Regulation of Ca\(^{2+}\) Sparks by Ca\(^{2+}\) and Mg\(^{2+}\) in Mammalian and Amphibian Muscle. An RyR Isoform-specific Role in Excitation–Contraction Coupling?

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Abstract Ca\(^{2+}\) and Mg\(^{2+}\) are important mediators and regulators of intracellular Ca\(^{2+}\) signaling in muscle. The effects of changes of cytosolic [Ca\(^{2+}\)] or [Mg\(^{2+}\)] on elementary Ca\(^{2+}\) release events were determined, as functions of concentration and time, in single fast-twitch permeabilized fibers of rat and frog. Ca\(^{2+}\) sparks were identified and their parameters measured in confocal images of fluo-4 fluorescence. Solutions with different [Ca\(^{2+}\)] or [Mg\(^{2+}\)] were rapidly exchanged while imaging. Faster and spatially homogeneous changes of [Ca\(^{2+}\)] (reaching peaks >100 \(\mu\)M) were achieved by photolyzing Ca NP-EGTA with laser flashes. In both species, incrementing cytosolic [Ca\(^{2+}\)] caused a steady, nearly proportional increase in spark frequency, reversible upon [Ca\(^{2+}\)] reduction. A greater change in spark frequency, usually transient, followed sudden increases in [Ca\(^{2+}\)] after a lag of 100 ms or more. The nonlinearity, lag, and other features of this delayed effect suggest that it requires increase of [Ca\(^{2+}\)] inside the SR. In the frog only, increases in cytosolic [Ca\(^{2+}\)] often resulted, after a lag, in sparks that propagated transversally. An increase in [Mg\(^{2+}\)] caused a fall of spark frequency, but with striking species differences. In the rat, but not the frog, sparks were observed at 4–40 mM [Mg\(^{2+}\)]. Reducing [Mg\(^{2+}\)] below 2 mM, which should enable the RyR channel’s activation (CICR) site to bind Ca\(^{2+}\), caused progressive increase in spark frequency in the frog, but had no effect in the rat. Spark propagation and enhancement by sub-mM Mg\(^{2+}\) are hallmarks of CICR. Their absence in the rat suggests that CICR requires RyR3 para-junctional clusters, present only in the frog. The observed frequency of sparks corresponds to a channel open probability of 10\(^{-8}\) in the frog or 10\(^{-8}\) in the rat. Together with the failure of photorelease to induce activation directly, this indicates a basal inhibition of channels in situ. It is proposed that relief of this inhibition could be the mechanism by which increased SR load increases spark frequency.

Keywords: sarcoplasmic reticulum • excitation–contraction coupling • Ca channels • ryanodine receptors • calcium photorelease

Introduction

After the action potential depolarizes the transverse (T) tubular membrane in muscle, a sequence of events takes place that result in rapid activation of Ca\(^{2+}\) release from the SR, followed by fast and synchronous closing of the release channels and removal of released Ca\(^{2+}\). The speed of the resulting Ca\(^{2+}\) transient’s onset and termination is crucial for a rapid contraction–relaxation cycle (Rome et al., 1996). Although a reasonably complete picture of the major aspects of excitation–contraction coupling has emerged, many aspects, especially those controlling the speed of activation and termination of Ca\(^{2+}\) release, remain to be understood. The presence of different RyR isoforms in variable proportions in mammals and amphibians suggests that mechanisms may be different in these two taxonomic classes. In both, channels of the “skeletal” isoform (RyR1 in mammals, α in amphibians) are located in two parallel rows facing the T tubular membrane. In this membrane, tetrads of dihydropyridine (DHP) receptors face and perhaps make contact with the large cytoplasmic domain of every other RyR, in a precise skipping pattern (Block et al., 1988). In the frog, in addition to functional α RyRs, there are β isoforms, homologous to RyR3 of mammals (for review see Sutko and Airey, 1996; Ogawa et al. 1999). β receptors are located in a parajunctional region close to the triad. Although they are ordered, the geometric arrangement of their clusters is different from that of their junctional counterparts (Felder and Franzini-Armstrong, 2002). Most importantly, they seem to be too far from DHP receptors for any physical interaction with them.

Studies of cell-averaged Ca\(^{2+}\) transients have found differences in the waveform of Ca\(^{2+}\) release in response
to pulse depolarization. In both mammals (rats) and amphibians (frogs), the response consists of an early peak, which is followed after 10–20 ms by a quasi-steady stage. The peak is much more prominent in the frog, and the ratio of amplitudes of these phases (peak/steady) is strongly voltage dependent in the frog, but not the rat (Shirokova et al., 1996).

\( \text{Ca}^{2+} \) release is entirely controlled by plasma membrane voltage, via DHPRs. But the extent to which this control relies on mediation by \( \text{Ca}^{2+} \) itself remains unclear, and appears to be different in these two taxonomic classes. As early as 1988, the peak of \( \text{Ca}^{2+} \) release was proposed to reflect CICR, activation of channels by \( \text{Ca}^{2+} \) (Ríos and Pizarro, 1988). While tests of this idea have been inconclusive (compare Jacquemond et al., 1991; Csernoch et al., 1993; Jong et al., 1993; Pape et al., 1993; Brum et al., 2003), it has the potential to rationalize the functional differences in terms of diversity of structure and isoform endowment. If indeed the frog’s large peak of \( \text{Ca}^{2+} \) release (and the voltage dependence of the peak/steady ratio) were a manifestation of CICR, then it would follow that parajunctional \( \beta \) receptors are the effectors of CICR.

Additional relevant data come from the study of local release events. In amphibian skeletal muscle, \( \text{Ca}^{2+} \) sparks constitute the elementary form of \( \text{Ca}^{2+} \) release, and they are believed to require CICR (Klein et al., 1996). In adult rat fibers, spontaneous \( \text{Ca}^{2+} \) sparks can be observed but at a much lower rate and under special conditions, including most effectively the chemical or mechanical disruption of the plasmalemma (Kirsch et al., 2001; Zhou et al., 2003a). The latter study demonstrated significant differences in the morphology of sparks of rats and frogs, which were attributed to differences in the properties of the release sources. While in the rat sources seem to be elongated parallel to the triad, which suggests that large portions of the junctional channel arrays fire together, in the frog the sources remain largely below the spatial resolution of the microscope (Zhou et al., 2003a). Sources seem to be not just distinct in structure but respond differently to the voltage stimulus. Csernoch et al. (2004) recently reported that voltage clamp pulses, which in frog elicit \( \text{Ca}^{2+} \) release constituted by sparks, in the rat only elicited “embers,” caused by the opening of individual channels. Ward et al. (2001) studied sparks in myotubes of the dyspedic 1B5 cell line. While myotubes expressing RyR1 showed almost no spontaneous release events, those expressing the RyR3 isoform had frequent spontaneous sparks. The sum of the structural and functional information indicates that different activation mechanisms are operating in mammalian and amphibians.

\( \text{Mg}^{2+} \) operates as a crucial brake on \( \text{Ca}^{2+} \) release by multiple mechanisms, including competition at channel sites where \( \text{Ca}^{2+} \) binding causes activation, binding at inhibitory sites, and partial block of \( \text{Ca}^{2+} \) current within the permeation pathway (for review see Meissner, 1994). In fact, the inhibitory effects of \( \text{Mg}^{2+} \) were so prominent in subcellular preparations that M. Endo used the presence of \( \text{Mg}^{2+} \) in the cytosol as basis for his influential dismissal of CICR as a contributor to physiological \( \text{Ca}^{2+} \) release (most forcefully presented in Endo, 1977). The interest in \( \text{Mg}^{2+} \) only increased upon the finding by Kirsch et al. (2001) that the frequency of spontaneous sparks in the mammal did not increase upon reduction of [\( \text{Mg}^{2+} \)\(_{\text{cyto}} \)] below 1 mM, in startling contrast with observations of Lacampagne et al. (1998) in frog muscle.

To further understand the control mechanisms, we compared the modulation of \( \text{Ca}^{2+} \) release by cytosolic \( \text{Ca}^{2+} \) and \( \text{Mg}^{2+} \) in these species. Although the effects of \( \text{Ca}^{2+} \) were qualitatively similar in rat and frog, a quantitative study revealed substantial differences. More strikingly, and in agreement with the result of Kirsch et al. (2001), the [\( \text{Mg}^{2+} \)] dependence of \( \text{Ca}^{2+} \) release was found to differ greatly, which complements the emerging picture of functional diversity in these taxonomic classes. Additionally, aspects of the \( \text{Ca}^{2+} \) effects suggest that intra-SR \( \text{Ca}^{2+} \) may play a role in release modulation as robust as the one it serves in cardiac muscle (Györke, et al., 2002). The present results have been communicated preliminarily (Zhou et al., 2002; Brum et al., 2004).

