Local Control Models of Cardiac Excitation–Contraction Coupling
A Possible Role for Allosteric Interactions between Ryanodine Receptors

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ABSTRACT In cardiac muscle, release of activator calcium from the sarcoplasmic reticulum occurs by calcium-induced calcium release through ryanodine receptors (RyRs), which are clustered in a dense, regular, two-dimensional lattice array at the diad junction. We simulated numerically the stochastic dynamics of RyRs and L-type sarcolemmal calcium channels interacting via calcium nano-domains in the junctional cleft. Four putative RyR gating schemes based on single-channel measurements in lipid bilayers all failed to give stable excitation–contraction coupling, due either to insufficiently strong inactivation to terminate locally regenerative calcium-induced calcium release or insufficient cooperativity to discriminate against RyR activation by background calcium. If the ryanodine receptor was represented, instead, by a phenomenological four-state gating scheme, with channel opening resulting from simultaneous binding of two Ca\(^{2+}\) ions, and either calcium-dependent or activation-linked inactivation, the simulations gave a good semiquantitative accounting for the macroscopic features of excitation–contraction coupling. It was possible to restore stability to a model based on a bilayer-derived gating scheme, by introducing allosteric interactions between nearest-neighbor RyRs so as to stabilize the inactivated state and produce cooperativity among calcium binding sites on different RyRs. Such allosteric coupling between RyRs may be a function of the foot process and lattice array, explaining their conservation during evolution.

KEY WORDS: sarcoplasmic reticulum • Monte Carlo • calcium-induced calcium release • dihydropyridine receptor • diad junction

INTRODUCTION

The intracellular signal that triggers the contraction of cardiac muscle is a transient rise in intracellular free calcium. The majority of this calcium (50–95%, depending upon species and conditions) is released from the sarcoplasmic reticulum (SR). This calcium is released by a process of calcium-induced calcium release (CICR) (Endo et al., 1970; Fabiato, 1985) via release channels, which have been shown to be type-2 ryanodine receptors (RyR2). It is generally believed that the major stimulus for CICR is calcium entering the cell via sarcolemmal dihydropyridine-sensitive L-type calcium channels. The rate and amount of calcium release from the SR is tightly controlled by the magnitude and duration of the L-type calcium current (See Ríos and Stern, 1997, for a review of the evidence). This graded control is paradoxical because the released calcium, which is roughly 10\(^x\) larger in amount than the trigger, would be expected to stimulate further CICR, leading to a regenerative, nearly all-or-none release. Several years ago (Stern, 1992), we proposed that this paradox of control might be explained if the stimulus for release of calcium by RyRs were actually the local nanodomains of [Ca\(^{2+}\)] generated by nearby L-type channels, rather than the global cytosolic [Ca\(^{2+}\)]. According to this local control hypothesis, the graded control of macroscopic SR calcium release would actually be achieved by graded statistical recruitment of individual, autonomous, stochastic release events.

Recent ultrastructural studies (Franzini-Armstrong, C., F. Protasi, and V. Ramesh, manuscript in preparation) show that, depending on species, from a few tens to \(~200\) RyRs are clustered in a two dimensional crystal-lattice array on the surface of SR release terminals, apposed, across the 15-nm cleft of the diad junction, to clusters of sarcolemmal dihydropyridine receptors (DHPRs). RyR2 is a homotetramer of a polypeptide of roughly 5,000 amino acids. The transmembrane calcium-sensitive channel is formed by the COOH-termini-
nal ~600 amino acids (Tunwell et al., 1996; Bhat et al., 1997a), while the remainder of the molecule forms a 30-nm quatrefoil “foot process” that spans the diadic cleft, and is required for interaction of the channel with a variety of modulators (Bhat et al., 1997b). The DHPRs in cardiac muscle, whose number varies in different studies from 10–100% of the number of RyRs (Wibo et al., 1991; Bers and Stiffel, 1993; Sun et al., 1995), are localized at the junctions, but are randomly positioned relative to the ryanodine receptor lattice (Franzini-Armstrong, C., F. Protasi, and V. Ramesh, manuscript in preparation). This contrasts with the regular arrangement of DHPR tetrads found in skeletal muscle.

We carried out numerical simulations of this system of channels, interacting stochastically via calcium diffusing in the diadic cleft. These simulations have revealed a new paradox. Local control succeeds if the gating of the RyR is represented by a simple, phenomenological, four-state scheme. However, published schemes derived from actual gating statistics of single RyR2 channels incorporated into lipid bilayers give rise to unacceptable instabilities when used in the local control model. These instabilities are traceable to two deficiencies in bilayer-derived gating schemes. The absence of strong inactivation prevents termination of locally regenerative release by clustered RyRs. Activation by binding of a single Ca\(^{2+}\) ion, in some bilayer-derived schemes, does not provide adequate discrimination against activation by global (rather than microdomain) Ca\(^{2+}\). This suggests that the gating of RyR2 in situ may differ significantly from its behavior in bilayers. One possible explanation of this difference would be the existence of allosteric interactions between the large foot processes of adjacent RyRs, which appear anatomically to be in contact. We show by simulation that RyR–RyR allosteric interaction energies can be chosen in such a way as to remedy both the inactivation and the cooperativity deficiencies of RyR2 gating schemes. Such interactions may be one of the important functions of the foot process, which has been highly conserved in evolution.

**METHODS**

Simulation

Monte Carlo simulations were carried out using a modification of an algorithm previously reported (Stern et al., 1997). In brief, the set of channels at each diad was treated as a single stochastic system whose state (“macrostate”) is defined by specifying the Markov state of each of the individual channels. State transitions of individual channels are considered to take place instantaneously so that a macrostate transition corresponds to a transition of exactly one channel. The transition rates of the individual channels were determined from their gating schemes as a function of the local [Ca\(^{2+}\)] at the position of each channel. The dwell time and destination of each transition were selected by use of appropriately distributed random numbers, while the local [Ca\(^{2+}\)] was determined concomitantly by solving the partial differential equations governing diffusion and binding reactions in the diadic cleft. The aggregate calcium fluxes produced by 1,000–10,000 such diads were taken as input to a conventional lumped-compartment model of global cytosolic and SR calcium dynamics. This model (a set of ordinary differential equations) was solved to update the values of [Ca\(^{2+}\)]\(_{\text{cytosol}}\) and [Ca\(^{2+}\)]\(_{\text{SR}}\)

These were used, in turn, to determine unitary currents of the channels and to set boundary conditions at the edges of the diadic clefts. The global model consisted of seven differential equations governing the following dynamical variables: [Ca\(^{2+}\)]\(_{\text{cytosol}}\) and [Ca\(^{2+}\)]\(_{\text{SR}}\) luminal calcium (in rapid equilibrium with calsequestrin), and calcium bound to five types of buffer sites, representing calcium-sensing dye (fuo-3), troponin, calmodulin, SR membrane sites, and low affinity sarcoplasmic sites. The SR calcium pump was modeled as a steady uptake, represented by a Hill function of [Ca\(^{2+}\)]\(_{\text{endo}}\), balanced (in the resting state) by a leak proportional to luminal [Ca\(^{2+}\)]. Compartmental volumes and the kinetic parameters of all buffers and pumps were taken from the literature (Table I) and were not treated as adjustable parameters of the model. The details of the cytosolic calcium removal model have very little effect on the short term simulations presented in this paper.

The extreme computational demands of such a simulation required the use of approximations to the reaction–diffusion equations in the diadic cleft. Because of the large aspect ratio of the cleft (diameter 100–400 nm, height 15 nm), diffusion was treated in two dimensions only, and the equations were discretized on a 10-nm mesh. This coarse approximation can be justified because the locations of the exit pore(s) and calcium sensing site(s) of the ryanodine receptor, as well as the diffusion paths of Ca\(^{2+}\) through the space occupied by the RyR foot processes, are not known with any greater precision. L-type channels were located at random on the mesh, while RyRs were located regularly at 30-nm center-to-center spacing (Fig. 1). Effective values of the cleft height and calcium diffusion coefficient were used to approximate the effects of surface charge, viscosity and tortuosity, as estimated by Soeller and Cannell (1997). Diads containing up to 121 RyRs were simulated; most of the results shown here were computed for 25-RyR diads containing an average of five randomly located DHPRs that display the qualitative features and save computation time.

