**Perspective**

**Ryanodine Receptor Adaptation**

M. Fill,* A. Zahradníková,‡ C.A. Villalba-Galea,§ I. Zahradník,‡ A.L. Escobar,§ and S. Györke§

From the *Department of Physiology, Loyola University Chicago, Maywood, IL; ‡Institute of Molecular Physiology and Genetics, Bratislava, 833334 Slovak Republic; and §Department of Physiology, Texas Tech University Health Science Center, Lubbock, Texas 79430

In the heart, depolarization during the action potential activates voltage-dependent Ca\(^{2+}\) channels that mediate a small, localized Ca\(^{2+}\) influx (I\(_{Ca}\)). This small Ca\(^{2+}\) signal activates specialized Ca\(^{2+}\) release channels, the ryanodine receptors (RyRs), in the sarcoplasmic reticulum (SR). This process is called Ca\(^{2+}\)-induced Ca\(^{2+}\) release (CICR). Intuitively, the CICR process should be self-regenerating because the Ca\(^{2+}\) released from the SR should feedback and activate further SR Ca\(^{2+}\) release. However, the CICR process is precisely controlled in the heart and, consequently, some sort of negative control mechanism(s) must exist to counter the inherent positive feedback of the CICR process. Defining the nature of this negative control has been a focus of investigation for decades. Several mechanisms have been suggested including all of the following: Ca\(^{2+}\)-dependent inactivation, adaptation, stochastic attrition, “fateful” inactivation, SR Ca\(^{2+}\) depletion, and coupled RyR gating. These mechanisms are generally regarded as being mutually exclusive (i.e., alternative). An emerging and more sophisticated view is that the regulation of CICR is not a single mechanism, but a complex phenomenon that involves multiple negative control mechanisms. One important mechanism is that of conventional inactivation (Fabiato, 1985; Nabauer and Morad, 1992). More recently, Lukyanenko and Györke (1999) showed that elevation of resting Ca\(^{2+}\) levels increased the frequency of spontaneous Ca\(^{2+}\) sparks, which are thought to arise from bursts of SR Ca\(^{2+}\) release channel activity. The conclusion was that SR Ca\(^{2+}\) release (RyR) channel activity in cells is not depressed by a high affinity Ca\(^{2+}\)-dependent inactivation. There are also no signs of high affinity Ca\(^{2+}\)-dependent inactivation (e.g., occurring at <0.1 mM Ca\(^{2+}\)) in studies of steady-state single RyR channel behavior (e.g., Györke and Fill, 1993; Zahradníková and Zahradník, 1995). Interestingly, there is evidence supporting the existence of low affinity inactivation that occurs at very high Ca\(^{2+}\) concentrations (>1 mM; Laver et al., 1995). It is not clear whether such high cytoplasmic Ca\(^{2+}\) concentrations are ever reached in the cell. 

Ca\(^{2+}\)-dependent Inactivation

Fabiato (1985) was the first to propose the existence of Ca\(^{2+}\)-dependent inactivation. He proposed that inactivation of SR Ca\(^{2+}\) release is due to slow Ca\(^{2+}\) binding to a high affinity inactivation site on the SR Ca\(^{2+}\) release channel. He showed, in a skinned cardiac cell preparation, that SR Ca\(^{2+}\) release was substantially inactivated at steady-state Ca\(^{2+}\) concentrations as low as 60 nM. He argued that this conventional absorbing Ca\(^{2+}\)-dependent inactivation leaves the SR Ca\(^{2+}\) release process refractory. Recovery from this refractory state requires removal of the Ca\(^{2+}\) stimulus and time. One important net of conventional Ca\(^{2+}\)-dependent inactivation is that it should be evident in both the stationary and nonstationary behavior of the channel. Another important net of conventional inactivation is that it will be essentially an all-or-none process (i.e., any channel will be either inactivated or not) at the single channel level. Thus, a second incremental Ca\(^{2+}\) stimulus would not be expected to reactivate an inactivated (i.e., refractory) channel regardless of stimulus amplitude or duration.

Historically, studies on intact and permeabilized cells present contradictory evidence concerning the existence of Ca\(^{2+}\)-dependent inactivation (Fabiato, 1985; Nabauer and Morad, 1992). More recently, Lukyanenko and Györke (1999) showed that elevation of resting Ca\(^{2+}\) levels increased the frequency of spontaneous Ca\(^{2+}\) sparks, which are thought to arise from bursts of SR Ca\(^{2+}\) release channel activity. The conclusion was that SR Ca\(^{2+}\) release (RyR) channel activity in cells is not depressed by a high affinity Ca\(^{2+}\)-dependent inactivation. There are also no signs of high affinity Ca\(^{2+}\)-dependent inactivation (e.g., occurring at <0.1 mM Ca\(^{2+}\)) in studies of steady-state single RyR channel behavior (e.g., Györke and Fill, 1993; Zahradníková and Zahradník, 1995). Interestingly, there is evidence supporting the existence of low affinity inactivation that occurs at very high Ca\(^{2+}\) concentrations (>1 mM; Laver et al., 1995). It is not clear whether such high cytoplasmic Ca\(^{2+}\) concentrations are ever reached in the cell.