**Materials and Methods**

Experiments were performed in singly dissected fiber segments from the semitendinosus muscle of *Rana pipiens* or in fiber segments separated enzymatically from the extensor digitorum longus (EDL) muscle of the rat (*Rattus norvegicus*, Sprague-Dawley) as described in detail in González and Ríos (1993) and Zhou et al. (2003a). In brief, frogs were deeply anaesthetized and killed by decapitation followed by pithing, a procedure approved by the Institutional Animal Care and Use Committee of Rush University. Fibers were dissected in a relaxing solution (Kovacs et al., 1983), fixed to the glass bottom of a 100-μl Lucite chamber, and moderately stretched to sarcomere lengths of 2.5–3.2 μm. 5-mo-old male rats were killed by CO₂ inhalation, and the EDL muscles of both legs were dissected and incubated for 1 h at 37°C in a modified Krebs solution with 2 mg/ml collageenate type I and 10% FBS, without added Ca²⁺ or Mg²⁺. Dystrophic muscles were washed in Krebs solution plus FBS; after 30 min, fiber segments of ~1 cm were removed gently and fixed slightly stretched (2.2–2.5 μm per sarcomere) to the glass bottom of the Lucite chamber. On the stage of the microscope, fibers were permeabilized by replacing the relaxing solution with a glutamate-based internal solution containing 0.002% saponin and 50 μM fluo-3; nominal [\( \text{Mg}^{2+} \)] was 1 mM for frog or 2 mM for rat. After 1–2 min, membrane permeabilization was complete, as judged from the increase in fluorescence within the fiber observed with the confocal microscope (Launikonis and Stephenson, 1997). At this time, saponization was stopped by replacing the solution in the chamber with the corresponding saponin-free internal solution.
Table I

Composition of Internal Solutions

|        | Cs Glutamate$^a$ or K$_2$SO$_4$$^b$ | Ca$_t$ | [Ca$^{2+}$] | Mg$_t$ | [Mg$^{2+}$] |
|--------|-----------------------------------|--------|-------------|--------|-------------|
| **Frog** | Varied [Ca$^{2+}$] | 100 | 0.177 | 100 | 4.51 | 0.3 |
|         |                                   | 100 | 0.304 | 200 | 4.51 | 0.3 |
|         |                                   | 100 | 0.474 | 400 | 4.51 | 0.3 |
|         |                                   | 100 | 0.659 | 800 | 4.51 | 0.3 |
| **Rat**  | Varied [Mg$^{2+}$] | 100 | 0.177 | 100 | 4.96 | 0.4 |
|         |                                   | 100 | 0.176 | 100 | 5.34 | 0.5 |
|         |                                   | 100 | 0.174 | 100 | 6.24 | 0.8 |
|         |                                   | 100 | 0.172 | 100 | 7.84 | 1.45 |
|         | Varied [Mg$^{2+}$] | 80 | 0.178 | 100 | 9.80 | 2.0 |
|         |                                   | 80 | 0.306 | 200 | 9.79 | 2.0 |
|         |                                   | 80 | 0.403 | 300 | 9.78 | 2.0 |
|         |                                   | 80 | 0.443 | 350 | 9.78 | 2.0 |
|         |                                   | 80 | 0.479 | 400 | 9.78 | 2.0 |
|         | Varied [Mg$^{2+}$] | 80 | 0.189 | 100 | 3.99 | 0.2 |
|         |                                   | 80 | 0.186 | 100 | 5.53 | 0.5 |
|         |                                   | 80 | 0.183 | 100 | 7.13 | 1.0 |
|         |                                   | 73 | 0.168 | 100 | 14.3 | 4.0 |
|         |                                   | 60 | 0.146 | 100 | 25.2 | 10.0 |
|         |                                   | 28 | 0.091 | 100 | 59.8 | 40.0 |

All internal solutions contained (in mM) HEPES or trismaleate (10), EGTA (1), phosphocreatine (5), ATP (5), glucose (10), dextran 8% and Fluo-4 (0.1). ATP was added as Mg salt up to 5 mM. When low Mg was required, the difference to 5 mM was replaced by the Na salt. pH was adjusted to 7 using CsOH, and osmolality to 265 mOsm/kg for frog or 320 mOsm/kg for rat, using cesium glutamate or potassium sulfate (for frog or rat) or water as needed.

$^a$Frog.

$^b$Rat.

Solutions

The composition of the solutions used to dissect and mount the fibers is the same as described in González et al. (2000) and Zhou et al. (2003a). Internal solutions used were also as described therein but with varying Ca$^{2+}$ and Mg$^{2+}$ concentrations (Table I). The nominal cation concentration is indicated in every experiment described. The actual [Ca$^{2+}$] of the internal solutions was measured with fura-2 as indicator in a spectrofluorimeter. The excitation spectrum of the solution between 320 and 420 nm was fitted to a linear combination of the saturated and Ca$^{2+}$ ter. The excitation spectrum of the solution between 320 and 420 nm was measured with fura-2 as indicator in a spectrofluorimeter (MRC-1000; Bio-Rad Laboratories) equipped with a 40×, 1.2 NA water immersion objective (Carl Zeiss MicroImaging, Inc.). The confocal scanning system (MRG-1000; Bio-Rad Laboratories) used for fluorescence excitation a 100-mW Ar laser attenuated to 3% of its power (Ríos et al., 1999).

Line scan images consist of 512 rows of 768 values of fluorescence intensity $I(x,t)$ at pixel spacing of 0.1428 μm, rows acquired at 2-ms intervals along a line parallel to or perpendicular to the fiber axis. Bleaching was evaluated and corrected as described by Zhou et al. (2003a). The bleaching-corrected version of fluorescence was normalized to the baseline intensity ($I_0(x,t)$), derived as an average of $I(x,0)$ in the nonsparking regions of the line scan image. To avoid photodamage, line scan images were not taken repeatedly at the same location.

Sparks were identified automatically by an objective detection program (Cheng et al., 1999, González et al., 2000), which also determined spark amplitude (maximum of $F/F_0 - 1$), full width at half magnitude (FWHM, or “half width,” defined as the spatial extent at the time of the peak where $F/F_0$ was $>1 + 0.5$ amplitude), and rise time (time lapse of rise between 10% and peak, measured on a 3-pixel central average). Every event detected was included in the statistics, provided that its amplitude was $>0.3$ and its half width $>0.53$ μm.

Solution Changes

In some experiments, Ca$^{2+}$ and Mg$^{2+}$ concentrations were changed by rapid perfusion in a specially designed chamber (Shirokova and Ríos, 1996), using inlet and suction valves electronically switched by a computer that also synchronized confocal image acquisition. The chamber, schematically depicted in the inset of Fig. 6, had a total capacity of ~100 μl. It was covered by a Lucite lid, leaving in and outflow passages at opposing ends. The solution exchange time, determined by the change in fluorescence in a line scan (Fig. 6, inset b), was <100 ms.
**Photorelease of Caged Calcium**

In other experiments, cytosolic Ca$^{2+}$ concentration was changed by flash photolysis of Ca-bound NP-EGTA. The internal solutions contained 1–5 mM NP-EGTA in frog experiments or 0.25–2 mM in rat experiments, with CaCl$_2$ added for a nominal [Ca$^{2+}$] of 100–150 nM, and no EGTA. [CaCl$_2$] was calculated first using parameters of Ellis-Davies et al. (1996), and then adjusted to match the fluorescence of the NP-EGTA solution with the fluorescence of the corresponding EGTA-containing reference solution.

Photolysis was achieved by UV light pulses, "flashes," of adjustable intensity, delivered via a tapered optic fiber (Fiberguide Industries). The 355-nm light pulses generated by a pulsed, frequency-tripled Nd:YAG laser (Quanta-Ray DHP [Spectra-Physics]), for experiments with identifiers MMD001, or Surelute III [Continuum], for ID MMDD03) were synchronized with image acquisition.

Fig. 1 illustrates calibrations of the technique in solution. Shown is a line scan image, acquired at 2 ms per line and 0.1428 μm per pixel, of an internal solution containing 50 μM fluo-3 and 5 mM NP-EGTA with 150 nM nominal [Ca$^{2+}$]. While scanning in progress, one pulse of 355-nm laser light was delivered through a 200-μm diameter optic fiber. The increase in fluorescence was sudden and brief, corresponding to a Ca$^{2+}$ "spike" that is spatially homogeneous within a cylindric region ∼150 μm in diameter in front of the optic fiber. Fig. 1 B is the profile of fluorescence, averaged over the box in the homogeneous region of the image, and its fit (a scaled [CaDye]$_0$) using the model of photolysis is diagrammed in the inset (Ellis-Davies et al., 1996) with parameter values listed in the figure legend.

