“Eventless” InsP$_3$-dependent SR-Ca$^{2+}$ Release Affecting Atrial Ca$^{2+}$ Sparks

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KEY POINT SUMMARY

- Inositol 1,4,5-trisphosphate receptors (InsP$_3$Rs) are functionally expressed in cardiac myocytes.
- Influence of inositol 1,4,5-trisphosphate-induced SR-Ca$^{2+}$ release (IP3ICR) on atrial excitation-contraction coupling under physiological and pathophysiological conditions remains elusive.
- The present study focuses on local IP3ICR and its functional consequences for ryanodine receptor (RyR) activity and subsequent Ca$^{2+}$-induced Ca$^{2+}$ release (CICR) in atrial myocytes.
- Here we show significant SR-Ca$^{2+}$ flux but eventless SR-Ca$^{2+}$ release through InsP$_3$ receptors.
- We suggest a new mechanism based on eventless and highly efficient InsP$_3$-dependent SR-Ca$^{2+}$ flux as a crucial mechanism of functional crosstalk between InsP$_3$Rs and RyRs, which may be an important factor in the modulation of excitation-contraction coupling sensitivity.

Word count: 121
ABSTRACT

Augmented inositol 1,4,5-trisphosphate receptor (InsP₃R) function has been linked to a variety of cardiac pathologies including cardiac arrhythmia. The contribution of inositol 1,4,5-trisphosphate-induced Ca²⁺ release (IP3ICR) in excitation-contraction coupling (ECC) under physiological conditions, as well as under cellular remodeling remain controversial. Here we test the hypothesis that local IP3ICR directly affects ryanodine receptor (RyR) function and subsequent Ca²⁺-induced Ca²⁺ release (CICR) in atrial myocytes. IP3ICR was evoked by UV-flash photolysis of caged InsP₃ under whole-cell configuration of the voltage-clamp technique in atrial myocytes isolated from C57/BL6 mice. Photolytic release of InsP₃ was accompanied by a significant increase in the Ca²⁺ release event frequency (4.14±0.72 vs. 6.20±0.76 events*100µm⁻¹s⁻¹). These individual photolytically triggered Ca²⁺ release events were identified as Ca²⁺ sparks which originated from RyR openings. This was verified by Ca²⁺ spark analysis and pharmacological separation between RyR and InsP₃R-dependent SR-Ca²⁺ release (2-aminoethoxydiphenyl borate, xestospongin C, tetracaine). Significant SR-Ca²⁺ flux but eventless SR-Ca²⁺ release through InsP₃R was characterized using SR-Ca²⁺ leak/SR-Ca²⁺ load measurements. These results strongly support the idea that IP3ICR can effectively modulate RyR openings and Ca²⁺ spark probability. We conclude that eventless and highly efficient InsP₃-dependent SR-Ca²⁺ flux is the main mechanism of functional crosstalk between InsP₃Rs and RyRs, which may be an important factor in the modulation of ECC sensitivity.

ABBREVIATIONS

<table>
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<th>Term</th>
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<tr>
<td>Inositol 1,4,5-trisphosphate-induced Ca²⁺ release</td>
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<td>Inositol 1,4,5-trisphosphate receptor</td>
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<td>Ca²⁺-induced Ca²⁺ release</td>
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<td>Cyclopiazonic acid</td>
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INTRODUCTION

In cardiac myocytes two mechanisms for SR-Ca\(^{2+}\) release have been established, Ca\(^{2+}\)-induced Ca\(^{2+}\) release (CICR) as the central mechanism for excitation-contraction coupling (ECC) and inositol 1,4,5-trisphosphate-induced Ca\(^{2+}\) release (IP3ICR). However, although InsP\(_3\)Rs are functionally expressed in cardiac preparations (Mackenzie et al., 2004; Zima & Blatter, 2004; Domeier et al., 2008; Harzheim et al., 2009), their role under physiological as well as under pathophysiological conditions in ECC and CICR is still a matter of an ongoing debate.

It has been suggested that SR-Ca\(^{2+}\) release via InsP\(_3\)R may effectively modulate ECC in ventricular cardiomyocytes, even though the expression of InsP\(_3\)Rs is only a fraction of RyR expression levels (Lipp et al., 2000; Kockskämper et al., 2008). These aspects gain even more importance in atrial myocytes, where InsP\(_3\)Rs are expressed 3-10 times more compared to ventricle myocytes (Lipp et al., 2000; Domeier et al., 2008). In addition, this hypothesis is supported by two key observations. First, under pathophysiological conditions, during cellular remodeling, InsP\(_3\)R expression is upregulated and may cause delayed after depolarizations (DADs), which lead to arrhythmogenicity (Zima & Blatter, 2004; Bootman et al., 2007). Second, InsP\(_3\)R deficient hearts are largely protected from pro-arrhythmogenic stress (Li et al., 2005). Supporting data that encourage a significant contribution of IP3ICR to ECC and CICR are colocalization studies of RyRs and InsP\(_3\)Rs (Lipp et al., 2000; Mackenzie et al., 2002), the microarchitecture of atrial myocytes (Yamasaki et al., 1997) and functional studies that have been carried out (Mackenzie et al., 2002; Zima & Blatter, 2004; Li et al., 2005).

In atrial myocytes normal Ca\(^{2+}\) transient propagation by concentric Ca\(^{2+}\) waves, which are not pro-arrhythmogenic, may be facilitated by IP3ICR (Mackenzie et al., 2004). Local Ca\(^{2+}\) signaling between InsP\(_3\)Rs / RyRs and vice versa might mediate a functional Ca\(^{2+}\) crosstalk between the two types of SR-Ca\(^{2+}\) release channels. This was conceptualized for smooth muscle cells, where local RyR Ca\(^{2+}\) release events, “Ca\(^{2+}\) sparks”, were activated via CICR subsequent to InsP\(_3\)R activation (Gordienko & Bolton, 2002).

