Control of Maximum Sarcoplasmic Reticulum Ca Load in Intact Ferret Ventricular Myocytes

Effects of Thapsigargin and Isoproterenol

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ABSTRACT In steady state, the Ca content of the sarcoplasmic reticulum (SR) of cardiac myocytes is determined by a balance among influx and efflux pathways. The SR Ca content may be limited mainly by the ATP-supplied chemical potential that is inherent in the gradient between SR and cytosol. That is, forward Ca pumping from cytosol to SR may be opposed by energetically conservative reverse pumping dependent on intra-SR free \( [Ca] \). On the other hand, SR Ca loading may be limited by dissipative pathways (pump slippage and/or pump-independent leak). To assess how SR Ca content is limited, we loaded voltage-clamped ferret ventricular myocytes cumulatively with known amounts of Ca via L-type Ca channels \( (I_{\text{L}}) \), using Na-free solutions to prevent Na/Ca exchange. We then measured the maximal resulting caffeine-released SR Ca content under control conditions, as well as when SR Ca pumping was accelerated by isoproterenol (1 \( \mu \)M) or slowed by thapsigargin (0.2–0.4 \( \mu \)M). Under control conditions, SR Ca content reached a limit of 137 \( \mu \)mol-liter cytosol\(^{-1} \) (nonmitochondrial volume) when measured by integrating caffeine-induced Na/Ca exchange currents \( (I_{\text{NaCaX}}) \) and of 119 \( \mu \)mol-liter cytosol\(^{-1} \) when measured using fluorescence signals dependent on changes in cytosolic free Ca \( ([Ca]_c) \). When Ca-ATPase pumping rate was slowed 39% by thapsigargin, the maximal SR Ca content decreased by 5 \% \( (I_{\text{NaCaX}}/dt \) method) or 23% \( (I_{\text{NaCaX}}/dt \) method); when pumping rate was increased 74% by isoproterenol, SR Ca content increased by 10% \( (I_{\text{NaCaX}}/dt \) method). The relative stability of the SR Ca load suggests that dissipative losses have only a minor influence in setting the SR Ca content. Indeed, it appears that the SR Ca pump in intact cells can generate a \( [Ca] \) gradient approaching the thermodynamic limit.

KEY WORDS: cardiac myocytes • sarcoplasmic reticulum • isoproterenol • thapsigargin • excitation–contraction coupling

INTRODUCTION In cardiac myocytes, many properties of the Ca-induced Ca release (CICR) and excitation–contraction coupling (ECC) processes depend on the quantity of Ca stored in the sarcoplasmic reticulum (SR). Among these are the gain or efficacy (Janczewski et al., 1995) and related factors such as the fraction of stored Ca released during twitches (Han et al., 1994; Bassani et al., 1995), the amount of Ca-dependent negative feedback on Ca influx through L-type Ca channels (Sipido et al., 1995; Grantham and Cannell, 1996), and the probability of local or propagated spontaneous Ca release events (Cheng et al., 1993; Satoh et al., 1997). Secondary to changes in CICR or ECC are, for example, inotropy with increased SR Ca loads (e.g., in phospholamban knockout mice; Wolska et al., 1996; Li et al., 1997), as well as rest potentiation or decay related with load changes (Bassani and Bers, 1994). Accordingly, mechanisms setting and otherwise controlling the maximum SR Ca load are of interest.

A particular SR Ca load is maintained in steady state when influx and efflux of Ca from the SR are equal. The maximal SR Ca load could be set by the limiting chemical potential gradient of free \( [Ca] \) between the SR lumen and the cytosol, \( \Delta G_{\text{SR}} = 2RT \ln ([Ca]_c/\text{[Ca]}) \). Efflux from the SR could be mediated by a combination of reverse pump flux (regenerating ATP; Makinose and Hasselbach, 1971) and dissipative leakage. In isolated rat SR vesicles, with dissipative leak through ryanodine receptor/release channels blocked, Shannon and Bers (1997) found a gradient of 7,000, corresponding to an apparent efficiency of 74%, based on \( \Delta G_{\text{ATP}} = 58.9 \text{kJ/mol} \) (Allen et al., 1985). Under this limiting condition, the flux balance should be set mainly by the forward and backward pump rates, and the energy cost incurred due to incomplete reversibility of the Ca-ATPase (Inesi and de Meis, 1988) should be minimal (Takenaka et al., 1982; Feher and Briggs, 1984).
In contrast, in intact cells, Ca must also leak from the SR through ryanodine receptors, dissipating the SR [Ca] gradient. One aim of this study was to determine whether enough Ca leaks from the SR in intact myocytes to reduce the steady state SR Ca load below the limit set by $\Delta C_{\text{SR}}$. If sufficient, leak flux, rather than reverse pumping, would be the main factor opposing forward pumping (SR uptake) and thereby determining steady state SR Ca load.

In intact cells, existing estimates of SR Ca pump and leak rates should help predict SR Ca loading, but appear disparate. Pump parameters estimated from cytosolic $[\text{Ca}]$, transients ($K_m = 0.27 \, \mu\text{mol-liter cytosol}^{-1}$ and $V_m = 210 \, \mu\text{mol-liter cytosol}^{-1} \cdot \text{s}^{-1}$; Sipido and Wier, 1991; Balke et al., 1994; Bassani et al., 1994) predict a forward rate of 27 $\mu\text{M} \cdot \text{s}^{-1}$ at cytosolic $[\text{Ca}] = 100 \, \text{nM}$. By contrast, direct estimates of unidirectional Ca leak from the SR of ventricular cells of rabbit and rat with normal loads amount to only 0.5 $\mu\text{mol-liter cytosol}^{-1} \cdot \text{s}^{-1}$ (Bassani and Bers, 1995). Since this Ca pump independent leak is much slower than the unidirectional SR Ca uptake, pump backflux may limit SR Ca content in intact cells, as appears to be the case in permeabilized myocytes with leak blocked as described above (Shannon et al., 1997).

Ca leak from the SR may increase with increasing SR Ca load (Bassani and Bers, 1995). Consistent with this, and consistent with the view that the ryanodine receptors are the source of the leak, Ca spark frequencies increase when SR Ca load is increased; e.g., by high external [Ca] or isoproterenol (ISO) treatment (Cheng et al., 1993; Gómez et al., 1996; Satoh et al., 1997). In intact rat cells, high SR Ca loading is associated with spontaneous waves that may indicate a sharply accelerating leak rate (Cheng et al., 1996; Diaz et al., 1997).

To assess the importance of leak in limiting SR Ca load, we loaded voltage-clamped cells cumulatively with known amounts of Ca via L-type Ca channels ($I_{\text{Ca}^l}$). During loading, Na-free solutions were used to prevent Na/Ca exchange current ($I_{\text{NaCaX}}$). We then measured the maximal resulting caffeine-released Ca content of the SR, using both Na/Ca exchange currents and fluorescence-based measurements of cytosolic free Ca ($[\text{Ca}]_c$). We perturbed SR Ca pump by accelerating it with ISO (1 $\mu\text{M}$; Tada et al., 1974) or slowing it (blocking it partially) with thapsigargin (TG; 0.2–0.4 $\mu\text{M}$; Sagara and Inesi, 1991; de Meis and Inesi, 1992; Hove-Madsen and Bers, 1993b). If Ca leak is minimal, stimulation or inhibition of the SR Ca pump rate should not alter maximal SR Ca load, but only change the time required to attain it. If leak is important in setting the SR Ca load, the load should change substantially when the pump rate is changed.

Under control conditions, SR Ca content reached a limit of 137 $\mu\text{mol-liter cytosol}^{-1}$ (nonmitochondrial volume) when measured via $\int I_{\text{NaCaX}} \cdot \text{dt}$ and 119 $\mu\text{mol-liter cytosol}^{-1}$ when measured using fluorescence. When Ca-ATPase pumping rate was slowed 39% by TG, the maximal SR Ca content decreased by 5% ($\int I_{\text{NaCaX}} \cdot \text{dt}$ method) or 23% (fluorescence method); when the pumping rate was increased 74% by ISO, SR Ca content increased by 10% (fluorescence method) or 20% ($\int I_{\text{NaCaX}} \cdot \text{dt}$ method). These modest changes indicate that leak flux has only a minor influence on maximal SR Ca content in intact ventricular myocytes.

**materials and methods**

**cell preparation**

Cells were isolated from adult ferret hearts by approved methods as previously described (Bassani et al., 1994). Animals were anesthetized with pentobarbital (50 mg/kg) and killed by exsanguination. Hearts were excised, rinsed of blood, mounted on a standard Langendorff apparatus, and retrogradely perfused at 30–70 mmHg with Ca-free Tyrode solution for 5 min, after which collagenase (0.5–0.8 mg/ml) and neutral protease (0.02–0.05 mg/ml) were added. When the heart was somewhat flaccid, it was removed, rinsed in Ca-free Tyrode solution, and the tissue was cut and teased to yield free cells. After washing, cells were put in MEM with 2 mM Ca, plated onto laminin-pretreated recording chambers, and secured to the stage of a Nikon TMD microscope equipped for epifluorescence (Photon Technology, Inc., South Brunswick, NJ).