Fig. 1 C is the evolution of free [Ca$^{2+}$], derived from the fit, together with the time course of [Ca dye]. It can be seen that the concentration briefly reaches 300 μM. Additional simulations included the buffers ATP at 5 mM, 1 mM parvalbumin, and 0.125 mM troponin, with conventional parameters for the reactions of these buffers (given in Zhou et al., 2003a). In this case, the peak [Ca$^{2+}$] was 220 μM and the kinetics of the transient changed only slightly. In experiments using 1 mM NP-EGTA, the peak [Ca$^{2+}$] was ∼70 μM (50 μM with buffers), but the kinetics were similar.

Fig. 1 D shows a simple simulation of the occupancy of a Ca$^{2+}$-binding site with kinetic parameters derived by Schiefer et al. (1995) for the activator site of the RyR, driven by a Ca$^{2+}$ spike like the one calculated for this photorelease event. The occupancy peaks at either 8 or 16%, depending on the concentration of free Mg$^{2+}$. When Ca$^{2+}$ spikes derived from 1 mM NP-EGTA drove the reaction, the occupancy peaked at 5.3 to 8.9%. Similar occupancies were found using parameter values of Zahradňková et al. (1999).

The changes in [Mg$^{2+}$] upon photolysis of NP-EGTA, both release from the cage and displacement of Mg$^{2+}$ by Ca$^{2+}$ from shared ligands, were also calculated (assuming a dissociation constant of 8.99 mM and a diffusion-limited binding rate for the reaction of NP-EGTA and Mg$^{2+}$; Ellis-Davies and Kaplan, 1994). Upon photolysis of 25% of the cage in a solution containing 1 mM NP-EGTA, 150 nM [Ca$^{2+}$], and 0.660 mM [Mg$^{2+}$], the calculation yielded a monotonic increase reaching a maximum of 0.612 mM Mg$^{2+}$, 6 ms after the flash. Based on the small magnitude of this change, any effects of the photolysis should be attributed to Ca$^{2+}$, rather than the Mg$^{2+}$ that it displaces.

An illustration of the technique applied to a frog fiber is in Fig. 2, a raw confocal image of fluorescence in an axon scan. The optic fiber is visible as a crescent at left. The black color represents the absence of fluorescence in the region occupied by the optic fiber. As illustrated in the inset diagram, the crescent shape results from the intersection of the optic fiber, a cylinder at ∼45°, and the horizontal plane of the confocal section, represented by the dashed line. Four flashes were applied at the times indicated in the plot on the right side (white trace). While the increase in fluorescence produced by the first flash was transient, the other flashes produced a sustained elevation because the main Ca$^{2+}$ removal agent, NP-
EGTA, was saturated with Ca\(^{2+}\) after the first flash. Additionally, the level of fluorescence reached inside the muscle fiber during the last flashes was close to saturation of the acquisition system. A rough normalization analysis demonstrated that the [Ca\(^{2+}\)] reached in front of the optic fiber was spatially homogeneous. In the example and almost every case, the fluorescence increased to a greater value inside the cell, but did so proportionally to the resting fluorescence, which was also greater inside due to binding of the Ca\(^{2+}\)-monitoring dye to cellular structures.

In control experiments without NP-EGTA, UV flashes of the highest intensity failed to cause any effect on Ca\(^{2+}\) release activity. When the more efficient Surelite III laser was used in later experiments, some bleaching of the Ca\(^{2+}\) monitoring dye was observed after high intensity pulses.

**Simulation of Ca\(^{2+}\) Sparks**

Computations in discussion require the “sampled volume” in a line scan: volume of a region close enough to the scanned line for an average spark originating there to be imaged with amplitude greater than the detection threshold. For this purpose, it was necessary to simulate sparks, and then simulate their scanning. Ca\(^{2+}\) sparks were simulated by solving diffusion–reaction equations of a model that included a discrete Ca\(^{2+}\) source, of known geometry and Ca\(^{2+}\) current, and a homogeneous surrounding space (a cytosol), with known concentrations of fixed or diffusing ligands. A detailed description of the model is in Zhou et al. (2005a). The parameter values used are described there as “model 1B.” While other parameter sets, explored in that paper, result in sparks of different sizes, the magnitude sought by the simulation here, which is the maximum distance between scan line and source of a detectable spark, was largely insensitive to model parameters. The simulation yields a “spark” (illustrated in Fig. 11 of Zhou et al., 2005a), which is the fluorescence associated with the calculated concentration of Ca\(^{2+}\)-bound fluo-4 at all values of the spatial coordinates and time. The spark is then imaged; that is, line scans at successive points in time are calculated by convolution with the point spread function of the imaging system, shifted according to the position of the source relative to the line of scanning. The measure of inter-
est, in this case amplitude, is made on the image formed by stacked line scans.

RESULTS

This section first presents results from experiments designed to study the role of cytosolic Ca\textsuperscript{2+} in shaping Ca\textsuperscript{2+} release events. This was explored by rapid solution exchanges or by Ca\textsuperscript{2+} photorelease in amphibian and mammalian muscle. A second set of results describes the effects of modifying [Mg\textsuperscript{2+}] by rapid solution changes, in both frogs and rats and under similar conditions.

Cytosolic Ca\textsuperscript{2+} Modulates Spark Frequency

Ca\textsuperscript{2+} sparks were affected by changes in cytosolic Ca\textsuperscript{2+} concentration. This was studied on spontaneous sparks of frog and rat fibers. Measurements were performed on single fibers with the plasmalemma made permeable by brief exposure to saponin. Experiments were started in a solution containing 100 nM [Ca\textsuperscript{2+}]. After recording two to four sets of 20–30 line scan images, the solution was changed rapidly to a different [Ca\textsuperscript{2+}] and more sets of images were acquired. Representative line scan images from a frog fiber are presented in Fig. 3. A–D correspond, respectively, to 100, 200, 400, and 800 nM [Ca\textsuperscript{2+}]. Images were acquired in the same cell in the order A, D, B. C is from a different cell of the same muscle.

The most obvious effect of the [Ca\textsuperscript{2+}] modification was a change in event frequency. A view of magnitude and time course of the effect is represented for frog and rat, diary plot-style, in Fig. 4. The number of events per frame was plotted (normalized to 1 s and 100 \(\mu\)m of scanned fiber length) as a function of time of image acquisition. Fig. 4 A shows that in frogs, spark frequency changed markedly upon solution changes. The effect was sometimes complex, with a rapid response followed by slower trends, and it was generally reversible. Fig. 4 B plots the [Ca\textsuperscript{2+}] dependence of frequency in nine experiments. Frequency increased in a linear manner with [Ca\textsuperscript{2+}]. However, while the dependence was as described up to 400 nM, at 800 nM [Ca\textsuperscript{2+}], the response was variable. When the high [Ca\textsuperscript{2+}] was reached after several solution changes, spark frequency was reduced. On the other hand, when the fiber was exposed to this concentration at early times during the experiment, the effect was a clear increase in spark frequency. The value shown in the graph is an average of frequencies observed during early exposures only.

A large number of events was detected in these experiments, giving us the opportunity to explore in detail the effect of [Ca\textsuperscript{2+}], on morphology of Ca\textsuperscript{2+} sparks. Table II summarizes the spark parameters at different cytosolic [Ca\textsuperscript{2+}], and a graphic view is in Fig. 5.
As already suggested by the images shown, amplitude of events (relative to resting fluorescence) decreased monotonically with $[\text{Ca}^{2+}]_{\text{cyt}}$. In the range of $[\text{Ca}^{2+}]$ explored, 100–800 nM, resting fluorescence was nearly proportional to $[\text{Ca}^{2+}]_{\text{cyt}}$. Thus it was possible to infer a measure of “absolute” amplitude of sparks, by multiplying the relative amplitude and the nominal $[\text{Ca}^{2+}]$ in the solution. Such absolute amplitude, represented by open symbols in Fig. 5 A, increased less than proportionally with cytosolic $[\text{Ca}^{2+}]$.