In atrial cells, RyR is the major Ca\(^{2+}\) release channel, organized in clusters and functionally coupled (Niggli & Shirokova, 2007). On a local scale, RyR openings give rise to “Ca\(^{2+}\) sparks”, the building blocks of global Ca\(^{2+}\) transients in cardiac cells (Cheng et al., 1993), while coordinated openings of clustered InsP\(_3\)Rs will form “Ca\(^{2+}\) puffs” with distinct properties (Yao et al., 1995).
Along with microscopic Ca$^{2+}$ release events (Ca$^{2+}$ sparks and Ca$^{2+}$ puffs) eventless Ca$^{2+}$ release has been discovered (“Ca$^{2+}$ quarks”, “Ca$^{2+}$ blips”) (Parker et al., 1996; Lipp et al., 1996; Brochet et al., 2011). However, these largely invisible Ca$^{2+}$ release events and concomitant Ca$^{2+}$ fluxes may have a significant impact on Ca$^{2+}$ signaling in sub-cellular microdomains and may be involved in the functional crosstalk between RyRs and InsP$_3$Rs, which is still enigmatic.

The present study focuses on local IP3ICR and its functional consequences for RyR activity in atrial myocytes. We examined local IP3ICR triggered by photoactivation of caged inositol 1,4,5-trisphosphate (InsP$_3$) in isolated atrial myocytes from mice under whole cell voltage-clamp conditions. This highly specific approach bypasses sarcolemmal membrane receptor activation and subsequent signaling cascades. Surprisingly, we found that although no individual IP3ICR events could be visualized, InsP$_3$ release significantly affected the Ca$^{2+}$ spark probability and foster that RyR Ca$^{2+}$ release can be modulated by Ca$^{2+}$ release via InsP$_3$Rs in intracellular microdomains. This suggested a new mechanism based on highly efficient but “eventless” InsP$_3$-dependent SR-Ca$^{2+}$ release. We conclude that invisible InsP$_3$-dependent SR-Ca$^{2+}$ events are the main mechanism of functional crosstalk between InsP$_3$Rs and RyRs.

MATERIAL AND METHODS

Atrial myocytes from adult male C57/BL6 mice, obtained from the Central Animal Facility, University Hospital, University of Bern, were freshly isolated by Langendorff perfusion technique. Hearts were removed after animals were killed by cervical dislocation. The number of animals (N) and myocytes (n) are given in the figure legends. All experiments were performed at room temperature and approved by the State Veterinary Office of Bern, Switzerland, according to Swiss Federal Animal protection law. Whole-cell voltage clamp was combined with confocal Ca$^{2+}$ imaging and UV-flash photolysis. The pipette solution contained (in mmol/L) 120 CsAsp, 10 HEPES, 20 TEA-Cl, 5 K-ATP, 1 MgCl$_2$, 0.1 K$_5$-fluoro-3 (Biotium), 2 GSH, (0.03, 0.06, 0.24) caged InsP$_3$-6Na (Sichem); pH 7.2 with CsOH. Cells were perfused with external solution containing (in mmol/L) 140 NaCl, 5 HEPES, 1.1 MgCl$_2$, 5.4 KCl, 10 Glucose, 1.8 CaCl, 0.5 BaCl, 1 CsCl; pH 7.4 NaOH. Pharmacological experiments used 5 µM 2-aminoethoxydiphenyl borate (Fluka), 5 µM xestospongin C (A.G Scientific Inc.), 1 mM tetracaine (Sigma) or 10 mM caffeine (Sigma) added to the external solution. For Ca$^{2+}$ leak/load experiments (Shannon et al., 2002) atrial myocytes were loaded with fluo-3 AM (Biotium).
Data are reported as mean ± SEM. Where the data was normally distributed, paired or unpaired, where appropriate, Student’s t-test was applied. For not normally distributed data the Wilcoxon Matched-Pairs Signed-Ranks Test was applied. A detailed Materials and Methods section is available in the Online Data Supplement.

RESULTS

ET-1 induced InsP₃R Ca²⁺ release in atrial myocytes

Figure 1 shows that rapid administration of 100 nM endothelin-1, an InsP₃ pathway activator, caused a significant increase in spontaneous local as well as global Ca²⁺ release events in resting mouse atrial cardiomyocytes. This increase was 2APB (InsP₃R blocker) sensitive (Fig. 1B) and was also absent in control (Fig. 1C). Observations agreed with previously performed experiments from other groups (Mackenzie et al., 2004; Zima & Blatter, 2004; Li et al., 2005), that ET-1 mediated IP3RICR could be obtained in various mammalian species including mouse atrial myocytes. ET-1 triggered IP3ICR is based on a complex signaling cascade which involves G-protein coupled phospholipase C activation followed by diacylglycerol and InsP₃ production. To bypass this complex signal transduction pathway, InsP₃Rs were activated by rapid and transient intracellular InsP₃ elevation induced by photorelease of InsP₃ from caged InsP₃.

InsP₃-induced Ca²⁺ release triggered by UV flash photolysis of caged InsP₃

Atrial myocytes were voltage-clamped in the whole-cell configuration of the patch-clamp technique and held at -80 mV. Caged InsP₃ in combination with the Ca²⁺ indicator fluo-3 were dialyzed into the cell through the patch pipette. A SR-Ca²⁺ loading protocol based on L-type Ca²⁺ current activation elicited by depolarization steps from -80 mV to 0 mV (up to 10 times) was applied to ensure comparable SR-Ca²⁺ content, before line scan images were recorded (Fig. 2A). Under these conditions spontaneous Ca²⁺ event activity of about 3.7±0.92 events*100µm⁻¹s⁻¹ was observed.