Even with these approximations, computation times of hours were required for the full simulation. For this reason, many of the results shown were computed using the approximation that calcium diffusion in the cleft is at steady-state during the dwell time in any given macrostate, which accelerates the computation by about two orders of magnitude. The accuracy of this approximation depends on the density of fixed calcium binding sites in the diadic cleft. In the absence of such sites, it agrees with the full simulation to within a few percent. In the presence of low affinity fixed Ca\(^{2+}\) binding sites at the density estimated by Post and Langer (1992), errors of 30–50% may occur with the steady state approximation, but the qualitative features of the results appear to be correct, and most of the quantitative results can be recovered by use of effective values of channel parameters that are well within the large experimental uncertainties (Fig. 2). The fixed buffer sites in the cleft do not appear explicitly in the steady state diffusion approximation since these sites are nonmobile and therefore have no effect on the steady state distribution of [Ca\(^{2+}\)], although they control the rate at which the steady state profile is established after a change in the sources.

**L-Type Channel**

There is no consensus on the gating scheme of the L-type sarcoplasmic calcium channel. In these simulations, it was modeled

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using a novel 24-state gating scheme that we developed as an improvement of the scheme recently proposed by Jafri et al. (1998). The new scheme combines the scheme of Imredy and Yue (1994) for calcium-dependent inactivation with that proposed by Shirokov et al. (1993) to explain gating charge movement and voltage-dependent inactivation. This scheme (shown in Fig. 3A) consists of four-gating modes (normal, calcium-inactivated, voltage-inactivated, and both calcium and voltage inactivated). This model is in the process of being validated and parametrized using measured whole cell and single channel calcium currents. For the purposes of this paper, it may be thought of as an empirical source of DHPR openings that act as input to the CICR process. Fig. 3, B and C, shows representative L-type calcium currents measured from a rat cardiac myocyte in the presence of 10 μM ryanodine. The parameters of the L-type gating model were adjusted manually to fit these data, as shown, and these parameter values were used for most of the simulations shown in the rest of the paper without any further adjustment. They should not be considered necessarily optimal or unique.

**RyR Gating Schemes**

Simulations were carried out using six different gating schemes for RyR2, shown in Fig. 4. Two of these are phenomenological

| Variable | Description/units | Value  |
|----------|--------------------|--------|
| B2T      | Total calmodulin Ca sites, mM | 0.024* |
| B3T      | Total troponin Ca sites, mM | 0.07* |
| B4T      | Total SR surface Ca sites, mM | 0.047* |
| B5T      | Total SL low-affinity Ca sites, mM | 1.124* |
| B7       | Total dye (fluo-3), mM | 0.1 |
| BCA0     | Starting cytosolic [Ca^{2+}], mM | 0.0001 |
| BCAIREST | Resting intracellular [Ca^{2+}], mM | 0.0001 |
| BCAO     | Extracellular [Ca^{2+}], mM | 1.0 |
| BCAOREST | Resting extracellular [Ca^{2+}], mM | 1.0 |
| BCA0R    | Starting SR calcium content, mmol/liter cytosol | 0.15* |
| BCA0RR   | Resting SR calcium content, mmol/liter cytosol | 0.15* |
| BCSQT    | [Calsequestrin sites], mM in SR lumen | 3.0 |
| BETA     | Na/Ca voltage splitting parameter | 0.35* |
| BKOFF    | Calmodulin Ca off rate, ms^-1 | 0.038* |
| BK20N    | Calmodulin Ca on rate, mM^-1 ms^-1 | 100.0* |
| BK20N    | Troponin Ca off rate, ms^-1 | 0.02* |
| BK30N    | Troponin Ca on rate, mM^-1 ms^-1 | 39.0* |
| BK40F    | SR surface site Ca off rate, ms^-1 | 0.1* |
| BK40N    | SR surface site Ca on rate, mM^-1 ms^-1 | 115.0* |
| BK50F    | SL surface site Ca off rate, ms^-1 | 1.0* |
| BK50N    | SL surface site Ca on rate, mM^-1 ms^-1 | 115.0* |
| BKC0     | Na/Ca exchanger K_i for Ca, mM | 1.38 |
| BKDCSQ   | K_i of calsequestrin sites, mM | 0.638* |
| BNKA     | Na/Ca K_i Na, mM | 87.5* |
| BKOFF    | Dye (fluo-3) Ca off rate, ms^-1 | 0.09* |
| BKON     | Dye (fluo-3) Ca on rate, mM^-1 ms^-1 | 80.0* |
| BKPOUM   | SR Ca pump affinity, mM | 0.0003* |
| BKPOUMR  | Resting SR Ca pump affinity, mM | 0.0003* |
| BKSAT    | Na/Ca saturation parameter | 0.1 |
| BNAI     | Cytosolic [Na^+], mM | 10.0 |
| BNAIRES  | Resting intracellular [Na^+], mM | 10.0 |
| BNAO     | Extracellular [Na^+], mM | 140.0 |
| BNAOREST | Resting extracellular [Na^+], mM | 140.0 |
| BNAP     | SR Ca pump cooperativity | 2.0 |
| BNRNC    | Na/Ca exchanger rate parameter, ms^-1 | 0.2 |
| BNSRF    | (Cytosol volume)/(SR volume) | 17.6 |
| BNVAC    | Relative volume density of diads | 1.0 |
| BNSVR    | SR Ca pump maximum rate, mM ms^-1/cytosolic volume | 0.00002 |
| BNSVR    | Resting SR Ca pump v_max, mM ms^-1/cytosolic volume | 0.00002 |
| BNVRREST | Resting potential, mV | -70.0 |

*Stern et al., 1998; †Bers, 1998; ‡Luo and Rudy, 1994; \$VFAC is proportional to the volume density of diads, scaled so that VFAC = 1 corresponds to the density of ryanodine receptors estimated by Bers (1999).
schemes representing, in simplest form, the combination of activation by binding of two calcium ions and either calcium-dependent or fateful inactivation. The other four are published gating schemes based on single-channel observations of purified RyRs in lipid bilayers.

Measurement of SR Release Flux

SR release flux was measured by the recently developed Oregon green/EGTA method, which has been described in detail by Song et al. (1998). In brief, adult rat ventricular cardiac myocytes were enzymatically isolated and studied under whole cell patch-pipette voltage clamp. The fast, low affinity fluorescent calcium indicator Oregon green 488 BAPTA 5N (1 mM) and EGTA (4 mM) were dialyzed into the cell from the pipette. As we have shown previously, this combination gives a fluorescence signal dominated by a term directly proportional to SR release flux. Myocytes were imaged in the line scan mode of an inverted confocal microscope (LSM-410; Carl Zeiss, Inc.), with the scan line oriented longitudinally. This method makes it possible to see the localized SR release at the Z line of each sarcomere. Whole cell calcium release rate was estimated by averaging over the length of the scan line.

