Interplay of Ryanodine Receptor Distribution and Calcium Dynamics

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ABSTRACT Spontaneously generated calcium (Ca^{2+}) waves can trigger arrhythmias in ventricular and atrial myocytes. Yet, Ca^{2+} waves also serve the physiological function of mediating global Ca^{2+} increase and muscle contraction in atrial myocytes. We examine the factors that influence Ca^{2+} wave initiation by mathematical modeling and large-scale computational (supercomputer) simulations. An important finding is the existence of a strong coupling between the ryanodine receptor distribution and Ca^{2+} dynamics. Even modest changes in the ryanodine receptor spacing profoundly affect the probability of Ca^{2+} wave initiation. As a consequence of this finding, we suggest that there is information flow from the contractile system *to* the Ca^{2+} control system and this dynamical interplay could contribute to the increased incidence of arrhythmias during heart failure.

INTRODUCTION

The control and loss of control of Ca²⁺ wave initiation (regenerative propagation of intracellular Ca²⁺ release) are important for both normal and pathological excitationcontraction coupling (E-C coupling) in cardiac muscle. Mammalian atrial cells lack an extensive transverse tubule system (t-tubules), so an action potential triggers Ca²⁺ release only from the ryanodine receptor clusters (or Ca^{2+} release units, CRUs) directly coupled to the L-type Ca²⁺ channels on the surface sarcolemma (1,2) and at interior sites associated with the less developed transverse axial tubular system (2). Nonpropagating Ca²⁺ release limited to the sarcolemma surface would activate only a fraction of the myofibrils near the surface. However, an action potential can also initiate a centripetally propagating Ca^{2+} wave (1–4) that allows rapid activation of the myofilaments in the central core of the cell (2). The mode of Ca^{2+} release (propagating or nonpropagating) determines the rate and magnitude of contraction in atrial myocytes.

Mammalian ventricular cells, due to their extensive and well-organized t-tubules, do not use Ca^{2+} waves for normal E-C coupling, because an action potential triggers synchronous Ca^{2+} release throughout the cell. Nevertheless, spontaneously generated Ca^{2+} waves do occur in ventricular cells as in atrial cells (4–6). Spontaneous Ca^{2+} release or Ca^{2+} waves can elicit delayed afterdepolarization as the excess Ca^{2+} is removed from the cell via the Na⁺-Ca²⁺ exchanger (7,8). A growing body of evidence also suggests that spontaneous waves may trigger cardiac arrhythmias (9–12). Hence, it is important to understand the factors that control the pathological generation of Ca^{2+} waves.

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This article examines the factors that influence wave initiation using mathematical modeling and large-scale computational (supercomputer) simulations. Of particular interest is the relationship between the three-dimensional (3-D) CRU distribution and Ca^{2+} wave initiation. We used antibody labeling and confocal imaging experiments to measure the spacing of CRUs along the longitudinal axis of cardiac myocytes (longitudinal spacing, equivalent to the Z-disk spacing), within the Z-disk (transverse spacing), and described a newly found intercalated CRU on the cell periphery (13). In this article, we describe and explain how even subtle changes in the CRU spacing along any of these dimensions can profoundly influence the Ca^{2+} wave initiation.

An important finding of these studies on CRU spacing and Ca^{2+} wave initiation is that the state of the contractile system can profoundly affect the dynamics of the Ca^{2+} control system. This flow of information from the contractile to the Ca^{2+} control system is complementary to the information flow from the Ca^{2+} to the contractile system in the canonical E-C coupling paradigm. Hence, this newly found feedback loop suggests a continuous and circular flow of information between the electrical, Ca^{2+} , and contractile systems, given the existence of feedback from the Ca^{2+} to the electrical excitation (see Bers (14) for review). Based on our results, we propose that the feedback from the contractile system to the Ca^{2+} control system may contribute, in part, to the increased incidence of arrhythmias in certain forms of heart failure and familial hypertrophic cardiomyopathy.

METHODS

In this section, a representation of the governing system of diffusion/reaction partial differential equations (PDEs) is presented along with a brief discussion of the numerical solution methods and the mathematical models that are used to implement the cardiac cell model. The PDEs that describe the diffusion and reaction of Ca^{2+} (*C*), the endogenous buffers (*B*), and fluorescent indicator (*F*) are essentially the same as described in our previous publications (15,16) except that now the number of spatial dimensions is three instead of two. We note that full 3-D simulations are required in this study to model the effects of

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the complex 3-D CRU distribution in cardiac myocytes measured by antibody labeling experiments (13). In addition, locally refined, highly resolved, unstructured meshes and temporal discretization are necessary for representation of the complex CRU distribution in space and time.

Briefly we present a generalized notation for the governing diffusion/ reaction systems of equations for the cardiac cell model. This general notation allows us to present the essence of the finite element approximation that follows very compactly. The governing PDEs for multi-component diffusion mass transfer and nonequilibrium chemical reactions are given by

$$\frac{\partial C_{i}}{\partial t} = \nabla \cdot \mathbf{D}_{i} \nabla C_{i} + S_{i} \quad \text{for} \quad i = 1, \dots N_{s}, \qquad (1)$$

where C_i is the concentration of species *i*, \mathbf{D}_i is a 3 × 3 diffusion mass transfer tensor that allows for anisotropic effects (in our model, this is a diagonal tensor), S_i is the volumetric source term for species *i*, and N_S is the total number of unknown concentrations in the simulation. A detailed description of the complete set of PDEs is presented in Appendix A. We present the equation for the free Ca²⁺ concentration here because it provides a useful reference for explaining the model and geometry.

$$\frac{\partial C(x, y, z, t)}{\partial t} = D_{Cx} \frac{\partial^2 C}{\partial x^2} + D_{Cy} \frac{\partial^2 C}{\partial y^2} + D_{Cz} \frac{\partial^2 C}{\partial z^2} + R_B(C, F_B) + R_D(C, F_D) + J_{pump} + J_{leak} + \sum_{i,j,k} \sigma S(x_i, y_j, z_k, t; T_{open}) \times \delta(x - x_i) \delta(y - y_j) \delta(z - z_k).$$
(2)

The Z-disks are in the *xy* plane and the longitudinal axis of the cardiac cell lies along the *z* axis. Diffusion (for both Ca²⁺ and fluorescent indicator) in the *xy* plane is assumed to be isotropic $(D_x = D_y)$ and the diffusivity is assumed to be half of that along $z (D_x = D_z/2 (17))$. $\delta(\cdot)$ is a Dirac delta-like function in the sense that the integral of the product of these functions and σ is the molar flux of Ca²⁺ (moles/s). $\delta(\cdot)$ is nonzero at the CRU and zero elsewhere. The exact specification of $\delta(\cdot)$ is given below in Numerical methods. *S* is a stochastic function that takes on the value of 1 (causing the CRU to "fire") or 0 (turning the CRU off). The probability that a CRU at position (x,y,z) will fire in time $t - \Delta t/2 < t < t + \Delta t/2$ is $P(C(x,y,z,t))\Delta t$, where *P* is the probability of firing per unit time. *P* is given by

$$P(C(x, y, z, t), K, n) = \frac{P_{\max}C^{n}}{K^{n} + C^{n}},$$
(3)

as measured by Lukyanenko and Györke (18).

The source strength σ is the Ca²⁺ molar flux density and is given by $\sigma = I_{SR}/2F$, where I_{SR} is the Ca²⁺ current (in pA) through the CRUs and *F* is the Faraday constant. In 3-D, σ has a very simple interpretation: it is the flux density of Ca²⁺. In 2-D, however, σ is a line source and has units of mol/s/ μ m and a reasoned "fudge" factor had to be introduced to relate σ to the current I_{SR} (16). No such factor is needed in 3-D.

The CRU currents, I_{SR}, used in our simulations range from 10 to 25 pA. This range is based on several studies using various methods to estimate I_{SR} . ISR, back-calculated from the size of sparks, ranges from 10 to 20 pA (15,19). Estimates based on ruthenium red inhibition of RyR2 (20) put the number of RyR2s underlying a spark at k > 10 and, therefore, $I_{SR} > 5$ pA, assuming single-channel current of 0.5 pA (21). Estimates based on spark amplitude variability give k > 18 and $I_{SR} \sim 10$ pA. (22). It is noted that the current magnitude underlying cardiac Ca²⁺ sparks remains a controversial issue to date. Based on the rate of rise of sparks, Cheng and co-workers estimated k between 1 and 3 (23,24). However, we have argued (13) that if k were small, then failure to trigger a spark would be high, which is at odds with the near certainty of eliciting sparks by depolarization (25). Moreover, sparks occurring under conditions that engender Ca²⁺ waves have larger amplitude and spatial size than those occurring under conditions where probability of Ca^{2+} waves is low (6). Since I_{SR} increases rapidly with spark size (15), I_{SR} values used for studying Ca²⁺ waves should be considerably larger than those for modeling sparks under quiescent conditions.

The model geometry of a single sarcomere, based on the experimentally measured RyR2 cluster distribution (13), is shown in Fig. 1 A. The dark blue circles mark the position of the Z-disks; the blue "bricks" are the CRUs on the Z-disk; and the cyan-colored bricks that lie between the Z-disks are the "intercalated" CRUs. Fig. 1 B shows the Z-disk in greater detail. The CRUs on the Z-disks are uniformly spaced $\Lambda \ \mu m$ apart (0.8 or 1 μm) and are radially arranged in layers. CRUs on the periphery (white arrows) lie in the circle of radius $r \equiv r_0 = 4.5 \ \mu m$. We refer to this layer of CRUs as the peripheral layer, or layer 0. The CRUs denoted by the yellow arrows lie on the circle whose radius is $r = r_1 = 4.5 \ \mu m - \Lambda$. We shall refer to this layer of CRUs $(r = r_1)$ as the first inner layer or layer 1. In many simulations, n contiguous CRUs on layer 0 will be triggered to fire. The central CRU on layer 0 is simply the one that is in the middle of the group. The central CRU on layer 1 is the CRU that is closest to the central CRU on layer 0. Note that there might be two CRUs on layer 1 that are equidistant to the central CRU on layer 0. The distance between Z-disks is L_z (2 μ m under control condition). For clarity, the plots along the z axis have been stretched out. The intercalated CRUs (cyan-colored bricks) are placed midway between the Z-disks and are present only on the periphery, at the same radial distance as the CRUs on layer 0 of the Z-disk. Their spatial separation is Λ , the same as the CRUs on the Z-disks. The central CRU on this ring of intercalated CRUs is the one that is closest to the central CRU on layer 0.

The simulation volume includes the two Z-disks and the region between them, plus flanking regions that extend an additional 1 μ m. Zero-flux boundary conditions are imposed on the cylindrical boundary and the flat end caps. The zero-flux boundary condition is a natural choice for the cylindrical boundary since the cell membrane is a barrier for free Ca²⁺ diffusion. We used this boundary condition for the end caps for computational efficiency; an absorbing boundary would be more natural. However, test simulations using a larger flanking domain showed that the conclusions are not materially affected by the edge effects arising from this boundary condition.

