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Publication Date

2021-12-01

DOI

10.1016/j.ceca.2021.102494

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Peer reviewed



HHS Public Access

Author manuscript *Cell Calcium.* Author manuscript; available in PMC 2022 October 12.

Published in final edited form as:

Cell Calcium. 2021 December ; 100: 102494. doi:10.1016/j.ceca.2021.102494.

Termination of Ca²⁺ puffs during IP₃-evoked global Ca²⁺ signals

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Abstract

We previously described that cell-wide cytosolic Ca²⁺ transients evoked by inositol trisphosphate (IP₃) are generated by two modes of Ca^{2+} liberation from the ER; 'punctate' release via an initial flurry of transient Ca²⁺ puffs from local clusters of IP₃ receptors, succeeded by a spatially and temporally 'diffuse' Ca²⁺ liberation. Those findings were derived using statistical fluctuation analysis to monitor puff activity which is otherwise masked as global Ca²⁺ levels rise. Here, we devised imaging approaches to resolve individual puffs during global Ca²⁺ elevations to better investigate the mechanisms terminating the puff flurry. We find that puffs contribute about 40% (~90 attomoles) of the total Ca²⁺ liberation, largely while the global Ca²⁺ signal rises halfway to its peak. The major factor terminating punctate Ca^{2+} release is an abrupt decline in puff frequency. Although the amplitudes of large puffs fall during the flurry, the amplitudes of more numerous small puffs remain steady, so overall puff amplitudes decline only modestly (~30%). The Ca^{2+} flux through individual IP₃ receptor/channels does not measurably decline during the flurry, or when puff activity is depressed by pharmacological lowering of Ca^{2+} levels in the ER lumen, indicating that the termination of punctate release is not a simple consequence of reduced driving force for Ca²⁺ liberation. We propose instead that the gating of IP₃ receptors at puff sites is modulated such that their openings become suppressed as the bulk $[Ca^{2+}]$ in the ER lumen falls during global Ca²⁺ signals.

Graphical Abstract

IP directed the project. IP and JTL performed experiments, analyzed data and wrote the manuscript.

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Competing Interests

The authors declare no competing interests.



Keywords

IP₃ receptor; IP₃ signaling; Calcium signaling; Ca²⁺ puffs; Calcium imaging

1. INTRODUCTION

Cytosolic Ca²⁺ signals generated by the liberation of Ca²⁺ ions sequestered in the endoplasmic reticulum (ER) through inositol trisphosphate receptor (IP₃R) channels regulate ubiquitous cellular processes as diverse as gene transcription, secretion, mitochondrial energetics, electrical excitability and fertilization[1,2]. Such unique repertories of Ca²⁺ dependent functions are achieved through a hierarchy of cytosolic Ca²⁺ signals with markedly different spatial scales and temporal durations, ranging from transient, subcellular Ca^{2+} signals called puffs[3–5] to more prolonged global Ca^{2+} elevations that engulf the cell[6]. This patterning is shaped by the functional properties of IP₃Rs together with their spatial arrangement in the cell. Biphasic modulation of IP₃R channel gating by cytosolic Ca^{2+} results in a phenomenon of Ca^{2+} -induced Ca^{2+} -release (CICR)[7,8]. Ca^{2+} diffusing from one open channel may thus trigger the opening of adjacent channels, with selfreinforcing CICR countered by inhibitory feedback at high $[Ca^{2+}]$. The clustered distribution of IP₃Rs further defines the extent of this regenerative process as puffs arise from localized CICR between IP₃Rs within stationary clusters, where they are preferentially 'licensed' to respond[4,9]. A widely-accepted model further posits that global Ca²⁺ signals arise through successive recruitment of puffs via cycles of CICR and Ca²⁺ diffusion between the IP₃R clusters at puff sites - in essence, that puffs are the basic 'building block' from which all IP₃-mediated cellular Ca^{2+} signals are constructed[5,10–15].