Ca\(^{2+}\)-dependent Adaptation

Györke and Fill (1993) were the first to explore the complex dynamics of RyR channel Ca\(^{2+}\) regulation. They showed that fast Ca\(^{2+}\) stimuli rapidly activated (τ ~1 ms) single RyR channels in planar bilayers to a high open probability (P\(_o\)) level. The P\(_o\) then slowly and spontaneously decayed. This was surprising because the Ca\(^{2+}\) stimuli generated a sustained Ca\(^{2+}\) elevation of ~1 μM, and there was no evidence of steady-state Ca\(^{2+}\)-dependent inactivation at that Ca\(^{2+}\) concentration. To test if the spontaneous decay was mediated by conventional Ca\(^{2+}\)-dependent inactivation, Györke and Fill
RyR deactivation following the very fast, large Ca$^{2+}$ decrease in channel activity might just reflect slow adaptation and modal gating was controversial. Lamb et al. (1994) suggested that inactivation, not adaptation is described later in the context of their intrinsic empirical evidence that adaptation is a manifestation of Ca$^{2+}$-dependent modal gating (for details see RyR Adaptation and Modal Gating).

The Györke and Fill (1993) RyR adaptation hypothesis was controversial. Lamb et al. (1994) suggested that the decrease in channel activity might just reflect slow RyR deactivation following the very fast, large Ca$^{2+}$ spike that was present on the leading edge of the applied Ca$^{2+}$ stimuli. The merits of each theoretical concern were experimentally addressed. These studies indicate that the fast Ca$^{2+}$ spike has no impact on the much slower (1,000-fold slower) adaptation phenomenon. They also indicate that the impact of the fast Ca$^{2+}$ spike is limited to accelerating the closed to open transition of the channel (i.e., essentially super-charging the Ca$^{2+}$ stimuli). The evidence supporting this conclusion is described later in Ca$^{2+}$ Spike Concern.

The nonstationary dynamics of RyR Ca$^{2+}$ regulation have also been explored using other methodologies (Schiefer et al., 1995; Sitsapesan et al., 1995; Laver and Curtis, 1996; Laver and Lamb, 1998). These studies also report a relatively fast Ca$^{2+}$-activation followed by a slow spontaneous decay in channel activity. However, the slow spontaneous decay is interpreted as conventional Ca$^{2+}$-dependent inactivation, not adaptation. Some investigators have evoked the original Lamb et al. (1994) concern to suggest that inactivation, not adaptation, warrants further investigation. We disagree and suggest that our focus should be on reconciling the diverse RyR data sets based on their intrinsic empirical value instead of simply defending a particular interpretation. The merits of each view are discussed later in Mechanical Solution Change Studies.

**RyR Adaptation and Modal Gating**

Under steady-state conditions, the opening of single RyR channels occurs in bursts. These bursts fall into two categories (i.e., high and low open probability; $P_o$). These bursts do not occur randomly, but, instead, they are temporally clustered into distinct modes of RyR channel gating (i.e., high- and low-$P_o$ modes). Several groups have now reported the existence of modal RyR channel gating (Zahradníková and Zahradník, 1995; Armisén et al., 1996; Villalba-Galea et al., 1998; Säftenku et al., 2000).

Stationary single RyR channel activity exhibits three distinct gating modes (Zahradníková and Zahradník, 1995). At intermediate Ca$^{2+}$ concentrations ($\sim$10 μM), the high $P_o$ mode (H-mode) is characterized by periods of frequent and long openings. The low $P_o$ mode (L-mode) is marked by periods of infrequent and short openings. The inactivated mode (I-mode) corresponds to periods where no opening events occur. A simple Markovian model of RyR modal gating was proposed by Zahradníková and Zahradník (1996) that predicts that upon a small fast Ca$^{2+}$ elevation the $P_o$ of a RyR channel residing in the H-mode will rapidly increase before the channel has time to relax to a new equilibrium between all three modes. The result is a rapid rise of channel activity that spontaneously decays over time. A second prediction is that the response of a single RyR channel to fast Ca$^{2+}$ elevation will depend on which mode the channel happens to be in at the moment the stimulus is applied. In other words, a single channel fluctuates spontaneously between modes with different capacities to respond to a Ca$^{2+}$ stimulus. For example, if a channel happens to be in the L- or I-mode (40% probability at 1 μM Ca), then the probability of a high $P_o$ response will be reduced to $\sim$17% because of the relatively long dwell times of these modes. Another prediction is that a single RyR channel will respond transiently and repeatedly to small fast incremental Ca$^{2+}$ elevations. This realization lead Zahradníková and Zahradník (1996) to propose that modal gating may explain the adaptation phenomenon observed by Györke and Fill (1993). For years, this proposal was based on theory, not experimental evidence. Recently, Zahradníková et al. (1999a) defined the evolving temporal correlations between the three RyR gating modes during the adaptation phenomenon and experimentally established that modal gating and the adaptation phenomenon are likely related.