Numerical methods

2-D simulations (finite-difference methods)

Simulations of the 2-D model equations were carried out using finite differences to approximate the derivatives, and solved using the method of lines, with the model as described previously (16).

3-D simulations (unstructured mesh finite-element methods)

Simulations of the full 3-D model with the complex distribution of CRUs was carried out using a Galerkin finite-element (FE) method as implemented in the MPSalsa transport/reaction simulation code (26). The system of $N_{\rm S}$ reaction/diffusion equation is solved in bounded open region Ω in \mathbb{R}^3 , with a sufficiently smooth boundary $\partial \Omega = \Gamma_{\rm m} U \Gamma_{\rm d}$ over the time interval (0,Y). The formal Galerkin weak form of the governing reaction/diffusion system is derived as follows. First, we multiply Eq. 1 by a test function ψ from the appropriate space $H_0^1(\Omega)$ (see Thomee (27)), which gives

$$\int_{\Omega} \psi \frac{\partial C_{i}}{\partial t} d\Omega = \int_{\Omega} \psi \nabla \cdot \mathbf{D}_{i} \nabla C_{i} d\Omega + \int_{\Omega} \psi s_{i} d\Omega$$

for $i = 1, \dots N_{s} \quad \forall \psi_{0} \in H_{0}^{1}.$ (4)

Using the divergence theorem we obtain the resulting weak form

$$\int_{\Omega} \psi \frac{\partial C_{i}}{\partial t} d\Omega = -\int_{\Omega} \psi \nabla \cdot \mathbf{D}_{i} \nabla C_{i} d\Omega + \int_{\Omega} \psi S_{i} d\Omega + \int_{\Gamma_{m}} \psi \, \hat{\mathbf{n}} \cdot \mathbf{D}_{i} \nabla C$$
for $i = 1, \dots, N_{s} \quad \forall \psi_{0} \in \boldsymbol{H}_{0}^{1}$ (5)

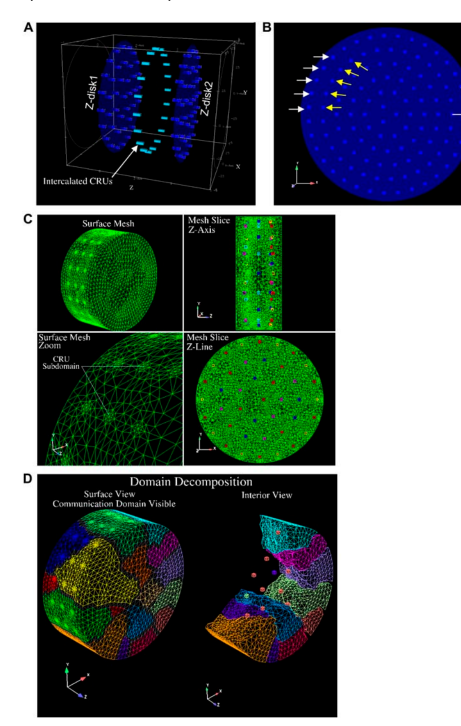


FIGURE 1 Three-dimensional model geometry of the sarcomere and finite element mesh geometry and domain decomposition. (A) Dark blue disks are the Z-disks into which the CRUs (blue bricks) are embedded. The Z-disks lie in the xy plane and the longitudinal axis of the myocyte is along the z axis. The distance between the Z-disks is L_z . The intercalated CRUs (cyan bricks) are only on the cell periphery, forming a ring that is located midway between two Z-disks. (B) Detail of a single Z-disk. CRUs are circularly arranged with spacing Λ indicated by the white lines. See text for meaning of arrows. (C) A composite view of the unstructured tetrahedral FE mesh of the cardiac cell with two z-planes and a set of intercalated CRUs is shown. The upper left image presents the surface mesh of the cell section and the lower left image shows a close-up view of the unstructured mesh used to resolve the individual CRUs. The upper right image shows a projection of the mesh on an xz-slice plane through the domain. The lower right image shows the projection of the mesh on an xy-slice plane. The individual CRUs in these two images have been colored to make them more visible. (D) Composite view of the subdomain-to-processor assignment used to partition the mesh on 16 processors is shown. The left image shows subdomain-to-processor assignments by color. The partitioning assigns FE nodes to subdomains and then maps these subdomains to processors. The gray intersubdomain region indicates boundaries between processors for which communication of information takes place in the parallel FE code. The image on the right is a partial exploded view with the intersubdomains removed.

of the governing equations. A Galerkin finite-element formulation for these equations restricts this system to a finite dimensional subspace $\psi^h \in H^1(\Omega)$ and $\psi^h_0 \in H^1_0(\Omega)$. In our implementation, we use linear hexahedral and tetrahedral elements that have a formal order of accuracy of $O(h^2)$. To complete the description of the method, we define the approximation to the time derivative by a first- or second-order approximation based on a backward Euler or trapezoidal rule method, respectively. This discrete approximation forms a large sparse system of nonlinear algebraic equations. In the paragraphs that follow, we give a brief overview of the critical issues of spatial and temporal resolution of the diffusion/reaction system and the parallel nonlinear and linear solution methods that are employed to solve the FE system of equations. A more detailed discussion of the spatial and temporal accuracy of these methods applied to diffusion/ reaction problems can be found in Ropp et al. (28).

Λ

Our implementation of the continuous Dirac delta function in space from Eq. 2 in our discrete model is a spatially localized source term applied in a CRU subdomain. This geometry requires resolution of multiple length scales for the representation of the small length scale distributed CRUs and the overall cardiac cell domain, which we accomplish via unstructured finiteelement tetrahedral meshes. An example of this type of mesh is shown in Fig. 1 *C* for a single saromere. Clearly, to adequately represent the smallest length scales, i.e., CRUs, local refinement about the CRUs is required.

In general, the governing diffusion/reaction equations and the discrete approximations to these equations produce a stiff system with multiple timescales. The stiffness of these equations is produced by the discrete approximation to the diffusion operator and by the source term operator. To integrate this stiff system in a stable and efficient way, we employ fully implicit solution methods. These techniques allow the simulation to resolve the dynamical timescale of interest (in this case the firing of the CRUs and the subsequent diffusion wave front) while not requiring, for stability, the simulation to run at the fastest stiff timescales of the system. It has been recently shown that fully implicit methods can have significant advantages in stability and accuracy when compared to some common operator split or semi-implicit methods for a subset of diffusion/reaction systems (28,29). In the time integration of the cardiac cell system, a second-order (trapezoidal rule) method is used and the time-step size is held constant to resolve the firing of the CRUs and the subsequent diffusion wave front. In the CRU model described below, the probability of firing was set at 30%/ms. To adequately resolve this release we selected a time-step size of 0.01 ms or a probability of firing per time step of 0.003. Numerical experiments were used to determine that this integration procedure was accurate (relative error <1%, utilizing equilibrium concentrations measuring total production of CRU source terms as a norm) for the simulations that are presented in this article.

As presented above, the fully implicit FE approximation of the diffusion/ reaction system produces a very large system of nonlinear algebraic equations. To solve these coupled systems we use Newton's method as a nonlinear solver and employ Krylov iterative solver techniques (conjugate gradient or generalized minimum residual (GMRES) method) with domain decomposition preconditioners to solve the linear systems at each substep. The decomposition and mapping of the FE domain (the cardiac cell) onto the parallel machine is accomplished by a graph partitioning method. Chaco (30), a general graph partitioning tool, is used to partition the FE mesh into subdomains and make subdomain-to-processor assignments. Chaco constructs partitions and subdomain mappings that have low interprocessor communication volume, good load balance, few message start-ups and only small amounts of network congestion. In Fig. 1 D, we present a sample partition of a cardiac cell mesh on 16 processors. In this figure, the color assignment of FE nodes is used to visualize the assignment to processors. In general, the partitioner assigns contiguous sets of elements to processors (see exploded view in B). Communication occurs across interprocessor boundaries to update information that is computed on neighboring processors. For a detailed description of parallel FE data structures and a discussion of the strong link between partitioning quality and parallel efficiency, see Hendrickson and Leland (30) and Shadid et al. (31). Finally, the domain decomposition preconditioners are based on parallel additive Schwarz domain decomposition methods (see, e.g., Smith et al. (32)) that use approximate incomplete LU solves on each subdomain. These preconditioners are used to efficiently solve the linear systems with a high degree of parallel scalability to thousands of processors. For a typical time step, this solution procedure requires one or two Newton steps and <10 linear iterations per Newton step.

The computations were performed on clusters at Sandia National Laboratories. A typical problem requires $\sim 200,000$ unknowns per time step (number of nodes in geometry \times number of species (calcium, fluorescent indicator, and protein buffers)). A 100-ms simulation requires 10,000 time steps. Depending on availability of compute nodes, where each node possesses two 1- to 3-GHz Pentium III or Pentium IV processors, we generally performed solutions on 4–16 nodes. Time to complete simulations ranged from 12 to 36 h, depending on the individual cluster's processor speed, excluding waiting time in the job scheduling queue.

Mathematical tools

The firing of a CRU is a stochastic process, so the outcome of a single simulation should not be viewed as representative of the archetypal behavior for a given set of parameters. An ensemble of repeated simulations is needed to obtain a more complete and accurate picture of the system's typical behavior. However, in some cases we can use the results of a single simulation to compute the probabilistic behavior of the system. This is possible in our model because the probability of CRU firing per unit time depends on the instantaneous cytosolic Ca²⁺ concentration. The probabilistic functions we will use are the waiting-time distribution $w(x,y,z,\tau,T)$, which is the probability of having to wait at least *T* ms starting from time τ before a CRU at (x,y,z) fires, and the probability density function $\phi(x,y,z,t,\tau)$. For brevity, we henceforth drop the (x,y,z) dependency. $w(T;\tau)$ is given by

$$w(T;\tau) = \exp\left(-\int_{\tau}^{\tau+\mathrm{T}} P(s)\mathrm{d}s\right),\tag{6}$$

where P(s) is the probability of spark firing per unit time at time *s* given in Eq. 3. This fundamental equation was derived in our earlier work (16), but here an explicit τ -dependence is included for studying synchronization. The probability density function (pdf) $\phi(T)$ is defined by

$$w(T;\tau) = \int_{\tau+T}^{\infty} \phi(s) ds, \quad \phi(T;\tau) = P(T+\tau)$$
$$\times \exp\left(-\int_{\tau}^{\tau+T} P(s) ds\right), \tag{7}$$

where the explicit expression for $\phi(T;\tau)$ comes from differentiation with respect to *T*.

