activation**

The effects of SERCA activity on exchange-mediated La\(^{3+}\) uptake (Fig. 10) indicate that local Ca\(^{2+}\) gradients contribute to maintaining the activated state of the exchanger. Thus, when a Ca\(^{2+}\) transient was elicited by ATP in the absence of Tg, the subsequent rate of exchange-mediated La\(^{3+}\) uptake was only about one-third of that seen in the presence of Tg. Even in the absence of Tg, however, La\(^{3+}\) uptake remained greater than that seen in the absence of a prior Ca\(^{2+}\) transient (Fig. 10, C and D), suggesting that SERCA activity did not completely abrogate persistent Ca\(^{2+}\) activation but merely accelerated its decay. Similar results were obtained with exchange-mediated Ba\(^{2+}\) uptake (unpublished data). Since global [Ca\(^{2+}\)], was well below \(K_{h}\) in these experiments, the results suggest that in the absence of SERCA activity, the local [Ca\(^{2+}\)] beneath the plasma membrane may have been high enough to maintain exchanger activation. Another possibility is that a local rise in Ca\(^{2+}\) activates a Ca\(^{2+}\)-dependent signaling process (e.g., calmodulin-dependent protein kinase) that secondarily maintains exchanger activation. In either case, SERCA activity would presumably reduce the local Ca\(^{2+}\) concentration and accelerate the relaxation of the activated state.

A gradient of free Ca\(^{2+}\) would be expected to dissipate by diffusion within a very brief period after a reduction in global [Ca\(^{2+}\)]. Therefore, we speculate that the most likely source for a local elevation in Ca\(^{2+}\) is the loss of Ca\(^{2+}\) from a membranous compartment that may be in close proximity to the plasma membrane. Mitochondria are unlikely to be involved since persistent Ca\(^{2+}\) activation was not blocked by CI-CCP (unpublished data). One candidate for such a compartment is the ER, consistent with the close apposition of the ER to the plasma membrane and the efficiency of the SERCA pump in accelerating the decay of the activated state (Fig. 10). For the experiments with ATP-induced Ca\(^{2+}\) release (Figs. 5–7, and 10), residual stores of un-released Ca\(^{2+}\) within the ER could be the source of a local Ca\(^{2+}\) elevation. For the experiments where the activating elevation of [Ca\(^{2+}\)], was initiated by store-operated Ca\(^{2+}\) channel activity (Figs. 8 and 9), some of the Ca\(^{2+}\) taken up by the cells may have been delivered to the ER by a SERCA-independent mechanism. Clearly, further investigation is needed to settle these speculative issues.

The persistence of Ca\(^{2+}\) activation suggests that exchange activity may monitor the integral of Ca\(^{2+}\) transients over multiple beats in functioning cardiac myocytes. Its integrating function would be markedly affected by conditions that influence the relaxation rate of the activated state. This possibility has profound implications with regard to the importance of exchange activity in mediating chronotropic and ionotropic alterations in cardiac function. For this reason, it will be important in future studies to determine if the Ca\(^{2+}\)-activated state of the exchanger persists in intact cardiac myocytes, and to identify potential mechanisms for regulating its relaxation rate.

**Appendix — Possible Sources of Error**

**Rate of Exchanger Activation**

In this study, we assumed that activation of the exchanger by Ca\(^{2+}\) is instantaneous and that the \(K_{h}\) for Ca\(^{2+}\) activation can be determined from the values of [Ca\(^{2+}\)], at each time point during the course of Ca\(^{2+}\) uptake. This assumption is compatible with the rapid rates of Ca\(^{2+}\) activation observed in giant patch studies.
Persistent Activation of Na\(^+\)/Ca\(^{2+}\) Exchange