In order to evoke IP3ICR in a highly specific manner, global UV-flashes were applied to photorelease InsP₃ in the entire cytosol, which was followed by an increase of local Ca²⁺ event frequency triggered in a dose dependent manner (Fig. 2C). The number of Ca²⁺ release events observed after photorelease of InsP₃ was normalized to the number of spontaneous Ca²⁺ releases counted before InsP₃ uncaging. Caged InsP₃ concentrations of 30 µM and 60 µM caused a slight but not significant increase in the frequency of Ca²⁺ release events (30 µM: 2.73±0.51 to 3.06±0.71 events*100µm⁻¹s⁻¹; 60 µM: 4.03±0.73 to 4.20±0.58 events*100µm⁻¹s⁻¹). However, photolysis of 240
µM caged InsP3 triggered a significant (p=0.02) increase in frequency of Ca2+ release events from 4.14±0.72 to 6.20±0.76 events*100µm^{-1}s^{-1} (Fig. 2B and 2C). It is to be mentioned, that the effective InsP3 concentration after UV-flash application is orders of magnitude lower but in proportion to the loading concentration of the caged compound in the pipette solution (see online data supplement). Figure 2C shows control experiments lacking the caged compound indicating that UV-flash application alone had no effect on the frequency of events. We expected, that at least part of the total number of photolytically triggered Ca2+ events were based on IP3ICR (e.g. Ca2+ “puffs”). Although Ca2+ sparks shows similarities with Ca2+ puffs, it has been shown that Ca2+ puffs exhibit significantly distinct spatio-temporal properties, that are different from Ca2+ sparks (Cheng et al., 1993; Yao et al., 1995; Tovey et al., 2001; Niggli & Shirokova, 2007). At maximal event activity (240 µM caged InsP3), local Ca2+ release events were analyzed in more detail for amplitude, full duration at half-maximum amplitude (FDHM), full width at half-maximum amplitude (FWHM), full duration (fulldur), full width (fullwidth) and rise time (Picht et al., 2007). Cells were divided into two groups: One group showed little (<1.3 events*100µm^{-1}s^{-1}) increase (frequency: 0.98±0.08 events*100µm^{-1}s^{-1}), a second group showed >1.3 events*100µm^{-1}s^{-1} increase in Ca2+ release events (frequency: 3.24±0.55 events*100µm^{-1}s^{-1}) after photolytic InsP3R activation. Ca2+ release events obtained in both groups showed no significant difference in their temporal characteristics. The fullwidth was significantly larger in the group of cells that showed little increase in the frequency of Ca2+ release events, from 3.80±0.29 to 4.51±0.38 µm (Fig. 2D), an indication for contribution of IP3ICR in Ca2+ spark formation. Nevertheless, Ca2+ event analysis did not show the anticipated two classes of local Ca2+ release events (i.e. Ca2+ sparks and Ca2+ puffs). The obtained parameters of Ca2+ events were comparable to those reported for Ca2+ sparks (Niggli & Shirokova, 2007) and are summarized in the supplementary Table 1 in the online material.

However, control measurements revealed photolytically triggered local InsP3 Ca2+ events (Ca2+ puffs) in neonatal rat cardiomyocytes. In other words, experimental conditions can be assumed to be appropriate for detecting microscopic subcellular Ca2+ events including Ca2+ puffs (see Fig. 2S, Supplementary Material). While photorelease of InsP3 lead to an increased Ca2+ spark frequency (Fig. 2), the precise mechanism by which this occurred still remained unclear. Therefore, pharmacological interventions were used to distinguish between Ca2+ sparks and Ca2+ puffs by selective inhibition of either RyRs or InsP3Rs. To inhibit IP3ICR, cells were incubated for at least 20 min in 5 µM Xesto, or 5 µM 2APB was acutely applied. TET (1 mM) was applied to inhibit RyR Ca2+ release (Fig. 3A).
Photolytic increase of Ca\(^{2+}\) release events was inhibited by 2APB and Xesto, without affecting the spontaneous Ca\(^{2+}\) release event activity.

Surprisingly, in the presence of TET a complete inhibition of all Ca\(^{2+}\) release events (Fig. 3B and 3C) was observed. This observation is in full agreement with the Ca\(^{2+}\) event analysis given in Fig. 2, showing that all provoked SR-Ca\(^{2+}\) release events can be classified as Ca\(^{2+}\) sparks. Taken together, photolytically activated IP3ICR efficiently facilitates RyR Ca\(^{2+}\) release.

**SR-Ca\(^{2+}\) load and invisible InsP\(_3\)R Ca\(^{2+}\) release (SR-Ca\(^{2+}\) leak)**

Ca\(^{2+}\) spark probability strongly depends on luminal Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_{SR}\)). RyRs and InsP\(_3\)Rs may share, at least in part, the same luminal Ca\(^{2+}\) pool, suggesting that [Ca\(^{2+}\)]\(_{SR}\) may have a potential role in the functional crosstalk of both Ca\(^{2+}\) release channels, which was examined using caffeine induced Ca\(^{2+}\) transients. InsP\(_3\)R block (Fig. 3D) did not significantly affect the SR-Ca\(^{2+}\) content (2APB: 1±0.18 to 1.09±0.08 a.u., p=0.8; and Xesto: from 1.13±0.07 to 0.78±0.09 a.u., p= 0.07), whereas TET caused a significant increase in the SR-Ca\(^{2+}\) content from 1±0.07 to 1.4±0.1 a.u., p=0.02, presumably by suppressing Ca\(^{2+}\) leak (Shannon et al., 2002). Besides the SR-Ca\(^{2+}\) content, SR-Ca\(^{2+}\) leak is an important determinant for Ca\(^{2+}\) spark frequency. Indirect evidence that InsP\(_3\)R stimulation may play a role in SR-Ca\(^{2+}\) leak came from the observation, that in the presence of TET subsequent InsP\(_3\) release caused a slow but significant increase in background fluorescence, which was not seen in the presence of 2APB (Fig. 3E). This result suggests that “eventless” Ca\(^{2+}\) flux through the InsP\(_3\)R, which cannot be resolved as local Ca\(^{2+}\) release events, may be more pronounced in atrial cardiomyocytes than previously expected.

In Figure 4 we addressed the question whether InsP\(_3\)Rs may contribute to the SR-Ca\(^{2+}\) leak and thus affect the Ca\(^{2+}\) spark appearance by performing a modified SR-Ca\(^{2+}\) leak protocol introduced by Shannon et al. (Shannon et al., 2002). Atrial myocytes were field stimulated for 45 seconds at 1 Hz, followed by a rapid solution switch from 1.8 mM extracellular Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_{o}\)) to a nominally [Ca\(^{2+}\)]\(_{o}\) free and [Na\(^{+}\)]\(_{o}\) free solution. After 10s 10 µM cyclosporin A (CPA, a potent SERCA blocker) was added and finally, 30 s later, the SR-Ca\(^{2+}\) content was measured with rapid caffeine application (Fig. 4A). Atrial myocytes were divided into two groups. The control group was recorded under conditions given above, the second group was pre-incubated with 100 nM ET-1 for at least 120 s before CPA was applied (Fig. 4A). CPA causes an inhibition of the SERCA function. Hence, in combination with the blocked Na\(^{+}/Ca^{2+}\) exchanger (NCX) function, any Ca\(^{2+}\) leaking from the SR is trapped in the cytoplasm and mirrored by an increase in the cytosolic Ca\(^{2+}\) concentration [Ca\(^{2+}\)]. This increase was measured and defined as SR-Ca\(^{2+}\) leak.
When CPA was added in the presence of ET-1 there was a significant increase in SR-Ca\(^{2+}\) leak (Δ[Ca\(^{2+}\)]\(_i\)) compared to the leak in control (0.16±0.03 to 0.38±0.09 ΔF/F\(_0\), p=0.02). The SR-Ca\(^{2+}\) leak/SR-Ca\(^{2+}\) load relationship increased accordingly, although not significantly (Δ[Ca\(^{2+}\)]/SR-Ca\(^{2+}\) load: 0.047±0.007 to 0.086±0.020, p=0.09). This may be due to the additional Ca\(^{2+}\) influx induced by ET-1 (He et al., 2000) causing an increased SR-Ca\(^{2+}\) content (3.30±0.24 to 4.53±0.26 F/F\(_0\), p=0.001). However, reduced SERCA activity unmasked a substantial contribution of InsP\(_3\)R activity to the total SR-Ca\(^{2+}\) leak (Fig. 4) which was not seen before. Taken together, eventless InsP\(_3\)R-Ca\(^{2+}\) release flux is part of a Ca\(^{2+}\) leak and facilitates SR-Ca\(^{2+}\) spark probability.
DISCUSSION