Except when studying synchronization of Ca^{2+} release (see below), we will take τ to be 0 and we will drop the explicit reference to τ . The meaning of $\phi(T)$ can be clarified by noting that $1 - w(T) \equiv q(T)$ is the probability that an event will occur in time 0 < t < T or

$$q(T) \equiv 1 - w(T) = 1 - \int_{T}^{\infty} \phi(s) \mathrm{d}s = \int_{0}^{T} \phi(s) \mathrm{d}s. \quad (8)$$

In other words, $\phi(T)$ is the pdf for q(T). We note that $\phi(T)$ is normalized; its integral from 0 to ∞ equals 1 follows from Eqs. 6 and 7. That w(0) = 1follows from Eq. 6 and, intuitively, from the certainty that one needs to wait more than zero ms before an event (a CRU firing) to occur.

The waiting-time distribution (WTD) is defined for a single CRU. The WTD for *N* CRUs, w(T,N), is the product of the individual WTDs because the CRUs are assumed to act independently. Thus, w(T,N) and its pdf $\phi(T,N)$ are

$$w(T,N) = \prod_{i=1}^{N} w_i(T), \quad \phi(T,N) = \left(\sum_{i=1}^{N} P_i(T)\right) \times \prod_{i=1}^{N} w_i(T).$$
(9)

Note that the assumption of independence does not imply that the firing probability of a CRU is unaffected by the history of firings of other CRUs. The assumption of independence only means that there is no direct effect of one CRU upon another except through the agency of changes in Ca^{2+} concentration.

The *N* CRUs in the product are those that are closest to the ones that have fired on the adjacent ring of CRUs. For example, in Fig. 1 *B*, the five CRUs that have fired are indicated by the white arrows. The N = 5 CRUs on the adjacent ring that go into the calculation of Eq. 9 are those pointed to by the yellow arrows. Note that "closest" is defined in terms of the Euclidean distance; when a CRU on the inner layer is equally close to two or more CRUs on the outer layer, other CRUs are chosen so that the number of CRUs in the calculation always equals the number that have fired. Although this might seem arbitrary, this choice does not materially change w(T,N). To see this, consider a CRU that does not see a substantial change in the Ca²⁺ concentration but is included in the product. Because its WTD is essentially unity, it makes little contribution to the product.

The mean waiting time before at least 1 CRU fires is MWT.

$$MWT = \int_{\tau}^{\infty} (s - \tau)\phi(s, N)ds$$
 (10)

Note that *MWT* can also be interpreted as the mean time of the first CRU to fire. From this viewpoint we can then ask what is the mean waiting time for the second, third, *N*th CRU to fire? These times characterize the synchronization of Ca²⁺ release. To calculate these times, we first note that, in this model, the probability that a CRU will fire depends only on its instantaneous ambient Ca²⁺ concentration and not on its history: CRUs have no memory of their previous states. Given *N* CRUs, the MWT for the first CRU to fire is given by Eq. 10, with $\tau = 0$. If the first CRU fires at $t = t_1$, then to calculate the MWT before the next CRU fires, we use Eq. 10, but now with $\tau = t_1$ and ϕ in Eq. 9 replaced by $\phi(s, N - 1)$, because the first CRU is no longer available to fire. We can apply this procedure iteratively to find the MWT between CRU firings for all *N* CRUs. An example of synchronization is given in Fig. 4 *B* (see Fig. 4).

Studying Ca²⁺ wave initiation using "canonical pulses"

Computing the WTD requires an explicit expression of *P* (see Eq. 6) or, equivalently, the spatial and temporal distribution of Ca^{2+} . We used the Ca^{2+} distribution generated by a canonical pulse (16). The canonical pulse is the simplest idealization of a distribution of Ca^{2+} . The leftmost panels in Fig. 2 show the canonical pulse of Ca^{2+} generated by triggering eight contiguous CRUs on the periphery. Using this canonical Ca^{2+} distribution, we can calculate the average behavior of the system, in effect getting the equivalent of an infinite number of simulations from one. By using this canonical pulse, we can compute w(T,N) and $\phi(T,N)$, which we can readily compare for different parameters (such as I_{SR} and Λ). These functions provide important insights into the mechanisms of Ca^{2+} wave initiation. The canonical pulse shown in Fig. 2 is meant to mimic the triggering of peripheral CRUs by an action potential. Later, we will use a canonical distribution to mimic a well-formed Ca^{2+} wave. To calculate w(T,N) and $\phi(T,N)$, we prevent other CRUs from firing by setting their P_{max} to zero (see Eq. 3)

RESULTS

In this section, we will study the important factors that influence the Ca^{2+} wave generation and examine the relationship between Ca^{2+} waves and the CRU distribution in 3-D. First, we will develop the important concepts and tools along with analyzing the effect of transverse spacing of CRUs on Ca^{2+} waves. Next, we will use these concepts and tools to analyze the effect of intercalated CRUs on Ca^{2+} waves. Finally, we will apply the analysis method to examine the influence of CRU longitudinal spacing on Ca^{2+} waves and study the effect of shortened sarcomere length on altering the Ca^{2+} dynamics, possibly linking these to increased propensity for spontaneous Ca^{2+} waves and arrhythmias in certain forms of heart failure and familial hypertrophic cardiomyopathy. Please see Figs. 2, 6, 7, 9, and 10 for snapshots of evolving Ca^{2+} dynamics. Movies of these simulations are available in Supplemental Materials.

Effect of transverse spacing of CRUs on Ca^{2+} waves: a surprise

Our previous modeling study (15) has shown that the initiation and propagation of Ca waves depends critically on the spacing of the CRUs. Based on the work of Parker et al. (17), our earlier 2-D modeling work on ventricular cells (16) assumed a CRU transverse spacing (Λ) of 0.8 μ m. Our initial 3-D simulations also used $\Lambda = 0.8 \ \mu m$. Using our novel method for getting improved resolution of CRU spacing in the transverse plane, we found that the transverse spacing of CRUs is 1.05 μ m in the ventricular and 0.97 μ m in the atrial myocytes (13). We did not expect to see qualitatively different dynamics in making the seemingly small change from $\Lambda = 0.8 \ \mu m$ to $\Lambda = 1.0 \ \mu m$. We were, however, very surprised. Fig. 2 shows two simulations that were identical except for the spacing of the CRUs. The leftmost panels show nine peripheral CRUs on the left-hand Z-disk firing at t = 0 to simulate triggering by an action potential; this is the canonical pulse. For $\Lambda = 0.8 \ \mu m$ (Fig. 2 A), the activation spreads to the next Z-disk and a Ca²⁺ wave is initiated and

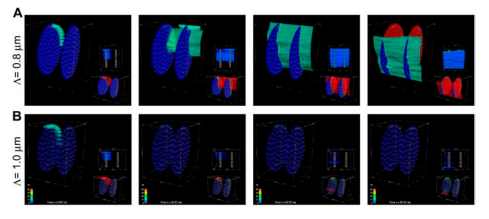


FIGURE 2 Qualitative difference in Ca²⁺ dynamics resulting from a small change in CRU spacing. These snapshots are of two simulations that used identical parameters and initial conditions, except that the CRU spacing is $\Lambda = 0.8 \ \mu m$ in A and 1.0 μm in B. A Ca^{2+} wave is initiated when $\Lambda = 0.8 \ \mu m$ but not when $\Lambda = 1.0 \ \mu m$. In this and subsequent figures, each image is divided as follows. The large image on the upper left is an oblique view of the sarcomere, which includes a $15-\mu M$ Ca2+ isoconcentration surface. The smaller image near the bottom right shows the Ca²⁺ concentration over the

smaller range of $0.1-2 \mu$ M, and shows the isoconcentration surface at 2μ M. The small image near the middle of the right-hand side is a side view of the sarcomere. A suite of isoconcentration surfaces at 1, 2, 5, 10, and 15 μ M are shown. The side view is especially helpful for understanding how CRUs on different Z-disks interact. The color bar ranges from 0.1 to 50 μ M for the large image and from 0.1 to 2 μ M for the lower right image. Simulation times are 0.92, 45.92, 65.92, and 85.92 ms (*left to right*). Simulation parameters: $I_{SR} = 13$ pA, $L_z = 1.6 \mu$ m; other parameters are given in Appendix A.

propagates across the two Z-disks. However, when $\Lambda = 1.0 \ \mu m$ (Fig. 2 *B*), the canonical pulse fails to initiate a Ca²⁺ wave. (Ca²⁺ waves are defined operationally based on the shape of the time-to-target plots described in Kirk et al. (2).) To understand how a seemingly small change in CRU spacing has such a large effect on Ca²⁺ wave initiation, we studied how waves are initiated and propagated in a simpler 2-D system.

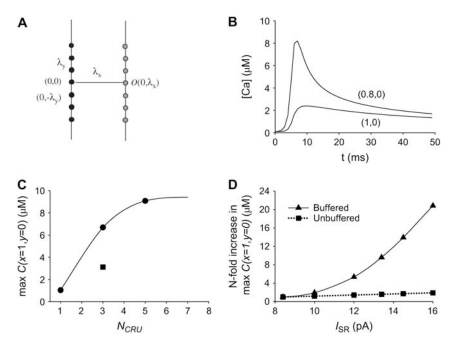
Important factors that influence Ca²⁺ wave initiation

Our efforts to understand the causes for this great sensitivity to Λ have given us new insights into the important factors that influence Ca²⁺ wave initiation and propagation. The basic insights gained here will reappear later when we examine the effects of CRU distributions in other dimensions, the intercalated CRUs, for example. Therefore, the next few sections will be devoted to developing the general strategy that we used for examining how Λ influences Ca²⁺ wave initiation.

Separation distance

To understand the basis for this great sensitivity of wave initiation on Λ , we needed to be able to examine the Ca²⁺ concentration at any point in space. We could do this much more easily in 2-D than in 3-D because of the much simpler 2-D computational mesh (uniform square cells versus tetrahedral or hexahedral elements). As will be shown below, the 2-D and 3-D systems share the same qualitative behavior.

A schematic of the 2-D simulation geometry is shown in Fig. 3 *A*. The CRUs on the boundary (x = 0, *black circles*) are spaced ℓ_v apart and the Ca²⁺ concentration is measured at



observation points (*gray circles*) at $x = \ell_x$, y = 0, a horizontal distance of ℓ_x from the boundary. The simulations start off with N = 7 CRUs on the boundary firing at t = 0. Fig. 3 *B* shows the Ca²⁺ concentrations at observation points (x = 0.8, y = 0) and (x = 1, y = 0). Note the fourfold decrease in peak Ca²⁺ when the observation point increases by just 0.2 μ m.