There are certain aspects of the Zahradníková and Zahradník (1996) model, however, that do not entirely reproduce the adaptation phenomenon. Our view is that these discrepancies indicate that the modal gating model needs refinement. Others have viewed these discrepancies as evidence against adaptation. Villalba-Galea et al. (1998) refined and expanded the original Zahradníková and Zahradník (1996) modal gating model to better reproduce various aspects of RyR gating behavior. The Villalba-Galea et al. (1998) form of the model incorporates an additional Ca$^{2+}$-dependent transition that shifts channel gating to the H-mode as the amplitude of the Ca$^{2+}$ perturbation becomes greater. In addition, the model includes a Ca$^{2+}$-dependent transition to the inactivated state to better reproduce high Ca$^{2+}$ (>1 mM) inhibition. The specific Markovian model presented by Villalba-Galea et al. (1998) is illustrated in Fig. 1 A. The simulated response of a single RyR channel to Ca$^{2+}$ stimuli like those applied by Györke and Fill (1993) is pre-
The Ca\textsuperscript{2+} stimuli (Fig. 1 B) did not contain the fast Ca\textsuperscript{2+} spike. The model predicts that single RyR channels display rapid activation followed by a slow, spontaneous decay in channel activity. The spontaneous decay occurs at Ca\textsuperscript{2+} levels (\(<1 \mu M\)) that are well below those known to induce Ca\textsuperscript{2+}-dependent inactivation under steady-state conditions (e.g., \(>1 \text{mM}\)). A second incremental Ca\textsuperscript{2+} stimulus induces a second transient of channel activity. Thus, a specific kinetic scheme (Fig. 1 A) can reproduce the fundamental hallmarks of the adaptation phenomenon (Györke and Fill, 1993) in the absence of the fast Ca\textsuperscript{2+} spike.

Recently, the Williams’ group has reported highly variable Ca\textsuperscript{2+}-dependent modal RyR gating (Saftenku et al., 2000). They found that elevating the steady-state Ca\textsuperscript{2+} concentrations increases the \(P_o\) within the L-mode and also increases the probability of the channel being in the H-mode. They suggest that this observation is inconsistent with the adaptation hypothesis. However, it is not clear, how this behavior might preclude the possibility of...
adaptation. The Villalba-Galca et al. (1998) model described above, in agreement with the results of Sretenku et al. (2000), clearly predicts a Ca\(^{2+}\)-dependent increase in H-mode probability. It also predicts a Ca\(^{2+}\)-dependent increase in P\(_o\) during the L-mode but found that this increase was significant but minor (\(<5\%\) P\(_o\)) at Ca\(^{2+}\) concentrations where the adaptation phenomenon occurs (i.e., between 0.5 and 20 \(\mu\)M Ca). It is the Ca\(^{2+}\)-dependent increase in P\(_o\) during the H-mode, not during the L-mode, that drives the adaptation phenomenon.

How does Ca\(^{2+}\)-dependent modal gating work? At low Ca\(^{2+}\) concentrations (100 nM), the states with the highest probability of being occupied are C1, C2, and C3 and, consequently, the channel will be closed most (97%) of the time. After a small Ca\(^{2+}\) elevation, the channel will move initially into the C4 and O1 states, inducing long openings and short closures (H-mode), and then into the C5, O2, and C6 states, inducing short openings and long closures (L-mode). Interestingly, the larger the Ca\(^{2+}\) stimulus, the longer the channel spends in the H-mode. The Ca\(^{2+}\)-dependent transition that connects C5 with O1 shifts the model to the H-mode during large Ca\(^{2+}\) stimuli. At high Ca\(^{2+}\) concentrations (e.g., >1 mM), the Ca\(^{2+}\)-dependent transition between the O1 and I states will move the channel into the nonconducting I state (I-mode).

At any steady-state Ca\(^{2+}\) level, there will be a dynamic equilibrium between the high P\(_o\), low P\(_o\), and inactivated modes. This dynamic equilibrium generates the well-known bell-shaped steady-state Ca\(^{2+}\) dependence of the RyR channel (Laver et al., 1995). As described above, small, fast Ca\(^{2+}\) elevations from a low Ca\(^{2+}\) concentration will upset the existing dynamic equilibrium momentarily in favor of the high P\(_o\) mode. Consequently, the P\(_o\) rises to a level above that predicted by steady-state measurements before activity spontaneously decays as the three modes reequilibrate at the new higher Ca\(^{2+}\) concentration. Such suprasteady-state P\(_o\) followed by a spontaneous decay has been experimentally observed previously (Györke and Fill, 1993; Valdivia et al., 1995; Laver and Curtis, 1996; Laver and Lamb, 1998). Application of a second incremental Ca\(^{2+}\) elevation will induce another transient modal gating shift and a second transient of channel activity like that reported by Györke and Fill (1993). The magnitude of the transient modal gating shift (i.e., adaptation) will depend on Ca\(^{2+}\) stimulus speed and magnitude. Slow and large Ca\(^{2+}\) stimuli will be less effective at generating adaptation. Slow Ca\(^{2+}\) stimuli are less effective because mode reequilibration is time dependent. Large Ca\(^{2+}\) stimuli are less effective because the channel begins to spend time in the inactivated mode. Thus, a relatively large slow Ca\(^{2+}\) stimulus will induce inactivation, not adaptation. This is consistent with the experimental results of Sitsapesan et al. (1995) and Schiefer et al. (1995). If refractory behavior was a characteristic of the inactivated mode, then this modal gating scheme would predict refractory behavior only at high Ca\(^{2+}\) levels. Interestingly, Schiefer et al. (1995) reported clear refractory behavior at 1 mM (following a 72% spontaneous decay), but found no evidence of refractory behavior at 10 \(\mu\)M (after \(\sim30\%\) spontaneous decay). In cells, the RyR channel is subjected to very brief (<1 ms) large (\(>10\) \(\mu\)M) Ca\(^{2+}\) stimuli (Stern, 1992). The modal gating scheme predicts that such large, fast Ca\(^{2+}\) stimuli will trigger one or two brief opening events followed by fast deactivation. These predictions are also quite consistent with the experimental record (Zahradniková et al., 1999b).