(see discussion). However, if an appreciable number of exchangers had entered an inactive state that is induced by cytosolic Na\(^+\) (Na\(^+\)-dependent or I\(_1\) inactivation, see Hilgemann, 1996), the rate of Ca\(^{2+}\) activation would be very much slower (Hilgemann et al., 1992b). In this case, K\(_h\) could be overestimated since activation would occur at times earlier than the time point identified by the matching procedure, i.e., at lower \([\text{Ca}^{2+}]\). Na\(^+\)-dependent inactivation is most pronounced at high cytosolic Na\(^+\) concentrations (~100 mM) and is counteracted by ATP, which acts by effecting the synthesis of phosphatidylinositol-4,5-bisphosphate (Hilgemann and Ball, 1996). Since \([\text{Na}^{+}]\), under normal conditions is low and ATP (and presumably phosphatidylinositol-4,5-bisphosphate) levels are high, we infer that most exchange carriers are not in the Na\(^+\)-inactivated state and that Ca\(^{2+}\) activation occurs rapidly.

This inference is supported by experiments with cells expressing a mutant exchanger that is resistant to I\(_1\) inactivation (Condrescu and Reeves, 1999); these cells behave essentially identically to cells expressing the wild-type exchanger and display a K\(_h\) of ~290 nM (unpublished data). Other lines of evidence consistent with a rapid activation of the exchanger by Ca\(^{2+}\) include the following: (a) There is no correlation between \(V_{\text{max}}\) and K\(_h\) among the individual cells in a population or under experimental conditions resulting in differing exchange rates (e.g., 1.0 vs. 0.1 mM external Ca\(^{2+}\)) (unpublished data). If Ca\(^{2+}\) activation were slow, K\(_h\) would be progressively overestimated as the rate of Ca\(^{2+}\) uptake increased and a positive correlation between K\(_h\) and exchange rates would be observed. (b) No systematic variations in K\(_h\) values were observed in experiments where gramicidin was used to stabilize cytosolic [Na\(^+\)] between 5 and 20 mM. If I\(_1\) inactivation were a factor in our experiments, we would expect...
more inactivated exchangers at the higher [Na\(^+\)], and a corresponding increase in \(K_h\). Together, our results are consistent with the assumption that little I\(_1\) inactivation occurs under our experimental conditions and that activation of the exchanger by Ca\(^{2+}\) therefore occurs rapidly.

**Mitochondrial Ca\(^{2+}\) Uptake**

Mitochondrial Ca\(^{2+}\) uptake during reverse exchange activity (Opuni and Reeves, 2000), could lead to an underestimate of the true rate of Ca\(^{2+}\) influx under the conditions of these experiments. When assays were conducted in the presence of 5 \(\mu\)M Cl-CCP, an uncoupling agent that depolarizes the mitochondrial membrane and blocks active mitochondrial Ca\(^{2+}\) uptake, we found \(K_h\) values slightly higher than those obtained in the absence of uncouplers (unpublished data). To explore the effect of mitochondrial Ca\(^{2+}\) uptake more precisely, we performed the experiment presented in Fig. 11. Fig. 11 A presents the average \([Ca^{2+}]_i\) values for 53 cells from a single coverslip in which K-PSS + 0.1 mM Ca\(^{2+}\) was applied twice, once in the absence of Cl-CCP (closed circles) and again, after a 10-min interval in Na-PSS + 0.1 mM EGTA, in the presence of 5 \(\mu\)M Cl-CCP (open circles). The two traces are superimposed in Fig. 11 B, where \(t = 0\) corresponds to the application of K-PSS + CaCl\(_2\) for each trace. \([Ca^{2+}]_i\), attained higher values in the presence of Cl-CCP, consistent with a substantial accumulation of Ca\(^{2+}\) by the mitochondrion.