These plots illustrate the important point that the free Ca²⁺ concentration is extremely sensitive to changes in separation distance between the source(s) and observation point. This extremely steep sensitivity cannot be understood in terms of linear diffusion. Making the approximation that M mol of Ca²⁺ are instantaneously deposited at the source point, then the Ca²⁺ concentration at a distance r (see Appendix A of Izu et al. (16) for the general case, where Ca²⁺ is delivered at a finite rate) is $C(r,t) = (M/(4\pi Dt^{1/2}))\exp(-r^2/(4Dt))$, which achieves its maximum value at $t = r^2/(2D)$. This $1/r^2$ dependency is not strong enough to account for the sensitivity we observe. For example, in a linear system the ratio of peak Ca²⁺ concentrations at r = 0.8 and 1 μ m would only be $(1/0.8)^2/(1/1)^2 = 1.56$, not the observed factor of 4 in the buffered system.

Superadditivity

The steep dependence of the free Ca^{2+} concentration on separation distance arises from the presence of large quantities of Ca^{2+} buffers, both endogenous and exogenous (including the fluorescent Ca^{2+} indicator). Ca^{2+} buffers give rise to the phenomenon that we call "superadditivity" (see Izu et al. (16) and Appendix B). In the absence of buffers or when the buffers are completely saturated, the change in the free Ca^{2+} concentration is linearly related to the number of moles of Ca^{2+} added. In an unsaturated buffered system,

> FIGURE 3 Effect of CRU spacing, number of firing CRUs, and current on Ca²⁺ concentration in 2-D. (A) Geometry of the 2-D lattice. The firing CRUs (*black dots*) are on the line x = 0; they are spaced λ_{v} apart. The observation point *O* (gray dot), where the Ca²⁺ concentration is measured, is at $x = \lambda_x$, y = 0. The Ca²⁺ concentrations at the other observation points are used in a later figure. (B) Ca^{2+} concentration when the observation point is $\lambda_x = 0.8$ or 1 μ m from the line of seven firing CRUs, which are spaced $\lambda_v = 0.8 \ \mu m$ apart. The modest 20% reduction in λ_x results in a fourfold increase in peak Ca^{2+} concentration at O. (C) Maximum Ca^{2+} concentration at O as a function of the number of firing CRUs. The square gives the Ca2+ concentration that would be seen if no \mbox{Ca}^{2+} buffers were present and a single CRU produced the concentration shown. (D) *n*-fold change in peak Ca^{2+} concentrations at O as a function of Ca^{2+} current. The squares show the *n*-fold change in Ca^{2+} concentration that would be seen if Ca²⁺ buffers were absent.

with each addition of δn moles of Ca²⁺, there is an ever greater increase in the change of the free Ca²⁺ concentration because of the ever decreasing number of free-buffer binding sites available to bind Ca²⁺. A formal definition of superadditivity is given in Appendix B and we show how this nonlinear behavior can be understood from an equilibrium analysis. The superadditive behavior of buffered systems means that the free Ca^{2+} concentration increase due to two Ca^{2+} sources is larger than a simple addition from two sources. To illustrate this phenomenon, Fig. 3 C shows the maximum concentration at $(x = 1 \ \mu m, y = 0)$ for different numbers of CRUs. Note that for $N_{CRU} = 3$, the maximum concentration is more than six times the maximum concentration obtained for a single CRU. In a linear system, the corresponding maximum concentration is <3 times that for a single CRU (*square*). (The concentration is less than threefold greater because addition is weighted by the distance between the source and the observation point; see Eq. 3 of Izu et al. (16).)

Superadditive effects apply not only to the number of CRUs but also to the current amplitude (I_{SR}) of the CRU. Fig. 3 *D* shows that the ratio of the maximum value of *C* at ($x = 1 \mu m, y = 0$) relative to its value for $I_{SR} = 8.4$ pA increases nonlinearly as I_{SR} varies in a buffered system (*triangles*). In a linear system (*squares*), the ratio is directly proportional to the magnitude of the current (see Eq. 3 of Izu et al. (16)). When I_{SR} changes from 8.4 to 20 pA, the approximately twofold change in maximum Ca²⁺ concentration in the linear unbuffered system is almost imperceptible on the scale needed to accommodate the 20-fold change in the nonlinear buffered system. The important point illustrated by this graph is that even small changes in source current can effect large changes in the free Ca²⁺ wave initiation and propagation.

Synchronization

A general definition of a Ca²⁺ wave front is an approximately synchronous firing of CRUs that recapitulates the firing pattern at an earlier time occurring on a well defined locus of points. Therefore, if the firing of *N* CRUs on the periphery of the Z-disk is to initiate a wave, then $\sim N$ CRUs on the adjacent layer should fire at about the same time for a wave to progress. This notion of synchronicity is captured by the iterative computation of the WTD, w(T,n), as explained in Methods (note that *n* is not necessarily *N*).

In our previous work (16), we used $\phi(t,N)$ as a predictor of wave initiation. The idea behind its use is that wave propagation implies a modicum of predictability of Ca²⁺ spark firing, and the breadth of $\phi(t,N)$ is inversely related to the uncertainty of when a Ca²⁺ spark will occur. When $\phi(t,N)$ is sharply peaked (Fig. 4 *A*, solid line) there is little uncertainty about when a Ca²⁺ spark will occur (in this case, ~6 ms after the canonical pulse), suggesting that a Ca²⁺ wave will be initiated. However, sometimes $\phi(t,N)$ is not sensitive enough to accurately predict wave initiation. For example, the two

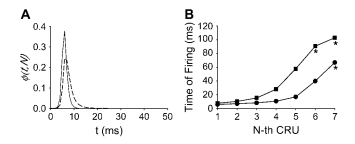


FIGURE 4 Understanding why a wave is initiated when $I_{SR} = 16$ pA but not when $I_{SR} = 13.4$ pA. (A) Probability density function $\phi(t,N = 7)$ for $I_{SR} = 13.4$ pA (*dashed line*) and 16 pA (*solid line*). $\phi(t,N)$ is not sufficiently sensitive to explain why a wave is initiated with $I_{SR} = 16$ pA but not when $I_{SR} = 13.4$ pA. (B) Synchronization curves show that five CRUs will fire within 11 ms of each other when $I_{SR} = 16$ pA (*circles*) but it takes ~50 ms for five CRUs to fire when $I_{SR} = 13.4$ pA (*squares*). This long delay reduces the likelihood of wave initiation. The accuracy of the firing times in the synchronization curve depends on the accuracy of estimating the integral of $\phi(t,n)$ over infinite time. Since our simulation of the canonical Ca²⁺ distribution is limited to 100 ms, $\phi(t,n)$ is known only to 100 ms. When the Ca²⁺ concentration is high, the integral converges quickly to 1 and the firing time is accurate. When the integral is <0.8 the computed firing time is only a crude lower bound. Firing times that are not known accurately (integral <0.8) are marked with asterisks.

 ϕ curves in Fig. 4 A are similar, yet a wave is initiated with $I_{SR} = 16 \text{ pA}$ (solid line) but not with $I_{SR} = 13.4 \text{ pA}$ (dashed *line*). The reason that ϕ , by itself, is not sensitive enough is that it generates the WTD for only the first Ca^{2+} spark occurrence. Since the Ca^{2+} wave front involves many Ca^{2+} sparks, a better indicator of wave initiation is the synchronization curve. The synchronization curve is the WTD of the first, second, ... *n*th Ca^{2+} spark occurrence (Fig. 4 *B*). For both currents, the first spark occurs at about the same time, \sim 6 ms after the canonical pulse, in agreement with the time the ϕ curves attain their peak. However, for $I_{SR} = 16$ pA (solid line), the synchronization curve is initially almost flat because the first five CRUs fire within 10 ms of each other; they are almost synchronous. On the other hand, when $I_{SR} =$ 13.4 pA (dashed line), CRU firing is asynchronous; the fourth and fifth CRUs fire 20 and 50 ms, respectively, after the first. It is this lack of synchronous firing that causes the failure of a wave to be initiated.

Large-scale cooperative CRU interactions

One of the most important insights derived from this study is that cooperative interactions of CRUs on adjacent Z-disks and intercalated CRUs significantly increase the probability of Ca^{2+} wave initiation. To see how important these largescale cooperative CRU interactions are, we first establish the reference state: the isolated Z-disk.

Initiation of Ca²⁺ waves on a single Z-disk: 3-D simulations

We wanted to determine whether the factors we found to be important for the initiation of waves from the 2-D simulations are also important in 3-D. The 2-D simulations are analogs for events on a single Z-disk. Therefore, in the following simulations only a single Z-disk is active (P_{max} for all of the CRUs on the second Z-disk are set to zero) and intercalated CRUs are absent.

As in the 2-D simulations, we find that there is a very steep Ca^{2+} concentration gradient as Λ changes from 0.8 to 1.0 μ m. Fig. 5 *A* shows the Ca^{2+} concentration measured at the CRU that sees the largest peak Ca^{2+} . There is an 1100% increase (0.48 vs. 5.95 μ M) in the maximum Ca^{2+} concentration as Λ decreases 20% from 1.0 to 0.8 μ m. Note that although the plots in Figs. 5 *A* and 3 *B* are qualitatively similar, they are not exactly comparable, because in Fig. 3 *B*, the CRU packing density was fixed ($\ell_x = 0.8 \,\mu$ m) and only the distance between adjacent layers ($\ell_x = 0.8 \,\sigma$ 1.0 μ m) was changed. In the 3-D calculations, Λ changes both the packing density and interlayer spacing. Nevertheless, the fact remains that the WTDs are very sensitive to changes in the spacing of CRUs.

Also similar to the 2-D case is the rapid increase in the maximum Ca^{2+} concentration seen by the central CRU with I_{SR} (Fig. 5 *B*, *solid line*). The effect of superadditivity is evident from the comparison with the expected values for an unbuffered linear system (*dotted line*).

To estimate the current needed to initiate a Ca^{2+} wave, we computed synchronization curves for $I_{SR} = 32$ pA (Fig. 5 *C*, *circles*) and $I_{SR} = 20$ pA (Fig. 5 *C*, *squares*). When $I_{SR} = 32$ pA, the MWT for the first CRU to fire (after the start of the canonical pulse) is ~6 ms and that the first five CRUs fire within 13 ms. By contrast, when $I_{SR} = 20$ pA, it takes ~22 ms before the first CRU fires and subsequent CRUs fire much later. Although the values in the figure are lower bounds for CRU number >1 (see figure caption for explanation), it is sufficient to show that for $I_{SR} = 20$ pA, the firing is asynchronous, so wave initiation is unlikely. The full simulations bear out these predictions. We find that a wave is readily initiated for $I_{SR} = 32$ pA, but not when $I_{SR} = 21$ pA (data not shown).

Because we have set up these 3-D simulations to be analogous to the 2-D simulations, it is not surprising that these results reinforce what we found in the 2-D studies, namely, that Ca^{2+} wave initiation is sensitive to CRU separation, superadditive effects, and synchronization of firing. Henceforth, we will focus on 3-D simulations that allow us to model the 3-D distribution of the CRUs in cardiac cells more realistically and reveal more complex interactions between CRUs.