The physiological role of atrial myocytes is to ensure ventricular blood refilling especially under conditions of physical stress and activity. This involves stimulation by direct autonomic nervous system innervation, neuro-humoral effects and interactions with hormones released by the vasculature which boost ECC on a cellular level. Similar to ventricular myocytes, the fundamental mechanism for ECC in atrial myocytes is driven by CICR. However, the contribution of IP3ICR in initiation, propagation and amplification of local and global SR-Ca\(^{2+}\) release and thus CICR in atrial myocytes remains controversial (Blatter et al., 2003; Mackenzie et al., 2004).

Our results obtained in atrial myocytes showed significant alterations in Ca\(^{2+}\) sparks (width and frequency) during InsP\(_3\)R activation. This emphasizes that local Ca\(^{2+}\) release from InsP\(_3\)Rs may play a significant role in the modulation of Ca\(^{2+}\)-dependent RyR activity in atrial myocytes. Our approach, based on the combination of different biophysical methods, revealed some discrepancies between InsP\(_3\)R activation and the appearance of local SR-Ca\(^{2+}\) release events such as apparent invisible, “eventless” Ca\(^{2+}\) release although InsP\(_3\)R activation occurred. From these discrepancies we derived a novel conclusion regarding functional crosstalk between RyRs and InsP\(_3\)Rs in atrial myocytes based on highly efficient but “eventless” IP3ICR.

Identification of Ca\(^{2+}\) sparks

We analyzed individual Ca\(^{2+}\) release events triggered by photorelease of InsP\(_3\). RyRs Ca\(^{2+}\) sparks are characterized by amplitudes of up to 2 \(\Delta F/F_0\), widths \(\approx 2\ \mu m\), FDHM \(\approx 30\ \text{ms}\), whereas InsP\(_3\) Ca\(^{2+}\) puffs are characterized by smaller amplitudes, increased widths \(\approx 6\ \mu m\) and a prolonged FDHM of about 100-600 ms (Zima & Blatter, 2004; Niggli & Shirokova, 2007; Cheng & Lederer, 2008). Our spark analysis revealed Ca\(^{2+}\) sparks but did not show a second distinct class of Ca\(^{2+}\) release events which could be \textit{ad hoc} assigned to Ca\(^{2+}\) puffs, respectively. In addition, selective pharmacological inhibition of either InsP\(_3\)Rs or RyRs let us conclude that the detected InsP\(_3\) triggered Ca\(^{2+}\) release events can be classified as Ca\(^{2+}\) sparks. However, the absence of photolytically triggered IP3ICR events was surprising.

Functional crosstalk of InsP\(_3\)R and RyR2 via invisible SR-Ca\(^{2+}\) release

Despite the evidence that IP3ICR occurs in cardiac cells (Zima & Blatter, 2004), reports of elementary Ca\(^{2+}\) signals arising from InsP\(_3\)R openings have been rare, which may be due to the Ca\(^{2+}\) spark dominance. In atrial myocytes, InsP\(_3\)Rs are colocalized with RyRs, preferentially in the subsarcolemmal space and around the nucleus (Lipp et al., 2000; Mackenzie et al., 2002). The
expression of InsP₃Rs is known to be regulated in a highly dynamic manner, notably during development and cellular remodeling in pathophysiological situations (Janowski et al., 2006).

RyRs are arranged in functional arrays (10–300 RyRs) and subject to stochastic cluster assembly processes (Baddeley et al., 2009). The synchronized openings of clustered RyRs result in local Ca²⁺ sparks. Under resting conditions, in diastole, spontaneous Ca²⁺ sparks reflect the finite open probability of RyRs which is controlled by [Ca²⁺], luminal Ca²⁺ ([Ca²⁺]₅R), phosphorylation state, and numerous other factors (Eisner et al., 1998). A comparable configuration of spatially segregated clusters of 10–100 InsP₃Rs is believed to be responsible for the generation of “Ca²⁺ puffs”, activated by InsP₃ and Ca²⁺ (Foskett et al., 2007), but direct ultrastructural evidence is yet missing (Shuai et al., 2007). In addition, there are species differences human/rat/rabbit vs. mouse in the InsP₃R cluster dynamics and microarchitecture which shape the InsP₃ Ca²⁺ event characteristics (Diambra & Marchant, 2011). It has been suggested, that InsP₃Rs are mobile in the SR membrane and can form clusters upon stimulation (Pantazaka & Taylor, 2011). There is also evidence that InsP₃R Ca²⁺ release sites represent pre-established, stable clusters of InsP₃Rs (Smith et al., 2009). Under this view, brief exposure to InsP₃ should trigger Ca²⁺ puffs, which was not the case in our study. We suggest, that InsP₃Rs in intact mouse atrial myocytes may not form sufficient cluster size to facilitate Ca²⁺ puffs due to low level of InsP₃R expression and/or intercluster distance. InsP₃Rs may rather be distributed near RyRs to facilitate Ca²⁺ sparks. This view is supported by our immunohistochemistry study showing that InsP₃Rs seem to be rather homogeneously distributed (see Fig. 3S, Supplementary Material). According to our data, expression of functional InsP₃Rs in atrial myocytes may not necessarily lead to the formation of distinguished Ca²⁺ puffs.