We recently challenged that notion[16], utilizing a statistical technique of fluctuation analysis of Ca^{2+} imaging data to monitor bulk puff activity during global responses, even when individual puffs were obscured by large cell-wide Ca^{2+} elevations. We found that although a flurry of puff activity was prominent during the initial portion of the global Ca^{2+} response and peaked early during the rising phase, it then terminated around the midpoint of the rising phase. Notably, global Ca^{2+} levels continued to climb after puffs had largely ceased, and cytosolic Ca^{2+} remained elevated for several seconds following the apex of the global signal, during which time local Ca^{2+} fluctuations were substantially suppressed. The flurry of puffs did not cease because IP₃Rs became inhibited by the rising cytosolic Ca^{2+} levels during the spike, but instead appeared to terminate because of falling levels of

 Ca^{2+} in the ER lumen[16]. Indeed, by prior partial depletion of ER Ca^{2+} we were able to evoke global Ca^{2+} responses in the absence of puffs. We thus proposed the existence of two distinct modes of IP₃-mediated Ca^{2+} liberation: 'punctate' liberation in the form of transient, local Ca^{2+} puffs, and a second 'diffuse' mode of liberation that is more homogeneous in space and time[16].

Although our fluctuation analysis technique provided a gross measure of puff activity across the cell it did not resolve individual puffs, leaving important questions unanswered. How is the termination of the puff flurry reflected in the behavior of individual puff sites? Does the decline in punctate Ca^{2+} liberation arise simply because of reduced driving force as ER levels decline, or is the gating of IP₃Rs modulated? Here, we address these issues by applying experimental and image-processing techniques that enable the resolution of individual puffs during the rise of global Ca^{2+} signals.

2. MATERIALS and METHODS

2.1. Cells and cell-loading

HEK293 wild-type cells were kindly provided by Dr. David Yule (University of Rochester) and were cultured in Eagle's Minimum Essential Medium (EMEM; ATCC #30-2003), supplemented with 10% fetal bovine serum (Omega Scientific #FB-11). Cells were grown in plastic 75 cm flasks and maintained at 37°C in a humidified incubator gassed with 95% air and 5% CO₂. For imaging, cells were collected using 0.25% Trypsin-EDTA (ThermoFisher #25200-056) and grown on poly-D-lysine coated (1mg/ml; Sigma #P0899) 35-mm glass bottom imaging dishes (MatTek #P35-1.5-14-C).

Cells were incubated with the membrane-permeant fluorescent Ca²⁺ indicator Cal520/AM (1-5 µM; AAT Bioquest #21130) for 1 hr at room temperature in a Ca²⁺-containing HEPES buffered salt solution (Ca²⁺-HBSS) together with membrane permeant ester of the caged IP₃ analogue ci-IP₃/PM [D-2,3,-O-Isopropylidene-6-O-(2-nitro-4,5 dimethoxy) benzyl-myo-Inositol 1,4,5,-trisphosphate hexakis (propionoxymethyl) ester] (1 µM; SiChem #cagiso-2-145-10). Except where indicated, cells were additionally loaded by incubation with EGTA/AM (15 µM; Thermo Fisher Scientific #E1219) for a further 1 hr at room temperature in Ca²⁺-HBSS to attenuate global Ca²⁺ elevations. Cells were finally maintained for 30 min at room temperature in Ca²⁺-HBSS before imaging. The genetically encoded, ERtargeted Ca²⁺ indicator R-CEPIA_{FR}[17] was transfected into cells using lipofectamine 3000 (ThermoFisher #L3000008) according to the manufacturer's instructions, and imaged 24-48 hrs later. Cal520/AM, i-IP₃/PM, and EGTA/AM were all solubilized with DMSO/20% pluronic F127 (ThermoFisher #P3000MP). Carbachol (CCH; #C4382) and cyclopiazonic acid (CPA; #C1530) were purchased from Sigma and dissolved in deionized water and 100% ethanol, respectively. Ca²⁺-HBSS contained (in mM) 135 NaCl, 5.4 KCl, 2 CaCl₂, 1 MgCl₂, 10 HEPES, and 10 glucose (pH=7.4). Ca²⁺-free HBSS consisted of the same formulation except that CaCl₂ was omitted and 300 µM EGTA was added.