Cooperative Z-disk interactions

The next level of complexity that we introduce is a second Z-disk. Fig. 6 shows two simulations that are identical ($I_{SR} = 21 \text{ pA}$) except that the CRUs on the second Z-disk (the one on the right, which we will call Z-disk 2) were either inactive (Fig. 6 *A*) or active (Fig. 6 *B*). When the CRUs are inactive, the Z-disk is functionally absent as it presents no diffusional

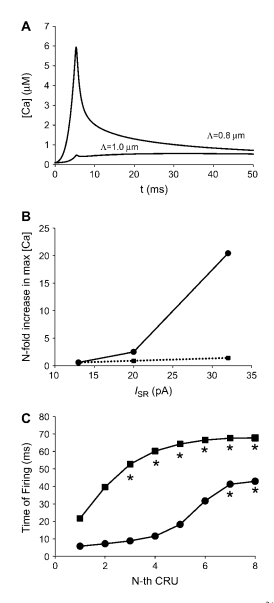


FIGURE 5 Effect of CRU spacing and current in 3-D. (A) Ca²⁺ concentration measured at the central CRU on layer 1 (see Methods for definition) when eight contiguous CRUs fire simultaneously on layer 0. The peak Ca²⁺ concentration is 12-fold higher when $\Lambda = 0.8 \ \mu\text{m}$ than when $\Lambda = 1.0 \ \mu\text{m}$. (B) *n*-fold increase in peak Ca²⁺ concentration at central CRU for different I_{SR} . The *dashed line* shows the increase that would be seen in an unbuffered system. (C) Synchronization curves for $I_{SR} = 20 \ \text{pA}$ (*squares*) and 32 \ pA (*circles*). Asterisks have the same meaning as in Fig. 4.

barrier to Ca^{2+} ; in the figure, Z-disk 2 is present just for illustration. No wave is initiated when the CRUs of Z-disk 2 are inactive (Fig. 6 *A*). When the CRUs of Z-disk 2 are present, however, a wave is initiated (Fig. 6 *B*). In this case, the firing CRUs on the left-hand Z-disk (Z-disk 1) elevate the ambient Ca^{2+} concentration of the CRUs on Z-disk 2, increasing their firing probability. The firing of CRUs on Z-disk 2, in turn, increases the ambient Ca^{2+} concentration of Z-disk 1's CRUs, thereby increasing their firing probability. This regenerative feedback between CRUs of the two neigh-

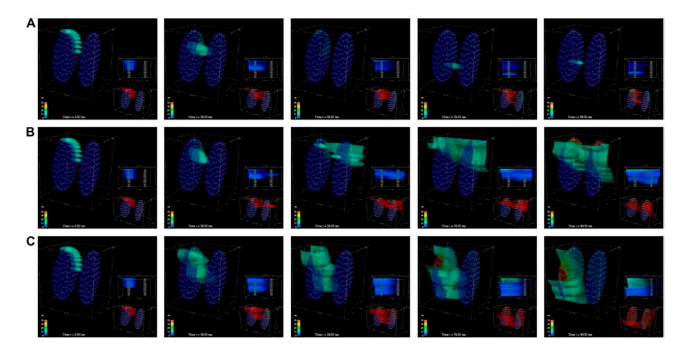


FIGURE 6 Cooperative Z-disk interactions. CRUs on Z-disk 2 (on the right) are inactive ($P_{max} = 0$) in A and active ($P_{max} = 0.3/ms$) in B. Otherwise, the simulations are identical. Because Ca²⁺ released from CRUs on one Z-disk can increase the probability of firing of a CRU on the other Z-disk, this regenerative feedback reinforces the nascent wave and a stable wave propagates across the Z-disks in B. In A, this cooperative interaction is absent and no wave forms. (*C*) The CRUs on Z-disk 2 are inactive but a wave forms because the current has been increased from $I_{SR} = 21$ pA (in A and B) to 25 pA. All simulations are started with the canonical pulse. The color bar ranges from 0.1 to 50 μ M for the large image and from 0.1 to 2 μ M for the lower right image. Snapshots are taken at 0.92, 30.92, 50.92, 70.92, and 90.92 ms.

boring Z-disks stabilizes the nascent wave and increases the probability of wave initiation.

In the absence of cooperative Z-disk interactions (when CRUs of Z-disk 2 are inactive), the CRU current needs to be increased to ~ 25 pA to initiate a wave (Fig. 6 *C*). Also, although it is not evident in these snapshots, the wave in Fig. 6 *C* propagates in a jerky fashion, whereas the wave in Fig. 6 *B* propagates smoothly (see movies in Supplemental Materials). Thus, not having CRUs on the adjacent Z-disk exacts a cost of ~ 4 pA to initiate a wave. These results demonstrate the important point that the interaction of CRUs of adjacent Z-disks reduces the current required to initiate and support Ca²⁺ waves.

Intercalated CRUs

The "intercalated" RyR2s are located approximately midway between the Z-disks on the cell periphery (mean distance to Z-disk = $0.97 \pm 0.42 \ \mu\text{m}$ in ventricular cells and $0.92 \pm 0.38 \ \mu\text{m}$ in atrial cells) but not in the interior of the cell (13). To explore the effect the intercalated CRUs have on wave initiation we carried out simulations with and without the intercalated CRUs (Fig. 7). All simulations start off with the canonical pulse (0.92 ms) and CRUs on both Z-disks are active. In the absence of intercalated CRUs, a current of $I_{\text{SR}} = 20 \text{ pA}$ is not sufficient to initiate a wave (Fig. 7 *A*), and ~21 pA is needed to initiate a wave (Fig. 6 *B*). When the intercalated CRUs are present, however, a current of just 15 pA is enough to initiate a robust wave (Fig. 7 B). Therefore, the intercalated CRUs significantly reduce the current needed to initiate a wave.

To understand how the intercalated CRUs reduce the current needed to initiate a wave, let us compare Ca²⁺ signal propagation within a Z-disk and between Z-disks. Fig. 8 A shows the Ca²⁺ distribution at t = 5.92 ms, the end of the canonical pulse (*left*). At t = 6.92 ms, two intercalated CRUs fire (*middle*), and at t = 9.92 ms, the first CRU on Z-disk 2 fires (right). Surprisingly, the first nontriggered CRU on Z-disk 1 fires much later, at 16.92 ms. These particular firing times are not statistical anomalies as they are close to the calculated firing times given in Fig. 8 B. The mean time for the first intercalated CRU firing (circles) is 4.8 ms, whereas the corresponding time for CRUs on Z-disk 1 is 21.7 ms (squares). Notice that the synchronization curve for the intercalated CRUs is initially almost flat, meaning that two or even three intercalated CRUs would fire almost simultaneously. This happened to be the case in this simulation where two intercalated CRUs fired at t = 6.92 ms. The intercalated CRUs are activated earlier because they see a larger and earlier rise of Ca²⁺ (Fig. 8 C, solid line) than the layer-1 CRUs on Z-disk 1 (Fig. 8 C, dashed line) because of the 2:1 anisotropic diffusion favoring faster flow along the z axis (longitudinally) over the transverse plane (within the Z-disk). (See Methods for definitions of layers and distances.)

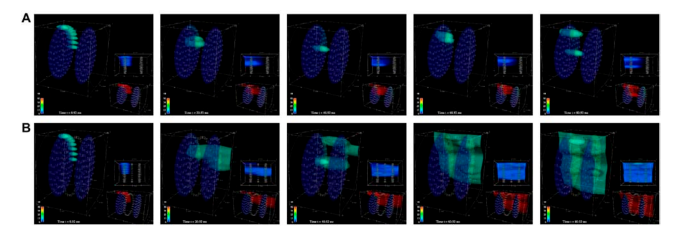


FIGURE 7 Effect of intercalated CRUs (i-CRUs). i-CRUs are absent in the simulations of A. When the current is $I_{SR} = 20 \text{ pA}$ (A), a wave is not initiated. In the absence of i-CRUs the minimum current needed to initiate a wave is ~21 pA (not shown). The current needed to initiate a wave is significantly reduced (to ~15 pA) when i-CRUs are present (B). The color bar ranges from 0.1 to 50 μ M for the large image and from 0.1 to 2 μ M for the lower right image. Snapshots are taken at 0.92, 20.92, 40.92, 60.92, and 80.92 ms. Simulation parameters: $L_z = 2.0 \ \mu$ m, $\Lambda = 1.0 \ \mu$ m, others standard.

Due to the fast diffusion in the *z*-direction, the firing of the intercalated CRUs can rapidly trigger peripheral CRUs on Z-disk 2. Indeed, the first peripheral CRU on Z-disk 2 fires at 9.92 ms, 7 ms before a CRU in layer 1 of Z-disk 1 fires, despite the fact that the CRUs on Z-disk 2 are about twice as far away. Therefore, the intercalated CRUs serve to link the dynamics of the adjacent Z-disks even more strongly than the cooperativity described in the previous section. The presence of intercalated CRUs causes Ca^{2+} waves to be initiated more rapidly and increases the velocity of the waves across the plane of the Z-disks (data not shown).

Z-disk interactions: sarcomere spacing variation

We have seen how CRUs on the different Z-disks reinforce each others' activities, making it easier for a wave to be initiated. In this section, we again consider how the activities of CRUs on one Z-disk affect those on an adjacent Z-disk but now focus on examining the strength of interaction as the sarcomere spacing varies. Of interest here is the relationship between sarcomere spacing and the probability of wave initiation. It is important to understand this relationship because it tells us how the mechanics of contraction affects Ca^{2+} dynamics.

Effect of shortening sarcomere length. To study the relation between sarcomere length and Ca²⁺ wave initiation, we forced all CRUs on one Z-disk (Z-disk 1) to fire, and, using the resultant Ca²⁺ concentration distribution, computed the probability of CRU firing on the adjacent Z-disk (Z-disk 2), that is, 2.0, 1.8, or 1.6 μ m from Z-disk 1. The synchronization curves (Fig. 9 A) show the strong effect of sarcomere spacing on CRU activity. The synchronization curve for $L_z = 1.6 \mu$ m (solid line) starts off almost flat, meaning that many CRUs fire almost simultaneously. The first 10 CRUs fire within 4.5 ms of each other. When $L_z = 1.8$,

the first 10 CRUs fire within 6.4 ms of each other (*dashed line*). However, when $L_z = 2.0 \ \mu$ m, the first 10 CRUs to fire are spread out over 23 ms (*dotted line*). In other words, the firings become asynchronous. This broad distribution in firing times is reflected in the steep slope of the $L_z = 2.0 \ \mu$ m synchronization curve.