Thus, how can the increase in Ca²⁺ spark frequency after InsP₃ photorelease be explained? The prevalent view that Ca²⁺ sparks and Ca²⁺ puffs are elementary events, becomes more comprehensive, since for both types of local Ca²⁺ release events, even smaller elementary Ca²⁺ release events have been confirmed. Compared to Ca²⁺ sparks, openings of single RyRs may result in smaller but more frequent events (“Ca²⁺ quarks”) (Lipp et al., 1996; Brochet et al., 2011). At the same time, similar to Ca²⁺ quarks, InsP₃R-dependent elementary Ca²⁺ release events arising from single InsP₃R openings, termed “Ca²⁺ blips”, have been discovered (Parker et al., 1996). Opposed to triggered events, which are experimentally easy to detect, “blips” and “quarks” are largely invisible because these Ca²⁺ events are below detection threshold (Sobie et al., 2002; Williams et al., 2011). This is due to the low signal-to-noise ratio given by fluorescent Ca²⁺ measurements. The absence of visible Ca²⁺ puffs in our measurements, together with the observed increase in spark frequency mediated by InsP₃Rs, suggests a significant contribution of “invisible” or “eventless”
IP3ICR to the occurrence of Ca\(^{2+}\) sparks (and possible Ca\(^{2+}\) wave propagation). We propose that these eventless InsP\(_3\)R openings may be responsible for microdomain [Ca\(^{2+}\)]\(_i\), increase that either sensitize RyRs for CICR or lead to direct RyR activation. Functional crosstalk between InsP\(_3\)R and RyR can operate in both directions. Hence, microdomain [Ca\(^{2+}\)]\(_i\) elevations could sensitize InsP\(_3\)R for InsP\(_3\) which would favor InsP\(_3\)R openings (Foskett et al., 2007).

Thus, the contribution of eventless SR-Ca\(^{2+}\) release via InsP\(_3\)R could be more substantial for the regulation of Ca\(^{2+}\) signaling in cellular microdomains than previously thought. This is consistent with a number of not entirely explained experimental reports regarding functional interactions of InsP\(_3\)R and RyR. It has been reported that InsP\(_3\)R activity increases [Ca\(^{2+}\)]\(_i\) in the vicinity of RyRs and thus facilitates CICR during ECC in adult cat atrial myocytes (Zima & Blatter, 2004) and that InsP\(_3\)-dependent Ca\(^{2+}\) release has a positive inotropic effect on ECC by facilitating Ca\(^{2+}\) release through RyR clusters in rabbit ventricle myocytes (Domeier et al., 2008). Furthermore, there is evidence that Ca\(^{2+}\) leak through InsP\(_3\)Rs is present at sites where RyRs are located and that this Ca\(^{2+}\) leak can modulate RyR Ca\(^{2+}\) release events (Gordienko & Bolton, 2002).

This “invisible” SR-Ca\(^{2+}\) leak, respectively SR-Ca\(^{2+}\) flux via InsP\(_3\)R, was further examined in a series of SR-Ca\(^{2+}\) leak measurements. Here, InsP\(_3\)R function was stimulated with ET-1. Compared to control, ET-1 may stimulate various Ca\(^{2+}\) influx pathways (e.g. via TRPC) (Treves et al., 2004), which may have led to an increased SR-Ca\(^{2+}\) load before the nominally Ca\(^{2+}\) and Na\(^+\) free solution was added. This would lead per se to a transient increase in SR-Ca\(^{2+}\) leak until a new steady-state with matched leak/load relationship is reached. Nevertheless, increased InsP\(_3\)R open probability will then concomitantly produce an “excess” SR-Ca\(^{2+}\) leak in the Ca\(^{2+}\) and Na\(^+\) free solution compared to control as seen in Figure 4A. This is largely independent of the total SR-Ca\(^{2+}\) content and results in an increased leak/load relationship.

Taken together, experiments using photorelease of InsP\(_3\) and SR-Ca\(^{2+}\) leak measurements suggest that “eventless” InsP\(_3\) dependent SR-Ca\(^{2+}\) leak is the main mechanism of functional crosstalk between InsP3Rs and RyRs.

**Pathophysiological implications**

A change in the SR-Ca\(^{2+}\) leak/load relationship can effectively change [Ca\(^{2+}\)]\(_{SR}\) transiently or even under steady-state conditions. It is well established that [Ca\(^{2+}\)]\(_{SR}\) is involved in various forms of cellular instability and affected during cellular remodeling in response to pathophysiological stress (Bers, 2003; Shannon et al., 2003; Wehrens et al., 2005). Under chronic atrial fibrillation, InsP\(_3\)Rs are targeted during cellular remodeling and were found to be up-regulated, which may have
positive inotropic effects on the global level (Zima & Blatter, 2004; Li et al., 2005; Bootman et al., 2007; Harzheim et al., 2009). Recently, subcellular mechanism(s) linking InsP₃R activity to the development of atrial fibrillation and cardiac hypertrophy has been demonstrated (Higazi et al., 2009; Nakayama et al., 2010). Cardiac specific blockage of the InsP₃R pathway, therefore, could offer a new therapeutic strategy for treatment of atrial arrhythmogenicity.

AUTHOR CONTRIBUTIONS

All experiments were performed at the Dept. of Physiology, University of Bern, Switzerland. M.E. conceived and designed experiments. T.H. collected and analyzed the experimental data. M.E., T.H. and N.D.U. drafted and revised the article. All authors approved the final version for publication.

ACKNOWLEDGEMENTS

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FIGURE LEGEN

FIGURE 1
ET-1 induced InsP₃R Ca²⁺ release in atrial myocytes
Time series of InsP₃ induced Ca²⁺ release events expressed as F/F₀ of fluo-3 fluorescence obtained in atrial myocytes. Cells were loaded with 5 µM fluo-3 AM (Biotium) for 20 min and left for de-estrification for 15 minutes before time series were collected in resting cells. A. InsP₃ pathway was stimulated with 100 nM ET-1 causing a substantial increase in SR-Ca²⁺ release event activity. B. Inhibition of ET-1-triggered Ca²⁺ events with 5 µM 2APB. C. Spontaneous Ca²⁺ event activity under control conditions. SR-Ca²⁺ load was measured at the end of each experiment and found not significantly different in the various conditions (CTRL: 3.45±0.28 F/F₀; ET-1: 3.89±0.48 F/F₀; 2APB+ET-1: 3.84±0.53 F/F₀). Bar graphs represent averaged F/F₀ of the first 20 s (Ctrl) and 160-200 s; scale bars represent 10 µm, N=3, n=4-9.