Fig. 11, C–E, present the data for three individual cells from this experiment. Note that the peak \([Ca^{2+}]_i\), values in Fig. 11 C were very high. Under these conditions, fura-2 is very close to saturation, and small variations in the fluorescence intensity at 380 nM excitation (the denominator of the fura-2 ratio) produce large fluctuations in the calculated \([Ca^{2+}]_i\). These very high \([Ca^{2+}]_i\), values and their fluctuations should not be taken as precise measurements. The \(K_h\) values obtained by the matching procedure with and without Cl-CCP are given in the figure and the data points corresponding to the estimated \(K_h\) are indicated by the diagonal arrows. For these cells, and most (but not all) of the other cells that were monitored, the apparent \(K_h\) was higher in the presence of Cl-CCP. Comparing the cells that responded under each condition (\(V_{max} \approx 5\) nM/s; \(n = 44\)), the average \(K_h\) (±SD) values were 257 ± 77 nM under control conditions and 336 ± 124 nM in the presence of Cl-CCP (\(P < 10^{-4}\), paired \(t\) test). When the results from four coverslips were pooled, the average \(K_h\) values (±SEM) were 242 ± 6.0 nM and 347 ± 7.4 nM in the absence and presence of Cl-CCP respectively (\(P \sim 0.003\), paired \(t\) test; \(n = 4\)). Thus, blocking mitochondrial Ca\(^{2+}\) accumulation increased the average \(K_h\) values for exchanger activation by 44 ± 5.5%.

**Rates of Ca\(^{2+}\) Uptake**

Another potential source of error in our analysis involves the influence of cytosolic Ca\(^{2+}\) buffering on the velocity of Ca\(^{2+}\) uptake. In our experiments, transport rates were measured as \(d[Ca^{2+}]_i/dt\) and expressed as nM/s. Conversion to units reflecting net Ca\(^{2+}\) uptake (nmol/s) requires knowledge of cytosolic buffering by cellular constituents as well as by the fura-2 present within the cell. Previous results (Fang et al., 1999) suggest that the fura-2 concentration within the cytosol is \(\sim 30\) \(\mu\)M.

Neher and Augustine (1992) defined the calcium binding capacities of endogenous buffering systems in the following manner:

\[
\kappa_S = d[S\text{Ca}]/d[Ca^{2+}],
\]

where S\text{Ca} represents the total amount of Ca bound to the endogenous buffers. An analogous relation was used to define the binding capacity of the exogenous buffer, i.e., fura-2:

\[
\kappa_S = d[B\text{Ca}]/d[Ca^{2+}].
\]

The equilibrium relation for the interaction between Ca\(^{2+}\) and fura-2 is

\[
K_D = \frac{[Ca^{2+}]B}{[B\text{Ca}]},
\]

where \(K_D\) is the dissociation constant for the Ca\(^{2+}\)/ fura-2 complex, \([B]\) is the concentration of unbound fura-2, \([B\text{Ca}]\) is the concentration of Ca\(^{2+}\)-bound fura-2 and \(B_T = [B] + [B\text{Ca}]\) is the total concentration of fura-2. With this relation, the expression for \(\kappa_B\) becomes (Neher and Augustine, 1992)

\[
\kappa_B = \frac{[B_T]}{K_D([Ca^{2+}] + 1)^2}
\]

For \([Ca^{2+}]_i \sim K_B\), \([B_T] = 30\) \(\mu\)M and \(K_B = 0.274\) \(\mu\)M, \(\kappa_B \sim 27\). For bovine chromaffin cells, Neher and Augustine (1992) found a mean value for \(\kappa_S\) of 70, which was relatively constant over the range \([Ca^{2+}]_i = 0.2–1\) \(\mu\)M. Thus, fura-2 makes an important contribution to the buffering power of the cytosol. Moreover, the buffering capacity of fura-2 changes significantly with \([Ca^{2+}]_i\), becoming progressively less as \([Ca^{2+}]_i\) increases. Changes in the buffering capacity of fura-2 during the course of Ca\(^{2+}\) uptake are therefore likely to alter the relation between \(d[Ca^{2+}]_i/dt\) and the rate of Ca\(^{2+}\) influx.

