Functional Coupling of Ryanodine Receptors to $K_{Ca}$ Channels in Smooth Muscle Cells from Rat Cerebral Arteries

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ABSTRACT The relationship between $Ca^{2+}$ release (“$Ca^{2+}$ sparks”) through ryanodine-sensitive $Ca^{2+}$ release channels in the sarcoplasmic reticulum and $K_{Ca}$ channels was examined in smooth muscle cells from rat cerebral arteries. Whole cell potassium currents at physiological membrane potentials (−40 mV) and intracellular $Ca^{2+}$ were measured simultaneously, using the perforated patch clamp technique and a laser two-dimensional (x–y) scanning confocal microscope and the fluorescent $Ca^{2+}$ indicator, fluo-3. Virtually all (96%) detectable $Ca^{2+}$ sparks were associated with the activation of a spontaneous transient outward current (STOC) through $K_{Ca}$ channels. A small number of sparks (5 of 128) were associated with currents smaller than 6 pA (mean amplitude, 4.7 pA, at −40 mV). Approximately 41% of STOCs occurred without a detectable $Ca^{2+}$ spark. The amplitudes of the $Ca^{2+}$ sparks correlated with the amplitudes of the STOCs (regression coefficient 0.8; $P < 0.05$). The half time of decay of $Ca^{2+}$ sparks (56 ms) was longer than the associated STOCs (9 ms). The mean amplitude of the STOCs, which were associated with $Ca^{2+}$ sparks, was 33 pA at −40 mV. The mean amplitude of the “sparkless” STOCs was smaller, 16 pA. The very significant increase in $K_{Ca}$ channel open probability (>104-fold) during a $Ca^{2+}$ spark is consistent with local $Ca^{2+}$ during a spark being in the order of 1–100 μM. Therefore, the increase in fractional fluorescence ($F/F_o$) measured during a $Ca^{2+}$ spark (mean 2.04 $F/F_o$, or $\sim$310 nM $Ca^{2+}$) appears to significantly underestimate the local $Ca^{2+}$ that activates $K_{Ca}$ channels. These results indicate that the majority of ryanodine receptors that cause $Ca^{2+}$ sparks are functionally coupled to $K_{Ca}$ channels in the surface membrane, providing direct support for the idea that $Ca^{2+}$ sparks cause STOCs.

KEYWORDS: $Ca^{2+}$ sparks • ryanodine receptor • sarcoplasmic reticulum • potassium currents • smooth muscle

INTRODUCTION Cytoplasmic $Ca^{2+}$ ions, which enter the cell’s cytoplasm through $L$-type voltage-dependent channels in the plasma membrane and through $Ca^{2+}$ release channels in the sarcoplasmic reticulum (SR),1 regulate contraction of arterial smooth muscle by activating myosin light chain kinase. Under most circumstances, this activating $Ca^{2+}$ will be broadly (“globally”) distributed throughout the cytoplasm and be sufficiently long lasting for $Ca^{2+}$ to approach equilibrium with myosin light chain kinase and major calcium buffers. Thus, global $Ca^{2+}$ regulates contraction of arterial smooth muscle. However, in the immediate vicinity of open $Ca^{2+}$ channels, “local” $Ca^{2+}$ concentrations are not in equilibrium with $Ca^{2+}$ buffers, and therefore could reach much higher levels than global concentrations (Neher, 1998). Local $Ca^{2+}$ transients (“$Ca^{2+}$ sparks”) caused by opening of ryanodine-sensitive $Ca^{2+}$ release channels (RyR) in the SR have been detected in arterial smooth muscle, using a laser scanning confocal microscope and the fluorescent indicator fluo-3 (Nelson et al., 1995; see also Bonev et al., 1997; Jaggar et al., 1998; Porter et al., 1998). Calcium sparks have been subsequently measured in myocytes from portal vein (Mironneau et al., 1996), airway (Sieck et al., 1997), esophagus (Kirber et al., 1997), and ileum (Gordienko et al., 1998).