**fluorescence measurement of [Ca],**

Cells were incubated in a normal Tyrode solution containing (mM): 140 NaCl, 6 KCl, 1 MgCl$_2$, 1 CaCl$_2$, and 10 HEPES, with pH 7.4 using NaOH. Dye loading with Indo-1/AM (10 $\mu\text{M}$, 20 min, 20°C) was done in this same solution, after which 30 min in dye-free solution was allowed for deesterification. In some cells, instead of Indo-1/AM, $K_C$-Indo (50 $\mu\text{M}$) was loaded through the patch pipette (see below). Indo-1 fluorescence was excited at 365 nm, detected at 405 ± 10 and 485 ± 10 nm by photomultipliers with current-to-voltage converters and recorded using PClamp6 software (Axon Instruments, Foster City, CA), which also acquired the voltage clamp data (see below) and controlled the application of bathing solutions. Autofluorescent background at each wavelength, measured each day in separate cells that were not dye loaded, was adjusted for apparent surface area of each cell and subtracted from the corresponding total fluorescence. In a few cells, a time-dependent trend for individual fluorescence was observed, but did not affect any conclusions.

**electrophysiology**

Perforated patch voltage clamp was used for most cells to record $I_{\text{Ca}^l}$ or $I_{\text{NaCaX}}$. Recording electrodes of resistance 1–2 $\Omega$ were tip-filled with (mM): 40 CsCl, 80 Cs-methanesulfonate, 0.1 EGTA, 1 MgCl$_2$, and 10 HEPES, with pH 7.2 using CsOH. The same solution, with amphotericin added at 120 $\mu\text{g/ml}$, was used to backfill the electrodes. In a few cells, ruptured patch recording was used, in which case $K_C$-Indo-1 (50 $\mu\text{M}$), Mg-ATP (5 $\mu\text{M}$), and Li-GTP (0.3 $\mu\text{M}$) were added to the pipette solution, and EGTA and amphotericin were omitted. During recording, cells were bathed in a standard solution containing (mM): 140 TEA-Cl, 4 CaCl$_2$, 1
MgCl₂, 10 HEPES, and 2 CaCl₂ with pH 7.4 using CsOH. Bathing solutions were substituted during appropriate periods by means of a fast switching device under computer control as set by the experimental protocol (see below and Fig. 1).

**Experimental Protocol**

Cells were maintained in the Na-free Ca-containing solution described above, so that I_{Ca} could be recorded without interference from I_{NaCaX}. To prevent cells from Ca overloading via reverse Na/Ca exchange in this solution, they were initially depleted of Na\(_2\) before experiments were begun. This was accomplished by incubating for 15 min in K-containing Na- and Ca-free solution containing (mM): 140 LiCl, 6 KCl, 1 MgCl₂, 1 EGTA, and 10 glucose, with pH 7.4 using LiOH.

As shown in Fig. 1, a cell was first initialized to deplete the SR Ca load by applying caffeine. Next, to Ca-load the cell progressively, we evoked I_{Ca} by depolarizing repeatedly from \(V_{hold} (-55\) mV) to 0 for 200 ms, at 0.25 Hz. The number of pulses was 1, 2, 5, 10, 20, or 40. Finally, the resulting SR Ca load was measured. Caffeine was applied in Na-free solution to release the SR Ca load. The initial increase in [Ca\(_i\)] was sustained long enough to use as a first measure of SR Ca load, the only routes for Ca to leave the cytosol being extrusion on the sarcolemmal Ca pump and/or uptake into mitochondria (Bassani et al. 1994a). 2 s after the start of caffeine application, caffeine-containing standard solution with 140 mM NaCl instead of TEA-Cl was substituted, which rapidly activated inward Na/Ca exchange current (I_{NaCaX}) and also caused [Ca\(_i\)] to decline steeply. After 10 s, caffeine-free standard solution was restored. During caffeine applications, \(V_{hold}\) was reset to \(-70\) mV to enhance I_{NaCaX}. Integration of I_{NaCaX} provided data for a second measure of SR Ca content (Callewaert et al., 1989; Varro et al., 1993). This initialization, loading, and measurement sequence was repeated for the various numbers of loading pulses, in random order. Thereafter, we added either ISO (1 μM sustained application) or TG (0.2–0.4 μM, 1–2 min application), and repeated the entire set of sequences (again with 1, 2, 5, 10, 20, or 40 pulses in random order) on the same cell.

**Calculations**

Cumulated molar Ca influx via I_{Ca} in each loading sequence was calculated by integrating I_{Ca} and summing over all pulses during the sequence. Since the depolarization for I_{Ca} was to 0 mV (eliminating leak current) and since K and Na were absent, all current was considered to be due to I_{Ca}, and the baseline for integration was set to 0. The molar influx was divided by the nonmitochondrial cytosolic volume to get Ca concentration influx. Cytosolic volume was found by dividing the cell’s capacitance by the ratio of capacitance to cytosolic volume for ferret myocytes (7.96 pF/pl cytosol; Satoh et al., 1996).

Fluorescence transients recorded as the ratio (R) of background-corrected fluorescence at 405 vs. 485 nm during Ca loading and subsequent release of SR Ca were converted to [Ca\(_i\)], according to [Ca\(_i\)] = \(K_β(R - R_{min})/(R_{max} - R)\) (Grynkwicz et al., 1985). Limiting ratio parameters \(R_{max}\) (corresponding with [Ca\(_i\)] < 10\(^{-9}\) M) and \(R_{min}\) (corresponding with [Ca\(_i\)] = 2 nM) were measured in vivo, separately for Indo-1/AM and K\(_2\)-Indo, as described in Bassani et al. (1995b). The value of \(β\), the ratio of free ([Ca\(_i\)] < 10\(^{-9}\) M) to bound ([Ca\(_i\)] at saturation) Indo-1 fluorescence measured in vitro at 485 nm, was 3.0. The dissociation constant (\(K_β\)) for intracellular Indo-1 was taken as 750 nm (reflecting higher \(K_β\) values seen in vivo vs. in vitro; see Hove-Madsen and Bers, 1992; comparable with Bassani et al., 1995b). [Ca\(_i\)] was used to follow the time course of SR Ca uptake during loading. To measure SR Ca content, resting [Ca\(_i\)], after loading but before caffeine application and peak [Ca\(_i\)], which occurred during the first 2 s of caffeine applications, were converted to total cytosolic Ca ([Ca\(_i\)]\(_T\)). The difference between these [Ca\(_i\)]\(_T\) values (Δ[Ca\(_i\)]\(_T\)) was taken to indicate the added SR Ca load, referred to the cytosolic volume. [Ca\(_i\)]\(_T\) was determined from [Ca\(_i\)], using [Ca\(_i\)]\(_T\) = [Ca\(_i\)] + 232 μM/(1 + (455 nM/[Ca\(_i\)]) + 50 μM/(1 + (750 nM/[Ca\(_i\)])], in which the second term reflects passive cellular Ca buffering (Bers and Berlin, 1995; based on data in Hove-Madsen and Bers, 1993a), and the third term accounts for Ca buffering by Indo-1.

Fluorescence-based SR Ca content measurements depend on numerous parameters, including those for cytosolic Ca buffering (\(K_β\) and \(B_{max}\)) and for calibration (\(R_{min}\), \(R_{max}\), \(K_β\), and \(B\)), and could be affected by non–Ca-related background fluorescence. We assessed the sensitivity of SR Ca content to these factors by putting typical observed values in the defining equations and noting how much the apparent SR Ca content changed when we varied each parameter ±20% around its typical value. \(B_{max}\) for cytosolic Ca binding affects apparent content quite strongly (±19.6% for ±20% change), compared with cytosolic binding \(K_β\) (±3.6%). However, these parameters are well established (Berlin et al., 1994; Hove-Madsen and Bers, 1993a). Apparent SR Ca content is sensitive to changes in background fluorescence (±1–10%). Although we could not measure the background for each individual cell, we did base our background correction on 10 or more cells each day with compensation for cell size. SR Ca content was relatively less sensitive to changes in Indo-1 AM calibration parameters \(R_{max}\) (±8%), \(K_β\) (±4.6%), and \(R_{min}\) (±4.4%).