2.2. Microscopy

Total internal reflection fluorescence (TIRF) imaging of Ca²⁺ signals was accomplished using a home-built microscope system, based around an Olympus (Center Valley, PA) IX50 microscope equipped with an Olympus 60X oil immersion TIRF objective (NA 1.45), and an Evolve EMCCD camera (Photometrics; Tucson, AZ) with a bit depth of 16 bits, using 2×2 binning for a final field at the specimen of 128×128 binned pixels (one binned pixel = 0.53 μ m) at rates of 125 or 227 frames s⁻¹. All references to 'pixel' in this paper refer to the binned pixel. Image data were streamed to computer memory using Metamorph v7.7 (Molecular Devices; San Jose, CA) and stored on hard disk for offline analysis. Cal520 fluorescence was excited using a 488 nm laser and detected through a 510 nm long-pass (LP) emission filter; R-CEPIAer imaging utilized a 561 nm excitation laser and a 568 nm LP emission filter. Photorelease of i-IP₃ was evoked by UV light from a xenon arc lamp filtered through a 350-400 nm bandpass filter and introduced by a UV-reflecting dichroic in the light path to uniformly illuminate the field of view. The amount of i-IP₃ released was controlled by varying the flash duration by an electronically controlled shutter. Carbachol was locally applied to cells through glass micropipettes with tip diameters of $\sim 1-2 \,\mu m$ using a pneumatic picospritzer. All imaging was performed at room temperature while cells were bathed in zero Ca²⁺-HBSS containing 0.3 mM EGTA and no added Ca²⁺.

2.3. Image processing

Image data in 16 bit integer multi-plane TIF format were imported into FLIKA (http://flikaorg.github.io), a freely available open-source image processing and analysis software in the Python programming language[18,19]. Internal processing and data output were performed using 64 bit floating-point arithmetic. Processing was performed on image stacks following subtraction of camera black offset level (500 camera units). F/F_0 ratio stacks were generated by dividing each frame in the offset-corrected raw image stack by an average of 100 baseline frames acquired before stimulation. F/F_0 image stacks were generated by subtracting 1 from F/F_0 ratio stacks.

To derive image stacks representing the pixel-by-pixel standard deviation (SD) of temporal fluctuations in fluorescence of the Ca²⁺ indicator dye we used a custom FLIKA script as described previously[16,19]. In brief, following subtraction of camera black offset level, the raw image stack from the camera was first spatially filtered by a Gaussian blur function with a standard deviation (sigma) of 2 pixels (~1µm at the specimen). To attenuate high-frequency photon shot noise and low frequency changes in baseline fluorescence, stacks were temporally filtered by a bandpass Butterworth filter with low and high cutoff frequencies of 3 and 20 Hz. A running variance of this temporally filtered movie was calculated, pixel by pixel, by subtracting the square of the mean from the mean of the square of a moving 20 frame (160 ms) boxcar window of the movie. The running standard deviation was calculated by taking the square root of the variance image stack to create a standard deviation (SD) stack. Finally, to remove the mean predicted photon shot noise (whose variance increases in linear proportion to the mean fluorescence intensity) the SD stack was corrected, pixel-by-pixel, by subtracting the square root of a running mean of the spatially filtered fluorescence image stack multiplied by a scalar constant derived as described previously [16,19]. Traces showing changes in SD signal throughout Ca^{2+}

responses were measured from a region of interest (ROI) encompassing the entire cell or part of the cell within the imaging field, and from small (4×4 pixel; 2×2 µm) ROIs. The locations of active puff sites were readily visualized by forming maximum intensity SD projection images where the intensity of each pixel is the maximum at that pixel across a range of frames acquired during the rising phase of Ca²⁺ responses.

To enable analysis of fluorescence signals generated by individual puffs we applied a custom FLIKA routine (scaled peak subtraction; SPS) to strip off the slow bulk rise in fluorescence during global Ca²⁺ responses to achieve a substantially flat baseline. This operated by forming an averaged image of frames (usually 100) at the time of peak global response and then subtracting a scaled version of this, frame-by-frame, from the original F/F_0 image stack. The scaling factor was derived as a running mean (usually 20 frames) of the whole-cell fluorescence at each time, expressed as a fraction of the change in F/F_0 from pre-stimulus baseline to the mean peak F/F_0 . The resulting image stack was then processed using the 'detect puffs' algorithm in FLIKA to measure peak amplitude, duration (from 20% rise to 80% fall from peak), and integral of each puff. Measurements of dwell-state amplitudes were made by visual inspection of SPS-processed F/F_0 traces from 4×4 pixel regions placed on puff sites identified from maximum-intensity projections of SD images.