These results show that even a modest reduction in sarcomere spacing from 2.0 to 1.8 μ m causes a tremendous increase in the ability of CRUs on adjacent Z-disks to increase each others' activities. The consequence of this reciprocating influence is that it is easier to initiate waves as illustrated in Fig. 9 B. Three simulations were carried out that are identical except for the distance between the Z-disks. Eight contiguous CRUs on the periphery of Z-disk 1 are forced to fire at t = 0 (*leftmost images*). With the Z-disks separated by $L_z = 2 \ \mu m$, the Ca²⁺ concentration increase seen by CRUs on Z-disk 2 is sufficiently low that no CRU fired and no wave was initiated (upper panel). When the Z-disk spacing is reduced to $L_z = 1.8 \ \mu m \ (middle \ panel)$, the rise in the Ca²⁺ concentration resulting from CRUs on Z-disk 1 firing is now sufficiently high at Z-disk 2 that CRUs on Z-disk 2 fire. Their firing, in turn, increases the likelihood of CRUs on Z-disk 1 to fire. This reciprocal interaction gives rise to a well formed wave front emerging at around t = 60ms. The interactions are even stronger when $L_z = 1.6 \ \mu m$ (lower panel) and the wave front begins forming even earlier, at ~ 40 ms.

Currents needed to initiate a wave as Z-disk spacing changes. Of further interest is the interplay between the Z-disk spacing and the Ca²⁺ current in initiating a wave. We determined the relationship between the minimum current needed to trigger a wave and the Z-disk spacing is roughly linear (Fig. 9 C). The current versus *z*-distance curve in Fig. 9 *C* separates this parameter space to stable and unstable zones. In the unstable zone where the current is large and *z*-distance is small, the system becomes unstable and a wave is initiated.

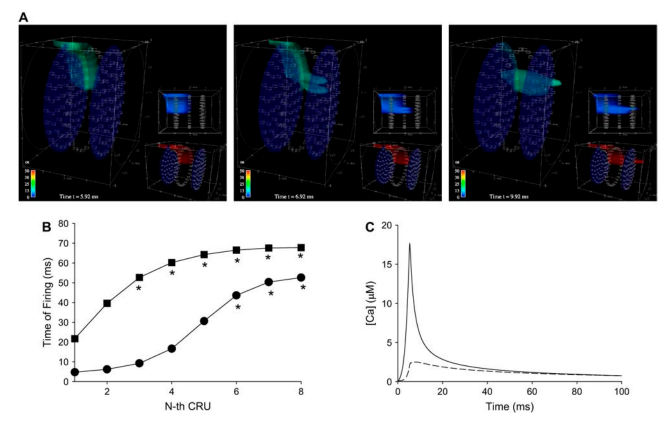


FIGURE 8 Intercalated CRUs act as relay stations. (*A*) The canonical pulse ends at 5.92 ms (*left panel*), and 1 ms later (at 6.92 ms, *center panel*), two intercalated CRUs fire. Their firing is seen more clearly in the side view of the sarcomere (*center panel*, *inset*, *right-hand side of image*). The first CRU on Z-disk 2 fires at 9.92 ms (*right panel*). (*B*) Synchronization curves for the i-CRUs (*circles*) and for the CRUs on Z-disk 1 (*squares*). Although the CRUs on layer 1 of Z-disk 1 and the i-CRUs are the same distance from the triggered CRUs (1 μ m), the expected times for the i-CRUs to fire are substantially shorter than for the layer-1 CRUs (for example, 4.8 ms vs. 21.7 ms for the first CRU to fire). The reason for this difference is shown in the curves of *C*, which show the Ca²⁺ concentration seen by the central i-CRU (*solid line*) and the central CRU on layer 1 (*dashed line*). Simulation parameters: $L_z = 2.0 \ \mu$ m, $\Lambda = 1.0 \ \mu$ m, $I_{SR} = 15 \ pA$, others standard.

Conversely, when the current is small and *z*-distance is large, the system is stable and wave initiation is unlikely.

Initiation versus support of wave propagation

It is important to distinguish between conditions that can initiate wave propagation from those that can support wave propagation. The simulations in Fig. 10 illustrate this distinction. Both simulations use identical parameters but differ in the initial conditions. In the first simulation (Fig. 10 A), the Ca^{2+} concentration is 0.1 μ M everywhere, with endogenous and exogenous buffers in equilibrium. As an initial stimulus, all CRUs on the left edge are forced to fire at t = 1 ms. We see that this stimulus fails to initiate a wave (time flows from left to right). In Fig. 10 B, we set up the initial stimulus as a well formed wave by raising the Ca^{2+} concentration between $0 \le$ $x \le 1.5 \ \mu \text{m}$ to 10 μ M, whereas "in front" of the wave (x >1.5 μ m) the Ca²⁺ concentration is 0.1 μ M, with all buffers in equilibrium with the Ca²⁺ at every point. In this case, the wave continues to propagate throughout the domain (to x = 16 μ m), showing that these conditions can support wave propagation but, by themselves, cannot initiate wave propagation. The reason wave propagation can be sustained after the wave front is formed is because there is a large reservoir of Ca^{2+} behind the wave that can keep the Ca^{2+} concentration at the front high long enough to trigger Ca^{2+} release from the next set of CRUs and thereby maintain propagation.

The explanation for propagation might seem circular positing the existence of a wave to continue a wave—but this circularity is broken if there is heterogeneity in the cell. For example, there may be a region where the sarcomeres are close together (say 1.6 μ m apart), which can be a site for wave initiation. Once the wave is initiated, it can spread to regions of the cell that can support but not initiate waves. In atrial cells, an action potential triggers the firing of surface CRUs, which may initiate Ca²⁺ waves. After its formation, the wave may propagate under less stringent conditions than that required for wave initiation.

DISCUSSION

 Ca^{2+} waves triggered by an action potential serve the normal physiological role of rapidly activating Ca^{2+} release

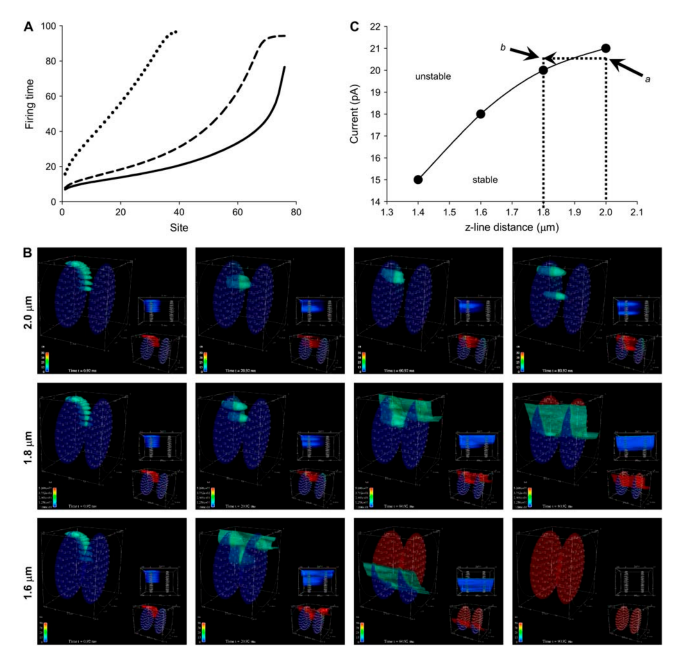


FIGURE 9 A small decrease in sarcomere length greatly increases Z-disk interaction. (A) Synchronization curves as functions of sarcomere spacing (L_z) for $L_z = 2.0 \ \mu m$ (*dotted line*), $L_z = 1.8 \ \mu m$ (*dashed line*), and $L_z = 1.6 \ \mu m$ (*solid line*). Simulation parameters: $I_{SR} = 20 \ pA$, $\Lambda = 1.0 \ \mu m$, others standard. (B) These three simulations used identical parameters and initial conditions, except that the Z-disk spacing was 2.0 μm (*upper*), 1.8 μm (*middle*), and 1.6 μm (*lower*). Closer Z-disk spacing allows greater regenerative interaction of CRUs on adjacent Z-disks, thereby increasing the likelihood of Ca²⁺ wave generation. The color bar ranges from 0.1 to 50 μ M for the large image and 0.1 to 2 μ M for the lower right image. Snapshots are taken at 0.92, 20.92, 60.92, and 80.92 ms. Simulation parameters: $I_{SR} = 20 \ pA$, $\Lambda = 1.0 \ \mu$ m, others standard. (C) Currents needed to initiate a Ca²⁺ wave as a function of Z-disk spacing. All simulations were started with the canonical pulse (eight contiguous CRUs on layer 0 firing at t = 0). Currents were varied until a Ca²⁺ wave was initiated within 100 ms of the start of the pulse. For a given Z-disk separation, fi I_{SR} is below the curve the probability of wave initiation is low (e.g., point *a*) and the system is stable. If I_{SR} remains constant but the Z-disk separation shortens to the extent that I_{SR} moves above the curve then wave initiation is likely (e.g., point *b*) and the system becomes unstable. Simulation parameters: $\Lambda = 1.0 \ \mu m$, the rest are standard.

throughout the atrial myocyte to ensure rapid and robust contraction. On the other hand, spontaneously generated Ca^{2+} waves in atrial and ventricular myocytes could contribute to generating arrhythmias. This article examines the factors that affect Ca^{2+} wave initiation in cardiomyocytes.

Our main focus is to understand the effect that the spatial distribution of CRUs has on Ca^{2+} dynamics. The most important finding is that the probability of Ca^{2+} wave initiation is exquisitely sensitive to the spatial distribution of the CRUs. This suggests that the contractile state of the myocyte

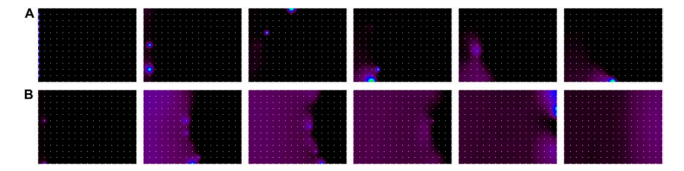


FIGURE 10 Initiation versus support of Ca^{2+} waves. In these 2-D simulations the initial conditions were chosen to mimic either an action-potential-triggered CRU firing (A) or a well-formed Ca^{2+} wave (B). White dots mark the positions of the CRUs. Simulation parameters: $\ell_x = \ell_y = 1 \mu m$ (see Fig. 6 A for definition of spacing parameters), $I_{SR} = 11 \text{ pA}$, others standard.

directly influences the dynamics of the Ca²⁺ control system. This important and novel concept is explored below.

In our effort to understand the physical basis for the sensitivity of Ca^{2+} dynamics to CRU spatial distribution, we also examined how the CRU Ca^{2+} current (I_{SR}) and the number of neighboring CRUs that fire synchronously affect the ambient Ca^{2+} concentration about a CRU. The large effect that these factors—spatial distribution of CRUs, number of CRUs firing, and I_{SR} —have on wave initiation probability all share a common physical basis: saturation of Ca^{2+} buffers. Intuitively, this is expected since the spread of the Ca^{2+} concentration at the CRU on the wave front. What the analyses and simulations reveal is the surprising magnitude of the nonlinear effect these factors have on the free Ca^{2+} concentration. We call this nonlinear dependence superadditivity.