As shown in Fig. 5, the other morphologic parameters were remarkably insensitive to changes in $[\text{Ca}^{2+}]_{\text{cyt}}$, with spatial width unchanged, and rise time perhaps increased by 1 ms. This stability of morphology indicates that the changes in amplitude must have been due to changes in unitary $\text{Ca}^{2+}$ current, rather than numbers of channels or the time course of their open probability.

While the general approach was similar in mammalian skeletal muscle fibers to that followed in the frog muscle experiments, the lower frequency of spontaneous events dictated that all experiments be performed in sulfate-based internal solutions. While this substitution of convenience may weaken the comparison, we believe that the anion change should not significantly change the modulation by cations. Indeed, Zhou et al. (2003a) demonstrated that the morphologic spark parameters barely change when glutamate is replaced by sulfate. Moreover, the effects of the changes in $[\text{Ca}^{2+}]$ were remarkably similar in both species, in spite of the different main anion in the solution. Finally, the observations on modulation by $\text{Mg}^{2+}$, communicated later, are consistent with earlier results by Kirsch et al. (2001) using glutamate as main anion. One additional difference in the protocol applied to mammalian fibers was the omission of 800 nM $\text{Ca}^{2+}$ solutions, which were not tolerated well.

Fig. 4 C is a diary plot of event frequency in successive line scan images of a complete experiment. $[\text{Ca}^{2+}]$ in the perfusing solution, starting at 400 nM, was changed several times, as indicated in the upper trace. The changes resulted in modifications of event frequency, which appeared to follow $[\text{Ca}^{2+}]_{\text{cyt}}$ roughly in the same manner as observed for frog muscle. The plot in Fig. 4 B, which includes data from 15 rat fibers, reproduces the observation made in frog cells, that spark frequency increases with $[\text{Ca}^{2+}]_{\text{cyt}}$. 

Figure 5. Dependence of $\text{Ca}^{2+}$ spark parameters on $[\text{Ca}^{2+}]_{\text{cyt}}$. Data are from all events detected in the experiments listed in Table II (for frogs, A–C) or Table III (rats, D–F). Parameter values were averaged first in individual fibers (including repeated passages at the same $[\text{Ca}^{2+}]_{\text{cyt}}$). Then fiber values were averaged with equal weight. Amplitude is the peak value of $\Delta F/F_0$. Absolute amplitude (open symbols, dashed lines in A and D) was obtained by multiplying amplitude by nominal $[\text{Ca}^{2+}]$. 

(A–C)
The onset of the effects after changes in $[\text{Ca}^{2+}]$ is sought in photorelease of $\text{Ca}^{2+}$ from NP-1. A reproducible observation was an apparent delay in the frequency usually occurred after 1 or 2 s. Moreover, scanning was performed at <10 $\mu$m from the fiber surface, so that the diffusion delays would be minimal. Finally, the changes in these cases seemed disproportionately large compared with the change in $[\text{Ca}^{2+}]$ in the solution. By analogy with the nonlinearity of the effects of intra-SR $\text{Ca}^{2+}$ reported in cardiac muscle (e.g., Shannon et al., 2000), this hints at a locus inside the SR as effector for the delayed increase in frequency.

Upon solution changes, however, we could not rule out significant delays in the actual changes of $[\text{Ca}^{2+}]_{\text{cyto}}$. A faster, more homogeneous change in $[\text{Ca}^{2+}]$ was observed within 100 ms (both in actual experiments and in calibration situations, with no fiber in the chamber, as shown in the inset). In contrast, the change in event frequency was often occurred after 1 or 2 s. Moreover, scanning was performed at <10 $\mu$m from the fiber surface, so that the diffusion delays would be minimal. Finally, the changes in these cases seemed disproportionately large compared with the change in $[\text{Ca}^{2+}]$ in the solution. By analogy with the nonlinearity of the effects of intra-SR $\text{Ca}^{2+}$ reported in cardiac muscle (e.g., Shannon et al., 2000), this hints at a locus inside the SR as effector for the delayed increase in frequency.

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**Frequency of Events Changes After a Lag**

A reproducible observation was an apparent delay in the onset of the effects after changes in $[\text{Ca}^{2+}]$ solution. Fig. 6 represents $xt$ scans obtained during a solution change in a frog (A) or a rat fiber (B). The upper traces schematically depict the time course of $[\text{Ca}^{2+}]$ in the solution. After the small artifact produced by the early flow of solution in the chamber, there is a lag that precedes the increased spark activity. This delay, which was always present, appeared to be unrelated to diffusion or mixing times, because the fluorescence increase expected from the increased $[\text{Ca}^{2+}]$ was observed within 100 ms (both in actual experiments and in calibration situations, with no fiber in the chamber, as shown in the inset). In contrast, the change in event frequency usually occurred after 1 or 2 s. Moreover, scanning was performed at <10 $\mu$m from the fiber surface, so that the diffusion delays would be minimal. Finally, the changes in these cases seemed disproportionately large compared with the change in $[\text{Ca}^{2+}]$ in the solution. By analogy with the nonlinearity of the effects of intra-SR $\text{Ca}^{2+}$ reported in cardiac muscle (e.g., Shannon et al., 2000), this hints at a locus inside the SR as effector for the delayed increase in frequency.

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**Frequency of Events Changes After a Lag**

A reproducible observation was an apparent delay in the onset of the effects after changes in $[\text{Ca}^{2+}]$ solution. Fig. 6 represents $xt$ scans obtained during a solution change in a frog (A) or a rat fiber (B). The upper traces schematically depict the time course of $[\text{Ca}^{2+}]$ in the solution. After the small artifact produced by the early flow of solution in the chamber, there is a lag that precedes the increased spark activity. This delay, which was always present, appeared to be unrelated to diffusion or mixing times, because the fluorescence increase expected from the increased $[\text{Ca}^{2+}]$ was observed within 100 ms (both in actual experiments and in calibration situations, with no fiber in the chamber, as shown in the inset). In contrast, the change in event frequency usually occurred after 1 or 2 s. Moreover, scanning was performed at <10 $\mu$m from the fiber surface, so that the diffusion delays would be minimal. Finally, the changes in these cases seemed disproportionately large compared with the change in $[\text{Ca}^{2+}]$ in the solution. By analogy with the nonlinearity of the effects of intra-SR $\text{Ca}^{2+}$ reported in cardiac muscle (e.g., Shannon et al., 2000), this hints at a locus inside the SR as effector for the delayed increase in frequency.

Upon solution changes, however, we could not rule out significant delays in the actual changes of $[\text{Ca}^{2+}]_{\text{cyto}}$. A faster, more homogeneous change in $[\text{Ca}^{2+}]$ was sought in photorelease of $\text{Ca}^{2+}$ from NP-1.
EGTA previously equilibrated in the cytosol. The photorelease approach also offered the possibility to achieve much greater increases in \([\text{Ca}^{2+}]_{\text{cyto}}\).

**Event Frequency Follows Homogeneous Changes in \([\text{Ca}^{2+}]_{\text{cyto}}\) After a Lag**

The photorelease technique, illustrated in materials and methods, consisted in delivery of high intensity pulses of UV light to a cell immersed in a solution with NP-EGTA, ~80% saturated with calcium. Each flash resulted in an abrupt and fairly homogeneous change in \([\text{Ca}^{2+}]\) within the irradiated fiber volume (Figs. 1 and 2). The presence of NP-EGTA itself resulted in substantial reduction of the frequency of spontaneous sparks. Use of scavengers of free radicals (like ascorbate, at 5 mM) did not change the effect. The inhibition of sparks was reversible upon removal of NP-EGTA, but it precluded use of cage concentrations >1 mM in the frog, or 0.5 mM in the rat for studies of the control of sparks.

Fig. 7A illustrates a line scan synchronized with a UV flash in a frog fiber. Upon application of the flash...
The spatially averaged fluorescence increased abruptly and then decayed to a steady value (B). As shown in materials and methods, this increase corresponds to a [Ca\(^{2+}\)] spike of \(~60\) nM, followed by a steady increase of \(~80\) nM above resting level. During the scanning time before the flash a few events were observed in the image. After the flash, the event frequency increased markedly. Sparks were located by the automatic detection program. A modified image, with markings made by the detector is in Fig. 7 C. The frequency of events detected in successive time bins is plotted in Fig. 7 D. Following the fluorescence transient, there was an interval when no increment in frequency was apparent. This was observed in every application of single flashes in six frog fibers. The delay was typically 80 ms, while that in solution change experiments was 1 s or greater.