RESULTS

Local Control Produces Graded Release of SR Calcium

Fig. 5 shows representative measurements of calcium release flux from the SR of a rat cardiac myocyte, compared with simulations using the phenomenological RyR gating Scheme 5 in Fig. 4. Fig. 5 (left) shows the time course of SR release flux during a single voltage clamp depolarization to 0 mV from a holding potential of −70 mV. The figure demonstrates that the local control model can simulate the observed time course of release, particularly the fact that release declines during sustained depolarization. The fact that release in this model is dependent on local geometry is illustrated by the much smaller dotted curve, which shows the release flux that would be computed assuming a diadic cleft height of 30 nm instead of the correct 15 nm. Fig. 5 (right) shows the peak release flux as a function of the voltage of the clamp pulse. $I_{Ca}$, the macroscopic current of the L-type channels, shows the bell-shaped voltage dependence typical of voltage-controlled ion channels, while the SR flux recapitulates (approximately, see below) the voltage dependence of $I_{Ca}$, but at an absolute magnitude 10-fold larger. The simulation reproduces these features well. This demonstrates the central point of the local control hypothesis: the complicated stochastic interactions of calcium-coupled channels can produce high amplification with stable, graded control by trigger calcium. As shown in Fig. 6, if the bell shaped $I_{Ca}$ vs. V and $I_{SR}$ vs. V curves are normalized to their respective peaks, there is a small difference between their positions. When these curves are then divided to give a plot of the gain of excitation–contraction (EC) coupling (peak $I_{SR}$ divided by peak $I_{Ca}$), the resulting gain diminishes with voltage. This was one of the original predictions of
the local control theory, which has been confirmed in several laboratories (Wier et al., 1994; Cannell et al., 1995; Janczewski et al., 1995; Santana et al., 1996). It stems from the fact that local activation of the RyR depends on the unitary current of the triggering L-type channel(s) rather than on the macroscopic ensemble-average current. As pointed out previously (Cannell et al., 1995; Santana et al., 1996), the decline of gain at positive voltages is a manifestation of the cooperativity of RyR activation by calcium, because the diminution of L-type unitary current as the calcium reversal potential is approached reduces $I_{SR}$ disproportionately in comparison to its linear effect on macroscopic $I_{CaL}$. As argued by those authors, a quadratic dependence of RyR activation on local calcium should result in a macroscopic gain that is roughly proportional to the L-type unitary current. The decline of gain with voltage in the simulations is steeper than that of the L-type current over the midrange of voltages (Fig. 6 B), even though activation of the RyR was assumed to require two Ca$^{2+}$ ions. This “elbow” in the voltage-gain relation results from the depletion of SR calcium (by release to the cytosol) during the depolarization, as indicated by the fact that it is prevented if SR depletion is minimized by reducing the number of diads per unit cytosolic volume, or reducing the number of available L-type chan-
Figure 3. (A) Structure of the scheme of L-type channel gating. The scheme consists of four modes. Each mode (above) consists of a cascade of five closed states connected by movement of four gating charges (horizontal transitions). Openings occur from the rightmost closed state. Each mode has its own center potential and barrier symmetry constants ($V_{ctr}$ and $\eta$). Inactivation is due to mode switching (vertical transitions) to modes with bound calcium, left-shifted $V_{ctr}$, or both (below). “Voltage-dependent” inactivation is actually due to voltage-independent transitions to left-shifted modes, which are strongly favored in the rightward charge states because of the requirements of microscopic reversibility, as suggested by Shirokov et al. (1993). (B and C) Measured L-type currents from a cardiac myocyte in the presence of 10 μM ryanodine are superimposed on simulations with SR release disabled, in the time (B) or voltage (C) domain. Simulated and measured currents were normalized so as to make the peak value of both equal to 1 at 0 mV. The L-type model parameters (Table II) were adjusted “by hand” to achieve the fit.

\[
\alpha = k \exp(\eta (V-V_{ctr})/K)
\]

\[
\beta = k \exp((\eta - 1)(V-V_{ctr})/K)
\]
nels (not shown). Such an elbow has been observed experimentally in intact cardiac myocytes (see Fig. 3 in Wier et al., 1994). It was not seen when $I_{Ca}$ (and therefore $I_{SR}$ and SR depletion) was reduced by a factor of 20 with nifedipine in order to study calcium sparks (Cannell et al., 1995).

**SR Calcium Release Is Locally Regenerative**

The number of RyRs that contribute to SR calcium release at a single diad is not known. As an index of that number, we computed the mean number of RyR openings per diad during a clamp pulse to 0 mV for 50 ms, averaged over those diads at which at least one RyR opening occurred. Neither the unitary current $i_{RyR}$ of the RyR nor its calcium sensitivity in situ is well known. We therefore varied the RyR permeability over two orders of magnitude, adjusting the sensitivity $k_i$ in each case to give the observed macroscopic gain of 10 at 0 mV and standard SR loading, to determine whether there could be a regime in which an L-type channel communicates with a single RyR without regenerative local recruitment of other RyRs. As shown in Fig. 7, the mean number of RyR openings per diad has a U-shaped dependence on $i_{RyR}$, with a nadir of $\sim 11$ (achieved, coincidentally, near the estimated physiologic $i_{RyR}$ of 0.4 pA; Mejia-Alvarez et al., 1999). This shows that, at least in the context of this model, RyR activation is always locally regenerative. This result can be understood qualitatively as follows. If $i_{RyR}$ were small, the RyR sensitivity would have to be high, since many RyRs would need to be recruited to achieve the macroscopic gain of 10. If $i_{RyR}$ were high, on the other hand, the sensitivity must be low, as a result of which the coupling from DHPR to RyR frequently fails. The release in this case is made up of rare diadic release events, each of which is highly regenerative. The fact that the DHPR-RyR communication fails before RyR-RyR recruitment is basically a consequence of geometry: if the DHPRs are located randomly at the junction, they cannot, on average, be much closer to RyR sensing sites than the latter are to the release pores of neighboring RyRs. If DHPRs were located in registry with RyRs, and if the calcium-sensing site of the latter were located on the “top” of the foot process, within a few nanometers of the DHPR, then nonregenerative 1:1 communication would be possible, as originally envisioned in the model of Györke and Palade (1993).  

**Local and Global Stability**

Before discussing simulations using empirically determined RyR gating schemes, we define two different concepts of stability. Local stability refers to the fact that there must be some mechanism to terminate the regenerative release at an individual diad. Global stability refers to the fact that the resting state of the myocyte must be stable against regenerative build up of global calcium, and the cell must eventually relax to this resting state after stimulation. The average diadic release rate triggered by global cytosolic calcium must, therefore, be less than the removal rate (SR pump + Na/Ca exchange) in the vicinity of the resting state. These two concepts of stability are distinct, though related. If there were no calcium removal, $[Ca^{2+}]_{cyt}$ would eventually build up to cause global regenerative release, no matter how well behaved the RyRs. On the other hand, it is evident that a cell cannot return to the resting state unless local release terminates. Both forms of stability are adversely affected by the clustering of RyRs. Clustered RyRs will have a more regenerative local release, which will be more difficult to terminate. Release triggered by background calcium will also be enhanced by clustering, since if a single RyR in the cluster opens in response to background calcium, this will trigger opening of most of the other RyRs in the cluster.

There are three mechanisms that contribute to the termination of local release: (a) depletion of SR calcium, (b) inactivation of RyRs, and (c) stochastic attrition. The last refers to the fact that for a finite-size cluster of RyRs, gating stochastically, there is always a nonzero probability that all RyRs will be closed at the same time, and this event will interrupt the positive feedback and extinguish the activity of the cluster, as long as there are no further DHPR openings to re-ignite it. SR depletion cannot be the principal mechanism of local release termination, since release terminates after voltage steps to very negative or positive voltage, which release very little of the SR calcium stores, and release in calcium sparks (Cheng et al., 1993) also terminates despite the fact that there is no global SR depletion. Conversely, greatly prolonged sparks occur in the presence of ryanodine (Cheng et al., 1993), indicating the absence of local SR depletion. Inactivation and stochastic attrition must, together, be capable of terminating release at a single diad.

The contribution of stochastic attrition may be roughly estimated from a simple, analytical model. Consider a “cluster” of $n$ identical channels, each of which has only two states, open and closed. Start with each channel in equilibrium, with an open probability $P_o$ and an opening duration $\tau_o$. The channels are assumed to gate independently, except that, if all close at once, the cluster is considered to be extinct. So long as the cluster is “alive,” the number of open channels, $n_o$, will be binomially distributed, except that $n_o = 0$ is excluded. The probability that a cluster becomes extinct during a time interval $dt$ is given by

$$\frac{dt}{\tau_{\text{attrit}}} = (\text{probability that only one channel is open}) \times (\text{probability that last channel closes}),$$

where $\tau_{\text{attrit}}$ is defined to be the time constant for extinction of clusters by stochastic attrition. Using this re-
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...relationship together with the binomial distribution, it is straightforward to determine the attrition time constant:

\[ \tau_{\text{attrit}} = \frac{\tau_0[1 - (1 - P_o)^n]}{n(1 - P_o)^{n-1}P_o}. \] (1)