SR Ca content was also calculated by integrating the current that flowed during caffeine application after Na was added (J_{NaCaX3} method; Varro et al., 1993; Terraciano and MacLeod, 1994; Delbridge et al., 1996). We believe this current is entirely I_{NaCaX}, because no change in current flow occurs in association with caffeine applications when Na is absent. To measure the amount of Ca added to the SR during loading, the I_{NaCaX} integral should run until [Ca\(_i\)] has decayed to the resting value it had attained at the completion of SR Ca loading. Even after [Ca\(_i\)] has returned to this resting value, I_{NaCaX} can be nonzero because [Ca\(_i\)] is still determined by a balance among Ca transport processes (excluding the SR Ca-ATPase). To predict resting I_{NaCaX} we first subtracted the leakage current flowing at the start of caffeine application before Na was added. We used linear regression (with zero asymptote) to predict how I_{NaCaX} depended on [Ca\(_i\)], selecting only the

![Figure 1](https://example.com/figure1.png)
decaying part of the caffeine-evoked [Ca], transient. A linear dependence of $I_{\text{NaCaX}}$ on [Ca], applies when $[\text{Na}], (= 0)$, [Ca], and $[\text{Na}],_0$ are constant, as applies here (Kimura et al., 1987; Barcenas-Ruiz et al., 1987). Using as a baseline the predicted $I_{\text{NaCaX}}$ at resting [Ca], we directly integrated the rising and decaying phases of the current. In some cases, especially where $I_{\text{NaCaX}}$ was small, the later part of the decaying phase was affected by noise or by transient changes in seal leakage during contraction (downward deflections in the $I_{\text{NaCaX}}$ record in Fig. 2 D). Since our linear model could predict the entire decaying portion of the $I_{\text{NaCaX}}$ record, we replaced any uninterpretable late segments of the actual $I_{\text{NaCaX}}$ record by the prediction, before integrating. The direct integral always included at least the first few seconds (sometimes all) of the $I_{\text{NaCaX}}$ decay, and accounted for 87 ± 4% (SEM, $n = 117$) of the total integral. The integral was divided by 0.75 to correct for the parallel removal of cytosolic Ca by slow transport processes (mitochondrial uptake and sarcolemmal Ca pump extrusion) in the ferret (Bassani et al., 1994b) and also by 0.65 to exclude the fraction of total cell volume occupied by mitochondria (Page et al., 1971), yielding content in micromoles-liter cytosol$^{-1}$.

The $\Delta[\text{Ca}]_T$ and $\int I_{\text{NaCaX}} dt$-based SR Ca content estimates are complementary. In our case, they are not truly independent because the baseline value for integration of $I_{\text{NaCaX}}$ and in some cases parts of the integrals themselves were determined from [Ca],. However, the respective calculations have independent bases: $\Delta[\text{Ca}]_e$ estimates rely on prior knowledge of cytosolic Ca binding but not on knowledge of Ca removal by non-$I_{\text{NaCaX}}$ processes, and vice versa for the $\int I_{\text{NaCaX}} dt$-based estimates.

RESULTS

Ca Loading and SR Ca Content Data

Fig. 2 (left) shows $I_{\text{Ca}}$ and corresponding fluorescence transients recorded during moderate Ca loading of the SR by a sequence of 10 pulses. For clarity, only pulses 1, 5, and 10 are shown. A shows control records, while B shows the loading of the same cell after ISO treatment. $I_{\text{Ca}}$ was larger with ISO. $I_{\text{Ca}}$ became progressively smaller and decayed faster from its peak as loading progressed in both control and ISO cases. However, the decay rate changed more within the first few pulses in the ISO case. The twitch [Ca], transient amplitude increased as loading progressed, but in the ISO case it also declined faster than in control.

Fig. 2 (right) shows the corresponding responses to caffeine after loading with 10 pulses. With the moderate total Ca influx during 10 pulses, the SR Ca load was higher in ISO (Dvs. C), whether measured from the Ca transient or the integral of $I_{\text{NaCaX}}$. In this example, caffeine-evoked [Ca], increased during the interval in Na and Ca-free solution. In most records, however, [Ca], was stable or decreased slightly during this period. On average, SR Ca content measured at the start of Na application was 0.916 ± 0.014 (SEM, $n = 142$) of the content at the peak of the [Ca], transient, which occurred 442 ± 56 ms (SEM) earlier. This slight loss of content was very similar in control, TG, and ISO conditions.

The effects of TG treatment on loading $I_{\text{Ca}}$ and [Ca], transients were complementary to those of ISO. Although $I_{\text{Ca}}$ still decayed faster with loading in TG-treated cells, the progression rate of this change was drastically reduced with TG. That is, the decay rate changed less after a given number of pulses after TG treatment than in control. Similarly, the [Ca], transient amplitude increased as loading progressed, but in the TG case this increase was smaller, and the loading transients were as a group smaller and decayed much more slowly with TG.

Decay of $I_{\text{Ca}}$ and [Ca], Transients During Loading

Fig. 3 shows that the $I_{\text{Ca}}$ decay time constant $\tau$ (single exponential fit) shortened as SR Ca loading increased. This is presumably caused by Ca-dependent inactivation. For this illustration, $\tau$ values were measured for cells under each treatment at numbers of pulses where corresponding measurements of the SR Ca load were available. For example, the SR Ca load before the first loading pulse was 0, the load associated with pulse 2 was that measured after a single pulse, and so on.
given cell, at each number of pulses, any repeated measurements of either \( I_{\text{Ca}} \) decay \( \tau \) or SR load were averaged. The single regression line was fitted to all of the data. For this illustration, SR Ca loads were calculated using the \( \Delta [\text{Ca}]_T \) method. \( I_{\text{Ca}} \) was larger and decayed faster with ISO (not shown), even before SR loading. This may explain why data points with ISO fell above the regression line (and conversely for TG).

**Degree of SR Ca Pump Alteration by ISO and TG**

We selected conditions (ISO and TG application) either to stimulate or partially inhibit the SR Ca-ATPase rate. In Fig. 4, the relative changes in SR Ca pump rates are assessed by evaluating rate constants of twitch \([\text{Ca}]_i\) decline under ISO and TG. These rate constants were calculated from half-decay times of fluorescence ratios (using \( \lambda = 1/\tau = \ln(2)/t_{1/2} \)). Fig. 4 A shows that the rate of \([\text{Ca}]_i\) decline slowed with loading and was slower with TG and faster with ISO, relative to control. As each cell was studied successively under control and treatment, paired comparisons of the decay rate constants are appropriate. Fig. 4 B shows these comparisons for the five-pulse-loading case. Decay was 39 ± 16% (SEM, \( n = 5 \)) slower in TG or 74 ± 5% (SEM, \( n = 5 \)) faster in ISO. These changes in Ca transient decay rate correspond approximately with changes in the SR pump rate, since SR uptake is the dominant process reducing \([\text{Ca}]_i\) during loading. We infer that the concentrations of TG and ISO we used nearly halved and nearly doubled, respectively, the SR Ca pumping rate.

Changes in decay rate do not always indicate changes in SR Ca pumping properties, because the pump rate depends intrinsically on \([\text{Ca}]_i\). It may increase, as found by Bers and Berlin (1995) or decrease (as discussed further below). Fig. 5 shows that TG- and ISO-dependent decay rate changes were not artifacts of simultaneous changes in \([\text{Ca}]_i\). Fig. 5 (top) shows \([\text{Ca}]_i\) transients recorded during the 10th loading pulse, with control and TG records from the same cell compared in A and control and ISO records from a different cell compared in B. Fig. 5 (bottom) shows how the decay rates of \([\text{Ca}]_i\) depend on \([\text{Ca}]_i\). TG slowed the decay rate (C) and ISO sped the decay rate (D), relative to the respective rates in the control condition, for all \([\text{Ca}]_i\), above rest.
Ca entering the cell in Na-free conditions must distribute between the cytosol and the SR, although some may be transported into mitochondria or extruded by sarcoclemmal Ca pumping. In Fig. 6, original records of fluorescence ratio show that diastolic $[\text{Ca}]_i$ increases substantially during our loading protocol in a rabbit ventricular myocyte (representative of $n = 4$). A much smaller corresponding increase of diastolic $[\text{Ca}]_i$ is apparent in the ferret, which was used for all analysis in this study. This greater recovery of $[\text{Ca}]_i$ may be due to the unusually high sarcoclemmal Ca-ATPase-dependent Ca removal flux in this species (Bassani et al., 1995a).

In fact, diastolic $[\text{Ca}]_i$ increased in ferret myocytes during loading by only $0.12 \pm 0.018 \text{ mmol/\mu mol}$ added Ca, resulting in a typical increase of only 50 nM after moderate to large loadings. Sarcolemmal Ca extrusion or mitochondrial uptake may limit our ability to account for the fate of all entering Ca in SR or cytosol. However, the relatively small variations of diastolic $[\text{Ca}]_i$ represent nearly constant driving conditions for the SR Ca pump, even with varying numbers of loading pulses and under control, ISO, or TG conditions. Further, the increase in $[\text{Ca}]_i$ with loading, though small, means that sufficient Ca has entered cells to load the SR to maximal capacity.