Fig. 8 documents the time course of the frequency change in a different cell, diary plot-style, over a more extended time scale. Each bar corresponds to an image acquired at 5-s intervals. The frequency was significantly increased by the flash. Unlike the effect of a solution change, where typically the first image in the new solution had a less-than-complete change in frequency, the effect of the flash was usually maximal in the first image; again, a consequence of the briefer lag. The effect was transient, typically reversing in minutes.

Another aspect of the effect in frog fibers is presented in Fig. 9, where a flash of low intensity was applied. In these cases, the rise in event frequency induced by the flash was rather small, and the delay was greater. This observation indicates that the low frequency interval after a UV flash was not due to inactivation by photoreleased Ca\(^{2+}\).

The lower intensity flashes also allowed for a better separation of post-flash sparks, and better resolution of their morphology. Visual examination showed numerous events like those magnified in insets in the figure, characterized by a slanted “front end.” As first shown for adult muscle by Brum et al. (2000) and for primary myotubes by Shirokova et al. (1999), the slanted front is evidence of physical migration of the Ca\(^{2+}\) source. It is noteworthy that the slant angles are essentially equal in all propagating sparks, consistent with a stereotyped propagation velocity of 156 \(\mu\)m/s.

As stated above, the presence of NP-EGTA inhibited the production of spontaneous Ca\(^{2+}\) sparks. When concentrations >1 or 2 mM were present, no sparks were observed. The effect of photorelease in those cases, though not on sparks, was still dramatic, and provided additional evidence of an intra-SR mechanism. An example using 5 mM NP-EGTA is in Fig. 10. Panel A is an xy scan image of fluorescence. Photorelease by a laser flash, applied at the arrow, had no immediate effects other than the expected, quite homogeneous spike in [Ca\(^{2+}\)]\(_{cyt}\). This spike, which as estimated in materials and methods peaked at \(~300\) \(\mu\)M, caused no immedi-
ate effects. After a lag of 250 ms, SR Ca\(^{2+}\) release started and propagated rapidly across the fiber. That this response required the increase in SR load was confirmed by the failure to elicit response by an identical flash, applied 60 s later (Fig. 10 B). Recovery from depletion requires 3 min or longer in frog fibers at room temperature (Pizarro and Ríos, 2004); hence the second flash found a partially depleted SR. Fig. 10 C is a line scan image (normalized to resting fluorescence), obtained 4 min later, with the delayed response now shown as a function of time. It evidences reproducibility of the effect on a rested SR, showing additionally that the response may originate more or less simultaneously at several locations. The activation of other areas may then be locally generated or propagated by CICR.

Similar photorelease experiments were performed in rat fibers. Due to the basal reduction of event frequency by NP-EGTA, cage concentration was limited to 0.5 mM. In the example of Fig. 11, the flash-induced transient did not have an immediate effect, but in the next frame, the number increased. Responses in rat fibers were qualitatively similar to those in the frog, but the magnitude of the changes was extremely variable, always less pronounced than in the frog, and often absent. Thus, while individual experiments like the one shown leave no doubt of the existence of an effect, the aggregate result of five flashes applied in the same fiber, shown in Fig. 11 B as an average diary plot of frequency before and after a flash, showed a barely significant change (the average frequency over four images after the flash was not different from the reference in a
two-sided $t$ test; it was significant at a $P < 0.05$ in the one-sided test. As shown in Fig. 8 for the effects in the frog, the increase in frequency appeared to decay rapidly. The difficulty to demonstrate significance may have been due to the use of a lower concentration of NP-EGTA, combined with the rapid restoration of resting frequency. An additional difference between the results in the two species was that propagating sparks were never observed in the rat, before or after the flash. As noted for the frog, even the most intense UV flashes acting on the highest concentrations of NP-EGTA used did not directly induce Ca$^{2+}$ release in rat muscle fibers.

**Changing Cytosolic [Mg$^{2+}$] in Frog Fibers**

[Mg$^{2+}$] in the cytosol was varied in the range 0.2–10 mM through rapid solution exchange. In experiments in frog muscle, all solutions were glutamate based and had 100 nM [Ca$^{2+}$]. Fig. 12 illustrates an experiment in a frog fiber. The diary plot shows that increasing [Mg$^{2+}$] produced a reduction in event frequency and vice versa, in agreement with the observations of Lacampagne et al. (1998). A plot of average frequency versus [Mg$^{2+}$] in several experiments, Fig. 14 (green symbols and lines), shows a monotonic increase as [Mg$^{2+}$] was lowered. The changes were restricted to concentrations <1.45 mM, as the frequency of events decayed precipitously above this range.

**In Rat Fibers, Changing Cytosolic [Mg$^{2+}$] Has Different Effects than in the Frog**

The changes of [Mg$^{2+}$] in the rat, however, had strikingly different effects, depending on the range of concentrations explored. Fig. 13 A illustrates the effects of changing [Mg$^{2+}$] in the 2–10 mM range. To maximize event frequency, in this and all other experiments the nominal [Ca$^{2+}$] was 300 nM. There is clear evidence of concentration dependence of frequency, which is qualitatively similar to that in the frog (i.e., an increase with decreasing [Mg$^{2+}$]). There were important quantitative differences, namely the ability to observe sparks in the rat at concentrations of 4 mM and higher. Because events could be observed even at 40 mM [Mg$^{2+}$], the range 0.2–40 mM was explored thoroughly in several fibers. Each experiment typically focused on two or three concentrations, which were visited repeatedly. The effects were fully reversible and the frequencies were calculated averaging data in all repeated passages. Table IV summarizes the results.

At variance with the frog, when [Mg$^{2+}$] was changed in the rat in the range <2 mM, the event frequency became essentially independent of concentration (Fig. 13 B). The changes in concentration often resulted in a transient increase in frequency, which occurred in the first few images after the change, but then settled to a frequency that was not statistically different from that before the change. Averages and SEM are plotted versus [Mg$^{2+}$] in Fig. 14 (dark red). The continuous lines

![Figure 12](image12.png)

**Figure 12.** Dependence of spark frequency on cytosolic [Mg$^{2+}$] in frog. Frequency in successive images (obtained as described for Fig. 4) vs. time. Dashed trace: nominal [Mg$^{2+}$] in perfusing solution. Image identifier 073003a.

![Figure 13](image13.png)

**Figure 13.** Dependence of spark frequency on cytosolic [Mg$^{2+}$] in the rat. Frequency in successive images vs. time. Dashed trace: nominal [Mg$^{2+}$] in perfusing solution. (A) Changes in a high [Mg$^{2+}$] range. (B) Changes in a low [Mg$^{2+}$] range. Note absence of systematic changes of frequency in B. Image identifiers: (A) 011301d; (B) 011701b.
(a and b) represent fits of functions considered in discussion.

Fig. 15 plots morphologic sparc parameters versus \([\text{Mg}^{2+}/\text{H}^{10001}]\) in the rat. There were only slight changes in morphology as a function of \([\text{Mg}^{2+}/\text{H}^{10001}]\). This is generally consistent with the observation in the frog of essentially invariant morphologic properties (Lacampagne et al., 1998).

**Table IV**

| \([\text{Mg}^{2+}/\text{H}^{10001}]\), mM | 40.0 | 10.0 | 4.0 | 2.0 | 1.0 | 0.5 | 0.2 |
|----------------------------------------|------|------|-----|-----|-----|-----|-----|
| 073001a                                | 0.83 |
| 073101a                                | 2.28 |
| 073101c                                | 3.43 |
| 091101a                                | 0.40 | 1.33 | 5.65|
| 091301b                                | 0.39 | 2.67 | 2.81|
| 091301c                                | 0.25 | 0.75 | 4.71|
| 091301d                                | 0.73 | 5.11 |
| 091701a                                | 3.32 | 2.90 | 1.83|
| 091701b                                | 0.45 | 2.85 | 1.48 | 2.65 |
| 092001a                                | 0.10 | 0.32 |
| 093005a                                | 5.02 | 6.18 |
| 093005b                                | 3.83 | 4.94 |
| 100903a                                | 4.06 | 3.36 |
| Averages                               | 0.10 | 0.32 | 1.19 | 3.84 | 2.22 | 3.99 | 4.3 |
| SEM                                     | 0.04 | 0.40 | 0.45 | 0.41 | 0.67 | 0.95 |

Scans were parallel to the fiber axis. Frequencies are events per second per 100 \(\mu\text{m}\) of fiber. In all fibers, at least one of the concentrations was applied twice. In most cases, several passes were made through two or more concentrations and all images at the same concentration were pooled. Averages are of equally weighted frequencies for individual experiments. Nominal \([\text{Ca}^{2+}/\text{H}^{10001}]\) was in all cases 100 nM.