FIGURE 2
InsP₃-induced Ca²⁺ release by UV-flash photolysis of caged InsP₃ in atrial myocytes.
A. SR-Ca²⁺ loading protocol performed by I_{Ca,L} (10 times). B. Representative line scan images of voltage clamped atrial myocytes in whole-cell configuration before and after UV-flash photolysis in control and with 30, 60, 240 µM caged InsP₃. C. Frequency of Ca²⁺ release events counted after UV-flash photolysis normalized to the spontaneous Ca²⁺ release events. D. Averaged spatio-temporal Ca²⁺ spark properties after uncaging of InsP₃ (240 µM) normalized to control. Cells were divided into 2 groups, white bars represent cells that showed little (<1.3) increase in Ca²⁺ release frequency (0.98±0.08 events*100µm⁻¹s⁻¹), black bars represent cells that showed >1.3 increase in Ca²⁺ release frequency (3.24±0.55 events*100µm⁻¹s⁻¹). Amplitude (ΔF/F₀), full width at half-maximum amplitude (FWHM; µm), full duration at half-maximum amplitude (FDHM; ms) showed no significant difference after flash photolysis in both groups (See supplementary Table 1). Full width (µm) showed a significant increase (from 3.80±0.3 µm to 4.50±0.4 µm) in cells with low Ca²⁺ event frequency in response to photolysis. *P<0.05 vs. control, N=6-7, n=6-17, mean±SEM.
**FIGURE 3**

**Identification of InsP$_3$ evoked Ca$^{2+}$ sparks.**

**A.** Experimental protocol, SR-Ca$^{2+}$ loading protocol performed by $I_{Ca,L}$ (10 times); atrial myocytes were voltage clamped under whole-cell conditions, the pipette solution contained 240 µM caged InsP$_3$; line-scans are numbered 1-3: 1) control, 2) pharmacological interventions and 3) pharmacological interventions combined with photolytically released InsP$_3$. Modified protocols were used for the application of 2APB and TET. **B.** Representative data showing line-scan images of fluo-3 fluorescence under whole-cell conditions taken at selected time points (1) control, (2) in the presence of 2APB or TET, (3) pharmacology and photolytic release of InsP$_3$. **C.** Frequency of Ca$^{2+}$ release events evoked by uncaging of caged InsP$_3$ normalized to control (spontaneous frequency in rest). The increase in Ca$^{2+}$ spark frequency was inhibited by the application of either 2APB or Xesto (InsP$_3$R inhibitors) and by TET (RyR inhibitor). **D.** SR-Ca$^{2+}$ content measured by rapid caffeine (10 mM) application. **E.** Change in background fluo-3 fluorescence F/F$_0$ after InsP$_3$ uncaging, in the presence of TET (dark grey) or 2APB (red). Fluorescence F/F$_0$ was normalized to background fluorescence before InsP$_3$ release (right). Background fluorescence increase after UV-flash photolysis of caged InsP$_3$ in the presence of TET was blocked by 2APB. *P<0.05 vs. control, N=2-5; n=6-13, mean±SEM.

**FIGURE 4**

**Invisible InsP$_3$ dependent SR-Ca$^{2+}$ release unmasked by SR-Ca$^{2+}$ leak-load relationship**

**A.** Representative data showing averaged F/F$_0$ of fluo-3 fluorescence changes (control: red, ET-1: black) and **B.** corresponding line-scan images of fluo-3 fluorescence recorded in atrial myocytes. Cells were field stimulated for at least 45 seconds at 1 Hz before line scan images were recorded. After pre-pulse protocol, superfusion was rapidly switched from 1.8 mM [Ca$^{2+}$]$_o$ to a nominally [Ca$^{2+}$]$_o$ and [Na$^+$]$_o$-free extracellular solution for 15 s, then 10 µM CPA was added for 20 s, before 10 mM caffeine was applied. The protocol was repeated in the presence of 100 nM ET-1 after pre-incubation with 100 nM ET-1 for at least 120 s. The increase in background fluorescence when CPA was applied represents SR-Ca$^{2+}$ leak ($\Delta$[Ca$^{2+}$]). **C.** Averaged data, SR-Ca$^{2+}$ leak ($\Delta$[Ca$^{2+}$]), SR-Ca$^{2+}$ content and SR-Ca$^{2+}$ leak/SR-Ca$^{2+}$ load relationship ($\Delta$[Ca$^{2+}$]/SR-Ca$^{2+}$ load) compared to control (no InsP$_3$ pathway stimulation). In the presence of ET-1 there was a significant increase in the SR-Ca$^{2+}$ leak ($\Delta$[Ca$^{2+}$]) when CPA was added unmasking a substantial contribution of the InsP$_3$R opening to the total SR-Ca$^{2+}$ leak. *P<0.02-0.009 vs. control, N=3; n=12-13, mean±SEM.
REFERENCES


ONLINE SUPPLEMENTS

Material and Methods

Atrial myocytes from adult male C57/BL6 mice, obtained from the Central Animal Facility, University Hospital, University of Bern, were freshly isolated by Langendorff perfusion technique. All experiments were approved by the State Veterinary Office of Bern, Switzerland, according to Swiss Federal Animal protection law. The authors have also read, and the experiments comply with, the policies and regulations of *The Journal of Physiology* described by Drummond, 2009 (Drummond, 2009). Adult male C57/BL6 mice (3-5 months old) were heparinized (50 U ip) and subsequently killed after 5-15 minutes by cervical dislocation. Hearts were rapidly removed and mounted on a Langendorff column and perfused retrogradely for 8-16 minutes with collagenase type II (14 U/ml, Worthington type 2) and protease type IV (0.2 U/ml, Sigma, type XIV) in a Ca\(^{2+}\) free solution (in mmol/L: 140 NaCl, 5.4 KCl, 1.1 MgCl\(_2\), 1 Na\(_2\)HPO\(_4\), 5 HEPES and 10 glucose at pH 7.4, 37°C). Atria (auricle) were removed and rinsed in 1% BSA solution before cells were dissociated mechanically from the tissue in Ca\(^{2+}\) free solution. The free [Ca\(^{2+}\)]\(_o\) was increased progressively to 500-700 µM over 1-2 hours before cells were used for experiments within 4 hours after isolation. All experiments were performed at room temperature.