This time constant varies roughly exponentially with the product \(nP_o\), the expected number of open channels. Some numerical values are instructive. For a cluster of 25 channels with a mean opening duration of 10 ms and \(P_o = 10\%\), \(\tau_{\text{attrit}}\) is \(\sim 46\) ms. For 75 channels with a 50\% open probability, it becomes 160 billion years, \(\sim 10^9\) the age of the universe! This makes it clear that stochastic attrition alone cannot be relied upon to terminate local release robustly if the number of RyRs in a diad is as large as present ultrastructural data indicate. There must also be an inactivation process that is capable of substantially reducing \(P_o\). For an opening duration of 10 ms, one finds numerically that to extinguish clusters with a time constant of 50 ms requires \(nP_o < 2.6\). Therefore, for RyR clusters of the size found in car-

| Variable | Description/units | Value |
|----------|------------------|-------|
| BPERM | L-type permeation constant (effective Ca activity coeff.) | 0.341* |
| CA1 | L-type permeation constant (equivalent channel mouth \([Ca^{2+}]\)) | 0.001* |
| ETAM01 | L-type mode-normal mode-V energy splitting factor | 0.84* |
| ETAM02 | L-type mode-normal mode-Ca energy splitting factor | 0.5* |
| ETAM13 | L-type mode-Ca mode-VCa energy splitting factor | 0.5* |
| ETAM23 | L-type mode-Ca mode-VCa energy splitting factor | 0.84* |
| ETAV0 | L-type voltage sensor barrier symmetry, normal mode | 0.5* |
| ETAV1 | L-type voltage sensor barrier symmetry, V-inact. mode | 0.5* |
| ETAV2 | L-type voltage sensor barrier symmetry, mode-Ca | 0.5* |
| ETAV3 | L-type voltage sensor barrier symmetry, mode-VCa | 0.5* |
| F0 | L-type transition rate into open state, ms\(^{-1}\) | 0.3* |
| F1 | L-type transition rate into open state, mode-V, ms\(^{-1}\) | 0 |
| F3 | L-type transition rate into open state, mode-VCa, ms\(^{-1}\) | 0 |
| G0 | L-type transition rate into closed state, mode-normal, ms\(^{-1}\) | 2.0* |
| G1 | L-type transition rate into closed state, mode-V, ms\(^{-1}\) | 100.6* |
| G2 | L-type transition rate into closed state, mode-Ca, ms\(^{-1}\) | 20.0* |
| G3 | L-type transition rate into closed state, mode-V-Ca, ms\(^{-1}\) | 100.6* |
| KC0 | L-type voltage sensor transition rate, normal mode, ms\(^{-1}\) | 0.2* |
| KC1 | L-type voltage sensor transition rate, mode-V, ms\(^{-1}\) | 0.1* |
| KC2 | L-type voltage sensor transition rate, mode-Ca, ms\(^{-1}\) | 0.2* |
| KC3 | L-type voltage sensor transition rate, mode-VCa, ms\(^{-1}\) | 0.1* |
| KIV | L-type V-inactivation rate constant, ms\(^{-1}\) | 0.005\* |
| KIVCA | L-type V-inactivation rate constant from mode-Ca, ms\(^{-1}\) | 0.005\* |
| KIVM | L-type repriming rate constant, mode-V \(\rightarrow\) normal, ms\(^{-1}\) | 0.00056\* |
| KK | L-type voltage sensor steepness, mV | 10.0\* |
| KVIOFF | L-type Ca off-rate in VCa-inactivated mode, ms\(^{-1}\) | 0.0001\* |
| KVION | L-type Ca binding rate in V-inactivated mode, mM\(^{-1}\) ms\(^{-1}\) | 0.2* |
| KVION | L-type Ca binding rate in mode-V open state, mM\(^{-1}\) ms\(^{-1}\) | 0 |
| KVOFF | L-type Ca off-rate in mode-Ca, ms\(^{-1}\) | 0.0001\* |
| KVOON | L-type Ca binding rate in normal mode, mM\(^{-1}\) ms\(^{-1}\) | 0.2* |
| KVORN | L-type Ca binding rate in mode-N open state, mM\(^{-1}\) ms\(^{-1}\) | 9.0* |
| VCTR0 | L-type voltage sensor center voltage, normal mode, mV | -24.0* |
| VCTR1 | L-type voltage sensor center voltage, mode-V mV | -104.0* |
| VCTR2 | L-type voltage sensor center voltage, mode-Ca, mV | -34.0* |
| VCTR3 | L-type voltage sensor center voltage, mode-VCa, mV | -114.0* |
| UIV | L-type unitary current at 0 mV, pA | 0.1* |

*Jafri et al., 1998; ‡Values estimated from Shirokov et al., 1993, and Imredy and Yue, 1994, and adjusted manually to give a good fit to the data in Fig. 3 while preserving semiquantitative agreement with charge movement results of Shirokov et al. (1993). Transition rates between modes are specified for the proopen state of the voltage sensor C4; intermode transition rates in other voltage-sensor states are determined by detailed balancing. §Mode-V and mode-VCa of the L-type channel are assumed not to open.
diac muscle, local stability requires an inactivation pro-
cess that, while it need not be completely absorbing,
can reduce the RyR open probability to a few percent
or less. It needs to be kept in mind, however, that sto-
chastic attrition is not an alternative to inactivation, but
an intrinsic feature of multichannel local control mod-
els, which always contributes to the cluster extinction
process regardless of what other mechanisms are
present.

Fateful Inactivation of RyR Gives Stable EC Coupling

Since the stability of local control EC coupling depends
critically on the inactivation process of the RyR, it is
reasonable to ask whether the calcium-dependent inac-
tivation mechanism employed in the phenomenologi-
gal gating of Fig. 4, Scheme 5, is essential to the success
of the model. The answer is no; as shown in Fig. 8, qual-
itatively comparable results can be obtained with Fig. 4,
Scheme 6, in which inactivation is fatefully linked to ac-
tivation, but not explicitly dependent on calcium (as
suggested by Pizarro et al., 1997). In this scheme, un-
like Scheme 5, the activation and inactivation “gates”
are not independent. The inactivation and repriming
rate constants (vertical transitions) are different in the
calcium-free and -activated states of the channel (sub-
ject to the constraint of microscopic reversibility, which
requires that the products around the loop of forward
and reverse rates be equal). By proper choice of these
rate constants, it can be arranged that, on exposure to
Ca²⁺ the channel first activates and then inactivates,
while on removal of Ca²⁺, channels predominantly de-
activate before repriming, avoiding another passage
through the open state. This arrangement requires
some tuning to secure sufficiently rapid decay of re-
lease both on depolarization and repolarization.

Bilayer-derived RyR Gating Schemes Give Unstable
EC Coupling

We now turn to consideration of models based on the
empirically derived RyR gating Schemes 1–4 in Fig. 4.
Each of these schemes proves to have one or more defi-
ciencies that preclude local and global stability. We can
consider, first, Scheme 3 proposed by Keizer and Le-
vine (1996). This scheme was developed to explain the
“adaptation” of the RyR observed by Györke and Fill
(1993). When used in an ensemble-average sense, where [Ca²⁺] is taken to be a global value imposed by
the researcher, this scheme gives a reasonable descrip-
tion of RyR adaptation as seen in bilayers with Cs⁺ as
current carrier. It is clear, however, that it cannot be
considered as a description of microscopic events in a
local control model, because it has a calcium-depen-
dent transition between two open states. As shown in
Fig. 4, once the channel reaches state O1, it will be ex-
posed to the microdomain of high [Ca²⁺] created by its
own permeating calcium. This will immediately induce
a transition to O2, precluding any opportunity for inac-
tivation or adaptation. The rate and extent of inactiva-

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Local control will be minimal in a local control setting, preventing termination of local release, as was confirmed by carrying out the simulation with this gating scheme (not shown).