2.4. Simulations

A custom plugin 'simulate puffs' in FLIKA was used to add synthetic puffs to an imaging record of a CPA-treated HEK cell that produced a global rise in Ca^{2+} fluorescence in response to photoreleased i-IP₃ in the absence of any detectable puff activity[16]. The size and kinetic parameters of simulated events listed in the legends of fig. S5 and fig. S6 were chosen to approximate experimental observations of small Ca^{2+} puffs and single-channel blips[20–22]. Simulations were then processed in the manner described above for automated and visual analysis of experimental Ca^{2+} imaging records.

2.5. Statistics

Mean values are expressed throughout as standardized means ± 1 SEM, where *n* is the number of cells analyzed. Comparison of mean values between two groups was assessed by both Student's T-test (parametric) and Mann-Whitney test (nonparametric). Differences were considered statistically significant when indicated by both parametric and nonparametric approaches; the larger of the two *P* values are presented throughout. Statistical significance in figures is denoted as **P*< 0.05 and ***P*< 0.01.

3. RESULTS

3.1. Techniques to improve resolution of Ca²⁺ puffs during global Ca²⁺ responses

Localized, Ca^{2+} transients (puffs) are observed during the rising phase of IP₃-evoked global (cell-wide) cytosolic Ca^{2+} responses in many cell types^{10–16}. Because the puffs become difficult to resolve as the global fluorescence Ca^{2+} signal increases, we previously developed image processing and analysis routines to display and analyze local fluctuations in Ca^{2+} fluorescence signals[16,19]. That procedure is illustrated in Fig. 1A,B. The top panels in Fig. 1A show 'snapshots' of fluorescence ratio signals (F/F_0) imaged by TIRF microscopy

of an HEK cell loaded with the Ca²⁺ indicator Cal520 at different times during the rise of a global Ca²⁺ signal evoked by photorelease of i-IP₃ from a caged precursor. Localized puffs can be discerned as increases in Cal520 F/F_0 in the first three top panels of Fig. 1A, but are more apparent in the middle panels that show corresponding SD images depicting a running time-average of the standard deviation of Ca²⁺-dependent fluctuations in fluorescence. Fig. 1B shows measurements from the same cell of the global (cell-wide) Ca²⁺ signal (smooth black trace; Cal520 F/F_0) and global SD signal (noisy red trace). Measurements of global SD signals provide a population average of Ca²⁺ fluctuations associated with puff activity at multiple sites across the cell[16], and measurement of fluctuations from small ROIs at active sites reveal the activity of individual puffs (Fig. 1C).

However, although SD fluctuation measurements identify puffs with a better signal-tonoise ratio than the underlying fluorescence signal, they are not optimal to determine the properties of individual puffs because the magnitude of the SD signal is a function of both the amplitude and kinetics of the Ca²⁺ fluorescence signal (F/F_0), and its time course is constrained by the duration (typically 20 frames, 160 ms) of the running boxcar window over which the standard deviation is calculated. We thus devised two approaches that, in combination, enable enhanced resolution of Ca²⁺ puffs while flattening the baseline on which they are superimposed. First, to flatten the baseline we stripped off the global rise in bulk fluorescence, using an algorithm to subtract from each frame of the F/F_0 image stack an average image of frames acquired around the peak of the response that was scaled in proportion to the global fluorescence of the cell at each time. This procedure (scaled peak subtraction; 'SPS') substantially flattened the baseline, but alone could not entirely compensate for spatial inhomogeneities in Ca²⁺ during the global response (lower panels of Fig. 1A; Fig. 1D). Moreover, the increased photon shot noise resulting from the increase in fluorescence impaired the resolution of puffs during later phases of the response.