We proposed that these calcium sparks serve as local calcium signals to activate $K_{Ca}$ channels in the surface membrane (Nelson et al., 1995; see also Fay, 1995), based on a number of lines of evidence. (a) SR can make intimate contacts with the plasma membrane (Devine, et al., 1972; Somlyo, 1985). (b) Immunostaining indicating that RyRs are distributed along the cell membrane (Gollasch et al., 1998). (c) Calcium sparks occurred close to the cell membrane (Nelson et al., 1995). (d) Calcium sparks had similar kinetics and frequencies as transient $K_{Ca}$ channel currents (Nelson et al., 1995; Bonev et al., 1997; Jaggar et al., 1998; Porter et al., 1998). These latter $K_{Ca}$ currents were originally described by Benham and Bolton (1986) as “spontaneous transient outward currents” (STOCs). (e) Inhibitors of sarcoplasmic reticulum release (e.g., ryanodine) prevented both calcium sparks and STOCs (Nelson et al., 1995).

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‡Abbreviations used in this paper: ES, enzyme solution; $F_o$, baseline fluorescence; RyR, ryanodine receptors; SR, sarcoplasmic reticulum; STOC, spontaneous transient outward current.
Activation of $K_{Ca}$ channels ("a STOC") by a local calcium release event ("a calcium spark") would cause a global hyperpolarization of the membrane potential, closing voltage-dependent calcium channels, and, therefore, feed back to decrease global Ca$^{2+}$ and contraction (see e.g., Brayden and Nelson, 1992; Knot and Nelson, 1998; Knot et al., 1998). A single STOC can cause a very significant membrane potential hyperpolarization ($>$20 mV; Ganitkevich and Isenberg, 1990). In contrast, a single Ca$^{2+}$ spark, based on its spatial spread, would activate $<$1% of a cell's myosin light chain kinase. Therefore, the location (particularly relative to $K_{Ca}$ channels), timing, and extent of such Ca$^{2+}$-release events will determine whether they contribute significantly to global Ca$^{2+}$, leading to contraction, or to local stimulation of $K_{Ca}$ channels, leading to a reduction of that global Ca$^{2+}$. In support of the latter mechanism in arterial smooth muscle, inhibitors of Ca$^{2+}$ sparks or $K_{Ca}$ channels increase in a nonadditive manner global Ca$^{2+}$ of pressurized cerebral arteries by 50 nM, which leads to a 30% vasoconstriction (Knot and Nelson, 1998; Knot et al., 1998).

The goal of this study was to determine the quantitative relationship between Ca$^{2+}$ sparks and transient potassium currents in arterial smooth muscle cells. Local calcium release transients are monitored in two distinct ways: (a) as an optical change in fluorescence as calcium binds to dye molecules in the surrounding cytoplasm, and (b) as $K_{Ca}$ channel currents activated by Ca$^{2+}$. Much like the light and thunder that arise from the electrical discharge of lightning, these two observable events would have individual characteristics, reflecting the nature of the originating transient event. Definitive evidence for causality of local calcium release events activating $K_{Ca}$ channels would come from a quantitative analysis of simultaneous measurements of fluorescence and potassium currents in single myocytes. Furthermore, the previous nonsimultaneous measurements do not address the issue of whether all Ca$^{2+}$ sparks activate nearby $K_{Ca}$ channels, or whether some sparks fail to activate STOCs. The stoichiometry of Ca$^{2+}$ sparks and STOCs has important implications regarding the location of Ca$^{2+}$ spark sites and their role in the regulation of cell function.

**Materials and Methods**

**Cell Isolation**

Sprague-Dawley rats (12–14 wk old) of either sex were killed by peritoneal injection of pentobarbital solution (150 mg/kg). The brain was removed and placed into ice cold oxygenated (95% O$_2$/5% CO$_2$) physiological salt solution containing (mM) 119 NaCl, 4.7 KCl, 1.2 KH$_2$PO$_4$, 1.6 CaCl$_2$, 1.2 MgSO$_4$, 0.0023 EDTA, and 11 glucose, pH 7.4. Cerebral basilar arteries were carefully dissected in ice-cold physiological salt solution and cleaned of surrounding connective tissue. Arteries were then washed with a low Ca$^{2+}$ enzyme solution (ES) containing (mM) 55 NaCl, 80 sodium glutamate, 5.6 KCl, 10 HEPES, 2 MgCl$_2$, 10 glucose, pH 7.3, and then digested with papain and collagenase as follows. The artery was first incubated in ES with the same ionic composition as described above, containing 0.3 mg/ml papain and 1 mg/ml diithioerythritol for 20 min at 37°C, and then transferred to a second incubation in ES with 100 μM CaCl$_2$ containing 1 mg/ml collagenase (types F and H in a 70/30% mixture, respectively) and incubated for 10 min at 37°C. The enzyme digestion was stopped by placing the artery in ice cold low Ca$^{2+}$ ES for 15 min. The digested tissue was triturated with a fire polished glass Pasteur pipette to yield single smooth muscle cells. After trituration the cells were kept on ice and used within the same day.