The remaining three schemes of Fig. 4 share certain features. All have a slow inactivation process, which is calcium dependent only in Scheme 4. Each also has the possibility of opening the channel after binding only a single calcium ion, although only Scheme 2 of Zahradnikova and Zahradnik (1996) has only one calcium binding site. Each of these schemes was put forward with a set of rate constants determined by measurements in lipid bilayers, in the absence of Mg$^{2+}$ and ATP. To incorporate these schemes into the local control simulation, it is necessary to decide how (if) these constants should be modified to describe the channel in the intracellular milieu. Fig. 4, Scheme 1, for example, has an open state reachable after binding two Ca$^{2+}$, as well as one reachable with only a single calcium. Clearly, by completely rearranging the rate constants, it would be possible to make this scheme behave somewhat similarly to the phenomenological Scheme 6. But Scheme 1 was determined by very careful fitting to a series of data on transient activation after flash photolysis of caged calcium, taking into account the transient overshoot of [Ca$^{2+}$] after the flash. It makes little sense to discard this information. We therefore explored parameter values according to the following strategy. Since all lipid bilayer schemes studied in the absence of Mg$^{2+}$ and ATP gave activation rates that were much too

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**Figure 5.** L-type calcium current and SR calcium release as a function of time at 0 mV (left) and peak current as a function of voltage (right), measured in a rat myocyte (above) and simulated (below). All currents were normalized to the peak $I_{CaL}$ at 0 mV. The time-dependent simulation was done using either the full dynamic-diffusion algorithm or the steady state approximation with $k_o$ and $k_i$ (see also Table III) adjusted to match the magnitude and timing of peak release. The graded dependence of $I_{SR}$ on $I_{CaL}$ across the range of voltages is successfully reproduced by the local control model. The dotted line at bottom left shows the markedly different result of the simulation if the assumed height of the diadic cleft is doubled. The strong dependence on geometry is the hallmark of local control.
high for local control, we reduced the rates of calcium binding by about two orders of magnitude to account for the effect of Mg$^{2+}$ competition; the exact factor used was determined by requiring the gain (ratio of peak release current to peak $I_{\text{CaL}}$) to be 10 for a depolarization to 0 mV. This criterion was relatively independent of the values of rate constants related to adaptation/inactivation. When the resulting models failed to show adequate stability, we examined the effect of increasing the kinetic rates of adaptation/inactivation steps by a factor of $\sim$20, consistent with the observations of Valdivia et al. (1995). This also failed, whereupon we examined the effect of increasing only the forward rates of inactivating steps, thereby stabilizing the inactivated/adapted states. This strategy was limited by the onset of an unacceptable degree of inactivation in the resting ([Ca$^{2+}$] = 100 nM) condition, and moved the steady state activation curve rightward so that the steady state $P_o$ approached unity only at [Ca$^{2+}$] values over 100 mM. Even with these adjustments, all three schemes gave models that manifested local instability (failure of release termination after activation) or global instability (spontaneous activation by background [Ca$^{2+}$]). Because of these instabilities, [Ca$^{2+}$]$_{\text{cyto}}$ fails to relax to the resting value following repolarization, as shown in Fig. 9 A (Schemes 1 and 2). The process of parameter exploration is illustrated for Scheme 2 in B–D. Initially, the bilayer-derived rate constants were kept, except that the rate of calcium binding to the activating site ($K_{\text{RC1}}$) was decreased from 1,000 to 16 mM$^{-1}$ ms$^{-1}$, consistent with a competitive effect of Mg$^{2+}$, giving an apparent gain of 10 for a depolarization to 0 mV, starting with all channels in the resting state (Fig. 9 B). The apparent success of this parameter set is illusory, as shown in Fig. 9 C, where the depolarization is preceded by a 1-s conditioning run-in at the holding potential.

Table III

| Variable | Description/units | Value |
|----------|-------------------|-------|
| Kl       | RyR Ca-dependent inactivation rate, mM$^{-1}$ ms$^{-1}$ | 0.5* |
| KI       | RyR repriming rate, ms$^{-1}$ | 0.005* |
| KO       | RyR Ca-activation rate, mM$^{-2}$ ms$^{-1}$ | 35.0* |
| KOM      | RyR deactivation rate, ms$^{-1}$ | 0.06* |
| UIC      | RyR unitary current at CASR = CASRR, pA | 0.41 |
| DCA      | Effective Ca$^{2+}$ diffusion coefficient in cleft, m$^{-2}$ ms$^{-1}$ | 0.15* |
| DB       | Dye diffusion coefficient, m$^{-2}$ ms$^{-1}$ | 0.021 |
| Hj       | Effective cleft height corrected for surface charge, m | 0.0297925 |

*Default values of the four parameters of the phenomenological RyR gating scheme were chosen “by hand” to give the observed gain of 10 at 0 mV, the observed time to peak of SR release (15 ms at 0 mV), and plausible macroscopic inactivation and repriming behavior. ‡Mejia-Alvarez et al., 1998. §Soeller and Cannel, 1997. ¶Harkins et al., 1993. ¶The full three-dimensional electrodiffusion problem was solved for the case of steady state diffusion from a point source at the center of a cylindrical diad, using the Galerkin finite-element method as implemented in the program PDEase (Macsyma Corp.) with parameters from Soeller and Cannel (1997). An effective value of cleft height and a two-dimensional (2D) resistive boundary condition at the edge of the cleft were then determined so as to give the best simultaneous least-squares fit by the analytical solutions of the 2D cylindrical free-calcium diffusion problem for diad radii of 50, 100, 200, and 250 nm. These values were then used in the numerical 2D diffusion computation in the Monte Carlo simulations.

Exposure to background calcium triggers a spontaneous SR release, which continues indefinitely, partially inactivating the channels so that, in the true resting state, the gain at 0 mV is reduced to only $\sim$1. As shown in Fig. 9 D, variation of the activating rate constant ($K_{\text{RC1}}$) over a fourfold range and of the inactivating rate constants (for $K_{\text{O1C2}}$, $K_{\text{O2C2}}$, and $K_{\text{C2I}}$; refer to Fig. 4, Scheme 2) over a 100-fold range was powerless to recover the normal gain in the true resting state of the model. Extensive parameter variations of this kind failed to identify a satisfactory parameter set for Schemes 1 or 2. The authors of Fig. 4, Scheme 2, did not specify a complete set of kinetic constants for the scheme, but it suffers from the same generic problems as the first two: slow and incomplete inactivation (73% at pCa 3) and activation by only a single Ca$^{2+}$ ion.

The global stability problem that arises if the RyR can be activated by a single calcium ion can be demonstrated in a relatively model-independent way. The time-averaged local [Ca$^{2+}$] at a distance $r$ from an L-type channel with an open probability $P_o$ passing a unitary calcium current $i_{\text{CaL}}$ considered as a point source, is given by

$$[\text{Ca}^{2+}]_{av} = \frac{i_{\text{CaL}} P_o}{8 \pi D_{\text{Ca}} r^4}.$$  \hspace{1cm} (2)

The openings of the L-type channel are brief and sparse, with $P_o$ probably not in excess of 5% (Rose et al., 1992). Using this value, and assuming a unitary current of 0.1 pA and an (effective) calcium diffusion coefficient of $0.15 \times 10^{-5}$ cm$^2$ s$^{-1}$, the time-average local [Ca$^{2+}$] at a distance of 10 nm from the L-type channel is only 1.4 $\mu$M, which is comparable to the global [Ca$^{2+}$]$_{\text{cyto}}$ reached during a forceful beat, and only 14× resting [Ca$^{2+}$]$_{\text{cyto}}$. A RyR that opens as the first power of local [Ca$^{2+}$] cannot, therefore, discriminate against ac-
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Portional to the local \([\text{Ca}^{2+}]\), but the mean RyR \(P_o\) during repeated exposure to brief pulses of high \([\text{Ca}^{2+}]\) is larger, as a result of nonlinear steps downstream from the calcium binding reaction. The difficulty with this argument becomes apparent if one tries to construct such a model. For a single RyR in isolation, the binding site occupancy translates into the probability that the Markov channel is found in certain states. But how is one to make the open probability depend nonlinearly on this probability when the master equations governing Markov transitions are linear? For the case of a channel with a microreversible (i.e., thermodynamically passive) gating scheme exposed to a steady calcium level, we can be quite rigorous about this. The equilibrium law of mass action requires that the \(P_o\) be a rational function of degree 1; i.e., a Michaelis-Menten function. The question, then, is whether there could be a gating scheme in which the non–steady state effects of exposure to a train of brief, very high pulses of \([\text{Ca}^{2+}]\) could “pump” the channel into a higher \(P_o\) than predicted by linear extrapolation from the resting state. The experimental constraints on this problem are actually very strong. In a resting cardiac myocyte, calcium sparks occur at a rate of 100 s\(^{-1}\) (Cheng et al., 1993). If the spark corresponds to a release of 4 pA for 10 ms, and if the RyR unitary current is 0.4 pA, then the trigger and release currents from the simulation are normalized to their respective maximum values, it is seen that the current–voltage relations are centered at somewhat different voltages. (B) When release is then divided by trigger to get the gain function of EC coupling, the resulting gain decreases with voltage, because the efficacy of the L-type current to trigger release from the RyR depends on the unitary current of the L-type channel, which decreases as the calcium reversal potential is approached. This effect, one of the basic predictions of local control theory, has been confirmed experimentally by several laboratories. Although activation of the RyR requires only two \([\text{Ca}^{2+}]\) ions in this model, the gain declines more rapidly than the L-type unitary current (scaled to match the gain curve at \(-35\) mV) in the midrange of voltages, as a consequence of partial depletion of SR calcium during the depolarization. This “elbow” effect has been observed in whole-cell calcium recordings, but not when the majority of the L-type current (and therefore SR release) has been blocked with nifedipine in order to count calcium sparks (see text).