**Figure 14.** \([\text{Mg}^{2+}/\text{H}^{10001}]\) dependence of event frequency in frogs and rats. Green symbols and lines, frog events. Dark red, rat events. Continuous curves: a, obtained with Eq. 4, \(K_{c_{\text{Ca}}} = 1 \mu\text{M}, K_{c_{\text{Mg}}} = 25 \mu\text{M}, K_{i} = 1 \mu\text{M}, \kappa = 0.025\), and \(N = 39,000\); b, best fit with Eq. 6, \(K_{c_{\text{Ca}}} = 375 \mu\text{M}, K_{i} = 3.1 \mu\text{M}, \kappa = 0.025\), and \(N = 134,000\).

**Figure 15.** Dependence of rat \(\text{Ca}^{2+}\) spark parameters on \([\text{Mg}^{2+}/\text{H}^{10001}]\). Data are from all events detected in the experiments listed in Table IV. Parameter values were averaged first in individual fibers (including repeated passages at the same \([\text{Mg}^{2+}/\text{H}^{10001}]\)), and then fiber values were averaged with equal weight.

**Discussion**

The present study sought to describe the dependence of local events of \(\text{Ca}^{2+}\) release on the cytosolic concentrations of the main ionic modulators of RyRs. The studies, initially designed to look for steady effects, turned out an unexpected temporal dependence of the frequency of events. Striking differences were also found in the modulation of events in mammalian (rat) compared with amphibian (frog) muscle.

**Cytosolic \(\text{Ca}^{2+}\) Had Major Effects on Event Frequency**

In both species studied, the increase in \([\text{Ca}^{2+}/\text{H}^{10001}]\) between 100 and 400 nM resulted in a substantial increase in event frequency. This increase was not distinguishable from a linear function, consistent with a simple, single (noncooperative) action site. The effect on event frequency was quantitatively more important than any changes of morphologic spark parameters. The latter included a decrease in relative amplitude (i.e., \(\Delta F/F_0\)
Frequency of Events Followed Increases in $[Ca^{2+}]_{cyto}$ After a Time Lag

Diary plots of frequency in successive images often showed a trend of slow increase after a change to a greater $[Ca^{2+}]$. When the change in concentration was done while imaging was in progress, a time lag was found between intervention and response. While the lag time between intervention and frequency increase was of 1 or 2 s when the intervention was a change in bathing solution, it was as brief as 80 ms after a Ca$^{2+}$ photorelease-leasing flash. When low intensity flashes were applied, the latency for the increase in frequency became greater, indicating an inverse relationship between lag time and the magnitude of the Ca$^{2+}$ concentration change.

A simple interpretation of lag and its relationship to magnitude of the change in $[Ca^{2+}]_{cyto}$ is that the effect requires an increase in free calcium inside the SR, $[Ca^{2+}]_{SR}$, or total SR calcium, $C_{SR}$. An alternative hypothesis, that the lag is due to Ca$^{2+}$-dependent inhibition of RyRs, cannot explain the inverse relationship between lag and magnitude of the concentration change.

Likewise, the slow trends in frequency observed after a solution change may reflect slow changes in the SR Ca$^{2+}$ content. Immediate and reversible changes, which were also seen, may instead reflect $[Ca^{2+}]$ acting on the cytosolic side of the channels.

Studies of cardiac muscle have provided evidence of control of Ca$^{2+}$ release by luminal SR calcium (for review see Györke, et al., 2002). These effects feature a highly supralinear dependence on $C_{SR}$, suggesting threshold phenomena of high cooperativity (Lukyanenko and Györke 1999; Shannon et al., 2000; Trafford et al., 2000). If the increase in spark frequency observed in the present work after a photorelease spike is due to the increase in $C_{SR}$, then it must also have a highly supralinear dependence on concentration. Indeed, $C_{SR}$ probably rises slowly during the latency interval after cytosolic $[Ca^{2+}]$ increase, but the event frequency increase is sudden and substantial. Therefore, the postulated intra-SR regulation is consistent with the well-established nonlinearity of the homologous mechanism in cardiac muscle.

While the present study was mainly focused on the effects of $[Ca^{2+}]_{cyto}$ on Ca$^{2+}$ sparks, we also made a limited exploration of global effects of photorelease, when higher cage concentrations prevented the occurrence of spontaneous Ca$^{2+}$ sparks. The results in those cases were qualitatively similar: no immediate response to the Ca$^{2+}$ spike, followed by release from the SR after a lag. This response had highly nonlinear properties, a sudden massive response, clearly self-sustained and reinforced by propagation, demonstrated in Fig. 10. It was also dependent on the SR load (or overload), as evidenced by the strict requirement of several minutes of recovery between repeated responses.

As mentioned, the intra-SR actions of Ca$^{2+}$ demonstrated in cardiac myocytes, and now putatively here, have supralinear properties; they are threshold like, apparently related to high powers of $[Ca^{2+}]_{SR}$. This may reflect multiple cooperative binding sites of Ca$^{2+}$ inside the SR lumen, consistent with a proposed regulatory role of calsequestrin (Ikemoto et al., 1989; Hidalgo and Donoso, 1995; Györke et al., 2004) discussed below. Alternatively, an interaction of sites may be at work, whereby intraluminal Ca$^{2+}$ increases both $p_{o}$ of RyRs and their unitary current. Such dually increased “leak” feeds back on the (cytosolic) activation site of the channels and becomes self-sustaining at some point, thus explaining disproportionate all-or-none responses.

While Ca$^{2+}$ sparks are believed to require CICR, and the response to photoreleased $[Ca^{2+}]_{cyto}$ also appears to involve CICR, the fact that large, propagated responses with CICR characteristics can be elicited without sparks suggests a separation of these two phenomena. Sparks apparently have additional requirements, and can therefore be inhibited separately in cells whose RyRs remain robustly activatable by calcium. Zhou et al. (2003a) showed that Ca$^{2+}$ sparks of amphibian and mammalian muscle are morphologically different, and suggested that the latter may not require CICR. We have also communicated preliminary evidence of two mechanisms for the mutual interactions of Ca$^{2+}$ release channels required in the production of sparks in the frog (Zhou et al., 2003b). Only one of these would involve CICR, while the other would be similar to the allosteric coupling postulated to occur between channels in bilayers (Marx et al., 1998).

In summary, the consequences of changing cytosolic Ca$^{2+}$ are complex, with features of direct cytosolic side effects, secondary luminal actions, and positive feedback. To better separate these interlocked mechanisms, we are implementing a new technique that affords direct measurement of $[Ca^{2+}]_{SR}$ (Zhou et al., 2004). Without such measurements, any dissection of luminal and cytosolic mechanisms will remain incomplete.

Large Increases in $[Ca^{2+}]_{cyto}$ Did Not Directly Cause Ca$^{2+}$ Release

As illustrated in Fig. 1, peak $[Ca^{2+}]_{cyto}$ upon photorelease of caged Ca$^{2+}$ probably reached near 300 μM when 5 mM NP-EGTA was present, and 70 μM with 1 mM NP-EGTA. These large spikes never caused an immediate response, while always eliciting an increase in frequency after a lag. Fast steps of cis Ca$^{2+}$ are known to open channels reconstituted in bilayers (e.g., Zahradníková et al.,

422 Regulation of Ca$^{2+}$ Sparks by Ca$^{2+}$ and Mg$^{2+}$
1999; 2003; Zoghbi et al., 2004). The failure to observe an immediate effect here might reflect a basal inhibition of RyRs in situ. We calculated numerically the occupancy of a Ca$^{2+}$ binding site exposed to spikes of photoreleased Ca$^{2+}$, using for the site kinetic parameters defined in bilayers. The calculation illustrated in Fig. 1 D used parameters of Schiefer et al. (1995) and yielded peak occupancies ranging from 8 to 16% (depending on [Mg$^{2+}$]). When 1 mM NP-EGTA was present, the peak occupancy was 3.5–8%. If, as postulated in the bilayer models, Ca$^{2+}$ binding to the site translated directly to channel opening, the frequency of sparks would have increased immediately and enormously (see analysis of relationship between channel opening and spark frequency in next section). Our failure to observe any direct response indicates that RyRs are inhibited in situ, in rats and frogs, by comparison with their properties in reconstituted systems. The conclusion that CICR would require extremely high [Ca$^{2+}$]$_{cyto}$ under physiological conditions was first reached by Endo (1975), who predicted that a concentration of at least 300 μM would be necessary. We now show that not even those levels are capable of triggering Ca$^{2+}$ release directly. Consequently, additional inhibitory factors must be contemplated.