**Simultaneous Patch Clamp and Fluorescence Recordings**

The isolated myocytes were plated in the recording chamber and loaded with the Ca$^{2+}$-indicator fluo-3 by incubation in ES containing 10 μM fluo-3 acetoxymethylster and 2.5 μg/ml pluronic acid (Molecular Probes, Inc.) in the dark for 30 min at room temperature. The incubation was followed by a 30-min wash in the same low Ca$^{2+}$ buffer. Cells were then washed with physiological Ca$^{2+}$ bathing solution (see below) and used for simultaneous patch clamp and confocal recordings after a stabilizing period of ~10 min. All measurements were made 15–45 min after the stabilizing period. Synchronicity of the current and fluorescence
measurements was achieved by means of a light-emitting diode placed near the recording chamber and switched on and off for 1.8 ms from a D/A output of Digidata board during the acquisition protocol, usually 10 s at ~40 mV. This system allowed us to align the fluorescence and current traces with a precision only limited by the sampling frequency of the electrical recording (0.20 μs/point).

Confocal Microscopy

The cells were scanned with a laser scanning confocal system (OZ; Noran Instruments) hosted by an Indy workstation (Silicon Graphics, Inc.) and Intevision software package. The confocal system is mounted in an inverted Diaphot microscope with a 60× water immersion lens (NA, 1.2; Nikon Inc.). Fluo-3 fluorescence was excited with a krypton/argon laser at 488 nm and emitted light was detected by the confocal detector at wavelengths >515 nm. Images were typically of 48.4 × 50.6 μm (or 220 × 230 pixels) and acquired every 8.33 ms (120 images/s) during 10-s laser exposure triggered from the patch amplifier, and stored on writable CDs for future analysis. The confocal aperture was adjusted to expand the z axis so that recordings included a relatively large fraction of the cell. The measurement depth of our recordings was estimated to be 3.0–3.5 microns (z-axis width at half intensity, using a mirrored cover slip), corresponding to ~50% of the cell volume. Since these isolated cells are thin, the confocal exclusion volume was roughly cubic; i.e., 2.2 × 2.2 × 3 μm (see below). (d) The box size considered for correlation analysis was adequate to screen for extraneous sources of light while maximizing our signal quality. This recording arrangement conveyed several significant advantages for this study. (a) The images measured were derived from a much larger fraction of each cell’s volume (~50%), permitting quantitative comparison with a majority of the STOCs. (b) Variability in spark amplitude was minimized due to variations in the relative z-axis position of the spark and the focal plane. (c) The volume within which quantitative signals were averaged for further analysis was roughly cubic; i.e., 2.2 × 2.2 × 3 μm (see below). (d) The efficiency of light collection was much higher, permitting longer recordings from the same cell with less photodamage.

Electrophysiological Recordings

Potassium currents were measured in the whole cell, perforated patch configuration of the patch clamp technique (Horn and Marty, 1988), using an Axopatch 200A amplifier (Axon Instruments). The bathing solution contained (mM): 134 NaCl, 0 KCl, 1 MgCl₂, 2 CaCl₂, 10 glucose, and 10 HEPES, pH 7.4. To minimize contraction, in some cases the bathing solution also contained 5 μM Wortmannin or 20 μM cytochalasin D. The pipette solution contained (mM) 110 potassium aspartate, 30 KCl, 10 NaCl, 1 MgCl₂, 10 HEPES, and 0.05 EGTA, pH 7.2, and 250 mg/ml amphotericin B. Membrane currents were recorded while holding the cells at a steady membrane potential of ~40 mV. Currents were filtered at 500 Hz and digitized at 1.6 kHz (620 μs/point). Cells were simultaneously scanned for fluorescence changes as indicated above.