**[Ca]_i Changes during Loading**

Fig. 7 shows how SR Ca loading depends on added Ca for two different cells recorded in ruptured patch with K$_5$-Indo, one treated with TG (A and C) and the other with ISO (B and D). The SR Ca content data were fitted to single-site binding (Michaelis-Menten) isotherms for descriptive purposes. In these single experiments, the maximal SR Ca contents, although different between cells, were unaffected by treatment: using the $\int_{\text{NaCaXdt}}$ method, the SR Ca content was $114.7 \pm 8.7 \text{ mmol-liter cytosol}^{-1}$ (control, 2 df) vs. $107.2 \pm 9.3 \text{ mmol-liter cytosol}^{-1}$ (TG, 3 df, Fig. 7 A), and $108.4 \pm 31.1 \text{ mmol-liter cytosol}^{-1}$ (ISO, 10 df, Fig. 7 B). Using the $\Delta[\text{Ca}]/_T$ method, corresponding values were $164.0 \pm 6.8 \text{ mmol-liter cytosol}^{-1}$ (control, 4 df) vs. $156.2 \pm 14.8 \text{ mmol-liter cytosol}^{-1}$ (TG, 5 df, Fig. 7 C), and $136.1 \pm 47.5 \text{ mmol-liter cytosol}^{-1}$ (ISO, 10 df, Fig. 7 D). The error tolerances represent the standard errors of the estimates of parameter $B_{\text{max}}$ and are shown with the corresponding number of degrees of freedom in the fits.

Fig. 8 A shows how the SR Ca load measured using the $\int_{\text{NaCaXdt}}$ method depended on the cumulated amount of Ca added, for a set of several perforated patch experiments ($n = 6$, TG; $n = 4$, ISO). Fig. 8 B shows results from the same cells with SR Ca load measured from the increase in peak fluorescence ($\Delta[\text{Ca}]/_T$ method) early during caffeine application. To show the data more clearly, observed loads were grouped within Ca influx ranges of 0–50, 51–100, 101–200, and >200 mmol-liter cytosol$^{-1}$, respectively.

Fits to the purely descriptive function $[\text{Ca}]_{\text{ser(total)}} = B_{\text{max}}/(1 + K_{1/2}/\Delta[\text{Ca}]/_T)$, presented as curves in Fig. 8, and as numeric values in Table I, were made to pooled raw (not grouped) data. Under control conditions, the fitted maxima were $110.4 \pm 17.5 \text{ mmol-liter cytosol}^{-1}$ ($n = 10$, $\int_{\text{NaCaXdt}}$ method) and $113.1 \pm 8.3 \text{ mmol-liter cytosol}^{-1}$ ($n = 8$, $\Delta[\text{Ca}]/_T$ method). Parameter $K_{1/2}$ is in-

![Image](image-url)

**Fig. 5.** TG- and ISO-dependent rate changes were not artifacts of simultaneous changes in $[\text{Ca}]_i$. (A) Loading $[\text{Ca}]_i$ transients, control and TG conditions, from one cell. (B) Loading $[\text{Ca}]_i$ transients, control and ISO conditions, from another cell. Each transient was the last evoked in a 10-pulse sequence. (C and D) $d[\text{Ca}]_i/dt$ as dependent on $[\text{Ca}]_i$, corresponding with A and B, respectively. TG slowed the decay rate at all $[\text{Ca}]_i$ above rest; correspondingly, ISO sped the decay at all $[\text{Ca}]_i$. Noise in the rate plots was reduced by fitting the decaying phase of each transient in A and B to a single exponential before differentiating.

![Image](image-url)

**Fig. 6.** Diastolic $[\text{Ca}]_i$ increases during loading in ventricular myocytes of both species, but less so in ferret than in rabbit. Original chart records of fluorescence ratio during execution of protocol in Fig. 1. Solid bars mark caffeine applications.
increased by TG and decreased by ISO. Although this $K_{1/2}$ does not measure a physical characteristic of the SR Ca-ATPase, it changes with TG or ISO, as would be expected from the corresponding changes in the pump’s ability to compete for cytosolic Ca against the other Ca transport systems.

Under the hypothesis that SR Ca loading is limited mainly by chemical potential and not by leakage, fitted SR Ca loads under all conditions must reach similar maxima. In Fig. 8 A, with the $f_{\text{NaCa}}\text{dt}$ method, ISO increased $B_{\text{max}}$ by 36% and TG decreased it by 14%. With the $\Delta[\text{Ca}]_T$ method ISO decreased $B_{\text{max}}$ by 12.6% and TG decreased it by 30%. Although it is not possible to combine analytically the content measurements made by the two methods, the overall pattern of the curve fits is consistent with a modest increase under ISO treatment and a modest decrease with TG.

In Fig. 8, the loading clearly increased progressively as Ca influx increased, under each treatment condi-

| Parameters of Fits Predicting Maximum SR Ca Load | $f_{\text{NaCa}}\text{dt}$ method | $\Delta[\text{Ca}]_T$ method |
|-------------------------------------------------|---------------------------------|----------------------------|
| $B_{\text{max}}$ | $K_{1/2}$ | $B_{\text{max}}$ | $K_{1/2}$ |
| ISO | $150.7 \pm 14.9$ | $2.8$ | $98.9 \pm 4.9$ | ~0 |
| CTRL | $110.4 \pm 17.5$ | $33.5$ | $113.1 \pm 8.3$ | 34.9 |
| TG | $94.6 \pm 16.6$ | $56.6$ | $78.7 \pm 10.0$ | 41.2 |

All values given in micromoles·liter cytosol$^{-1}$.  

To measure TG or ISO effects, data from each experiment, under each condition, would ideally be fit separately and the parameters compared pairwise. However, the raw data had some scatter, and as a result load in some experiments did not increase monotonically with increasing influx, leaving individual fitting difficult. To reduce uncertainty, we therefore compared pairwise the absolute maximum loads in each cell, regardless of stimulation history, with and without TG or ISO treatment.

Fig. 9 shows how ISO and TG affected maximal SR Ca load. For illustration, loads under ISO and TG were normalized to the corresponding control loads in the particular cells. Absolute loads appear in Table II. Using the $f_{\text{NaCa}}\text{dt}$ method, the SR Ca load with TG was 95.3% of control, and the load with ISO was 119.4% of control. The load with TG was 76.7% of the control load, while the load with ISO was 110.0% of control, using the fluorescence method. In paired $t$ test comparisons of the loads (before normalization) before and after treatment, the deficit with TG was significant ($P < 0.05, \Delta[\text{Ca}]_T$ method only), but the increment with ISO was not (either $f_{\text{NaCa}}\text{dt}$ or $\Delta[\text{Ca}]_T$ method). When maximum loads under TG and ISO treatment were compared (unpaired $t$ tests before normalization, either method), the difference was also insignificant.

To clarify the paired comparisons, loads under control conditions are shown separately in Table II for cells later treated with ISO or with TG. As noted in the abstract and discussion, the average control load for all cells was $118.7 \pm 5.8$ μmol-liter cytosol$^{-1} (f_{\text{NaCa}}\text{dt}$
For in the SR.

Mark unity slope that would result if all added Ca were accounted for in the SR.

Figure 8. Composite data from several cells showing how SR Ca load saturated as cumulated amount of Ca added increased. (top) \( \int \text{NaCa} \times dt \) method \((n = 6, \text{TG}; n = 4, \text{ISO})\). (bottom) Peak fluorescence \( \Delta[\text{Ca}]_T \) method \((n = 6, \text{TG}; n = 4, \text{ISO})\). For clarity, data points were grouped in influx ranges of 0–50, 51–100, 101–200, and 201–500 \( \mu \text{mol} \cdot \text{liter cytosol}^{-1} \), but curves were fit to individual data. Parameters of fits appear in Table I. Heavy lines mark unity slope that would result if all added Ca were accounted for in the SR.

Model for Steady State SR Ca Loading Characteristics

A simple theoretical model may make clear how our experimental alterations in SR pump function would be expected to change maximal SR Ca content. In accord with the ability of the SR Ca-ATPase to translocate Ca bidirectionally with minimal dissipative loss (Makinose and Hasselbach, 1971; Takenaka et al., 1982; Feher and Briggs, 1984; Inesi and de Meis, 1988), pumping by the SR Ca-ATPase is assumed to behave as a reversible enzymatic reaction, in which the net pump rate \( V_{\text{pump}} \) is defined as (Segel, 1976; Shannon and Bers, 1997):

\[
V_{\text{pump}} = \frac{V_{m}^{\text{mf}} \left( \frac{[\text{Ca}]}{K_{\text{mf}}} \right)^{2} - V_{m}^{\text{mr}} \left( \frac{[\text{Ca}]}{K_{\text{mr}}} \right)^{2}}{1 + \left( \frac{[\text{Ca}]}{K_{\text{mf}}} \right)^{2} + \left( \frac{[\text{Ca}]}{K_{\text{mr}}} \right)^{2}}.
\]

(1)

In this equation, \( V_{m}^{\text{mf}} \) and \( V_{m}^{\text{mr}} \) are the forward and reverse maximum velocities, and \( K_{\text{mf}} \) and \( K_{\text{mr}} \) are the forward and reverse affinities, respectively. \([\text{Ca}] \), and \([\text{Ca}]_\text{SR} \) are the free Ca in cytosol and SR, respectively. If the pump reaches a thermodynamic steady state and there is no leakage, then the net flux \( V_{\text{pump}} \) is 0. We assumed, in accord with Takenaka et al. (1982), that the maximum possible forward and backward rates are equal; i.e., \( V_{m}^{\text{mf}} = V_{m}^{\text{mr}} = V_{m} \). This \( V_{\text{pump}} = 0 \) condition reduces Eq. 1 to \( K_{\text{mf}} / K_{\text{mr}} = [\text{Ca}]_{\text{SR}} / [\text{Ca}] \).