To estimate the influence of these factors on \(K_h\), we used the following relation to convert the rate of change in \([Ca^{2+}]_i\) to the rate of Ca\(^{2+}\) uptake (Neher and Augustine, 1992).
The impact of the correction for fura-2 buffering was dependent on the assumed concentration for fura-2. When the six cells displayed in Fig. 4 were reanalyzed by the curve-fitting approach assuming a value for \( B_T \) of 100 \( \mu \)M, the average \( K_b \) value (±SD) was 151 ± 60 nM compared with 191 ± 68 nM with \( B_T = 30 \) \( \mu \)M and 248 ± 89 nM with no correction for \( \text{Ca}^{2+} \) buffering. Thus, at the higher \( \text{Ca}^{2+} \) concentration, the apparent overestimate of \( K_b \) obtained with uncorrected velocities increased to ~40%. The \( \text{Ca}^{2+} \)-levels vary considerably among individual cells, to judge by the variation in individual cellular fluorescence intensities. Thus, a cell-by-cell correction for the effects of fura-2 buffering is not possible and we will assume that the effects of buffering by fura-2 result in an overestimate of \( K_b \) between 25–40%, depending on the amount of fura-2 within the cell.

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TABLE I

| Procedure | Buffering correction | \( K_b \) nM | \( n \) | \( V_{max} \) (nmol/L cell water/s) |
|-----------|---------------------|-------------|------|----------------------------------|
| Curve-fitting | No | 232 ± 18 | 2.8 ± 0.15 | 153 ± 22 |
| Curve-fitting | Yes | 201 ± 19 | 2.8 ± 0.31 | 12,854 ± 1,700 |
| Matching | No | 231 ± 14 | | |
| Matching | Yes | 171 ± 13 | | |

The buffering correction involved the use of equations 4 and 5 with \( B_T = 30 \) \( \mu \)M, \( K_b = 70 \) (Neher and Augustine, 1992) and \( K_b \) for fura-2 of either 224 nM (37°C) or 274 nM (22°C), as appropriate. \( n \), Hill coefficient. Parameters for the curve fitting approach were based on 21 cells; parameters for the matching approach were based on 6 coverslips. Results are mean values ± SEM.

\(^{a} P < 0.01 \) compared to no correction.

\(^{b} P < 0.05 \) compared to no correction.

\[
\frac{d[\text{Ca}^{2+}]}{dt} = \left( 1 + \frac{K_b}{[\text{Ca}^{2+}]} \right) \frac{d[\text{Ca}^{2+}]}{dt},
\]

where \([\text{Ca}^{2+}]_T\) refers to the change in total cytosolic \( \text{Ca}^{2+} \) (nmol/L cell water). This relation assumes \( \text{Ca}^{2+} \) buffering is rapid and neglects slower processes, including \( \text{Ca}^{2+} \) uptake by cellular organelles. We used Eq. 4 to define the \( \text{Ca}^{2+} \) dependence of \( K_b \) and assumed that \( K_b \) was independent of \([\text{Ca}^{2+}]_T\) and equal to the value of 70 obtained by Neher and Augustine (1992) for bovine chromaffin cells. Rates of \( \text{Ca}^{2+} \) uptake, computed using Eq. 5 with \( B_T = 30 \) \( \mu \)M, were analyzed by the curve-fitting procedure as described for Fig. 4 and the combined results for the 21 cells discussed in results (see text relating to Fig. 4) are compared with the uncorrected values in Table I. As shown, correcting for \( \text{Ca}^{2+} \) buffering decreased the value of \( K_b \) significantly, reflecting the reduced buffering power of fura-2 at high \( \text{Ca}^{2+} \) concentrations. The \( K_b \) values obtained by the matching procedure were also reduced when velocities were analyzed using Eq. 5. As shown in Table I, the average \( K_b \) decreased from 231 to 171 nM after correcting for \( \text{Ca}^{2+} \) buffering for the six coverslips analyzed by the matching procedure. Thus, with an assumed \( B_T \) of 30 \( \mu \)M, the effects of fura-2 buffering result in an overestimate of \( K_b \) by ~25%.
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