Thus, modal gating reconciles a large body of apparently contradictory experimental results and, thus, provides a useful comprehensive theoretical context for understanding RyR Ca\(^{2+}\) regulation. In this context, RyR inactivation and adaptation should not be viewed as mutually exclusive, but rather two different manifestations of the same underlying mechanism (i.e., modal gating). However, this modal gating scheme is still deficient because the potential impact of certain physiologically important ligands (e.g., luminal Ca, cytosolic Mg, and ATP) is not considered. This deficit, and others, will have to be addressed to correlate single RyR channel data with local Ca\(^{2+}\) release events in cells.

### The Fast Ca\(^{2+}\) Spike Concern

Here, the possibility that the fast Ca\(^{2+}\) spike drives the adaptation phenomenon is discussed. It is known that flash photolysis of certain caged Ca\(^{2+}\) compounds (e.g., DM-nitrophen and NP-EGTA) can generate a very fast Ca\(^{2+}\) overshoot (i.e., the Ca\(^{2+}\) spike) at the leading edge of a sustained Ca\(^{2+}\) elevation. Lamb et al. (1994) suggested that the adaptation phenomenon might reflect slow RyR deactivation following a fast Ca\(^{2+}\) spike. The suggestion was that the Ca\(^{2+}\) activation sites on the RyR channel are rapidly occupied during the brief
Ca$^{2+}$ spike (lasting $\sim 1$ ms), resulting in fast RyR activation to a high $P_C$ level. As the Ca$^{2+}$ concentration falls to a substantially lower level after the brief Ca$^{2+}$ spike, the RyR channel may simply “turn off” (i.e., deactivate) very slowly as occupancy of Ca$^{2+}$ activation sites falls. This concern was very legitimate considering the vacuum of knowledge that existed in 1994.

Lamb et al. (1994) estimated the properties of the fast Ca$^{2+}$ spike using the available published properties of the Ca$^{2+}$ cage complex. Györke and Fill (1994) began to experimentally evaluate the potential impact of the Ca$^{2+}$ spike on the adaptation phenomenon. They reported no compelling evidence to support the theoretical contenotions of Lamb et al. (1994). Subsequently, it became clear that the previous estimates of Ca$^{2+}$ spike properties were wrong. The experiments of Escobar et al. (1997) revealed that the Ca$^{2+}$ spike was considerably faster and smaller than previously thought. This supported the intuitive assumption of Györke and Fill (1994) that it was very unlikely that such a brief Ca$^{2+}$ signal (lasting $< 0.2$ ms) could drive the slow adaptation phenomenon (lasting $> 1,000$ ms). Another interesting insight gained from the Escobar et al. (1997) study was that the fast Ca$^{2+}$ spike is actually a reasonable representation of the fast Ca$^{2+}$ signals that may occur in the dyadic cleft during openings of single dihydropyridine receptor (DHPR) channels (Stern, 1992). This latter insight prompted Zahradníková et al. (1999b) to further explore the potential impact of the fast Ca$^{2+}$ spikes on single RyR channel activity. They improved the experimental recording conditions (i.e., increasing bandwidth, minimizing flash artifacts, etc.), so that they could define the impact of a Ca$^{2+}$ spike alone (with little or no sustained Ca$^{2+}$ elevation) during the first few milliseconds following its application. The rationale was that bandwidth and flash artifacts may have masked the impact of the Ca$^{2+}$ spike in previous studies (Györke and Fill, 1993; Györke et al., 1994; Valdivia et al., 1995). They showed that a fast Ca$^{2+}$ spike alone, albeit slightly smaller than those applied previously, induced one or two brief opening events during the first few milliseconds after the flash. When Zahradníková et al. (1999b) duplicated the Györke and Fill (1993) stimuli (i.e., spike plus sustained Ca$^{2+}$ elevation), the classical adaptation phenomenon was observed. Thus, the RyR channel’s response to a Ca$^{2+}$ spike alone is dramatically different than its response to an Ca$^{2+}$ spike followed by a sustained Ca$^{2+}$ elevation. Zahradníková et al. (1999b) concluded that Ca$^{2+}$ spikes are simply too brief to drive the very slow adaptation phenomenon.