The previous argument tacitly assumed that an RyR with only a single calcium binding site would have a time-averaged open probability roughly proportional to the time-averaged local \([\text{Ca}^{2+}]\). It might be argued that this is not necessarily so: there could be schemes in which the occupancy of the single binding site is proportion to the local \([\text{Ca}^{2+}]\), but the mean RyR \(P_o\) during repeated exposure to brief pulses of high \([\text{Ca}^{2+}]\) is larger, as a result of nonlinear steps downstream from the calcium binding reaction. The difficulty with this argument becomes apparent if one tries to construct such a model. For a single RyR in isolation, the binding site occupancy translates into the probability that the Markov channel is found in certain states. But how is one to make the open probability depend nonlinearly on this probability when the master equations governing Markov transitions are linear? For the case of a channel with a microreversible (i.e., thermodynamically passive) gating scheme exposed to a steady calcium level, we can be quite rigorous about this. The equilibrium law of mass action requires that the \(P_o\) be a rational function of degree 1; i.e., a Michaelis-Menten function. The question, then, is whether there could be a gating scheme in which the non–steady state effects of exposure to a train of brief, very high pulses of \([\text{Ca}^{2+}]\) could “pump” the channel into a higher \(P_o\) than predicted by linear extrapolation from the resting state. The experimental constraints on this problem are actually very strong. In a resting cardiac myocyte, calcium sparks occur at a rate of 100 s\(^{-1}\) (Cheng et al., 1993). If the spark corresponds to a release of 4 pA for 10 ms, and if the RyR unitary current is 0.4 pA, then

Figure 6. (A) If the trigger and release currents from the simulation are normalized to their respective maximum values, it is seen that the current–voltage relations are centered at somewhat different voltages. (B) When release is then divided by trigger to get the gain function of EC coupling, the resulting gain decreases with voltage, because the efficacy of the L-type current to trigger release from the RyR depends on the unitary current of the L-type channel, which decreases as the calcium reversal potential is approached. This effect, one of the basic predictions of local control theory, has been confirmed experimentally by several laboratories. Although activation of the RyR requires only two \([\text{Ca}^{2+}]\) ions in this model, the gain declines more rapidly than the L-type unitary current (scaled to match the gain curve at \(-35\) mV) in the midrange of voltages, as a consequence of partial depletion of SR calcium during the depolarization. This “elbow” effect has been observed in whole-cell calcium recordings, but not when the majority of the L-type current (and therefore SR release) has been blocked with nifedipine in order to count calcium sparks (see text).

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Figure 7. The unitary current of the RyR, and its calcium sensitivity \(k_o\), neither of which is known precisely in vivo, were varied over two orders of magnitude, keeping the gain of EC coupling at 0 mV equal to the experimental value of 10. The mean number of RyR openings in those diads in which at least one opening occurred is plotted on the abscissa; it is a measure of the degree of local regenerative recruitment of RyRs by CICR. This number was never <11, indicating that, for this model, there is no regime in which EC coupling operates with the observed gain without local regeneration.
resting spark release amounts to an average of only \( \sim 10 \) open RyRs per cell. If there are, conservatively, \( 10^5 \) RyRs per myocyte, the resting \( P_0 \) at \([Ca^{2+}] = 100 \text{ nM}\) is \( 10^{-4} \). Is it possible that a one-calcium gating scheme with this resting \( P_0 \) could, in response to a 200-\( \mu \text{s} \) pulse of \([Ca^{2+}] = 100 \text{ \mu M}\), open with a probability of, say, 0.1? This is a special case of a problem that has been solved previously in connection with thermodynamic constraints on RyR adaptation (Stern, 1996). Using the methods in that paper, we found that it is impossible, assuming that the Markov scheme follows the kinetic law of mass action. We can therefore be fairly confident that the difficulty in obtaining global stability in simulations with single-calcium RyR gating schemes is generic.

Local Control Can Be Stabilized by RyR–RyR Allosteric Interactions

The fact that many RyR gating schemes derived from lipid bilayer data fail to support stable EC coupling in simulations suggests that RyR gating in situ may differ from that in bilayers. Two features are required to achieve local and global stability: “strong” inactivation and cooperative activation by more than one \( \text{Ca}^{2+} \) ion. A possible clue as to how these features might arise comes from the ultrastructure and molecular biology of the RyR. In all striated muscles from crustaceans to man, including those that are activated purely by CICR (e.g., crayfish skeletal, mammalian cardiac) RyRs are found in dense two-dimensional crystalline arrays, although the details of the crystal lattice have varied somewhat (Loesser et al., 1992). Moreover, the amino acid sequence of the foot process has been strongly conserved. Tunwell et al. (1996) found 98.6% amino acid identity between rabbit and human RyR2. We have identified a sequence of 200 amino acids located in the foot region that is 100% identical between mouse and human RyR2, despite the presence of 82 synonymous mutations at the DNA level (data not shown). Since only \( \sim 29\% \) of random mutations are synonymous (Nei,
In the absence of selection is only $0.29^{82}$ ($8.2 \times 10^{-45}$). This implies that the foot has an important (though unknown) function even though the COOH-terminal region of the molecule suffices to form a calcium-sensitive channel (Bhat et al., 1997a). Could the difference between in situ and in vitro gating of the RyR be due to allosteric interactions between the foot processes of nearest-neighbor RyRs, which appear ultrastructurally to be in contact? To test this possibility, we modified the simulation algorithm to make the transition rates of each RyR dependent on the states of its (up to) four nearest neighbors. To satisfy the constraint of microscopic reversibility, the interactions were specified in the form of free energies of interaction between neighboring RyRs, depending (symmetrically) on the states of each. To determine the transition rate $K_{ij}$ of a given RyR from state $i$ to state $j$, the sum of the allosteric contact energies with its neighbors (in their present states) were added up in the initial and final states of the transition, and the exponential of the difference, divided by $kT$ was multiplied into the affinity $k_{ji}/k_{ij}$ of the transition. This still leaves one kinetic degree of freedom to determine how the effects of the free energy change are to be partitioned between the forward and backward rate constants. The transition rate therefore has the form

$$K_{ij} = k_{ij} \exp[\eta_{ij} \sum_{m=1}^{4} (E_{sm} - E_{cm})/(kT)],$$

(3)
was assumed, except for the case of a diffusion-limited 
that adding an allosteric energy 
requires stabilizing the inactivated state(s) relative to 
of inactivating transitions need to be increased, which 
two parts. To produce local stability, the forward rates 
and would decrease it only if it engendered a 
sufficient conformational change in the molecule to re-
strict access to the binding site, or a sufficient reduc-
tion in the magnitude of the binding energy to prevent 
capture of the ion.