Ca$^{2+}$ Spark Frequency Should Reflect Spontaneous Opening of Single Channels

While the effects of changes in cytosolic [Ca$^{2+}$] were similar in rats and frogs, the concentration profiles of the effects of Mg$^{2+}$ were qualitatively different, a fact that may afford new insights into diverse mechanisms that control their Ca$^{2+}$ release channels. To interpret the species differences within a theoretical framework of channel regulation, we used simple models of control, largely based on measurements in bilayer-reconstituted channels, to account for the steady frequency of Ca$^{2+}$ sparks and its changes.

The state of openness of RyRs can be affected by many ligands (for review see Meissner, 1994). Ca$^{2+}$ is known to exert its effect mainly by binding to two sites, named I and II by Laver et al. (1997), on cytosolic domains of the RyR. In the discussion that follows, and in the diagrams of Fig. 16, these sites are named instead “a” and “i”. Binding to site a, with dissociation constant of 1 μM in mammalian muscle, activates the channels (an action underpinning the CICR phenomenon). Binding to site i, with dissociation constant near 1 mM (Meissner et al., 1986), puts the channels in a closed state with features of inactivation (in this state the channel cannot be opened by binding to its activation site). Mg$^{2+}$ may bind to both sites, with dissociation constant ~30 μM at site a and one indistinguishable from that of Ca$^{2+}$ at site i (Laver et al., 1997). Bound to either site, Mg$^{2+}$ is inhibitory; at site a preventing activation by Ca$^{2+}$, and at site i causing inactivation.

**Binding at a High Affinity Site Explains the Effect of Mg$^{2+}$ on Frequency in the Frog**

The effect of reducing [Mg$^{2+}$] on spontaneous Ca$^{2+}$ sparks of frog muscle can be accounted for quantitatively on the basis of the interactions described above. The approach here generalizes one of Laver et al. (1997) introduced to interpret effects on unitary currents of channels in bilayers. Upon simplification, it is reducible to that used by Lacampagne et al. (1998) to explain the effect of [Mg$^{2+}$] on frequency of Ca$^{2+}$ sparks in the frog.

![Scheme 1](attachment://Scheme_1.png)

The assumed interactions of the RyR with Ca$^{2+}$ and Mg$^{2+}$ are represented in state Scheme 1. C, O, and I represent states of the channel. Because the analysis is only concerned with steady probabilities, states that are theoretically possible but populated very seldom (like O with no bound Ca$^{2+}$) are omitted. Therefore, all transitions require ion binding or unbinding, and all dissociation constants have dimensions of concentration. At the cytosolic [Ca$^{2+}$] used in the present experiments, Scheme 1 reduces to Scheme 2. The relevant dissociation constants are represented as $K$, with self-explanatory sub-indices.

![Scheme 2](attachment://Scheme_2.png)

Because it is not necessary in the argument that follows, these diagrams do not assume cooperativity. A simple cooperativity is introduced in Scheme 3 to interpret the rat data.

The four equilibrium equations of Scheme 2 and its conservation equation can be solved for the probability (occupancy) $O_Ca$ of the open state (occupancies and concentrations are in italics).

$$O_Ca = \frac{Ca K_{Mg} K_i}{K_{Ca}(Mg^2 + K_{Mg} + K_{Mg} K_i)}.$$  \hspace{1cm} (1)

**Translating Open Channel Probability to Event Frequency**

To compare Eq. 1 with data, it is necessary to relate observed frequency (sparks per second per 100 μm of scan) to $p_o$, which is identical to occupancy. This requires recognizing first that a line scan samples a nearby volume, where there are N channels, randomly gating at open probability $p_o$. 

423  \hspace{1cm} ZHOU ET AL.
N can be derived from either biochemical studies of ryanodine binding or morphometry of EM images. The density of RyRs was reported as 44 pmoles/g of muscle and 77 pmoles/g of muscle, respectively, for frog and rabbit (Anderson et al., 1994). This is $4.4 \times 10^{-11}$ moles/g $\times N_p$, molecules mole$^{-1} \times 4 \times 10^{-12}$ $\mu$m$^{-3}$ g, or 25 channels per cubic micron of muscle. Considering that isoforms 1 and 3 are present in similar densities, the number would be 12.5 channels of each isoform per $\mu$m. The density in the rabbit, used here as estimator for the rat, will be 43 $\mu$m$^{-3}$.

Different estimates are reached through morphometry. We combined the fractional volume of T tubules in frog muscle (0.0025; Mobley and Eisenberg, 1975), with an average sectional area of T tubules of 0.0025 $\mu$m$^2$ (Peachey, 1965) to derive length of T tubules per unit volume (1.25 $\mu$m$^{-2}$). Assuming that 80% of tubule length is junctional (Mobley and Eisenberg, 1975), and that in these regions there are 120 channels of each isoform per micron, a channel density of 120 $\mu$m$^{-3}$ is reached. While there are no comparable numbers for mammalian muscle, it is possible to use images and morphometry of stained T tubules in Franzini-Armstrong et al. (1999) to relate density of RyRs in rats and frogs. In similarly prepared thick sections, the average number of (junctional) channels derived by these authors from measured tubule length in a 0.5 $\mu$m$^3$ volume (selected to contain T tubules in the whole section) was 120 for rat EDL and 70 for frog iliofibularis. Considering that the number of T tubule-containing sections per unit fiber length is double for the rat, the ratio of densities, rat/frog, should be 240/70, or 3.43, which is actually the same as that derived from ryanodine binding measurements (43/1.5, or 3.44). In conclusion, the best estimates of receptor density from morphometry are 120 $\mu$m$^{-3}$ for frog and 412 $\mu$m$^{-3}$ for rat fast-twitch muscle.

The next step is to associate to the line scan a sampled volume, defined so that every spark of average magnitude that originates in it will be detected. In experiments on permeabilized frog fibers equilibrated with solutions of 100 nM [Ca$^{2+}$], the average amplitude of sparks, given in Table II, is 0.97. A basic result of the theory of spark scanning is that the “true amplitude” of sparks will be underestimated by 2–2.5-fold due to off-focus errors (Ríos et al., 2001). This implies that the modal amplitude of sparks, if they were all in focus, would be 1.9–2.4. Using a simulation method described in detail in Zhou et al. (2003a), whereby sparks are produced as a consequence of a known current of Ca$^{2+}$ release exiting a source of specified dimensions, we found that to have amplitude 2.1, a simulated spark, sampled in-focus by our confocal microscope (i.e., its fluorescence convolved with the PSF of our system), required a current of 9 pA flowing for 5 ms. This “modal” or “canonical” spark was then used to determine the sampled volume of a line scan, the volume of a region such that canonical sparks originating there will have in the scan image an amplitude greater than the detection criterion (0.3 $F_0$). For this purpose, the spark’s source was shifted relative to the scan line in the y or the z direction (both perpendicular to x, the scanning coordinate). It was found that a vertical (z) shift of 1.15 $\mu$m and a horizontal (y) shift of 0.90 $\mu$m reduced the scanned amplitude to 0.3; these marks the boundaries of the detection region.

The associated volume has therefore the length of the scan, 100 $\mu$m, and the cross area of an ellipse, of diameters 2.3 and 1.8 $\mu$m, or 325 $\mu$m$^3$. Using the density estimates from morphometry, the number $N$ of channels of each isoform contained in this volume in frog muscle is 39,000 and in the rat 134,000 of isoform 1. The number of openings in 1 s is $N p_o \tau^{-1}$ (where $\tau$ is the mean open duration).

Assume that a single channel opening results in the production of a spark in a fraction $\kappa$ of cases. Hence, the frequency of Ca$^{2+}$ sparks will be

$$f = N p_o \kappa \tau^{-1}.$$  \tag{2}

Assuming that the channel’s open time is equal to the spark’s rise time, i.e., 5 ms,

$$f = 200 N p_0 \kappa.$$  \tag{3}

The probability in Eq. 1 can be substituted in Eq. 3 to obtain the predicted frequency $f$ in line scan images

$$f = 200 N \kappa \frac{Ca K_{Mg} K_i}{K_{Ca} (Mg^2 + K_i Mg + K_{Mg} K_i)}.$$  \tag{4}

While the derivation of Eq. 4 required several approximations, its main virtue, to allow for an unbiased comparison of event frequencies in frogs and rats, should not be affected by its likely errors.