We thus applied a complimentary experimental approach to mitigate these effects, based on cytosolic loading of EGTA. This slow Ca^{2+} buffer selectively diminishes global baseline Ca^{2+} elevations evoked by IP₃[27–29], while having little effect on the amplitude of the transient fluorescence signals from puffs (fig. S1), 'sharpening' them in space and time so they become more localized and display kinetics that closely track the openings and closings of individual IP₃R channels[22,29,30]. Figs. 1E–H illustrate representative records, processed and displayed in the same way as for the cell in Fig. 1A–D, but from a cell loaded with EGTA by prior incubation with 15µM EGTA/AM for 1 hr; a protocol chosen to substantially reduce the peak amplitude of the global signal while minimally affecting the SD signal reflecting puff activity. As illustrated in Fig. 1H, the combination of EGTA-loading and SPS image processing to strip the bulk global rise in Ca^{2+} signal enabled recordings on which individual puffs were readily resolved on a stable, flat baseline, and with reduced photon shot noise because the overall fluorescence increase was smaller.

A possible concern with using EGTA to reduce the amplitude of global Ca^{2+} elevations was that, beyond buffering the rise in cytosolic free Ca^{2+} , it might depress the Ca^{2+} release process by impairing CICR between IP₃Rs. To discriminate between these possibilities we monitored ER Ca^{2+} levels during global responses evoked by bath application of carbachol (CCH; 100 μ M) using the ER-targeted, low-affinity Ca^{2+} sensor R-CEPIA_{ER}[17]. EGTA-

loaded cells showed a larger drop and slower refilling of ER Ca²⁺ (fig. S2A), mirroring the reduced amplitude and slowed falling phase of cytosolic Ca²⁺ signals (fig. S2B). The reduced amplitude of the global cytosolic Cal520 signal is readily explained by the action of EGTA to buffer free Ca²⁺ ions. Further, the greater fall in ER Ca²⁺ level indicates that Ca²⁺ release was actually potentiated (fig. S2C), likely because the reduced rise in free cytosolic Ca²⁺ both increased the concentration gradient for Ca²⁺ efflux from the ER and reduced inhibitory feedback of cytosolic Ca²⁺ on the IP₃R channels. Importantly, despite strongly attenuating the CCH-evoked rise in global cytosolic Ca²⁺ (fig. S2D), the distinct punctate and diffuse modes of Ca²⁺ liberation were preserved after EGTA loading[16] and their temporal relationship was retained, with puff activity declining abruptly as the global response reached about half-way toward its peak (Fig. 1).

Unless otherwise noted, all following results in this paper were obtained in EGTA-loaded HEK cells. Moreover, all experiments were done while cells were bathed in a solution containing no added Ca^{2+} and 300 μ M EGTA to exclude Ca^{2+} influx across the plasma membrane, so the responses arise exclusively from Ca^{2+} liberated from intracellular stores.

3.2. Patterns of puff activity during global i-IP₃-evoked responses

Fig. S3 illustrates representative spatio-temporal patterns of puff activity during a global i-IP₃-evoked response in an EGTA-loaded cell, derived using both SD analysis (fig. S3A–D) and SPS processing (fig. S3E,F) of fluorescence signals. Projection images from the SD image stack highlight the locations of multiple hot-spots during the rising phase of the global response (fig. S3A, middle panel), whereas these were absent in images generated from periods before simulation or at the peak of the global response (fig. S3A, left and right panels). Raster plots in fig. S3C,E summarize activity patterns at multiple hot-spots within a single cell displayed, respectively, in terms of local fluctuations (SD signal) and fluorescence ratio signals (F/F_0), throughout the time course of the global Ca²⁺ response. Corresponding traces in fig. S3D,F respectively illustrate SD and F/F_0 signals at five representative sites.

3.3. Changes in puff amplitudes and frequencies during global Ca²⁺ responses

Records like those in fig. S3 portray both a decline in puff amplitudes at some individual sites during the rise of the global Ca^{2+} response, and a marked attenuation in puff frequency and whole-cell SD signal around the time of half-rise to the peak. To quantify these changes we used FLIKA[18,31] to automate detection and measurement of puff parameters (frequency, peak amplitude, duration and integral), after first applying SPS processing to derive F/F_0 image stacks of puffs on a stable baseline fluorescence level. Because the rise time of the global Ca^{2+} signal varies strongly as a function of the amount of photoreleased i-IP₃[16,32], we separately pooled measurements from four HEK cells that gave relatively 'slow' responses (~50 s from photorelease to peak; Fig. 2, left panels) and from four cells that gave 'fast' responses (~20 s; Fig. 2, right panels). To further compensate for remaining variability in kinetics between the cells in each group (Fig. 2A,H, insets) we plot the data on a normalized time scale, with time t=0 corresponding to the time of the photolysis flash and t=1 corresponding to the time the global fluorescence signal rose to 50% of its peak value for each cell.