In our simulations, the free Ca²⁺ concentration between the Z-disks during a wave typically reaches ~15 μ M. This value is ~10 times higher than some estimates of experimentally measured [Ca²⁺]_i (e.g., 4). However, amplitudes of Ca²⁺ waves vary widely. For example, Kockskämper and Blatter (33) report *F*/Fo values of 5–13 in cat atrial myocytes using fluo-4. The Ca²⁺ concentrations corresponding to the *F*/Fo range of 5–11 are 0.79–12 μ M, calculated from the selfratio formula (34) assuming $K_d = 1.1 \ \mu$ M (35) and resting [Ca²⁺]_i of 0.1 μ M. Using the data from Lipp et al. (36) and their fluo-3/fura-red calibration curve (37), we calculated the peak averaged [Ca²⁺]_i in a wave to be ~10 μ M.

A wave having a Ca²⁺ concentration of ~10 μ M is not unexpected given Lukyanenko and Györke's (18) finding that the Ca²⁺ sensitivity of the CRUs is low, ~15 μ M. If the Ca²⁺ concentration of a wave were 1 μ M, then in 20 ms (the time it takes a wave traveling 100 μ m/s to traverse a sarcomere) only ~8% of the CRUs on a Z-disk would fire. (This calculation uses Eq. 10 and assumes for simplicity that the CRUs see the 1 μ M concentration throughout the 20-ms period. Thus, 8% is an upper bound of the fraction of CRUs that fire.) It is unlikely that such a low fraction of firing CRUs could support a wave. However, when a wave has a Ca²⁺ concentration of 10 μ M, 87% of the CRUs fire in that 20-ms period. Ca^{2+} waves with lower peak Ca^{2+} can exist when the Ca^{2+} sensitivity of CRUs is higher, for example, in the presence of low concentrations of caffeine.

Spontaneous Ca^{2+} spark frequency, I_{SR} , and Ca^{2+} waves—effects of superadditivity

An increase in the spontaneous Ca^{2+} spark frequency often precedes the appearance of spontaneous Ca^{2+} waves (6). Fig. 3 *C* provides a quantitative explanation for why this occurs. The ordinate is the *n*-fold increase of the maximum Ca^{2+} seen by a CRU (the "observer") 1 μ m away from 1, 3, 5, and 7 CRUs that fire. Notice that when three CRUs fire, the observer sees not a threefold increase in Ca^{2+} concentration but rather a sixfold increase. This disproportionate increase in peak Ca^{2+} is due to superadditivity of CRU release. The superadditive summation of Ca^{2+} sparks explains why a small increase in the spark frequency can produce an inordinate increase in the probability of more sparks and, by extension, a large increase in the probability of initiating a Ca^{2+} wave.

Superadditive effects also underlie the great sensitivity of the Ca²⁺ concentration to changes in CRU current. Figs. 3 *C* and 5 *B* show that the peak Ca²⁺ concentration increases nonlinearly with I_{SR} . An approximately twofold change of I_{SR} (from 8.4 to 16 pA) produces a 21-fold increase in the peak Ca²⁺ concentration measured 1 μ m away. Thus, even a modest increase in the Ca²⁺ flux from a spark can greatly increase the probability of Ca²⁺ waves.

Coupling between contractile dynamics and Ca²⁺ dynamics: large-scale cooperative CRU interactions

A surprising and important finding of this study is the remarkable sensitivity of the Ca^{2+} dynamics to the spatial distribution of the CRUs. A small decrease in the distance between CRUs in the plane of the Z-disk (Λ) can result in qualitatively different Ca^{2+} dynamics (Fig. 2). This sensitivity results from the rapid decline of the Ca^{2+} concentration as a function of distance from the source, as shown in Figs. 3 *B* (the 2-D case) and 5 *A* (the 3-D case). Note that a CRU sees a fourfold (2-D) or 12-fold (3-D) increase in the peak Ca²⁺ concentration as it moves just 0.2 μ m closer, from 1.0 to 0.8 μ m, to the Ca²⁺ source. Since the probability of CRU firing depends on the ambient Ca²⁺ concentration, a small change in CRU spacing will profoundly affect the firing probability.

 Ca^{2+} dynamics is also sensitive to large-scale cooperative CRU interactions. By this, we mean that the firing probability of a CRU on one Z-disk not only depends on the firing history of other CRUs on that Z-disk, but also on the firing histories of adjoining intercalated CRUs and of CRUs on adjacent Z-disks. The simulations in Fig. 6 illustrate the basic idea of large-scale cooperative CRU interactions. The simulations are identical except that cooperative interactions between CRUs on the two Z-disks are absent in Fig. 6 A but present in Fig. 6 B. In the absence of cooperative interactions the triggered firing of CRUs fails to initiate a Ca^{2+} wave (Fig. 6 A), but by allowing interaction of CRUs on adjacent Z-disks, the firing activity of CRUs on one Z-disk regeneratively reinforces the firing activity of CRUs on the other Z-disk. This mutual reinforcement results in Ca²⁺ wave initiation (Fig. 6 *B*).

Because of large-scale cooperative CRU interactions, the probability of wave initiation is influenced by the distance between Z-disks, that is, the sarcomere length. Fig. 9 *B* illustrates this point. Fig. 6*A* shows that when the Z-disk distance is 2.0 μ m a Ca²⁺ wave fails to initiate. However, when the Z-disk distance is reduced to 1.8 μ m (Fig. 6*B*) or 1.6 μ m (Fig. 6*C*), the same conditions allow a Ca²⁺ wave to form. The physiological implication is that the contractile state of the myocyte affects the dynamics of the Ca²⁺ control system. This novel feedback pathway arises solely from changes in the spatial distribution of CRUs as a result of contraction. This pathway is distinct from the coupling of contractile system to the Ca²⁺ control system via an NO-mediated mechanism (38) and also independent of the decrease in Ca²⁺ binding to troponin-C at shortened sarcomere lengths (39,40).

Physiological implications of large-scale cooperative CRU interactions

Ca²⁺ waves in atrial cells

Atrial myocytes, unlike ventricular myocytes, lack t-tubules, so Ca^{2+} release directly triggered by depolarization occurs mainly at the cell periphery (1–4,41,42) and, to a lesser extent, at sparse sites inside the cell (2). If Ca^{2+} release occurred only from CRUs directly triggered by depolarization, it would activate the myofilaments near the periphery, but myofilaments near the cell core would be only slightly activated and the slow rate of activation would reflect the passive diffusion of Ca^{2+} (2). On the other hand, centripetally propagating Ca^{2+} waves would greatly increase the rate and magnitude of myofilament activation at the core (2). For atrial cells to use

 Ca^{2+} waves, the Ca^{2+} control system must be delicately poised in a state of "controlled instability" wherein there is enough regenerative Ca^{2+} release to support wave propagation but not so much as to generate waves spontaneously.

The spacing of CRUs in the plane of the Z-disk (Λ) is important for determining the state of controlled instability. A small change of Λ from 1.0 to 0.8 μ m has a large effect on the probability of Ca²⁺ wave initiation (Fig. 3). Indeed, in the limit of $\Lambda = 0$ we would obtain the common pool model, whose failure to explain the graded Ca²⁺ release prompted Stem (43) to propose the local control theory of E-C coupling. On the other hand, our model could not produce Ca²⁺ waves with a transverse spacing of $\Lambda = 2 \,\mu$ m, as reported in cat atria (44). Our results suggest, then, that RyR2 clusters in species other than rat (which our model is based on), and in other cell types (e.g., neurons) where Ca²⁺ waves are found, would also be spaced ~1 μ m apart. Large departures from this value would indicate that compensatory changes (e.g., I_{SR} , RyR2 sensitivity to Ca²⁺) need to occur to maintain a state of controlled instability.

Possible function of intercalated CRUs to control wave initiation in atrial cells

A way of modulating the rate and magnitude of contraction in atrial myocytes would be to control the initiation of Ca^{2+} waves. When the intercalated CRUs are functional, both the probability of Ca^{2+} wave initiation and Ca^{2+} wave velocity greatly increase (Figs. 7 and 8). Thus, if there were a way of turning the intercalated CRUs on or off then this would be a means of controlling atrial myocyte contractility.

One possible way of modulating intercalated CRU activity is suggested by the finding that inositol-1,4,5 trisphosphate receptors (IP₃Rs) are colocalized with the RyR2s on the sarcolemma but not in the interior of rat atrial cells (45,46). Activation of the IP₃Rs results in an increase in the spark frequency in the subsarcolemmal region (45) and extra Ca²⁺ transients after action potential triggered Ca²⁺ release (46). A different pattern of IP₃R distribution is found in cat atrial cells. In these cells Zima and Blatter (47) observed that IP₃Rmediated Ca²⁺ release, similar to Ca²⁺ puffs (47), occurred throughout the cell.

We want to be clear that at this time we do not know whether the intercalated RyR2s are functional CRUs that release Ca^{2+} in response to depolarization or whether they release Ca^{2+} at all. Woo et al. (42) found that Ca^{2+} sparks on the cell periphery of rat atrial myocytes were spaced between 1.4 and 2 μ m apart. This result suggests that, at least under the conditions of their experiments (no IP₃R agonists were used), the intercalated CRUs were not functional.

Possible role of contractile Ca^{2+} system coupling in generating arrhythmias: "Short CRU spacing, more Ca^{2+} waves" concept

Spontaneous Ca^{2+} waves can induce sufficiently large depolarization to trigger an action potential (10,11), causing

delayed afterdepolarizations (49,50) and possibly cardiac arrhythmias (7,12,51). Known factors that predispose the cell to spontaneously generate Ca^{2+} waves include increased sarcoplasmic reticulum (SR) Ca^{2+} load (6,52–54) and increased RyR2 open probability (55–58). The study presented here suggests an additional important factor: shortened CRU spacing.

Shortened CRU spacing was observed in the ventricular myocytes from the spontaneously hypertensive rat (SHR) at the late-stage cardiac hypertrophy and heart failure. In comparison to the controls, the average longitudinal CRU spacing (which corresponds to Z-disk spacing or sarcomere length) in the SHR was 5% shorter in the isolated cells and 10% shorter in the tissue cross section (Y. Chen-Izu, C. W. Balke, and L. T. Izu, unpublished data) in comparison to the age-matched WKY controls. Consonant with the "short CRU spacing, more Ca²⁺ waves" idea, the hypertrophied SHR hearts had higher incidence of spontaneous Ca²⁺ waves (59) and higher susceptibility to arrhythmias in vivo (60) and in the isolated heart (61).