Assuming that the SR pump can generate a maximum gradient \(([\text{Ca}]_{\text{SR}} / [\text{Ca}] \)) of 7,000 in the absence of leak (based on data of Shannon and Bers, 1997), then \( K_{\text{mf}} = 7,000 \cdot K_{\text{mf}} \).

In intact cells, Ca must leak from the SR, and in steady state, \( V_{\text{leak}} = V_{\text{pump}} \). Defining \( V_{\text{leak}} \) as an independent variable and substituting 7,000 \( \cdot K_{\text{mf}} \) for \( K_{\text{mr}} \), we can make Eq. 1 explicit for free \([\text{Ca}]_{\text{SR}} \):

\[
[\text{Ca}]_{\text{SR}} = \frac{7,000}{\sqrt{V_{m}^{\text{mf}} - 1} - K_{\text{mf}}^{2}} \left( \frac{V_{m}^{\text{mf}}}{V_{\text{leak}}} \right)^{2} + \left( \frac{V_{m}^{\text{mf}}}{V_{\text{leak}}} \right)^{2}.
\]

(2)

Under control conditions, we assume \( V_{m} = 210 \mu \text{mol} \cdot \text{liter cytosol}^{-1} \cdot \text{s}^{-1} \) and \( K_{\text{mf}} = 270 \text{ nM} \) (Bassani et al., 1994a). With the SR loaded, we found \([\text{Ca}]_{\text{i}} = 150 \text{ nM} \), so that, in the absence of leak, \([\text{Ca}]_{\text{SR}} = 1,050 \mu \text{M} \).

The relationship between free \([\text{Ca}]_{\text{SR}} \) and total SR Ca \(([\text{Ca}]_{\text{SRT}}) \), determined predominantly by calsequestrin binding, is (Shannon and Bers, 1997):

\[
[\text{Ca}]_{\text{SRT}} = [\text{Ca}]_{\text{SR}} \cdot B_{\text{maxSR}} / \left[ 1 + (K_{\text{dSR}} / [\text{Ca}]_{\text{SR}}) \right].
\]

(3)

Using for convenience \([\text{Ca}]_{\text{SRT}} = 100 \mu \text{mol} \cdot \text{liter cytosol}^{-1} \), taking SR volume as 3.5% of the cytosolic volume (Page et al., 1971), and setting \( K_{\text{dSR}} = 600 \mu \text{mol} / \text{liter SR} \) (Shannon and Bers, 1997), \( B_{\text{maxSR}} \) is calculated.
as 1.3 mmol/liter SR when binding is at thermodynamic equilibrium.

Addition of TG is simulated by a 50% reduction of both forward and reverse $V_m$, to $105 \mu$mol·liter cytosol$^{-1}$·s$^{-1}$; i.e., a complete blockage of 50% of the pumps. ISO treatment is simulated by reducing both forward and backward $K_m$ by 50% to 135 nM and 945 mM, respectively. This is in accord with data of Kirchberger and Wong (1978) and leaves the energy efficiency unchanged and the maximal gradient at 7,000.

Fig. 10 shows how maximum SR Ca load is expected to decrease with increasing leak rates, as predicted by Eqs. 2 and 3. In Fig. 10 A, the leak rate is expressed as a fraction of the forward pump rate. When leak is expressed this way, the loss in SR Ca content as leak rate increases is the same in control, TG, and ISO conditions. For instance, a loss of 25% of the SR Ca content requires a leak rate that is 59% of the forward rate, regardless of treatment condition. Under control conditions, this leak would be $\sim 15 \mu$mol·liter cytosol$^{-1}$·s$^{-1}$, 50X as high as the value actually observed, 0.3 $\mu$mol·liter cytosol$^{-1}$·s$^{-1}$ (Bassani and Bers, 1995).

SR Ca content relative to absolute leak rate is shown in Fig. 10 B. For each condition, the model is defined at leak rates up to the forward pump rate, at which the SR Ca content is 0. Fig. 10 B shows that for a given leak rate, the calculated steady state SR Ca content is higher with ISO and lower with TG, compared with control. At the physiological leak flux, 0.3 $\mu$mol·liter cytosol$^{-1}$·s$^{-1}$, TG, or ISO treatment are predicted to leave the SR Ca content virtually unchanged. Even with a leak flux 10X

**FIGURE 9.** Absolute maximum SR Ca loads, regardless of stimulation history, with and without TG or ISO treatment, were similar. (top) $\int I_{NaCa} dt$ method; (bottom) Peak fluorescence ($\Delta [Ca]_r$) method. *Load with TG was significantly lower ($P < 0.05$) in paired comparisons with control data. For illustration, maximum load with ISO or TG treatment was expressed as a fraction of the maximum load in same cell before treatment.

**FIGURE 10.** Model based predictions of changes in SR Ca load with TG or ISO due to leakage. Maximum SR Ca load decreases with increasing leak flux, but at the experimentally observed leak rate of 0.3 $\mu$mol·liter cytosol$^{-1}$·s$^{-1}$, the effect is minor. (A) Loss of SR Ca load as leak is increased, with leak rate expressed as a fraction of forward pump rate. (B) Same, expressed using absolute leak rate, log scale. (C) Normalized and expanded version of B. The forward pump rate stays relatively constant as $V_{\text{leak}}$ increases, but the reverse rate decreases, maintaining $V_{\text{forward}} - V_{\text{reverse}} - V_{\text{leak}} = 0$. 

499  GINSBURG ET AL.
higher (3 µmol-liter cytosol$^{-1}$·s$^{-1}$), there is only a modest variation in SR Ca load, and it is not until the leak is 30–50× the observed value, well beyond the physiological range, that the model predicts substantial changes. Fig. 10C shows an expanded view of the likely physiological range of leak flux. Even with a leak of 3 µmol-liter cytosol$^{-1}$·s$^{-1}$, the SR Ca content remains 95% of maximum.

Reverse pump flux is critical to the model. Since in the steady state, $V_{\text{forward}} - V_{\text{reverse}} - V_{\text{leak}} = 0$, $V_{\text{leak}} = V_{\text{forward}}$ must hold in the absence of reverse flux. In fact, to obtain a total [Ca]$_{\text{SR}}$ of 100 µmol-liter cytosol$^{-1}$ with a model neglecting $V_{\text{reverse}}$, $V_{\text{leak}}$ would need to be 63.7 µmol-liter cytosol$^{-1}$·s$^{-1}$, an unreasonable amount, inconsistent with observation.

**DISCUSSION**

**SR Ca Content Reaches a Limit**

Using a cumulative loading protocol, we saw that under control conditions SR Ca content in ferret ventricular myocytes reached a maximum that we estimated as $118.7 \pm 5.8$ µmol-liter cytosol$^{-1}$ from $\int_{10^{-5}}^{10^{-4}} \text{d}t$ measurements and $136.7 \pm 15.1$ µmol-liter cytosol$^{-1}$ from $\Delta$[Ca]$_{1}$ (fluorescence) measurements. The steady state load in the SR must be determined by a balance among influx and efflux pathways that include both conservative (forward and reverse pumping) and dissipative (pump slippage and/or pump-independent leak) pathways (Inesi and de Meis, 1988). When TG slowed the SR Ca-ATPase pump rate by 39 ± 16% or ISO increased the rate by 74 ± 5%, the perturbations in maximum SR Ca load were systematically smaller. These minor perturbations are consistent with a minimal influence of dissipative losses on SR Ca content.

**Origin and Amount of Leak Flux**

Under steady state SR Ca loading, influx and efflux of Ca from the SR are equal. The maximal SR Ca load could be set by the limiting chemical potential gradient of free [Ca] between luminal and cytosolic compartments, $\Delta G_{\text{ATP}} = 2RT \ln ([\text{Ca}]_{\text{SR}}/([\text{Ca}]_{i}))$, but some Ca must leak from the SR through the ryanodine receptors, dissipating the SR Ca gradient. Experimentally, the question of whether enough Ca leaks from the SR in intact myocytes to reduce the steady state SR Ca load below the chemical potential-dependent limit can be answered by perturbing the SR Ca uptake rate. If leakage, rather than reverse pumping, is the dominant factor opposing forward pumping (SR uptake), steady state SR Ca load will change when the forward rate is perturbed as we have done.