Lamb et al. (2000) have a different interpretation of the Zahradníková et al. (1999b) data with adaptation again being driven by the Ca$^{2+}$ spike and not by the sustained Ca$^{2+}$ elevation. This interpretation is based, in part, on the fact that the Lamb et al. (1999b) Ca$^{2+}$ spikes were somewhat smaller ($\leq 30$ µM peak or estimated Ca$^{2+}$ time integrals of 2–7 µMms) than those applied by Györke and Fill (1993; 30–60 µM peaks or estimated Ca$^{2+}$ time integrals of 8–30 µMms). The Lamb group suggests that the Zahradníková et al. (1999b) data simply show that there is some sort of critical threshold for the Ca$^{2+}$ spike to induce the adaptation phenomenon. In other words, the RyR channel distinguishes a Ca$^{2+}$ spike peaking at 40 µM (with a Ca$^{2+}$ time integral of 8 µMms) from one peaking at 30 µM (with a Ca$^{2+}$ time integral of 6 µMms) and its response to each is different (slow or fast deactivation, respectively). They speculate that the action of large Ca$^{2+}$ spikes is mediated by some hypothetical very low affinity Ca$^{2+}$ binding sites that are preferentially occupied only during large Ca$^{2+}$ spikes. However, this new hypothesis is inconsistent with common kinetic principles of receptor–ligand interaction and directly contradicts existing experimental evidence regarding RyR behavior.

Common kinetic principles dictate that very low affinity Ca$^{2+}$ binding sites typically have fast kinetics (conversely high affinity sites typically have slow kinetics). It is difficult to understand how the hypothetical low affinity sites could mediate a very slow phenomenon like adaptation. In numerical terms, the low affinity sites would have a $K_d$ near 50 µM and an off-rate of 1 s$^{-1}$ (i.e., the published time constant of RyR adaptation). Consequently, the probability of that site being occupied during a fast 50-µM Ca$^{2+}$ spike lasting $< 1$ ms will be $< 0.001$; thus, the spike is unlikely to have a dramatic impact on such a site (Escobar et al., 1997). Additionally, the probability of a state of being occupied in any kinetic reaction is typically described by a probability distribution and not by a particular triggering level. Lamb et al. (2000; this issue) acknowledge the difficulty of explaining their new hypothesis in specific kinetic terms, and offer no specific kinetic scheme to support their point of view. Instead, they simply suggest that large Ca$^{2+}$ spikes must drive the RyR channel into some unusual configuration and/or induce some unknown mechanisms. On the other hand, we have forwarded a specific and relatively simple Markovian gating scheme (Fig. 1) that describes adaptive behavior arising from Ca$^{2+}$-dependent modal gating changes driven by the sustained component of the photolytic Ca$^{2+}$ stimuli. As described earlier, the Lamb et al. (2000) predicts that adaptation of a single RyR channel occurs when some critical Ca$^{2+}$ spike threshold is exceeded (i.e., $\geq 30$ µM peak or Ca$^{2+}$ time integral $\geq 8$ µMms). However, there is experimental evidence that suggest otherwise. Györke and Fill (1993) and Valdivia et al. (1995) used the same method but different caged-Ca$^{2+}$ compounds, DM-nitrophen and NP-EGTA, respectively. Consequently, the applied Ca$^{2+}$ stimuli had a similar sustained Ca$^{2+}$ elevation, but fast Ca$^{2+}$ spikes of dramat-
ically different amplitude (and time integral). The similarity of RyR adaptation in both works suggested that adaptation is not highly dependent on subtle features of the fast Ca$^{2+}$ spike. More recently, our group (Zahradníková, A., and S. Györke, unpublished results) has also explored the impact of large Ca$^{2+}$ spikes on single RyR channel function. In this work, large Ca$^{2+}$ spikes (80–150 μM, Ca$^{2+}$ time integrals of up to 15 μMms) were generated by photolysis of NP-EGTA (3 mM). The large Ca$^{2+}$ spikes triggered only solitary RyR openings, not the slow adaptation phenomenon. The openings were reminiscent of the spike-activated events previously reported by Zahradníková et al. (1999b). Thus, it does not appear that adaptation depends on the amplitude or Ca-time integral of the fast Ca$^{2+}$ spike.

The premise that the RyR channel undergoes slow deactivation (lasting seconds) after the fast Ca$^{2+}$ spike is also inconsistent with existing experimental evidence. The rate of RyR deactivation in response to a fast Ca$^{2+}$ reduction has been measured (Schief er et al., 1995; Vélez et al., 1997; Zahradníková et al., 1999b). These experimental data show that RyR deactivation from high Ca$^{2+}$ concentrations is fast ($\tau$s < 10 ms), not slow (lasting seconds). Thus, the slow adaptation process is not likely due to very slow RyR deactivation.