Figs. 2A,B show, respectively, overlays of global fluorescence ratio (F/F_0) signals and global SD signals from the four slow-rising cells. Fig. 2C overlays representative puff activity (F/F_0 after SPS processing) at 12 individual sites in a single cell, and histogram bars in Fig. 2D show the total numbers of puffs arising in the four cells during normalized time bins. Notably, about 75% of puffs occurred before the global Ca^{2+} signal had risen to half the peak value (normalized time t=1). Fig. 2E shows a scatter plot of all puff amplitudes (grey symbols) and mean amplitudes at binned intervals (black symbols) throughout the rise of the global responses. Many small amplitude events were observed throughout the course of the global signal, but there was a marked fall-off in large amplitude puffs as Ca^{2+} levels rose, resulting in a 35% reduction in mean puff amplitudes from the start of the puff flurry to the half-rise time in global Ca²⁺ fluorescence. The mean durations of puffs increased slightly throughout the rise of the global Ca²⁺ signal (Fig. 2F), roughly balancing the decline in amplitudes so that the mean integrated fluorescence signal during each puff ($F/F_0 * s$; a measure of the cumulative amount of Ca²⁺ released during a puff) remained about constant (Fig. 2G). Thus, the marked decline in overall puff activity around the half-rise time of the global Ca²⁺ response reported by the SD signal (Fig. 2B) can be attributed primarily to a reduced frequency of puffs (Fig. 2D), rather than to a diminution in the mean amount of Ca^{2+} liberated by each puff.

The right-hand panels in Fig. 2H–N show corresponding records and measurements from the four cells showing 'fast' rising global Ca²⁺ responses. Plotted on a normalized timescale the changes in frequency and amplitude of puffs closely match those during slow rising responses. Fig. S4 further compares the distributions of the puff parameters (frequencies, amplitudes, durations, and integrals) during slow- and fast-rising global signals. Despite their occurrence on a more compressed timescale during the fast-rising responses, the overall magnitudes and kinetics of puffs were closely similar to those during slow responses.

3.4. Heterogeneity between puff sites

Inspection of puff traces revealed considerable heterogeneity between puff sites within each cell (e.g. fig. S3E and Fig. 3A). Some sites displayed only small events involving openings of only one or two IP_3R channels, whereas others showed much larger puffs. We thus segregated the pooled measurements of puff amplitudes from Fig. 2 into two populations; 'big' sites where at least one puff with an amplitude > 0.5 F/F₀ was observed (Fig. 3B; n = 930 events, 158 sites), and 'small' sites where no events > 0.5 F/F₀ were detected (Fig. 3C; n = 367 events, 133 sites). At big sites, large puffs occurred predominantly during the initial part of the response and progressively fewer large puffs were observed as the global signal rose, although numerous small events continued (Fig. 3B). Sites that showed only small puffs did not show any consistent decline in puff amplitudes (Fig. 3C). These behaviors are summarized in Fig. 3D, plotting mean puff amplitudes from big (filled red squares) and small (filled blue squares) sites, binned at normalized time intervals during the global response. The mean amplitude of events at big puff sites declined overall by about 50% over the time course of the puff flurry. This reduction could be attributed almost entirely to a fall-off in larger (> 0.5 F/F₀), events because the mean amplitude of small (< 0.5 F/F₀) puffs at the same sites remained about constant during the flurry (open red squares, Fig. 3D), closely matching the behavior of puffs at small puff sites (filled blue squares). In contrast to

the difference in temporal evolution of puff amplitudes between big and small puff sites, the temporal evolution of overall event frequencies was similar at both types of sites (Fig. 3E,F).