Leakage from the SR may be due to infrequent spontaneous openings of ryanodine receptors, which are manifest as Ca sparks (Cheng et al., 1993). The rate of spark occurrence is consistent with the low leak rate observed by Bassani and Bers (1995). Some dissipative loss of Ca from the SR could also result from normal slippage of the Ca-ATPase (Wolosker and de Meis, 1985).

If the leak flux from the maximally loaded SR of ferret cells is comparable to the flux measured directly in rabbit and rat ventricular cells with a normally loaded SR, 0.3 µmol-liter cytosol$^{-1}$·s$^{-1}$ (Bassani and Bers, 1995), then modifying the SR pump rate with ISO or TG should hardly perturb the maximum SR Ca content at all. Considering the data at varying loads, we found that perturbations were very minimal in particular cells (Fig. 7), though readily detectable in the fits to the pooled data (Fig. 8). In the data at absolute maximum loads, the finding of significant variations in load depended on the particular method and statistical test used (see Fig. 9, legend). In summary, leak rates in our cells may have been several times higher than observed previously, but they are not comparable to the forward rate. This supports the idea that backflux, rather than leak, mainly limits SR loading.

Using [Ca], transient decay data, Balke et al. (1994) calculated a much larger SR leak flux in rat ventricular myocytes, 18 µmol-liter cytosol$^{-1}$·s$^{-1}$, than did Bassani and Bers (1995). The model of Balke et al. (1994) considered the forward SR uptake rate to be balanced entirely by leakage, not pump-mediated backflux.

**How Ca Influx from Outside the Cell Affects SR Ca Loading**

All Ca added to a cell must distribute among cytosolic, SR, and mitochondrial compartments or be extruded. Initially, nearly all Ca added to the cell goes into the SR. This corresponds with the lines of unity slope on Fig. 8. The SR Ca content reaches a limiting value below proportionality for large additions. It is not surprising that almost all Ca initially enters the SR, since the SR Ca-ATPase is the transporter of highest capacity and its most effective competitor, the Na/CaX, is inoperative. As the SR fills progressively, the net uptake rate will decline as reverse flux increases (see Fig. 4 and Eq. 1; Shannon et al., 1997). Then, more of the entering Ca is transported by the slow systems; e.g., mitochondria and the sarcolemmal Ca-ATPase. This may account for the deviation below unity slope in Fig. 8 as load increases.

Mitochondria may sequester some of the Ca that does not end up in the SR or cytosol. However, we have recently measured SR Ca content in rabbit myocytes using the same protocol described here, and could discern no difference in experiments done with and without a mitochondrial Ca uptake blocker, RU360, a kind gift of Dr. Abdul Matlib, University of Cincinnati (Cincinnati, OH; unpublished observations). Even in cells
having substantial SR Ca loads and both SR Ca uptake and Na/CaX-mediated Ca extrusion blocked, redistribution of Ca into mitochondria appears very minimal, although detectable (Bassani et al., 1994a).

Some Ca will be extruded on the sarcolemmal Ca pump. We have predicted the cumulated sarcolemmal pump-mediated Ca efflux during loading over the duration (80 s) of the 20-pulse sequences in Fig. 6. From \([\text{Ca}^2+]_i\), measured at each sample point, and using sarcolemmal Ca pump parameters estimated by Bassani et al. (1994a, 1995a) for rabbit and ferret myocytes, we calculated an integrated Ca efflux of 45–100 \(\mu\text{mol-liter cytosol}^{-1}\). Cumulated sarcolemmal pump extrusion seems only partially to explain the fate of Ca, which enters neither SR nor cytosol at high cumulative loads. However, slow Ca efflux via this route does have the capacity to maintain low diastolic \([\text{Ca}^2+]_i\), even with Na absent and SR function suppressed (Lamont and Eisner, 1996). SL Ca efflux could be accelerated manyfold by calmodulin or by ISO-mediated PKA activation (Dixon and Haynes, 1989). If this happened, we might not detect increases in SR pumping capacity (and hence loading) with ISO. Since initial Na-free caffeine-induced \([\text{Ca}^2+]_i\) transients do not decay any faster with ISO present (see Fig. 2, legend), SL pumping may already be at maximum under control conditions.

Although we cannot trace the fate of all admitted Ca in our cells, we do not believe any other cytosol-related processes have worked to limit SR loading in our experiments. In the ruptured patch experiments, the patch pipette also sinks Ca from the cell cytoplasm. This loss appears insignificant, as SR Ca contents measured in ruptured and perforated patch experiments were close. Ca efflux on an active Na/CaX could limit SR loading, but our use of Na-free solutions eliminates this possibility (contrast Ginsburg and Bers, 1996; Trafford et al., 1997; Bassani et al., 1995a) and so better drives the cells toward a maximal SR Ca load.

With ISO treatment, the SR Ca content often exceeded the Ca influx as measured by \(\int I_{\text{Ca}}\,dt\); that is, it fell above a line of unity slope in Fig. 8, particularly for small influxes. One possible explanation is leakage of Ca into the cell (Lamont and Eisner, 1996). SR Ca reloading without stimulation has been seen previously (Trafford et al., 1997), but is not expected in the absence of internal Na. In a 30-pl cell, each picoampere of leakage carried by Ca would add 0.16 \(\mu\text{mol-liter}^{-1}\cdot\text{s}^{-1}\) to the total cell Ca content. For a worst-case 40-pulse loading sequence of duration 160 s, 1 pA could increase the SR Ca content by \(\approx 23 \mu\text{mol-liter cytosol}^{-1}\) in the absence of removal processes (see Eq. 1). Under low Ca loading conditions, the SR Ca pump can compete quite effectively with other Ca transport processes, especially when \(K_m\) is reduced by ISO. For example, with external Ca absent, the SR can recover Ca from mitochondria without raising \([\text{Ca}^2+]_i\). (Bassani et al., 1993).

Another possible explanation for the supralinear Ca content in ISO is potential Ca loading via Na/Ca exchange. Since K was absent, Na entering cells during caffeine/Na measurements of load via INaCaX was not rapidly extruded via the sarcolemmal Na/K-ATPase. In a typical caffeine/Na record, 100 \(\mu\text{M} Ca\) was extruded by INaCaX, which could correspond to 0.3 mM Na gain/trial. Even this small growth of \([\text{Na}^+]_i\) would, however, be opposed by ongoing Na/H exchange (since external Na is absent except during the short periods that caffeine-induced INaCaX flows) and by diffusion into the patch pipette. This maximum change in \([\text{Na}^+]_i\) would also not be expected to significantly activate Ca entry via Na/Ca exchange because the exchanger’s \(K_m\) for Na is 30 mM (Reeves and Philipson, 1989).

**Decline of \(I_{\text{Ca}}\) and Increase of \([\text{Ca}^2+]_i\)**

Larger SR Ca loads corresponded with a faster \(I_{\text{Ca}}\) decline. We infer that this was caused by Ca-dependent inactivation after SR Ca release since the SR is a major source of inactivating Ca (Sham et al., 1995; Grantham and Cannell, 1996; Sipido et al., 1995). SR Ca load was a good predictor of the fractional decline of \(I_{\text{Ca}}\) during successive loading pulses, since increased loads lead to greater fractional Ca release. Ca admitted via \(I_{\text{Ca}}\) would also be expected to enhance \(I_{\text{Ca}}\) inactivation, as was seen by Hadley and Hume (1987) and Yuan and Bers (1995). Between no-load and maximum-load conditions, resting \([\text{Ca}^2+]_i\) increases, though variable, were on average 50 nM. This gradual increase in the average \([\text{Ca}^2+]_i\) may have enhanced \([\text{Ca}^2+]_i\)-dependent \(I_{\text{Ca}}\) inactivation and limited peak \(I_{\text{Ca}}\) as load progressed (Fig. 2). We assume that the \(I_{\text{Ca}}\) inactivation at pulse 1 (with no SR Ca load) was almost entirely due to Ca influx through the Ca channel. Since \(I_{\text{Ca}}\) decline was accelerated four-to fivefold with SR Ca loading (Fig. 3), we infer that \(\sim 75\%\) of the Ca-dependent inactivation is due to SR Ca release. Together, these observations suggest that Ca originating from \(I_{\text{Ca}}\) SR Ca release, and average diastolic \([\text{Ca}^2+]_i\) all contribute to Ca-dependent inactivation.