What then is the impact of the Ca$^{2+}$ spike on RyR channel function? The fast Ca$^{2+}$ spike essentially supercharged the Ca$^{2+}$ stimulus in the flash photolysis studies (Györke and Fill, 1993). Its impact is best illustrated using the RyR scheme presented above. Simulated RyR channel response to a fast Ca$^{2+}$ elevation with and without a fast Ca$^{2+}$ spike at its leading edge is shown in Fig. 1, E and D, respectively. The sustained component of the Ca$^{2+}$ stimulus elevated Ca$^{2+}$ from 0.1 to 1.0 μM. The fast Ca$^{2+}$ spike peaked at ~50 μM and lasted 0.2 ms. The rate and extent of the slow spontaneous decay (i.e., adaptation) is identical in the presence or absence of the fast Ca$^{2+}$ spike. The presence of the fast Ca$^{2+}$ spike, however, introduces a brief peak in channel activity that rapidly decays (i.e., deactivates). This brief peak is generated by the spike-activated events as described experimentally by Zahradníková et al. (1999b). In essence, the impact of the fast Ca$^{2+}$ spike is limited to accelerating the closed to open transition of the channel. It does not impact the rate or extent of the much slower adaptation phenomenon. This is consistent with the original assumption of Györke and Fill (1993) that the fast Ca$^{2+}$ spike is simply too fast to impact the adaptation phenomenon.

Mechanical Solution Change Studies

Several studies have explored the complex dynamics of single RyR channel Ca$^{2+}$ regulation using the flash photolysis methodology (Györke and Fill, 1993; Györke et al., 1994; Valdivia et al., 1995; Vélez et al., 1997; Zahradníková et al., 1999b). The data are quite consistent between studies, and are generally interpreted in terms of adaptation, not inactivation. Other studies have explored the dynamics of RyR Ca$^{2+}$ regulation using Ca$^{2+}$ stimuli generated by different mechanical solution change methodologies (Schief er et al., 1995; Sitsapesan et al., 1995; Laver and Curtis, 1996). These studies report RyR activation followed by either a slow spontaneous decay or no decay at all. When RyR channel activity spontaneously decayed, the decay was interpreted as inactivation, not adaptation. This interpretation and the absence of fast Ca$^{2+}$ spikes in these latter studies have been construed as evidence against the adaptation phenomenon.

The following questions are addressed below. First, are the mechanical solution change data internally consistent enough to dismiss data collected using a different methodology? Second, are the stimuli in the mechanical solution change studies “true” [Ca$^{2+}$] steps and the only ones suitable for defining the nonstationary behavior of RyR? Finally, is conventional Ca$^{2+}$-dependent inactivation the only mechanism consistent with the mechanical solution change data? We believe the answer to each of these questions is no.

Three different mechanical solution change strategies have been applied to study single RyR channel function in planar bilayers (Schief er et al., 1995; Sitsapesan et al., 1995; Laver and Curtis, 1996; Laver and Lamb, 1998). Schief er et al. (1995) used a piezo-based method and achieved complete solution exchange over a 1-ms period. Sitsapesan et al. (1995) used a solenoid-based method that exchanged solutions over a period of 10 ms. Laver and Curtis (1996) and Laver and Lamb (1998) used a “puffing” method that exchanged solutions over a period ranging from 20 to 110 ms. Each of these mechanical solution change studies also applied different size Ca$^{2+}$ stimuli. Some applied small Ca$^{2+}$ stimuli (<10 μM; Schief er et al., 1995, Laver and Curtis, 1996). All applied very large Ca$^{2+}$ stimuli (>100 μM; Schief er et al., 1995; Sitsapesan et al., 1995; Laver and Curtis, 1996; Laver and Lamb, 1998). The reported rates of Ca$^{2+}$ activation varied from 0.2 to 20 ms, whereas the reported rates of inactivation varied from 200 to 15,000 ms. Inactivation was highly voltage-dependent in one study (Sitsapesan et al., 1995), but was voltage-independent in another (Schief er et al., 1995). Two of the studies report inactivation only at Ca$^{2+}$ concentrations (≥100 μM) known to inactivate channels under steady-state conditions (Schief er et al., 1995; Sitsapesan et al., 1995). Two other studies report inactivation at Ca$^{2+}$ concentrations (1 μM) that do not inactivate channels under steady-state conditions (Laver and Curtis, 1996; Laver and Lamb, 1998). The variability in results between the different solution exchange studies probably reflects the different methodologies used and ultimately the types of Ca$^{2+}$ stimuli applied (see below).
Another potential contributing factor may be a limited sample size in some cases (Schiefer et al., 1995; Laver and Lamb, 1998). We suggest that the data presented in the solution exchange studies is important, but is not internally consistent enough to categorically dismiss results collected using a different methodology.