3.5. Dwell-state amplitudes

The amplitude of a puff is a function of both the number of IP₃R channels that are open at the peak, and the fluorescence signal arising from Ca^{2+} flux through each open channel[22]. To better assess how the Ca^{2+} flux through individual IP₃R channels might change during the rise of global Ca^{2+} signals we measured the amplitudes of dwell-state levels (Fig. 4A) - steps during puffs that reflect differing numbers of IP₃R channels open at different times during a puff - and the amplitudes of small, 'square' events (blips) that result from opening of a single channel [20–22]. Figs. 4B,C show, respectively whole-cell Cal520 F/F₀ and SD traces obtained from imaging records in five cells, acquired at a faster frame rate than in Figs. 2 and 3 to better resolve individual fluorescence step-levels. Fig. 4D shows a scatter plot of all dwell-state amplitudes during the rise of the global responses. Although there was a marked decline in the size of the largest events, an underswell of a greater number of smaller events continued. The mean fluorescence dwell-state amplitude late in the puff flurry (from 20-25 s) was thus only modestly lower compared to the beginning (5-10 s): $F/F_0 0.254 \pm 0.009$ vs. 0.409 ± 0.027 . The mean frequency of dwell-state events matched the time course of the global SD signal, rising to a maximum during the foot of the global Ca^{2+} response and declining abruptly at about the mid-point of the global Ca^{2+} rise (Fig. 4E).

Prompted by our analysis of peak puff amplitudes at big and small puff sites (Fig. 3), we further segregated measurements of dwell-state event amplitudes from big sites where at least one puff > 0.5 F/F₀ was observed (Fig. 4F; n = 787 events, 23 sites), and small sites where no events exceeded 0.5 F/F_0 (Fig. 4G; n = 424 events, 20 sites). At sites giving large puffs, high amplitude dwell-states were most apparent early during the puff flurry, whereas numerous small events continued with undiminished amplitude (Fig. 4F); as did dwell-state amplitudes at sites that showed only small puffs (Fig. 4G). These behaviors are summarized in Fig. 4H, plotting mean dwell-state levels binned at time intervals during the global response. The overall mean amplitudes of dwell-states at big puff sites declined to about one half over the time course of the puff flurry (Fig. 4H, filled red circles). Similar to the trends observed for peak puff amplitudes this decrease could be attributed almost entirely to a fall-off in larger (> 0.5 F/F₀) dwell-states, as the mean amplitudes of smaller (> 0.5 F/F₀) dwell-states at big sites remained steady (Fig. 4H, open red circles). The mean dwell-state amplitudes at sites generating only small puffs also remained almost constant (Fig. 4H, filled blue triangles). The frequencies of dwell-state events followed similar temporal patterns at both small and big sites (Fig. 4I,J).

3.6. Limits of resolution

To determine how our ability to detect and reliably measure puffs and blips might be impaired as photon shot noise increases during global Ca^{2+} responses we generated test data by synthesizing idealized local Ca^{2+} events of various amplitudes and added these throughout an experimental record of a global Ca^{2+} response in a cell pre-treated with cyclopiazonic acid (CPA) to inhibit puff activity[16] that was of similar amplitude to the

records in Fig. 2A,H but was substantially free of any local fluctuations. The resulting image stacks were processed identically to the experimental puff data and, with the observer blinded as to the random timing of simulated events, were analyzed by automated puff detection in FLIKA (fig. S5A–E), by visual inspection (fig. S5F–J), and by using our SD routine (fig. S6A–D).

Fig. S5A shows the whole-cell Ca²⁺ fluorescence signal from the image stack to which 'square' (120 ms duration) channel-like events were added at random times with fluorescence amplitudes (F/F_0) of: 0.08, 0.12, 0.16, and 0.24 (fig. S5B–E, respectively). Although many of the smallest events tested (0.08 F/F_0) were below the algorithm's detection limit, simulated events approximating that evoked by a single IP₃R channel[20] (0.12 F/F_0) were detected by the algorithm with a 58% success rate and identified equally well at the beginning of the global response as at the peak. Simulated events of larger amplitudes (F/F_0 0.16, and 0.24) were readily resolved (84% and 95% detected, respectively). In all cases FLIKA typically overestimated the true simulated event amplitude (fig. S5B–E, dashed lines). This was expected as the algorithm was designed to detect peaks and would be biased by baseline noise spikes superimposed on the synthetic Ca²⁺ signals. Visual inspection of the same simulations detected events with similar fidelity to FLIKA, but more closely matched the simulated event amplitudes (fig. S5G–J) because, as with the data in Fig. 4, we interpolated mean amplitude measurements within noise fluctuations during the events.