HeLa cells were a kind gift from N. Demaurex’s lab, Geneva. HeLa cells were cultured on glass cover slips in DMEM, 10% FBS, 1% HEPES, 0.1% penicillin / streptomycin until 80% confluent. Once 80% confluency was reached, cells were used for experiments. Ca\(^{2+}\) responses were detected with K\(_5\)-fluo-3 after flash photolysis of 30 µM caged InsP\(_3\) in the patch pipette.

*Patch Clamp Recordings with caged InsP\(_3\).* Electrophysiological recordings were combined with simultaneous confocal Ca\(^{2+}\) imaging. Caged InsP\(_3\) (Sichem) was dialyzed into the atrial myocytes via low resistance pipettes (1.5-3 MΩ) which were pulled from borosilicate glass micropipettes using a Zeitz DMZ puller (Zeitz instruments, Germany). The pipette solution contained (in mmol/L) 120 CsAsp, 10 HEPES, 20 TEA-Cl, 5 K-ATP,1 MgCl\(_2\), 0.1 K\(_5\)-fluo-3 (Biotium), 2 GSH, (0.03, 0.06, 0.24) caged InsP\(_3\) 6Na (Sichem), pH 7.2 with CsOH. Free Ca\(^{2+}\) was calculated using Patcher’s Power Tools (IgorPro plug-in, MPI Göttingen, Germany) and was approximately 59 nM. During the experimental procedure, cells were perfused with external solution containing (in mmol/L) 140 NaCl, 5 HEPES, 1.1 MgCl\(_2\), 5.4 KCl, 10 Glucose, 1.8 CaCl\(_2\), 0.5 BaCl, 1 CsCl, pH 7.4 NaOH. Pharmacological experiments used 5 µM 2-aminoethoxydiphenyl borate (2APB, Fluka), 5 µM xestospongin C (Xesto, A.G Scientific Inc), 1 mM tetracaine (TET, Sigma) or 10 mM caffeine (Sigma) added to the external solution.
Figure 2

A

+ 10 mV
- 40 mV
- 80 mV
UV flash
Caffeine

B

Before uncaging	After uncaging
Control
30 µM InsP₃
60 µM InsP₃
240 µM InsP₃

200 ms
10 µm

C

D

# of events normalized

normalized to control
Figure 3

A

UV flash

+ 10 mV
- 40 mV
- 80 mV

2APB

Caffeine

B

2APB

TET

1

2

3

200 ms

10 µm

F/F₀

1.0

3.0

C

# of events normalized

Control
InsP₃
InsP₃ + 2APB
InsP₃ + TET

D

SR-Ca²⁺ content normalized

InsP₃
InsP₃ + 2APB
InsP₃ + TET

E

UV flash

2APB + InsP₃

TET + InsP₃

normalized background fluorescence F/F₀
Figure 4

A

3.0
F/F₀
1.0

5 s

[Ca²⁺]ᵢ

1.8 mM Ca²⁺

Ca²⁺ and Na⁺ free

Ca²⁺ and Na⁺ free + CPA

Caffeine

Control

ET-1

100 nM ET-1

B

Control

ET-1

10 µm

5 s

C

SR-Ca²⁺ leak

( [Ca²⁺]ᵢ )

SR Ca²⁺ content

[Ca²⁺]ᵢ / SR-Ca²⁺ load

a.u.

0.4
0.2
0

ET-1

Control

ET-1

Control

ET-1

Control

* 12 13

F/F₀

3.0
1.5
0

* 12 13

a.u.

0.06
0.03
0

* 12 13
Cells were held at -80 mV and depolarized from -80 mV to 0mV ten times before line-scan images were taken; membrane voltages were controlled with an Axopatch 200 amplifier (Axon Instruments, Foster City, CA) driven by custom-written software developed under LabView software (National Instruments). Fluo-3 was excited with the 488 nm line of an argon laser alternatively with a solid-state laser (50 µW; Sapphire, Coherent) and fluorescence (> 515 nm) in line-scan mode (6 ms/line) was collected with a PMT on an MRC-100 or MicroRadiance confocal microscope (both BioRad, Glattbrugg, Switzerland). Line-scans of 1024 lines were collected in 2 equal, sequential segments. In line-scan recordings where InsP3 was uncaged, UV light from a xenon short-arc flash lamp (300-350 nm, pulse duration ~ 400 µs, discharge energy up to 230 J) was delivered and applied at the beginning of line-scan recordings. For more details see Del Principe et al. (DelPrincipe et al., 1999). Line-scan images were processed using customized Image SXM software (Steve Barrett, University of Liverpool, UK). 3000 ms were used for analysis of each line-scan. Sparkmaster plug-in (Picht et al., 2007) for ImageJ (Wayne Rassband, NIH, USA) was used to identify and analyze Ca2+ release events. IgorPRO (WaveMatrics, Portland, OR, USA) software was used to analyze caffeine transients.

**Ca2+ leak measurements.** Using the RyR2 blocker TET, Ca2+ leak was determined and using the SR-Ca2+ pump (SERCA) blocker cyclopiazonic acid (CPA), the increase in [Ca2+]i (Δ[Ca2+]i) was determined with an adapted protocol from Shannon et al. (Shannon et al., 2002). Line-scans were recorded at 50 l/s on a laser scanning confocal microscope (MicroRadiance, BioRad, UK). Atrial myocytes were loaded with 5 µM fluo-3 AM (Biotium) for 20 min and left for de-estrification for 15 minutes before confocal images were collected. Fluo-3 (Biotium) was excited using an argon ion laser. Cells were field stimulated for at least 45 seconds at 1 Hz before line-scan images were recorded. After the pre-loading protocol, superfusion was rapidly switched from (in mmol/L) 140 NaCl, 5 KCl, 5 HEPES, 1 MgCl, 1.8 CaCl2, 10 glucose, pH 7.4 with NaOH to a nominally Na+ and Ca2+-free solution (by substitution of Na+ with Li+ and addition of 0.5 mmol/L EGTA). A further solution switch was done after 10 s to a nominally Ca2+ and Na+ free solution containing CPA for 30 s, before caffeine was applied to measure SR-Ca2+ load. The same protocol was repeated in the presence of 100 nM ET-1 in all solutions. CPA application was accompanied with an increase in [Ca2+]i.