Chemicals

Unless otherwise stated, all chemicals used in this study were obtained from Sigma Chemical Co. and Calbiochem-Novabiochem International. All experiments were conducted at room temperature (20°C–22°C).

Data Analysis

To determine the amplitude of the STOCs, analysis was performed off line, using a custom analysis program. The threshold of STOCs was set at three times the single KCa channel amplitude at ~40 mV or at 6 pA. The activity of KCa channels in the absence of Ca²⁺ sparks is very low at ~40 mV (nF, ~10⁻⁵; see Bonev et al., 1997), with the probability of three simultaneous openings being exceedingly low. Image analysis was performed using custom written analysis programs using Interactive Data Language software (Research Systems Inc.). Baseline fluorescence (F₀) was determined by averaging the 30 images of (1,200) with no activity. Ratio images were then constructed and replayed for careful examination to detect active areas where sudden increases in F/F₀ occurred. F/F₀ vs. time traces were further analyzed in Microcal Origin (Microcal Software, Inc.), and represent the averaged F/F₀ from a box region of 2.2 × 2.2 μm centered in the active area of interest to achieve the fastest and sharpest changes. This box size (4.8 μm²) was determined empirically to be the best compromise between temporal and spatial precision of Ca²⁺ sparks and the signal to noise ratio.

Statistical Analysis

Results are expressed as means ± SEM where applicable. All the statistical analysis was made with SigmaStat 2.03 software (Jandel Scientific Software). Spearman rank order correlation test was used for correlation analysis and Mann-Whitney rank sum test for significant differences.

Results

STOCs Were Associated with Ca²⁺ Sparks

To minimize disruption of the cell’s cytoplasm, whole cell current was measured in isolated cerebral artery myocytes, using the perforated patch approach of the patch clamp technique (Hamill et al., 1981; Horn and Marty, 1988). Outward potassium currents (“STOCs”) have previously been shown to be through iberiotoxin-sensitive, large conductance potassium (KCa) channels in cerebral arterial myocytes (Nelson et al., 1995). Intracellular Ca²⁺ was measured using the Ca²⁺-sensitive fluorescent dye (fluo-3) and a Noran Instruments laser scanning confocal microscope, with images obtained every 8.3 ms. Under such conditions, Ca²⁺ sparks (n = 128) and STOCs (n = 208) were measured with an average spark frequency of 0.7 ± 0.1 Hz and an average STOC frequency of 1.4 ± 0.3 Hz at ~40 mV (10 cells from different arteries). Fig. 1 A illustrates the life cycle of a Ca²⁺ spark, which peaks in ~20 ms and decays over ~200 ms. Fig. 1 B illustrates simultaneous recordings of whole cell current (at ~40 mV) (in blue) and fluorescence measurements for two spark sites that were identified in the cell (indicated by green and red boxes and F/F₀ traces). Approximately 2.0 ± 0.3 (Table I) Ca²⁺ spark sites per cell were detected (n = 10). The Ca²⁺ sparks in the red site were relatively large (F/F₀ 2.8 ± 0.4), and the associated STOCs were also relatively large (57 ± 10 pA). This Ca²⁺ spark site generated four sparks over a 10-s recording period. The spark site indicated in green had much smaller sparks (mean 1.5 F/F₀) and much smaller associated STOCs (mean 7.9 pA).
Figure 1. Ca\textsuperscript{2+} sparks generate STOCs in myocytes from rat basilar cerebral artery. (A) Original sequence of two-dimensional confocal images obtained every 8.33 ms of an entire smooth muscle cell (top), followed by subsequent images of the region of interest (dotted box) illustrating the time course of the fractional increase in fluorescence (F/F\textsubscript{0}) and decay of a typical Ca\textsuperscript{2+} spark. The images are color coded as indicated by the bar. (B) Simultaneous STOC (pA) and spark (F/F\textsubscript{0}) measurements, at –40 mV, illustrating temporal association. (Blue) Current, (red and green) the F/F\textsubscript{0} average of the red and green boxes (2.2 μm/side) indicated in A, respectively. The pink bar indicates the segment of the trace illustrated in A.