For the allosteric model, we used gating Scheme 2 of 
Zahradnikova and Zahradnikov (1996; see Fig. 4), since 
it can be considered a “worst case,” having slow inacti-
vation and only a single calcium binding site. In this 
model, RyRs interact both allosterically and by CICR—
a process difficult to analyze intuitively. The strategy we 
used to choose the allosteric interaction energies had 
two parts. To produce local stability, the forward rates 
of inactivating transitions need to be increased, which 
requires stabilizing the inactivated state(s) relative to 
the noninactivated. A simple counting argument shows 
that adding an allosteric energy $E_{ij}$ that is equal to $-E/4$ 
if one of the states $i$ and $j$ is an inactivated state and 
$-E/2$ if both are, will contribute $-E$ to the free energy 
change associated with any inactivating transition of a 
channel that has four nearest neighbors. To produce 
global stability is somewhat more subtle, because it re-
quires creating, de novo, cooperativity of RyR activa-
tion by calcium. Since, in Scheme 2, each RyR has only 
a single Ca$^{2+}$ binding site, it is necessary to create posi-
tive cooperativity between binding sites on different 
RyRs. The strategy here is to add a large, positive allosteric 
contact energy between a RyR in an open state and one in 
the resting state. Initially, all RyRs in the diad are in 
the resting state $R$. Binding of calcium to one of these 
RyRs will move it to state $C_1$, from which it could open 
if the channel were in isolation. But, to move this RyR 
to states $O_1$ or $O_2$ when it is surrounded by resting 
channels would cause a large increase in allosteric en-
ergy, so this transition is effectively prevented by an en-
ergy barrier. If more of its neighbors acquire bound 
calcium, moving them out of state $R$, the energy barrier 
is decreased so that the RyR can open, effectively re-
sponding to the binding of several Ca$^{2+}$ ions by the ar-
ray as a whole. The important insight here is that, for 
this strategy to work, the underlying single-RyR gating 
scheme must have a closed state with bound calcium, so 
that binding of multiple Ca$^{2+}$ ions by the array can oc-
cur before any channel opens. If binding of the first 
Ca$^{2+}$ is tantamount to opening of the channel, then 
this event, however rare, will supply the local calcium to 
permit opening of nearby channels so that the whole 
array is effectively triggered by a single Ca$^{2+}$.

Once one RyR opens, a complicated interplay of al-
losteric and CICR interactions ensues, including some 
unexpected effects. For example, the energy barrier 
that prevented opening of a RyR surrounded by resting 
neighbors will, during the relaxation phase, prevent 
calcium from dissociating from RyRs that are in contact 
with one that is still open. We started with allosteric en-
ergies chosen by the two strategies described in the pre-
ceding paragraph, which succeeded in creating cooper-
tivity of activation and strong inactivation. We then 
made adjustments, guided by admittedly incomplete in-
tuition, until the performance of the model was satis-
factory. The final matrix of allosteric interaction en-
ergies is shown in Fig. 10 A; they are not necessarily ei-
ther unique or optimal. It should be noted that in this 
model we made no changes in the bilayer-derived pa-
rameters other than the introduction of the allosteric 
interactions. In Fig. 10 B, we show typical time courses 
of SR release in response to 50-ms depolarizations to the 
voltages shown. While the responses are not exactly 
the same as those of the phenomenological model, 
they serve to demonstrate that Fig. 4, Scheme 2, when 
augmented by the allosteric couplings, produces locally 
stable release. Fig. 10 C shows the rate of release stimu-
lated by global cytosolic Ca$^{2+}$, for Scheme 2 with and 
without allosteric couplings. The presence of the allo-
steric couplings converts the first order dependence of 
release flux on global [Ca$^{2+}$]$_{cys}$ into a cooperative re-
response. This markedly reduces the rate of release trig-
gered by global [Ca$^{2+}$]$_{cys}$ in the physiologic range, as 
required for global stability. Fig. 11, A and B, shows ex-
plicitly that the dynamics of [Ca$^{2+}$]$_{cys}$ after repolariza-
tion, and the spontaneous release rate when exposed to 
resting [Ca$^{2+}$]$_{cys}$, have been stabilized by the allosteric 
interactions. These allosteric interactions are basically 
inhibitory in character, as demonstrated in Fig. 11, C 
and D, which shows the effect of reducing the number of 
RyRs by leaving random vacancies in the RyR lattice. 
When the lattice is more than $\sim$60% filled, increasing 
the number of RyRs actually decreases the absolute re-
lease rate, because the effect of allosteric inhibitions 
more than compensates for the increased number of 
RyRs carrying the release flux. It is striking that even a

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few vacancies per diad markedly increase background calcium release, impairing global stability (Fig. 11 D).

**DISCUSSION**

In contrast to the situation in skeletal muscle, it is well established that mammalian cardiac excitation–contraction coupling is mediated by calcium-induced calcium release. This creates the paradox of control: why does the calcium released from the SR not feed back to trigger more release, causing an explosive chain reaction that would release all of the calcium in the SR? Previously (Stern, 1992), we analyzed some highly simplified and idealized models of stochastic CICR, with the conclusion that local control might be able to explain graded coupling. Since that time, a considerable amount of evidence has arisen in support of local control (see Ríos and Stern, 1997, for a review), but we have yet to see a single local calcium release event triggered by an identified L-type channel opening. However, considerably more information has become available about the ultrastructure of the diad junction and the gating of L-type channels and RyRs, and computer power has increased to the point where it is now practical to make use of this information in simulations.

Ideally, such simulations should be used to rule out erroneous mechanisms, prove the uniqueness of correct models, and identify the numerical values of their parameters. In practice, it is impossible to explore completely the multidimensional parameter space of these models, so the goal must be approached in stages. First, we ask whether local control, in a realistic setting of diad geometry, number, and location of channels, could explain the paradoxical gradedness of cardiac

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**Figure 10.** (A) Nearest neighbor allosteric interaction energies, as a function of the states of the interacting channels, for the model using Scheme 2 (from Fig. 4, shown at top left) augmented by allosteric interactions. (B) SR release currents produced by depolarizations to different voltages, showing a pattern similar to those produced by the phenomenological Schemes 5 and 6 (see Fig. 4), with termination of release upon repolarization. (C) “Spontaneous” release triggered by exposure to a constant global calcium level for Scheme 2 with and without the allosteric interactions. The interactions cause the activation of the diad to depend cooperatively on calcium, even though each RyR has only a single Ca$^{2+}$ binding site. This suppresses release at physiologic levels of global [Ca$^{2+}$], conferring global stability on the model.
EC coupling. This question is answered in the affirmative by the demonstration that a phenomenological local control model with numerous, regularly spaced RyRs and multiple randomly located DHPRs at each diad junction can display graded control of SR calcium release flux by a sarcolemmal calcium current only one tenth as large. In the second stage, we ask whether one or more of the proposed RyR gating schemes based on lipid bilayer data are compatible with local control. It is difficult to answer this question with complete rigor, because of the large parameter space of these schemes. The approach that we have taken is to begin with the parameters obtained in lipid bilayers and make a trial series of parameter modifications consistent with known effects of the intracellular milieu; e.g., competition of Mg$^{2+}$ at Ca$^{2+}$ binding sites and acceleration of adaptation/inactivation kinetics. Based on these explorations, we have drawn inferences about how and why RyR gating schemes fail in the local control model, and what properties a successful RyR must possess. We have tried to back these inferences with relatively model-independent arguments. The third stage would involve detailed fitting of models to large amounts of whole-cell EC coupling data, identifying parameter values and establishing

![Graph A](image1.png)

**Figure 11.** (A) Global cytosolic [Ca$^{2+}$] for the model using Scheme 2 (see Fig. 4) with and without allosteric interactions is shown. With Na/Ca release disabled, there is no relaxation of global [Ca$^{2+}$] after a stimulus; normal relaxation is produced when allosteric interactions between RyRs are included. (B) Global instability of the model using Scheme 2 is revealed by major calcium release when the diads are exposed to resting global background [Ca$^{2+}$] (100 nM) in the absence of depolarization. Inclusion of allosteric interactions eliminates this background sensitivity by requiring cooperative activation by Ca$^{2+}$ bound to different RyRs. (C) The effect of the allosteric interactions is basically inhibitory. As a result, if the number of RyRs per diad is reduced by leaving vacancies in the RyR lattice, there is a paradoxical increase in the rate of SR calcium release, until the number of RyR remaining is no longer sufficient to support the release. (D) The allosteric suppression of spontaneous release triggered by background calcium (C) is very sensitive to completeness of the RyR lattice. Introduction of small numbers of vacancies per diad unmasks the unstable release characteristic of Scheme 2 RyR activation by a single bound Ca$^{2+}$. The effect of reducing the number of RyRs by introducing vacancies in the lattice is much larger than the effect of simply reducing the size of a complete array, which has only a slight effect on global stability.
what experiments would discriminate between competing models. This stage goes beyond the scope of this paper, although the results obtained so far offer some tantalizing hints.