**Immunofluorescent staining.** Atrial myocytes were enzymatically isolated as described earlier. Without increasing Ca2+, 500 µl cell dispersion was left to settle for 2 hours at room temperature on previously prepared gelatin coated coverslips. Samples were washed twice before they were fixed in 4 % paraformaldehyde for 10 minutes. Prior to staining, samples were washed one more time with
PBS before cells were permeabilized and unspecific sites were blocked for 30 min in 0.1% Triton-X-100, 5 % BSA in PBS. Myocytes were stained against InsP3R2 and RyRs by incubating the samples with rabbit polyclonal InsP3R2 (Abcam, 1:500, Cat.# ab77838, see also Wang et al., 2012) and mouse monoclonal RyR (Abcam 1:200, Cat.# ab2827) antibodies overnight at 4ºC in 0.1% Triton-X-100, 5 % BSA in PBS. Cells were then again washed 3 times in 0.1% Triton-X-100, 5 % BSA in PBS for 10 minutes before incubating with the secondary antibodies (Alexa Fluo 488, 1:1000; Alexa Fluo 568, 1:1000) and DAPI, 1:20000 for 1.5 hours at room temperature. Cells were then washed 3 times with PBS before coverslips were mounted on slides with mounting medium (Fluoroshield). Cells were imaged using a fluorescent microscope (TMD300 microscope, Nikon) equipped with a high-sensitivity Visicam camera (Visitron Systems, Germany).

Verification of InsP₃R Ca²⁺ release

In a series of control experiments we ruled out flaws in the applied experimental settings. HeLa cells are known to have a highly developed InsP₃R pathway and were used as a bio-indicator for InsP₃ Ca²⁺ response. HeLa cells have several advantages, they express InsP₃R at high density and InsP₃R dependent elementary Ca²⁺ release events are well characterized. Although they express mainly type 1 and type 3 isoforms, InsP₃Rs type 2 are also present (Tovey et al., 2001). This workaround was inevitable because there is still a deficit in the availability of InsP₃ indicators which could be used to monitor rapid intracellular InsP₃-mediated signals. Indeed, by using UV-flash photolysis, robust InsP₃ Ca²⁺ transients were elicited (Fig. 1S). As shown in Fig. 1S, using identical settings and solutions containing 30 µM caged InsP₃, reliable global Ca²⁺ transients were photolytically triggered. This suggests that UV-flash triggered IP3ICR can be obtained similarly in cardiac cell preparations. The reliability of the experimental approach was further underscored by the dose dependency of InsP₃-induced increase in Ca²⁺ event frequency (Fig. 2C).

As a positive control for elementary Ca²⁺ release events we used neonatal rat cardiomyocytes that are known to exhibit both Ca²⁺ puffs and Ca²⁺ sparks (Luo et al., 2008). Fig. 2S A,B and C show individual photolytically triggered InsP₃ “Ca²⁺ puffs”, “Ca²⁺ sparks” and local mini-waves identified by their spatio-temporal properties (Niggli & Shirokova, 2007).

For a variety of caged compounds it is known that only a small fraction of the total amount of the compound is being uncaged during a single UV-flash. This essentially depends on the physicochemical properties of the compound (e.g. quantum yield), the power of UV-light and the duration of illumination (Ellis-Davies, 2007), which in our hands is in the range of 200 µs. In addition, the pipette solution contains an infinite reservoir for caged InsP₃ compared to the cell volume. Finally, we would expect to end up in a biologically relevant concentration range of approximatively 10 nM - 1 µM InsP₃.
Experimental approach to verify intracellular InsP$_3$ release.

Representative data showing averaged F/F$_0$ of fluo-3 and corresponding line-scan images of changes in [Ca$^{2+}$]$_i$ in HeLa cells under whole-cell conditions of the patch-clamp technique. InsP$_3$ was released by UV-flash photolysis. A. With 30 µM caged InsP$_3$ in the patch pipette there is an immediate global increase in Ca$^{2+}$ after UV flash photolysis observed in 100% of all approached cells. The inset shows normalized fluorescence F/F$_0$ before and after UV-flash (n=6); B. Control, with no InsP$_3$ in the patch pipette, shows no response to flash photolysis. n=5-12
Figure 2S. Representative data showing averaged ΔF/F₀ of fluo-3 fluorescence and corresponding line-scan images of changes in [Ca²⁺]ᵢ in neonatal rat myocytes in response to InsP₃ photorelease. Cells were loaded with 5 µM InsP₃-PM (D-2,3-O-Isopropylidene-6-O-(2-Nitro-4,5-Dimethoxy) benzyl-myo-Inositol-1,4,5-Trisphosphate-Hexakis Propionoxymethyl Ester, SiChem) together with 4 µM fluo-3-AM (Biotium) for 120 and 30 min at room temperature, respectively. InsP₃ was liberated by UV-flash photolysis as described in the Material and Methods section. **A. a), b)** Individually triggered Ca²⁺ puff consecutively followed by a Ca²⁺ spark. **B.** Expanded part of the line-scan given above, now presented in unprocessed raw data format. From FDHM and FWHM analysis (1) can be classified as Ca²⁺ puff and (2) as Ca²⁺ spark (Picht et al., 2007). **C.** Ca²⁺ mini-wave elicited in response of initial InsP₃ triggered local [Ca²⁺]ᵢ increase. This local Ca²⁺ response shows that elementary Ca²⁺ release via InsP₃R may be involved in the conditioning of CICR (e.g. Ca²⁺ mini-waves as given in this example) in cardiac cells.
Figure 3S. Colocalization of RyR2 and InsP₃R2 in an atrial myocyte. A. Atrial myocyte was stained against RyR2 (primary antibody from Abcam 1:200, secondary Alexa Fluo 568 1:1000) B. Atrial myocyte was stained against InsP₃R2 (primary antibody from Abcam 1:500, secondary Alexa Fluo 488 1:1000), intensity profiles of immunofluorescence across the cell (right). These profiles were obtained by sampling the immunofluorescence intensity of the InsP₃R2 isoform across a line one pixel wide. C. The nucleus of atrial myocyte was stained with DAPI 1:20000 D. Merged image.
Ca\textsuperscript{2+} Spark characteristics.

Ca\textsuperscript{2+} spark characteristics were analyzed as mentioned in the Material and Methods section. Myocytes were divided into two groups.

**Supplementary Table 1**

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<td>9 myocytes, 5 animals</td>
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**REFERENCES**