Figure 2. Almost every Ca\textsuperscript{2+} spark (96%) generates a STOC. Time course of whole cell current and F/F\textsubscript{0} from three different cells. Blue traces represent whole cell currents at −40 mV. F/F\textsubscript{0} traces represent the average from a square box of 2.2 μm/side centered at identified active sites in the cell. One site is displayed in cell 1, one site in cell 2, and four sites in cell 3 (A–D; maximum distance between box centers was 7.3 μm [A and D]; minimum distance between box centers was 2.9 μm [B and C]).
Figs. 1 and 2 illustrate a close temporal association of two very low frequency (<1.5 Hz) events, a Ca$^{2+}$ spark and a STOC. Virtually all (123 of 128, 96%) measured Ca$^{2+}$ sparks were accompanied by a STOC with amplitude >6 pA, occurring within the time required to acquire one image (8.3 ms). Even the smallest number (five) of Ca$^{2+}$ sparks that did not match our criteria for association with a STOC were, in fact, paired with small transient currents (mean amplitude 4.7 pA, Table I), which were below our STOC threshold of 6 pA.

The Ca$^{2+}$ sparks and STOCs occurred virtually simultaneously whenever they were paired. Using the photodiode synchronization method described above, we could precisely align the electrical recordings with our optical signals. The mean time difference between the onset (10% over baseline) of a Ca$^{2+}$ spark and a STOC was $-0.3 \pm 0.4$ ms ($n = 114$), with the negative number indicating that the sparks preceded the STOCs (Table I). For these low frequency events, the chances that such close temporal association could have been coincidental is essentially nil. The mean time difference of the peak of an associated STOC and Ca$^{2+}$ spark was $6.4 \pm 1.0$ ms ($n = 118$), with the peak of the STOC occurring earlier than the peak of the Ca$^{2+}$ spark (Table I). The decay half time of STOCs (9.3 $\pm$ 0.4 ms) was shorter than the decay half time of Ca$^{2+}$ sparks (55.9 $\pm$ 3.6 ms) (Table I). The Ca$^{2+}$ spark decay was well fitted by two exponentials (time constants were: $\tau_1 = 31.5$ ms and $\tau_2 = 274.8$ ms, Table I, with both components contributing similarly, 49 and 51%, respectively), whereas the STOC decay was fitted by one exponential ($\tau_1 = 13.4$ ms, Table I).

These observations indicate the close relationship between Ca$^{2+}$ sparks and STOCs. In addition, several previous lines of evidence suggested that Ca$^{2+}$ sparks cause STOCs, including the observations that inhibitors of Ca$^{2+}$ sparks (ryanodine and thapsigargin) block STOCs, and kinetic similarities between Ca$^{2+}$ sparks and STOCs (Nelson et al., 1995). Furthermore, the close temporal relationship between spark and STOC onset, and the observation that STOCs subsequently peak and decline before sparks, indicate that the majority of Ca$^{2+}$ spark sites are close enough to the cell membrane to activate K$_{\text{Ca}}$ channels quickly, before the full spatial spread of the Ca$^{2+}$-dye complex.

**Relationship between Ca$^{2+}$ Spark and STOC Amplitudes**

One apparent aspect of our recordings is that both the sparks and the STOCs varied in amplitude. Fig. 2 illustrates Ca$^{2+}$ sparks and STOCs from three different cells, with one cell having four spark sites. Fig. 2 as well as Fig. 1 illustrates that STOCs were associated with Ca$^{2+}$ sparks, and that larger amplitude sparks were associated with larger STOCs. Fig. 4 also shows the spread in amplitudes of the observed STOCs, ranging from events just above our threshold of 6 pA to 101 pA.

A causal relationship between the Ca$^{2+}$ release and the generation of STOCs would predict that if Ca$^{2+}$ spark amplitude varied due to differences in the amount of Ca$^{2+}$ locally released from the SR during each spark, then the amplitude of the associated STOC would also vary in relation to the release amount. In contrast, other possible sources of variability (e.g., uneven dye distribution, spark events at different positions in the confocal plane, differences in the distances...
between spark sites, and $K_{Ca}$ channels) would cause uncorrelated amplitudes between the two events. We have therefore studied the quantitative relationship between $Ca^{2+}$ spark and STOC amplitudes. Fig. 3 shows a plot of $Ca^{2+}$ sparks and STOC amplitudes from all the cells used in this study ($n = 10$). The amplitudes of $Ca^{2+}$ sparks and STOCs were strongly correlated (correlation coefficient 0.8). The mean amplitudes of associated $Ca^{2+}$ sparks ($F/F_o$) and STOCs (pA) at $-40$ mV were $2.04 \pm 0.05$ and $33.6 \pm 1.9$ pA, respectively ($n = 123$) (Table I). This correlation provides strong evidence that the common element (amount of $Ca^{2+}$ released) between the two observed events is the factor causing the common variation.