In the Frog, Spark Activity is Controlled by Mg$^{2+}$ at the Activation Site

Eq. 4 with conventional values for the dissociation constants, $K_{Ca} = 1 \mu$M, $K_{Mg} = 25 \mu$M, $K_i = 1 \text{mM}$, and $\kappa =$

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1This figure, derived from stereology of EM images, is substantially different from in vivo estimates by Launikonis and Stephenson (2002). They are used here for consistency, because the information regarding T tubule cross-sectional area was obtained from EM images as well (Peachey, 1965).

3The dwell times of single RyRs opening in situ, equated to the rise times of “q = 1” or single channel sparks, were shown by Wang et al. (2004) to be distributed exponentially, with $\tau = 11.6$ ms, in rat ventricular myocytes. In multichannel sparks, the rise time remained close to this value. By analogy, here we assume the mean open time to be equal to the average rise time, or 5 ms.
0.025, generated curve “a”, which describes well the frog values in Fig. 14. In the interesting range where \( f \) is measurable, the denominator of Eq. 4 is dominated by the first order term, implying that the channels are largely inhibited by \( \text{Mg}^{2+} \) at the activation site.

Because the present frog data are similar to those of Lacampagne et al. (1998), the simple function used in that study

\[
f = \frac{K}{\text{Mg}^n}
\]

also described the data well (with \( n = 1.24 \); unpublished data).

At the low \([\text{Ca}^{2+}]\) used here, the interactions at site \( a \) determine the changes in spark frequency when \([\text{Mg}^{2+}]\) changes. Therefore, the occupancy of site \( i \) is not reflected in the frequency versus \([\text{Mg}^{2+}]\) dependence, and cannot be characterized in the frog studies. No insights on site \( i \) were expected, given the impossibility to elicit sparks when site \( a \) is saturated.

The frog results can therefore be accounted for within existing theory (represented by Eq. 4), provided that an efficiency factor, \( \kappa \), is included. While we introduced \( \kappa \) as an efficiency, a success rate at which channel openings cause sparks, its definition must be revised. Indeed, fluorescence signals from single channel openings should be detectable as “embers” (for example see González et al., 2000; Zhou et al., 2003a), but under the present conditions, we did not observe many embers. Therefore \( \kappa \) should be understood as an ad hoc factor. Its low value, 0.025, underscores the basic mismatch between the activity of channels in the simplified preparations where the properties of the control sites were established, and their activity in situ, which appears to be much lower. A parsimonious speculation, previously made for cardiac muscle (for example see Györke et al., 2004), reconciles this observation with the evidence provided here of control from the SR lumen: perhaps the inhibition manifested in the low \( \kappa \) is relieved by intra-SR \( \text{Ca}^{2+} \). For example, a basal inhibition by calsequestrin could be eliminated by conformational changes associated with an increased \([\text{Ca}^{2+}]_{\text{SR}}\). This and other mechanisms are summarized graphically in Fig. 16.

**A Low Affinity \( \text{Mg}^{2+} \) Site Determines Spark Frequency in the Rat**

The rat data (dark red in Fig. 14) clearly cannot be accounted for by Eq. 4 or its simplified version, Eq. 5. Qualitatively, the frequency ceases to depend on \([\text{Mg}^{2+}]\) at concentrations <1 or 2 mM. This cannot be fitted by Eq. 5, or by Eq. 4 with consensus parameter values.

A straightforward change in the reaction scheme accounted for the difference. This is represented in Scheme 3.

\[\text{OCa} \xrightarrow{K} \text{ICaMg} \xrightarrow{K} \text{IMg} \xrightarrow{K} \text{ICaMg}_2 \xrightarrow{K} \text{ICaMg}_2 \]

Activation is still equated to transition to a bound state \( \text{OCa} \), but there is no more competition by \( \text{Mg}^{2+} \). \( \text{Mg}^{2+} \) binds as before to site \( i \), with low affinity, to inhibit the channel (driving it to states \( \text{IMg} \) or \( \text{ICaMg} \)). Scheme 4, which assumes the possibility of binding of a second \( \text{Mg}^{2+} \), providing cooperativity, improves the description slightly. Solving the system of equations of Scheme 4 for \( \text{OCa} \) and using Eq. 3, the following expression for event frequency is obtained:

\[
f = \frac{200NK}{K_{\text{Ca}}[\text{Mg}^2 + K_{\text{Mg}} + K_i^2] + \text{Ca}[\text{Mg}^2 + K_{\text{Mg}} + K_i^2]}
\]

The continuous curve “b” in Fig. 14 represents a best fit of Eq. 6, with \( K_{\text{Ca}} = 375 \mu\text{M} \), \( K_i = 3.1 \text{mM} \), and \( \kappa = 0.025 \). When the simpler Scheme 3 was used, a slightly worse fit was obtained, with \( K_{\text{Ca}} = 350 \mu\text{M} \) and \( K_i = 2.1 \text{mM} \). The agreement of these values of \( K_i \) with the dissociation constant of the inhibitory site \( i \) (Meissner et al., 1986) identifies \( i \) as the probable \( \text{Mg}^{2+} \) binding site.\

By comparison with the frog, spark activity in the rat was found to be much less sensitive to activation by \( \text{Ca}^{2+} \) (\( K_{\text{Ca}} = 375 \mu\text{M} \) instead of the 1 \muM expected for site \( a \)). Strangely as well, spark activity appears to occur in spite of the occupancy of site \( a \) by \( \text{Mg}^{2+} \). The seemingly inescapable conclusion is that the CICR site is deeply altered in the mammal, with reduced \( \text{Ca}^{2+} \) affinity and inure to \( \text{Mg}^{2+} \).

The conclusion is consistent with a body of evidence (for review see Ríos and Zhou, 2004) of limited or no involvement of CICR in mammalian muscle. The functional evidence includes the paucity of \( \text{Ca}^{2+} \) sparks in rats in conditions that elicit them abundantly in the amphibian (Shirokova et al., 1998), the smaller size in the rat of the peak of \( \text{Ca}^{2+} \) release elicited by voltage clamp (Shirokova et al., 1996), and the absence in the rat of voltage dependence of the ratio peak/steady release flux, which in the frog is steep and appears to reflect CICR. Additionally, there are clear differences in

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\( ^3 \)The parameters \( \kappa \) and \( K_{\text{Ca}} \) are dependent, as at low \([\text{Ca}^{2+}]\) the rhs of Eq. 4 depends on their product only. \( K_i \) instead can be fitted independently of \( \kappa \).
the ability of directly applied Ca$^{2+}$ to induce release in skinned mammalian fibers (for review see Lamb, 2000). Finally, CICR should, under some conditions, result in wave-like propagation of channel opening. Indeed, as reported by Brum et al. (2000) and shown in Fig. 9, Ca$^{2+}$ sparks may propagate in space, doing so at velocities close to those predicted by CICR models (Stern et al., 1997, 1999) and those of the CICR-driven Ca$^{2+}$ waves of cardiac muscle (Lukyanenko and Györke, 1999). After examining thousands of sparks, we did not find propagating ones in rat muscle (Zhou et al., 2003a and the present work).

The functional evidence of differences is consistent with the presence in frog muscle of two isoforms of the RyR. Indeed, the parajunctional arrays of RyR3 or β have been proposed as structural basis of the functional differences and main effectors of CICR (Rios and Zhou, 2004). In this view, these channels would underlie spontaneous and voltage-evoked Ca$^{2+}$ sparks, making a major contribution to the peak of Ca$^{2+}$ release, as well as other manifestations of CICR, including propagating sparks. Because of intrinsic properties or tissue-specific interactions, RyR1 would be much less able to participate in these manifestations of CICR, remaining largely under control of the voltage sensors.

The control actions discussed above are illustrated schematically in Fig. 16 (top). Action I is the excitatory effect of the voltage sensors on the (junctional, RyR1) channels. Action II is the promotion of channel opening by Ca$^{2+}$. III is the inhibition/inactivation by binding to site i. IV is the basal inhibition evidenced here by the low frequency of spontaneous sparks and the failure to respond to photoreleased Ca$^{2+}$, an effect possibly mediated by interactions in situ with intra-SR proteins like calsequestrin and triadin. The promotion of activity by intra-SR Ca$^{2+}$ could be a relief from this inhibition upon binding to sites c. High [Ca$^{2+}$]$_{SR}$ will also increase unitary current, which feeds back on the cytosolic sites upon channel opening. For some of these effects, apparent dissociation constants were derived, and are listed in the bottom panel of Fig. 16. Because high affinity parameters of the activation site were consistent with the properties of the activation site in the frog only, they are attributed to RyR3, present only in the frog. The values derived in the rat experiments are tentatively assigned to RyR1 and, as stated before, are inconsistent with a significant role of CICR in mammalian muscle.

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