Recently, studies of RyR Ca\(^{2+}\) dynamics performed with improved time resolution show that RyR Ca\(^{2+}\) activation is very fast (activation time constant 0.1 ms; Zahradníková et al., 1999b). Such a rapid response time is consistent with the local control theory (Stern, 1992). In vivo, the RyR channel may be activated by very fast, brief local Ca\(^{2+}\) changes associated with single openings of a DHPR channel. Considering the fast kinetics of RyR channel Ca\(^{2+}\) activation, the Ca\(^{2+}\) stimuli applied by the mechanical solution change studies described above cannot be regarded as true instantaneous Ca\(^{2+}\) steps. The possible exception is the Schiefer et al. (1995) study. However, the other studies applied relatively slow Ca\(^{2+}\) changes that required 10–110 ms to complete due to considering the large unstirred layers that exist immediately adjacent to the bilayer (Laver and Curtis, 1996). Consequently, the Ca\(^{2+}\) stimuli applied in all but one of the mechanical solution change studies can be more accurately described as Ca\(^{2+}\) ramps instead of steps. The point is that the same critical scrutiny that was applied to the photolytic Ca\(^{2+}\) stimuli has not yet been applied to the mechanical Ca\(^{2+}\) stimuli.

The dynamics of RyR channel behavior in response to a very fast Ca\(^{2+}\) step or a relatively slow Ca\(^{2+}\) ramp are likely to be different. The potential impact of Ca\(^{2+}\) stimulus speed on single RyR channel gating can be seen in simulated channel data presented in Fig. 1. Not surprisingly, RyR behavior to a fast Ca\(^{2+}\) step (rise time ≤100 μs) or a ramp-like Ca\(^{2+}\) stimulus (rise time 100 ms; Laver and Curtis, 1996) was quite different. Changes in Ca\(^{2+}\) stimulus speed impacted both Ca\(^{2+}\) activation and spontaneous decay (Fig. 1, D–F). The point is that Ca\(^{2+}\) stimuli with such slow rise times (10 to 100 ms) can have rather dramatic impact on RyR channel function. This may explain the tight parallel between the dynamic and stationary behavior of single RyR channels in certain mechanical solution change studies (Sitsapesan et al., 1995). The high sensitivity to Ca\(^{2+}\) stimulus speed (Zahradníková et al., 1999b) and the very fast Ca\(^{2+}\) stimuli thought to occur in vivo (Stern, 1992) suggest that the relatively slow Ca\(^{2+}\) stimuli available in most mechanical solution change studies may not be appropriate to study the dynamics of RyR regulation especially in the physiologically relevant range of Ca\(^{2+}\) stimuli rates (<1 ms).

We also believe that the mechanical solution change results provide evidence for both conventional inactivation and adaptation. Schiefer et al. (1995) and Sitsapesan et al. (1995) report a spontaneous decay in RyR activity at Ca\(^{2+}\) concentrations (i.e., ≥100 μM) where in-
Correlating Single RyR Behavior to Ca^{2+} Release Phenomena in Cells

The identity of the negative control mechanism(s) that counters the inherent positive feedback of CICR in heart has been debated over the last 15 years. Two candidate mechanisms are conventional Ca^{2+}-dependent inactivation and adaptation. However, it appears that neither of these mechanisms alone is sufficient to explain the properties of CICR in cells. It is clear that a more comprehensive understanding of the mechanisms that govern intracellular Ca^{2+} release is required.

Fabiato (1985) was the first to propose that Ca^{2+}-dependent inactivation is the negative control mechanism that regulates CICR in the heart. This proposal was controversial because early patch-clamp studies found no evidence of inactivation of cell-averaged (global) Ca^{2+} release in experiments using a conventional two-pulse protocol. Subsequent patch-clamp studies, using a sustained trigger Ca^{2+} stimulus, showed that SR Ca^{2+} release does, indeed, “turn-off” (Yasui et al., 1994). Interestingly, the “turned-off” Ca^{2+} release could be reactivated by a further increase in the trigger Ca^{2+} stimulus. Other groups have reported such paradoxical behavior in variety of preparations (e.g., Pizarro et al., 1997) and found it to be analogous to quantal or incremental Ca^{2+} release observed from IP_{3}-sensitive Ca^{2+} stores. This is interesting because the RyR and IP_{3}-receptor are homologous intracellular Ca^{2+} release channels. Each IP_{3} increment of an incremental IP_{3} stimulus induces a small transient Ca^{2+} release. The mechanism that turns-off Ca^{2+} release after an incremental IP_{3} stimulus is not desensitization because a second incremental IP_{3} stimulus could reactivate Ca^{2+} release. This property of IP_{3}-sensitive Ca^{2+} stores is interesting here for the following two reasons: (1) it is reminiscent of the RyR adaptation phenomenon; and (2) it could be related to the apparent quantal or incremental Ca^{2+} release observed from ryanodine sensitive Ca^{2+} stores (Yasui et al., 1994; Pizarro et al., 1997). The IP_{3} literature has focused on two potential explanations of quantal Ca^{2+} release. First, quantal release phenomenon could be generated by Ca^{2+} stores with heterogeneous IP_{3} sensitivity, so that each stimulus increment recruits a new segment of the Ca^{2+} store population. Second, a homogenous Ca^{2+} store population could generate quantal release if each release was governed by an adaptation-like phenomenon. The RyR-IP_{3} homology, the homogeneity of IP_{3} binding affinity, and the existence of RyR adaptation supports the second hypothesis. Therefore, it is possible that much of the complexity of global Ca^{2+} signaling in cells may be attributable to properties of individual Ca^{2+} release channels.