**STOCs that Were Not Associated with a Detectable $Ca^{2+}$ Spark**

Although virtually all $Ca^{2+}$ sparks observed were associated with a STOC (Figs. 1 and 2), we also observed many STOCs (41%) that were not associated with a spark above the detection threshold of $F/F_o > 1.2$ (Fig. 4, Table I). However, as shown in Fig. 4, STOCs that were not associated with $Ca^{2+}$ sparks had smaller amplitudes (mean, $16.3 \pm 1.3$ pA, $n = 83$ STOCs, see Table I) and had faster rise and decay times (Table I). If the amplitude correlation illustrated in Fig. 3 also held for these STOCs, then they would have been associated with small sparks. Since a significant fraction (~50%) of

the cell volume was scanned, it is unlikely that large calcium sparks outside the scan volume would have been missed, but it is likely that small and fast calcium sparks outside the scan volume could have been missed.

**Discussion**

$Ca^{2+}$ Sparks Cause STOCs: Implications for the Location and Nature of $Ca^{2+}$ Spark Sites

This study provides the first quantitative analysis of simultaneous $Ca^{2+}$ spark and potassium current measurements in smooth muscle. Mironneau et al. (1996) provided one simultaneous recording of a spark (line scan mode) and STOC in a portal vein myocyte (Mironneau et al., 1996), without quantitative information. In a preliminary report, evidence that some, but not all, $Ca^{2+}$ sparks cause STOCs in esophageal myocytes was provided (Kirber et al., 1997). Our results indicate that virtually all detectable $Ca^{2+}$ transients are associated with a detectable potassium current in arterial smooth muscle, that these events initiate virtually simultaneously, and that they have closely correlated amplitudes. These results therefore demonstrate a close functional coupling between $Ca^{2+}$ spark sites and $K_{Ca}$ channels in the surface membrane. In support of the close proximity of $Ca^{2+}$ spark sites (i.e., RyR) to the surface membrane, SR elements are within 20 nm of the sarcolemmal membrane (Devine et al., 1972; Somlyo, 1985), and RyR have been shown to colocalize with the plasma membrane (Carrington et al., 1995; Gollasch et al., 1998). Taken together with previous pharmacological evidence, we can conclude that $Ca^{2+}$ sparks cause STOCs in arterial smooth muscle.
Sparkless STOCs

Although we establish here that essentially all detectable Ca\(^{2+}\) sparks cause a STOC in cerebral artery myocytes, many STOCs were observed that were not associated with a measurable Ca\(^{2+}\) spark. These “sparkless” STOCs appeared to result from small and fast Ca\(^{2+}\) sparks in regions of the cell that were not scanned. The sparkless STOCs had a smaller mean amplitude (16 pA) than STOCs associated with Ca\(^{2+}\) sparks (33.6 pA) (Table I). Since the lower amplitude STOCs were generally caused by smaller Ca\(^{2+}\) sparks (Fig. 3), the missed Ca\(^{2+}\) sparks are likely to be small as well. Given that the scanned volume covers only about half the cell, and falls off in a graded way for events outside the optimum recording plane, small events were more likely to be missed. In addition, the optical sampling rate (8.33 ms per frame) also restricts the detection limits in the time domain. We show that the sparkless STOCs were faster (both rise and fall time, Table I) than those that were associated with sparks, indicating that the corresponding Ca\(^{2+}\) transients were also likely to be faster. For such small, fast events, it is increasingly likely that the peak of the transient fell between frames, and was therefore further underestimated. These combined “spatio-temporal filtering” effects of the optical recording apparatus would increase the likelihood that small, rapid spark events outside the recording plane would be missed, giving rise to a population of sparkless STOCs. Although our recording system provides significant advantages over other imaging alternatives, it seems clear that K\(_{Ca}\) channels are far superior to the fluorescent dye fluo-3 in sensing local and rapid calcium changes.