Shortened sarcomere spacing was also reported in two transgenic mouse models of familial hypertrophic cardiomyopathy (FHC). One model with the troponin T (R92Q) mutation displayed 10% shortening of the sarcomere length in the ventricular myocytes (62). Another model with mutations in the actin-binding domain of α -myosin heavy chain displayed 5% shortening of the sarcomere length (63). Humans with the same mutations suffer high risk of sudden cardiac death (64,65)

The above data from SHR and FHC models support the plausibility of the "short CRU spacing, more Ca^{2+} waves" concept. However, it should be noted that these measurements were done in unloaded cells and in tissue sections from unloaded hearts. It is unknown whether the degree of shortening seen in the unloaded cells and tissues is also present in the working myocardium. Thus, care must be taken in extrapolating the expected increase in spontaneous Ca^{2+} waves due to sarcomere shortening seen in unloaded cells and tissues to the working heart.

In both the SHR and the two FHC mutants described, the sarcomeres were shortened and the SR Ca2+ load was normal or even elevated compared to controls (63,66). This combination acts in synergy to increase the likelihood of spontaneous Ca²⁺ waves. By contrast, in human heart failure (HF) (67), in the dog (68,69), and in a rabbit model (8) of pressure-overload-induced HF, the SR Ca²⁺ load was reduced. Moreover, the mean sarcomere spacing remained the same in isolated myocytes from failing and nonfailing human left ventricle (70) and in isolated right ventricular myocytes from hypertrophied and normal rabbits (71). Thus, in human and rabbit HF, the combination of reduced SR Ca²⁺ load and normal sarcomere lengths would reduce the probability of spontaneous Ca²⁺ waves (assuming sarcomere lengths remain the same as the rabbit heart progresses from hypertrophy to HF). This conclusion, however, should be tempered, because it does not take into consideration the distribution of sarcomere lengths. In the human HF, the distribution of mean sarcomere lengths from normal hearts is Gaussian, but from hypertrophied hearts it is significantly left-skewed (71), meaning that there is a higher preponderance of cells from hypertrophied hearts than expected with short sarcomere lengths. This preponderance might be important, because spontaneous Ca²⁺ waves could be started in these cells with shortened sarcomeres and become foci for initiating Ca^{2+} waves. Fig. 9 C shows the relationship between sarcomere spacing and CRU current that determines whether the likelihood of spontaneous Ca²⁺ waves is high (unstable) or low (stable). Suppose point *a* is the initial state of the myocyte; the sarcomere length is 2 μ m and $I_{SR} \approx 20$ pA. Since this state is below the curve, the likelihood of spontaneously generated Ca²⁺ waves is low. If the sarcomere length is reduced to 1.8 μ m and the current is unchanged, then, as point b is above the curve, the system loses stability and there is a high probability that Ca^{2+} waves emerge spontaneously.

Limitations of the model and future developments

The model uses a number of simplifications dictated by a need to keep the computational burden within reason and the unavailability of experimental data to warrant a more detailed model. The steady-state response of the RyR2 opening probability to Ca^{2+} was used (Eq. 3), which allows us to analytically compute the WTD for CRU firing based on a canonical distribution of Ca^{2+} (a pulse or wave), enables us to rapidly obtain the archetypal behavior of this stochastic system. Without this simplification the mean behavior of the system would need to be found by averaging the results of many simulations. However, the use of steady-state behavior is likely to be inadequate when simulating the response of the CRUs to the opening of L-type Ca^{2+} channels, as the RyR2 opening probability is higher when Ca^{2+} changes rapidly than under steady-state conditions (72,73).

We have proposed a novel explanation of the heightened risk of arrhythmias in some forms of HF based on our finding of the strong coupling between the contractile and Ca²⁺ control systems in tandem with the finding that the sarcomere length is significantly shortened in the SHR and two FHC mutants (62,63). This explanation, however, is based on a static view of the sarcomeres. The cooperative interactions between CRUs on adjacent sarcomeres still operate during a heartbeat, but the magnitude of the cooperativity will depend also on the changing SR Ca²⁺ load and changing sarcomere distance. The current model does not include the dynamics of the lumenal SR Ca^{2+} . We have explored how the interplay of a reduced SR Ca^{2+} load and sarcomere shortening affect wave initiation (Fig. 9 C), but a dynamic description of the SR Ca²⁺ content is needed to fully understand the interaction between the contractile and Ca²⁺ control systems. New developments in dynamic measurement of lumenal Ca²⁺

content (74–76) will provide information about how SR load affects Ca^{2+} release and the next generation of our model will incorporate these new insights.

SUMMARY

Simulations and analytic results presented here show that the probability of Ca^{2+} wave initiation is extremely sensitive to even small changes in RyR2 distribution, CRU current, and Ca^{2+} spark frequency. This insight suggests that there is a strong, hitherto underappreciated, coupling between the contractile system—which dynamically modulates the RyR2 distribution—and the Ca^{2+} control system. Cardiac arrhythmias have a complex etiology and we propose that the coupling between the RyR2 distribution and the Ca^{2+} control system may contribute, in part, to the generation of cardiac arrhythmias.

APPENDIX A

The model equations are nearly identical to those used previously (15,16) except that for most simulations they were extended to three dimensions; the results shown in Figs. 3 and 10 are from 2-D simulations. The coordinate axes for the 3-D simulations are shown in Fig. 1. The Z-disk lies in the *xy* plane and the long axis of the cell is along the *z* axis. The partial differential equation (PDE) for Ca²⁺ is given by Kirk et al. (2). The PDEs for the free endogenous ($F_{\rm B}$) and exogenous ($F_{\rm D}$) buffers are

$$\frac{\partial F_{\rm B}}{\partial t} = R_{\rm B}(C, F_{\rm B}, H_{\rm B}) \tag{A1}$$

$$\frac{\partial F_{\rm D}}{\partial t} = \nabla \cdot \left(D_{\rm D} \nabla F_{\rm D} + R_{\rm D} (C, F_{\rm D}, H_{\rm D}) \right) \tag{A2}$$

where (j = D or B)

$$R_{j}(C,F_{j},H_{j}) = -k_{j}^{+}C \cdot F_{j} + k_{j}^{-}(H_{j} - F_{j})$$
(A3)

and H_j is the total (free + Ca²⁺-bound) buffer concentration. The standard parameter values for these buffer reactions are given in Table 1 of our previous work (15), which also gives the sources for these values. For convenience, these parameters are listed here: $H_B = 123 \ \mu$ M; these values are lumped parameters for endogenous Ca²⁺ buffers (77). To reduce the computational load, the model did not differentiate between different endogenous buffers such as troponin-C and calmodulin. Other parameters are $D_{Cx} = D_{Cy} = 0.15 \ \mu$ m²/ms, $D_{Cz} = 0.30 \ \mu$ m²/ms, $D_{Dx} = D_{Dy} = 0.01 \ \mu$ m²/ms, $D_{Dz} = 0.02 \ \mu$ m²/ms; $k_B^+ = 100/\mu$ M/s, $k_B^- = 100/s$; $k_D^+ = 80/\mu$ M/s, $k_D^- = 90/s$ (78). The initial Ca²⁺ concentration is $C(0,x,y,z) = 0.1 \ \mu$ M (except for the simulation in Fig. 10). The parameters describing the probability of opening of the CRUs given in Eqs. 2 and 3 have the following standard values: $T_{open} = 5 \ ms$, $P_{max} = 0.3/ms$, $K = 15 \ \mu$ M, and n = 1.6 (see Izu et al. (16) and references therein). The parameters for the SERCA2 pump,

$$J_{\text{pump}} = \frac{V_{\text{p}} C^{n_{\text{p}}}}{K_{\text{p}}^{n_{\text{p}}} + C^{n_{\text{p}}}},\tag{A4}$$

are $V_p = 200 \ \mu$ M/s, $K_p = 0.184 \ \mu$ M, and $n_p = 4$. J_{leak} is adjusted such that $J_{pump} - J_{leak} = 0$ when $C = 0.1 \ \mu$ M.

APPENDIX B

The idea of superadditivity captures the effect that multiple CRUs or increasing CRU current have on the Ca^{2+} concentration at an observation

point. If the firing of a CRU raises the Ca²⁺ concentration at the observation point by $\Delta C(1)$, what would the concentration change ($\Delta C(2)$)) be if another CRU fires? We chose the term superadditivity because buffer systems share the defining property of superadditive functions: $f(x + y) \ge f(x) + f(y)$ for all x and y. For example, x and y could be the amount of Ca²⁺ released by two CRUs and the concentrations change at an observation point because the combined effect, f(x + y), is greater than the linear combination, f(x) + f(y), of each CRU by itself. To sensibly compare the changes that occur in buffered and unbuffered systems we consider the changes in concentration relative to some reference state instead of the absolute changes. That is, we are interested in $\Delta C(2)/\Delta C(1)$. In the special case where the chemical reactions are in equilibrium, the superadditivity effect is readily calculated. Consider the following bimolecular reaction occurring in a volume V:

$$B + C \rightleftharpoons P.$$
 (B1)

Assume that $\overline{C} \cdot V$ moles of *C* are added. The constraints on the number of moles of *C* are

$$V \times C + V \times P = V \times C$$
$$V \times B + V \times P = V \times \overline{T}$$
(B2)

where \overline{T} is the total concentration of buffer (free and bound). Note that the volume cancels, and these constraints together with the equilibrium expression for $B \times C/P = K_d$, provide the three equations needed to solve for *C*, *P*, and *B* uniquely. The free concentration is

$$C = ((\bar{T} - \bar{C} + K_{\rm d}) + \sqrt{(\bar{T} - \bar{C} + K_{\rm d}) + 4K_{\rm d}\bar{C})/2}.$$
 (B3)

Superadditivity is measured by the ratio S,

$$S = \frac{\Delta C(2)}{\Delta C(1)} \approx \frac{\partial C/\partial C|_{\bar{C}_2} \times \delta C}{\partial C/\partial \bar{C}|_{\bar{C}_1} \times \delta \bar{C}}$$
(B4)

The reference state is that which gives a free ligand concentration of $C = 1.0 \mu$ M. In the unbuffered system this reference state is $\bar{C}(1) = 1.0 \mu$ M and for the buffered system, $\bar{C}(1) = 51 \mu$ M, assuming $\bar{T} = 100 \mu$ M and $K_d = 1 \mu$ M. In the absence of a buffer, $\bar{T} = 0$, *S* is identically 1. This means the concentration change is always in the same proportion to the number of moles added. For the buffered system, *S* increases monotonically, indicating that as more of the ligand *C* is added and fills up the binding sites, each additional bolus of *C* would cause a larger increase in the free concentration. Doubling \bar{C} increases the free Ca²⁺ concentration 10-fold.

SUPPLEMENTARY MATERIAL

An online supplement to this article can be found by visiting BJ Online at http://www.biophysj.org.

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