**Comparison with Previous Measures of SR Ca Content**

Our estimates of the SR Ca load (either \(\int \Delta [\text{Ca}^2+]_i\,dt\) or \(\Delta [\text{Ca}^2+]_i\) methods) agree well with most previous estimates. Our preliminary Na/Ca exchange–based estimate in ferret, which was based on steady state loading in Na-containing solutions was somewhat higher (170 \(\mu\text{mol-liter cytosol}^{-1}\); Ginsburg and Bers, 1996). In saponin-treated ferret ventricular muscle strips, maximal SR Ca content with \([\text{Ca}^2+]_i\) \(= 100 \text{nM} \) was 260 \(\mu\text{mol-liter cytosol}\) (Kawai and Konishi, 1994). Previous \(\int \Delta [\text{Ca}^2+]_i\,dt\)-
based estimates have varied somewhat with the species used (rat: 185 μmol-liter⁻¹ cytosol, Varro et al., 1993; 115 μmol-liter cytosol⁻¹, Terraciano and MacLeod, 1997; 169 μmol-liter cytosol⁻¹, Diaz et al., 1997; rabbit: 87 μmol-liter cytosol⁻¹, Delbridge et al., 1996; guinea pig: 58.8 μmol-liter cytosol⁻¹, Terraciano and MacLeod, 1997). Our fluorescence-based content measures are quite comparable with other fluorescence-based measures from our own laboratory in ferret (91 μmol-liter cytosol⁻¹; Bassani et al., 1995c), rat (114 μmol-liter cytosol⁻¹; Bassani and Bers, 1995), and rabbit (106 μmol-liter cytosol⁻¹; Bassani and Bers, 1995). For comparison, these measurements have been adjusted where needed to refer to nonmitochondrial cell volume (Page et al., 1971); however, not all previous measurements were made under conditions that maximized loading.

We combined both \( \int \Delta [Ca] \text{A}_T \) methods, for which calculations differ somewhat, to evaluate SR Ca load. The initial exposure to caffeine in 0 Na/0 Ca solution is valuable in the \( \Delta [Ca] \text{A}_T \) method because it prevents snubbing of the peak of the [Ca] transient (Bassani et al., 1994a). A minor potential disadvantage of the \( \int \Delta [Ca] \text{A}_T \) procedure is that any removal of Ca by the sarcolemmal Ca-ATPase (possibly boosted during ISO treatment) will have progressed somewhat by the time Na is added. We found, however, that the average loss of SR Ca content between peak [Ca], and the start of Na application was under 10% in each condition.

**Modeling**

Backflux via the SR Ca pump (Inesi and de Meis, 1988; Takenaka et al., 1982; Feher and Briggs, 1984) is an essential feature of our SR loading model, which has not been included in previous related models (Negretti et al., 1993; Balke et al., 1994). It allows us to explain the apparent disparity between previous measurements of forward pumping (Sipido and Wier, 1991; Balke et al., 1994; Bassani et al., 1994a) and backward leak fluxes (Kawai and Konishi, 1994; Bassani and Bers, 1995).

**Physiological Implications**

Some implications of a limiting SR Ca load have been discussed previously (Bassani et al., 1995c). As SR Ca content increases, the fraction that is free \([Ca]_\text{SR}\) increases, amplifying the inotropic tendency of increased loads. A limit to loading mitigates this factor, providing an additional control point that may help to optimize twitch force so as to maintain efficient excitation–contraction coupling.

Ca leak from the SR can be presumed to increase with increasing SR Ca load (Bassani and Bers, 1995) as Ca spark frequencies increase when SR Ca load is increased; e.g., by high external [Ca] or ISO treatment (Cheng et al., 1993; Gómez et al., 1996; Satoh et al., 1997). Leak may increase in proportion to the SR Ca content (rate constant 0.0027 s⁻¹ or 16% min⁻¹, Bassani and Bers, 1995; 0.005 s⁻¹, Kawai and Konishi, 1994). However, as we noted in the introduction, rat myocytes undergo spontaneous SR Ca release at high SR loads (Cheng et al., 1996; Diaz et al., 1997), indicating that leak from the SR, and hence \([Ca]_\text{SR}\), may increase disproportionately with increasing SR Ca load. \([Ca]_\text{SR}\) might then reach triggering threshold more often during random fluctuations. It is somewhat surprising that we seldom saw spontaneous SR Ca release or waves in these ferret cells even as they approached the maximum SR Ca load. This might be related to the particularly strong sarcolemmal Ca-ATPase in ferret ventricular myocytes (Bassani et al., 1995a).

A potentially important pathophysiological implication of our results is that \( \Delta G_{\text{ATP}} \) changes due to energetic compromise in ischemia or heart failure would reduce the maximal SR Ca load. Any loss of the maximal \([Ca]_\text{SR}/[Ca]_\text{i}\) gradient would reduce both the amount of SR Ca available for release and the fraction of that Ca released (Bassani et al., 1995c). These factors could thus contribute to systolic cardiac dysfunction in various pathophysiological states.

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**REFERENCES**

Allen, D.G., F.G. Morris, C.H. Orchard, and J.S. Pirolo. 1985. A nuclear magnetic resonance study of metabolism in the ferret heart during hypoxia and inhibition of glycolysis. *J. Physiol. (Camb.).* 361:185–204.

Balke, C.W., T.M. Egan, and W.G. Weir. 1994. Processes that remove calcium from the cytoplasm during excitation–contraction coupling in intact rat heart cells. *J. Physiol. (Camb.).* 474:447–462.

Barcenas-Ruiz, L., D.J. Beuckelmann, and W.G. Weir. 1987. Sodium–calcium exchange in heart: membrane currents and changes in [Ca], *Science.* 238:1720–1722.

Bassani, J.W.M., R.A. Bassani, and D.M. Bers. 1993. Ca2+ cycling between sarcoplasmic reticulum and mitochondria in rabbit cardiac myocytes. *J. Physiol. (Camb.).* 460:603–621.