Is this the case for RyR channels in heart cells? One caveat here is that the conditions that produce RyR adaptation in vitro may not exist in vivo. In the heart, the RyR channels operate in local Ca^{2+} microdomains, where even small Ca^{2+} stimuli are thought to elevate local Ca^{2+} concentrations to high levels (Stern, 1992). Additionally, RyR channels in vivo operate in essentially all-or-none clusters (Stern, 1992), where activation of one activates all. Thus, the small incremental Ca^{2+} stimuli, known to induce RyR adaptation in vitro, may not occur in the local control environment of the cell. A second caveat is that termination of local Ca^{2+} release events in cells is ~10 times faster than both RyR adaptation (Györke and Fill, 1993) and inactivation (Schiefer et al., 1995; Sitasapan et al., 1995), as defined in bilayers (Györke and Fill, 1993; Valdivia et al., 1995). Thus, the impact of RyR adaptation and/or inactivation in the heart is not yet clear.

Recently, Sham et al. (1998) extended the findings of Yasui et al., (1994) to the local Ca^{2+} release level. Not surprisingly, considering the caveats described above, they found only weak evidence for quantal/adaptive behavior of local Ca^{2+} release units. They activated a subset of local Ca^{2+} release units and applied a second incremental Ca^{2+} stimulus. The second incremental Ca^{2+} stimulus preferentially activated release units that had not been activated by the first stimulus. Their interpretation was that local Ca^{2+} release is terminated by an inactivation-like mechanism, not an adaptation-like process. However, there is one interesting problem here. Intuitively, the initial small, but sustained, Ca^{2+} stimulus should have continued to recruit (i.e., activate/inactivate) additional release units throughout its duration. In contrast, Sham et al. (1998) showed that the sustained Ca^{2+} stimulus activated/inactivated a subset of release units only at the beginning of the Ca^{2+} stimulus. This rather paradoxical behavior was unexplained. It appears as if local release units were sensi-
itive to the rate of Ca\(^{2+}\) change during the stimulus and not necessarily stimulus amplitude. Such rate-sensing behavior has been considered a hallmark of quantal Ca\(^{2+}\) release behavior (Pizarro et al., 1997). The point is that unconventional behavior is difficult to reconcile with conventional mechanisms. Consequently, the Sham et al. (1998) data clearly illustrate the complexity of local SR Ca\(^{2+}\) release regulation more than they identify the mechanism(s) that may underlie it. Interestingly, Callamaras and Parker (2000) recently showed that such rate sensing, or quantal behavior of local Ca\(^{2+}\) release events may occur during IP\(_3\)-dependent Ca\(^{2+}\) signaling in nonmuscle cells.

At this point, there is an insufficient understanding of the mechanisms that regulate local Ca\(^{2+}\) signaling in heart. It is becoming clear that none of the candidate negative control mechanisms by themselves is sufficient. In efforts to define the mechanisms that underlie complex Ca\(^{2+}\) signaling phenomena in cells, two general themes are developing. The first evolves around defining the extent to which single Ca\(^{2+}\) release channel gating mechanisms govern the complexity of local Ca\(^{2+}\) release events. We believe that defining RyR modal gating dynamics under relatively simple ionic conditions (i.e., no Mg\(^{2+}\) and ATP) is an important first step in this direction. The second evolves around defining how interactions between Ca\(^{2+}\) release channels contribute to the complexity of local Ca\(^{2+}\) signaling in cells. Recently, Stern et al. (1997) has forwarded a local control model of RyR-mediated Ca\(^{2+}\) release that predicts quantal behavior as a collective phenomenon of stochastically interacting RyR channels. This model describes Ca\(^{2+}\) release in the well defined spatial RyR geometry in skeletal muscle and, thus, may not fully apply to other less ordered systems. However, it shows that groups of interacting RyR channels can indeed exhibit collective phenomena (i.e., “group dynamics”) that are qualitatively different from those predicted solely from the properties of individual RyR channels.

Concluding Remarks

In this perspective, we attempt to forward the concept that there will be no simple answers to the complex questions concerning RyR-mediated Ca\(^{2+}\) signaling in the heart. An emphasis on reconciling and/or unifying apparently contradictory data sets is clearly the path that will advance knowledge. In this context, we believe there has been substantial progress. We suggest that Ca\(^{2+}\)-dependent modal gating may be the mechanism that underlies the complex dynamics of single RyR channel Ca\(^{2+}\) regulation, and we have presented a specific kinetic modal gating scheme that reconciles a large body of apparently contradictory results. This scheme suggests that RyR adaptation and inactivation are two distinct manifestations of the same underlying mechanism (i.e., modal gating).

Submitted: 21 August 2000
Revised: 30 October 2000
Accepted: 31 October 2000

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