The missed smaller sparks and their associated sparkless STOCs might conceivably represent a completely different source of transient Ca\(^{2+}\). For example, these smaller, faster Ca\(^{2+}\) transients might be caused by Ca\(^{2+}\) entering the cell through L-type Ca\(^{2+}\) channels (Wang et al., 1997). We believe, however, that these events are more likely to belong to the lower end of RyR-mediated Ca\(^{2+}\) release events. This latter hypothesis appears more likely since thapsigargin and ryanodine abolish all the STOC activity in these cells (Nelson et al., 1995).

Ca\(^{2+}\) Spark and STOC Amplitudes Were Correlated

The amplitudes of both the Ca\(^{2+}\) sparks and the STOCs reported here are distributed from just above the detection threshold to 3.6 F/F\(_{0}\) and 101 pA, respectively. A number of factors contribute to distribution of Ca\(^{2+}\) spark amplitudes. First, spark events might vary in z-axis position with respect to the confocal recording plane. Second, the release channel(s) that give rise to the sparks might have variable open duration, thus releasing more or less Ca\(^{2+}\) during each spark. Third, the number and distribution of release channels that participate in each spark might differ from one event or cell region to another. Each of these factors undoubtedly plays a role in our observed distributions. Similarly, there are several reasons why STOC amplitudes might be broadly distributed. Assuming STOCs are caused by local Ca\(^{2+}\) release, the distance of that release from the membrane, the local density of K\(_{Ca}\) channels, the quantity of Ca\(^{2+}\) released, and the spatial spread of the released Ca\(^{2+}\) are all factors that would influence the magnitude of the STOC. We report here, however, a strong correlation between the amplitude of individual spark and STOC events (Fig. 3). This observation reduces the possible explanation to those that would be common to both processes. We therefore expect that the correlation of Ca\(^{2+}\) sparks and STOCs amplitudes reflects a common, underlying variability in the quantity of Ca\(^{2+}\) released during one event.

On the Kinetics of Ca\(^{2+}\) Sparks and STOCs

The rise time of a Ca\(^{2+}\) spark (~20 ms) (Nelson et al., 1995; this study, Table I) may reflect in part the kinetics of fluo-3 (Escobar et al., 1997). The kinetics of a STOC did not appear to be affected by fluo-3 (time to peak in cells without fluo-3, 17 ms [Nelson et al., 1995]; time to peak in cells with fluo-3, 16 ms [this study, Table I]), suggesting that fluo-3 (~50 μM) does not buffer the local Ca\(^{2+}\) that activates the K\(_{Ca}\) channels. This observation is consistent with a very short gap (20 nm) (Devine et al., 1972; see also Neher, 1998 for a discussion of Ca\(^{2+}\) diffusion over short distances) between the SR with Ca\(^{2+}\) spark sites and the surface membrane. The relatively slow rise time of a STOC (10–20 ms) may reflect the activation time of K\(_{Ca}\) channels by Ca\(^{2+}\), and not diffusion (Markwardt and Isenberg, 1992). The rapid decay of STOCs may reflect the diffusion and the steep Ca\(^{2+}\) dependence of K\(_{Ca}\) channels.

The decay of Ca\(^{2+}\) sparks in arterial smooth muscle appears to be biexponential (Table I), and much slower than the decay of the associated STOCs. The faster decay time constant (31 ms) is similar to that measured for Ca\(^{2+}\) sparks in heart muscle (Santana et al., 1997), and consistent with the diffusion of the Ca\(^{2+}\)–fluo-3 complex. The nature of the slower time constant of decay is unknown. Although it is clear that a STOC is associated with a Ca\(^{2+}\) spark, the precise interpretation of timing of the onsets and peaks of STOCs and Ca\(^{2+}\) sparks (Table I) is much less certain. Our precision is limited by a number of factors including image sampling rate (120 Hz) and the kinetics of fluo-3.