Bassani, J.W.M., R.A. Bassani, and D.M. Bers. 1994a. Relaxation in...
rabbit and rat cardiac cells: species-dependent differences in cellular mechanisms. J. Physiol. (Camb.). 476:270–293.
Bassani, J.W.M., R.A. Bassani, and D.M. Bers. 1994b. Relaxation in ferret ventricular myocytes: unusual interplay among calcium transport systems. J. Physiol. (Camb.). 476:295–308.
Bassani, J.W.M., R.A. Bassani, and D.M. Bers. 1995. Calibration of Indo-1 and resting intracellular [Ca], in intact rabbit cardiac myocytes. Biophys. J. 68:1453–1460.
Bassani, J.W.M., W.-L. Yuan, and D.M. Bers. 1995c. Fractional SR Ca release is regulated by trigger Ca and SR Ca content in cardiac myocytes. Am. J. Physiol. 268:C1313–C1329.
Bassani, R.A., and D.M. Bers. 1994. Na–Ca exchange is required for rest-decay but not for rest-potentiation of twitches in rabbit and rat ventricular myocytes. J. Mol. Cell. Cardiol. 26:1335–1347.
Bassani, R.A., and D.M. Bers. 1995. Rate of diastolic Ca release from the sarcoplasmic reticulum of intact rabbit and rat ventricular myocytes. Biophys. J. 68:2015–2022.
Bassani, R.A., J.W.M. Bassani, and D.M. Bers. 1995a. Relaxation in ferret ventricular myocytes: role of the sarcolemmal Ca ATPase. Pflügers Arch. 430:573–578.
Berlin, J., J.W.M. Bassani, and D.M. Bers. 1994. Intrinsic cytosolic calcium buffering properties of single rat cardiac myocytes. Biochimie. 67:1775–1787.
Bers, D.M., and J.R. Berlin. 1995. Kinetics of [Ca], decline in cardiac myocytes depend on peak [Ca]. Am. J. Physiol. 268:C271–C277.
Callevaert, G., L. Cleeman, and M. Morad. 1989. Caffeine-induced Ca2+ release activates Ca2+-extrusion via Na+-Ca2+ exchanger in cardiac myocytes. Am. J. Physiol. 257:C147–C152.
Cheng, H., W.J. Lederer, and M.B. Cannell. 1993. Calcium sparks: elementary events underlying excitation–contraction coupling in heart muscle. Science. 252:740–744.
Cheng, H., M.R. Lederer, W.J. Lederer, and M.B. Cannell. 1996. Calcium sparks and [Ca2+] waves in cardiac myocytes. Am. J. Physiol. 270:C148–C159.
Delbridge, L.M.D., J.W.M. Bassani, and D.M. Bers. 1996. Steady-state twitch Ca2+ fluxes and cytosolic Ca2+ buffering in rabbit ventricular myocytes. Am. J. Physiol. 270:C192–C199.
de Meis, L., and G. Inesi. 1992. Functional evidence of a transmembrane channel within the Ca2+ transport ATPase of sarcoplasmic reticulum. FEBS Lett. 299:33–35.
Díaz, M.E., A.W. Trafford, S.C. O’Neill, and D.A. Eisner. 1997. Measurement of sarcoplasmic reticulum Ca2+ content and sarcosomal Ca2+ fluxes in isolated rat ventricular myocytes during spontaneous Ca2+ release. J. Physiol. (Camb.). 501:3–16.
Dixon, D.A., and D.H. Haynes. 1989. Kinetic characterization of the Ca pumping ATPase of cardiac sarcolemma in four states of activation. J. Biol. Chem. 264:13612–13622.
Fehér, J., and F.N. Briggs. 1984. Unidirectional calcium and nucleotide fluxes in cardiac sarcoplasmic reticulum II. Experimental results. Biochim. Biophys. Acta. 761:211–232.
Ginsburg, K.S., and D.M. Bers. 1996. Sarcoplasmic reticulum Ca content in intact ferret ventricular myocytes. Biophys. J. 70:4209.
Gómez, A., H. Cheng, J.W. Lederer, and D.M. Bers. 1996. Ca2+ diffusion and sarcoplasmic reticulum transport both contribute to Ca2+ decline during Ca2+ sparks in rat ventricular myocytes. J. Physiol. (Camb.). 496:571–581.
Grantham, G.J., and M.B. Cannell. 1996. Ca2+ influx during the cardiac action potential in guinea pig ventricular myocytes. Circ. Res. 79:194–200.
Grynkiewicz, G., M. Poenie, and R.Y. Tsien. 1985. A new generation of Ca2+ indicators with greatly improved fluorescence properties. J. Biol. Chem. 260:3440–3450.
Hadley, R.W., and J.R. Hume. 1987. An intrinsic potential-dependent inactivation mechanism associated with calcium channels in guinea-pig myocytes. J. Physiol. (Camb.). 380:205–222.
Hale, S., A. Schiefer, and G. Isenberg. 1994. Ca2+ load of guinea-pig ventricular myocytes determines efficacy of brief Ca2+ currents as trigger for Ca2+ release. J. Physiol. (Camb.). 480:411–421.
Hove-Madsen, L., and D.M. Bers. 1992. Indo-1 binding to protein in permeabilized ventricular myocytes alters its spectral and Ca binding properties. Biophys. J. 63:89–97.
Hove-Madsen, L., and D.M. Bers. 1993a. Passive Ca buffering and SR Ca uptake in permeabilized rabbit ventricular myocytes. Am. J. Physiol. 264:C677–C686.
Hove-Madsen, L., and D.M. Bers. 1993b. Sarcoplasmic reticulum Ca2+ uptake and thapsigargin sensitivity in permeabilized rabbit and rat ventricular myocytes. Circ. Res. 73:829–828.
Inesi, G., and L. de Meis. 1988. Regulation of steady state filling in sarcoplasmic reticulum. J. Biol. Chem. 264:5929–5936.
Janeczkowski, A.M., H.A. Spurgeon, M.D. Stern, and E.G. Lakatta. 1995. Effects of sarcoplasmic reticulum Ca2+ load on the gain function of Ca2+ release by Ca2+ current in cardiac cells. Am. J. Physiol. 268:H916–H920.
Kawai, M., and M. Konishi. 1994. Measurement of sarcoplasmic reticulum calcium content in skinned mammalian cardiac muscle. Cell Calcium. 16:123–136.
Kimura, J., S. Miyamae, and A. Noma. 1987. Identification of sodium–calcium exchange current in single ventricular cells in guinea pig. J. Physiol. (Camb.). 384:199–222.
Kirchberger, M.A., and D. Wong. 1978. Calcium efflux from isolated cardiac sarcoplasmic reticulum. J. Biol. Chem. 253:6941–6945.
Lamont, C., and D. Eisner. 1996. The sarcolemmal mechanisms involved in the control of diastolic intracellular calcium in isolated rat cardiac trabeculae. Pflügers Arch. 432:961–969.
Li, L., E.G. Kranias, and D.M. Bers. 1997. Calcium transport in phospholamban knockout mouse: relaxation and endogenous CaMKII effects. Biochim. Biophys. Acta. 132:4234.
Makinoose, M., and W. Hasselback. 1971. ATP synthesis by the reversal of the sarcoplasmic reticulum calcium pump. FEBS Lett. 12:271–272.
Negretti, N., S.C. O’Neill, and D.A. Eisner. 1993. The effects of inhibitors of sarcoplasmic reticulum function on the systolic Ca2+ transient in rat ventricular myocytes. J. Physiol. (Camb.). 468:35–52.
Page, E., L.P. McCallister, and B. Power. 1971. Stereological measurements of cardiac ultrastructure implicated in excitation–contraction coupling. Proc. Natl. Acad. Sci. USA. 68:1465–1466.
Reeves, J.P., and K.D. Philipson. 1989. Sodium–calcium exchange activity in plasma membrane vesicles. In Sodium–Calcium Exchange. T.J.A. Allen, D. Noble, and H. Reuter, editors. Oxford University Press, Oxford, UK. 27–53.
Sagara, Y., and G. Inesi. 1991. Inhibition of the sarcoplasmic reticulum Ca2+ transport ATPase by thapsigargin at subnanomolar concentrations. J. Biol. Chem. 266:12506–12613.
Satoh, H., L.A. Blatter, and D.M. Bers. 1997. Effects of [Ca2+]i, Sr2+ load, and rest on Ca2+ spark frequency in ventricular myocytes. Am. J. Physiol. 272:H657–H668.
Satoh, H., L.M. Delbridge, L.A. Blatter, and D.M. Bers. 1996. Surface-volume relationship in cardiac myocytes studied with confocal microscopy and membrane capacitance measurements: Species-dependence and developmental effects. Biochips. 26:1494–1504.
Segel, I. 1976. Biochemical Calculations. 2nd ed. John Wiley & Sons, New York. pp. 220–221.
Sham, S.K., L. Cleeman, and M. Morad. 1995. Functional coupling of Ca2+ channels and ryanodine receptors in cardiac myocytes. Proc. Natl. Acad. Sci. USA. 92:121–125.
Shannon, T., and D.M. Bers. 1997. Assessment of intrasR free [Ca]...
and buffering in rat heart. Biophys. J. 73:1524–1531.
Shannon, T., K.S. Ginsburg, and D.M. Bers. 1997. SR Ca uptake rate in permeabilized ventricular myocytes is limited by reverse mode of the SR Ca pump. Biophys. J. 72:A167.
Sipido, K.R., G. Callewaert, and E. Carmeliet. 1995. Inhibition and rapid recovery of Ca\textsuperscript{2+} current during Ca\textsuperscript{2+} release from sarcoplasmic reticulum in guinea pig ventricular myocytes. Circ. Res. 76:102–109.
Sipido, K.R., and W.G. Weir. 1991. Flux of Ca\textsuperscript{2+} across the sarcoplasmic reticulum of guinea pig cardiac cells during excitation-contraction coupling. J. Physiol. (Camb.). 435:605–630.
Tada, M., M.A. Kirchberger, D.I. Repke, and A.M. Katz. 1974. The stimulation of calcium transport in cardiac sarcoplasmic reticulum by adenosine 3\textsuperscript{′}-5\textsuperscript{′}-monophosphate-dependent protein kinase. J. Biol. Chem. 249:6174–6180.
Terraciano, C.M.N., and K. MacLeod. 1994. The effect of acidosis on Na\textsuperscript{+}/Ca\textsuperscript{2+} exchange and consequences for relaxation in isolated cardiac myocytes from guinea-pig. Am. J. Physiol. 267:H477–H478.
Terraciano, C.M.N., and K. MacLeod. 1997. Measurements of Ca\textsuperscript{2+} entry and sarcoplasmic reticulum Ca\textsuperscript{2+} content during the cardiac cycle in guinea pig and rat ventricular myocytes. Biophys. J. 72:1319–1326.
Trafford, A.W., M.E. Diaz, and D.A. Eisner. 1997. Enhanced calcium current and decreased Ca efflux restore sarcoplasmic reticulum Ca content following depletion. J. Physiol. (Camb.). In press.
Varro, A., N. Negretti, and D.A. Eisner. 1993. An estimate of the calcium content of the sarcoplasmic reticulum in rat ventricular myocytes. Pflügers Arch. 423:158–160.
Wolsker, H., and L. de Meis. 1985. Ligand-gated channel of the sarcoplasmic reticulum Ca\textsuperscript{2+} transport ATPase. Biosci. Rep. 15:365–376.
Wolska, B.M., M.O. Stojanovic, W. Luo, E.G. Kranias, and R.J. Solorio. 1996. Effect of ablation of phospholamban on dynamics of cardiac myocyte contraction and intracellular Ca\textsuperscript{2+}. Am. J. Physiol. 271:C391–C397.
Yuan, W.-L., and D.M. Bers. 1995. Protein kinase inhibitor H-89 reverses forskolin stimulation of cardiac L-type calcium current. Am. J. Physiol. 268:C651–C659.