The somewhat surprising result of our stage-2 explorations is that none of the published bilayer-derived gating schemes gives a successful local control model, even when plausible modifications are made to adjust for the effects of the intracellular milieu. While it is difficult to exclude rigorously the possibility that some alternative set of parameters might solve the problem, it appears from our explorations that this could not be done without sacrificing crucial features of the gating behavior seen in bilayers; e.g., turning an adaptive inactivation into an irreversible one. All the schemes failed by lack of sufficient stability. Based on our explorations, this failure appears to result from the absence of two critical features: cooperative activation by more than one calcium ion and a “sufficiently” rapid and complete inactivation mechanism. The isolated RyR in bilayer is a very artificial system, and there are many reasons why it might not demonstrate physiological gating behavior, including the absence of other proteins associated with the RyR at the junction (e.g., triadin, junctin, calsequestrin). However, the most intriguing possibility is that the missing factor is the densely packed regular array of other RyRs that has been so well conserved through hundreds of millions of years of evolution. Recently, it has been reported that pairs (and, on occasion, groups of three, four, or even five) of skeletal muscle RyRs in lipid bilayers can gate synchronously when the associated protein FKBP-12 is present (Marx et al., 1998). Similar behavior is sometimes observed with RyR2 (A.R. Marks, private communication). This positive cooperativity was seen even when barium, rather than calcium, was the permeant ion, indicating that it is probably not mediated by the CICR mechanism. Our simulations show that cooperative allosteric interactions between nearest neighbor RyRs, interacting with CICR, can lead to stable local control of SR calcium release by a RyR whose gating scheme in isolation lacks the properties required for stability.

Limitations of the Allosteric Gating Model

Our construction of a stable gating scheme using allosteric interaction energies should be considered only as a proof of principle. In particular, our use of gating Scheme 2 from Fig. 4 as a starting point should not be construed as an endorsement of that gating scheme—we chose it only because it displays the pathologies of both inactivation and cooperativity in a simple form, and because it has a calcium-bound closed state, needed for our construction of RyR–RyR cooperativity. Our particular choice of allosteric energies should also not be considered either unique or optimal. They were obtained by starting with a certain strategy and then “tinkering” to get improved activation and inactivation behavior. The resulting model may deviate significantly from observations in many areas that we did not examine carefully, such as repriming dynamics. Our choice of allosteric interactions, while it produces positive cooperativity between RyRs, does not give rise to the perfect synchronization of opening and closing of coupled RyRs observed by Marx et al. (1998). Bers and Fill (1998) have suggested that such synchronization among all the RyRs in a cardiac diad might be a mechanism to generate calcium sparks, as an alternative to stochastic CICR, which also produces spontaneous multi-channel release events consistent with observed sparks (results not shown). It is not likely that allosteric interactions are required for the production of sparks since removal of FKBP12, the immunophyllin that is associated with the native RyR and is required for synchronized gating (Marx et al., 1998), does not abolish sparks (McCall et al., 1996), but may actually increase their frequency and duration and induce global calcium oscillations (Xiao et al., 1997). This observation finds a natural place in our allosteric coupling scheme, which is fundamentally inhibitory in character, as shown in Fig. 11. In fact, for the model based on RyR gating Scheme 2, even partial relief of the allosteric inhibitions (Fig. 11 D) unmasks a much greater degree of instability than is observed in intact cardiac myocytes deprived of FKBP12 by either knockout or the drug FK506. In interpreting such findings, it is important to bear in mind that FKBP has major effects on the gating properties of RyRs even in isolation, perhaps by mediating coordination between monomers in the tetramer (Kaftan et al., 1996).

There are, moreover, fundamental reasons to doubt that allosteric coupling alone, unaided by CICR, could produce acceptable synchronous gating of all the RyRs at a diad. Marx et al. (1998) reported that all the components of the dwell time distribution of synchronously gating pairs of RyR1 (two open times and two closed times) were unchanged from those of single channels. As pointed out by Bers and Fill (1998), this is quite surprising on energetic grounds. One way to model such an array is to assume that the coupling gangs together the molecular degrees of freedom involved in gating so that the only available states of the array are those in which all channels are in the same state. If that were the case, then an array of $n$ channels would have all the state and, presumably, transition-state, energies $n$-fold larger than a single channel. For the large $n$ found in the cardiac diad, these increased energies, which appear as negative exponentials, would produce an overwhelming slowing of the gating process, as well as reduce the occupancy of the less probable states in the
gating scheme to negligible values. Another way to model a synchronized array, more in keeping with our approach in the simulations, is to assume that the gating of individual channels in the array remains well defined, but is modulated by allosteric interaction energies that are additive over all the points of RyR–RyR contact in the array. The fact that Marx et al. (1998) did not observe intermediate states (one channel open, one closed) in coupled channel pairs implies that occupancy of such states is suppressed by a positive allosteric energy, large compared with $kT$, at a contact between open and closed channels. In order for all the RyRs at a diad to go from closed to open, it would be necessary to pass, however briefly, through such intermediate states, which would have an allosteric energy proportional to the perimeter of contact between open and closed channels. This perimeter must, at some point in the process, be as large as the minimum diameter of the diad, 10 or more RyRs for large diads (Franzini-Armstrong, C., F. Protasi, and V. Ramesh, manuscript in preparation). This would represent an enormous energy barrier, which would, again, greatly slow the synchronous gating.

We are led to the conclusion that coupled gating, unaided by CICR or other exogenous sources of energy, should be greatly slowed compared with single channel gating. These arguments fall short of rigorous proof, because we have assumed that the states and transitions states of individual channels remain well defined in the array, and that allosteric interactions can be treated as an additive perturbation generated at the interface between nearest-neighbor RyRs. One could, instead, achieve coupled gating without change in the dwell times by assuming that ganging together $n$ channels has a catalytic effect that reduces the transition state energies of individual channels by a factor of $1/n$. Such a long range mechanical interaction in a lattice of huge macromolecules would be, at the least, ad hoc, although it might not be physically impossible. This argument shows that synchronous gating of many channels, with unchanged dwell times, is not a natural consequence of the interaction of multiple RyRs. It is therefore likely, as indicated by Bers and Fill (1998), that allosteric interactions collaborate with local CICR to produce the observed macroscopic and microscopic phenomena of cardiac EC coupling. Whether this collaboration is of the form modeled here is a matter presently beyond the resolution of direct measurement. In light of the observations by Marx et al. (1998), it is also likely that allosteric interactions would contribute substantially to the quantitative behavior and robustness of gating schemes (e.g., Fig. 4, Scheme 6) that are not intrinsically unstable, but require “tuning” to give acceptable behavior.

Although the possible role of allosteric interactions in cardiac EC coupling is the most novel phenomenon to emerge from the modeling, it should be put in perspective. To achieve stable local control, two characteristics are required: “strong” RyR inactivation and cooperative RyR activation by more than one Ca$^{2+}$ ion. Inactivation could be produced by any interaction of the RyR with its native environment that stabilizes the inactivated state—it is not necessary to invoke state-dependent allosteric interactions with neighboring RyRs for this purpose. In contrast, the creation of cooperative activation, starting from a gating scheme that permits RyR opening in response to a single Ca$^{2+}$, does require the exchange of information between RyRs. However, it is not entirely certain that the cooperativity problem is a real one. While published gating schemes based on bilayer data all permit single-Ca$^{2+}$ activation, some studies have shown the steady state open probability of a single RyR2 depending on [Ca$^{2+}$] with a Hill coefficient of $>2$ (Laver et al., 1995; Sitsapesan and Williams, 1994; Copello et al., 1997), suggesting that intramolecular cooperativity might be adequate to achieve global stability.

In any case, it is unlikely that any of the currently proposed RyR gating schemes captures all of the features of importance for physiologic EC coupling. One complicating factor that we have not considered here is the possibility that SR luminal calcium may directly influence gating kinetics. Another (possibly related to the first) is that increasing the SR calcium load of a rat myocyte modestly above normal provokes periodic calcium oscillations, whereas “latch up” of the cell in a stable state of high [Ca$^{2+}$]$_{cyt}$ and high RyR open probability is never observed, even when egress of calcium from the cell is prevented. None of the models described above reproduces this global behavior accurately. It is possible that “adaptation” or other modulation of RyR gating by sustained elevation of [Ca$^{2+}$] in the submicromolar range plays an important role in determining global stability. Further simulation and experimentation will be required to determine how this and other physiologic constraints translate into required properties of the RyR at the molecular level.
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