Fidelity of Fluo-3 Ca\(^{2+}\) Measurements

A quantitative consideration of the Ca\(^{2+}\) sensitivity of K\(_{Ca}\) channels also supports the idea that Ca\(^{2+}\) spark sites are close to the sarcolemmal membrane containing K\(_{Ca}\).
channels. The whole-cell activity \((n_P)\) of \(K_{Ca}\) channels, in the absence of Ca\(^{2+}\) sparks, has been directly measured using the perforated patch, whole cell configuration of the patch clamp technique, and is \(\sim 10^{-2}\) at 0 mV (Bonev et al., 1997; Porter et al., 1998). Therefore, taking into account the voltage dependence of \(K_{Ca}\) channels (e-fold per 10–20 mV; Carl et al., 1996), activity \((n_{cell}P_{o,base})\) of \(K_{Ca}\) channels in the absence of Ca\(^{2+}\) sparks would be \(\sim 10^{-3}\) at physiological membrane potentials (−40 mV), where \(n_{cell}\) is the total number of functional \(K_{Ca}\) channels in a single cell and \(P_{o,base}\) is the average open probability of these \(n_{cell}\) channels. The activity of \(K_{Ca}\) channels that were activated by a Ca\(^{2+}\) spark or \(n_{spark}P_{o,spark}\) can be determined by dividing the STOC by the unitary current (see Nelson et al., 1995), where \(n_{spark}\) is the number of \(K_{Ca}\) channels that have been significantly activated by a Ca\(^{2+}\) spark and \(P_{o,spark}\) is the mean \(P_o\) of these \(n_{spark}\) channels. \(n_{spark}P_{o,spark}\) was \(\sim 16\) (33/2 pA) at −40 mV for these experiments. Thus, the minimum increase of the open probability \((P_o)\) of \(K_{Ca}\) channels can be determined by dividing \(n_{cell}P_{o,base}\) by \(n_{spark}P_{o,spark}\) (or 16/10\(^{-3}\)).

This calculation shows that channel open probability increased at least 10\(^4\) during a Ca\(^{2+}\) spark, even assuming that the Ca\(^{2+}\) transients were felt equally by all the \(K_{Ca}\) channels in the membrane (i.e., \(n_{cell} = n_{spark}\)). However, since a Ca\(^{2+}\) spark (average size, 13 \(\mu m^2\), Table I) covers \(\sim 1\%\) of the surface membrane (1,300 \(\mu m^2\)), then only 1\% of homogeneously distributed \(K_{Ca}\) channels would be activated by a Ca\(^{2+}\) spark. The \(P_o\) of \(K_{Ca}\) channels near the Ca\(^{2+}\) spark would increase more than 10\(^4\) and closer to 10\(^6\), depending on the distribution of \(K_{Ca}\) channels. For channel \(P_o\) to increase by 10\(^4\)–10\(^6\), local Ca\(^{2+}\) would have to increase by at least 10–1,000-fold, assuming open probability increases to second to fourth power of [Ca\(^{2+}\)] (for review see McManus, 1991; Carl et al., 1996), with the beta subunit of the \(K_{Ca}\) channel increasing Ca\(^{2+}\) sensitivity (Tanaka et al., 1997). Therefore, the local intracellular Ca\(^{2+}\) that activates the \(K_{Ca}\) channels during spark ranges from 1 to 100 \(\mu M\) (assuming 100-nM resting Ca\(^{2+}\)). These estimates of relatively high local Ca\(^{2+}\) in the junctional space between the RyR in the SR and the \(K_{Ca}\) channels are in accord with simulations of local Ca\(^{2+}\) changes during a Ca\(^{2+}\) spark in the junctional space in cardiac muscle (Cannell and Soeller, 1997). In contrast, the mean increase in Ca\(^{2+}\) during a Ca\(^{2+}\) spark, as measured with fluo-3, would predict an increase in \(K_{Ca}\) channel \(P_o\) of no more than 16-fold, assuming [Ca\(^{2+}\)]\(^4\) relationship with channel \(P_o\). These estimates indicate that local Ca\(^{2+}\) sensed by the \(K_{Ca}\) channels is much higher (1–100 \(\mu M\)) than peak of the Ca\(^{2+}\) spark (~200–300 nM range at peak amplitude) measured with fluorescent Ca\(^{2+}\) indicator fluo-3.

In conclusion, our results support the idea that Ca\(^{2+}\) sparks cause STOCs, and that the RyR that generate Ca\(^{2+}\) sparks in cerebral arterial myocytes are located in SR elements that are close enough for local Ca\(^{2+}\) to cause significant activation (10\(^4\)–10\(^6\)-fold increase in open probability) of \(K_{Ca}\) channels in the plasma membrane.

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