Because these simulations indicated that a proportion of small ($F/F_0 < 0.2$) Ca²⁺ events might have been missed as global Ca²⁺ signals approached their peak we further tested the ability of the SD routine to report the activity of such small events. Using a similar base image stack (fig. S6A), we added randomly occurring events of two different amplitudes at 25 sites, beginning at the onset of the global response. Maximum-projection SD images clearly revealed simulated local events with amplitudes of $F/F_0 0.125$ (fig. S6C) and 0.25 (fig. S6D) during both the initial rise of the global signal and the subsequent rise to peak that were absent from simulations with no add events (fig. S6B). Moreover, corresponding traces of the SD signal measured from the whole cell showed a sustained elevation when simulated events were present throughout the global Ca²⁺ rise (fig. S6C,D), without the marked decline at the mid-point of the Ca²⁺ rise characteristic of IP₃-evoked responses like those in Fig. 2.

We thus conclude that the marked decline in puff activity around the time of half-rise of the global signal that we report does not arise artifactually from a failure to resolve putative events, or from an underestimate of event amplitudes.

3.7. Single-channel Ca²⁺ flux when ER Ca²⁺content is replete or partially depleted

In light of our finding that the rundown in puff activity during global responses is associated with declining levels of Ca^{2+} in the ER [16], we considered whether the decrease in puff amplitudes (Fig. 3D) might simply result from an attenuated driving force for Ca^{2+} efflux through IP₃R channels. We thus compared fluorescence signals arising from single channel openings (blips) under conditions when the ER Ca^{2+} levels were expected to be replete, or partially depleted. Prior treatment with the SERCA inhibitor CPA to partially deplete ER

 Ca^{2+} content activity substantially abolishes puff activity while preserving diffuse global Ca^{2+} responses[16]. In some cells treated with CPA we still observed weak punctate activity, evident as hot spots of local Ca^{2+} release in maximum intensity projections of SD image stacks (Fig. 5A) and as small whole-cell SD signals during the initial rise of the global responses (Fig. 5B). Fluorescence traces from hotspots showed discrete Ca^{2+} transients, most with amplitudes and 'square' time course (Fig. 5C) characteristic of single-channel IP₃R openings[21,22]. These events provided an opportunity to examine whether Ca^{2+} flux through single IP₃R channels was diminished when punctate activity was substantially suppressed by prior reduction of the ER Ca^{2+} content.

Fig. 5D shows the distribution of event amplitudes in three CPA-treated cells. The measurements largely cluster around a single peak and fit well to a Gaussian distribution with a mean of 0.135 F/F_0 . We interpret this value as a measure of single-channel Ca²⁺ flux in light of the 'square' appearance of these events and the similarity of their magnitudes to prior Cal520 measurements of single IP₃R channel signals in WT HEK cells (0.126 F/F_0)[20], as well as those obtained with a less sensitive indicator (0.11 F/F_0)[21,22].

To compare this estimate of single-channel Ca^{2+} flux to that when the ER Ca^{2+} content is replete or declining during IP₃-evoked global responses, we further plot in Fig. 5E,F dwell-state amplitude distributions from the cells in Fig. 4. The data in Fig. 5E were obtained during the first 8 s of the puff flurry, before the ER Ca^{2+} content would have been appreciably depleted, whereas the data in Fig. 5F were obtained during the last 16–32 s when puff activity had markedly declined. The initial peaks of these two distributions match closely with one another, and with the distribution of events in CPA-treated cells (Fig. 5D); as illustrated by the superimposed black curves which replicate the Gaussian distribution fitted to the distribution of event amplitudes in CPA-treated cells after scaling in the y-axis to match the differing numbers of observations.

A possible caveat in this comparison is that the CPA-treated cells were not loaded with EGTA. However, because buffering by EGTA is too slow to appreciably affect the amplitudes of the fast, local Ca^{2+} fluorescence transients[29] (fig. S1), we conclude that the similar fluorescence amplitudes in Fig. 5D–F indicate that single-channel Ca^{2+} flux is not appreciably diminished when the ER Ca^{2+} content is reduced by prior SERCA inhibition, or by release of Ca^{2+} during global IP₃-mediated responses.

3.8. Cumulative Ca²⁺ liberation by puffs

In EGTA-loaded cells the local fluorescence signal recorded by TIRF microscopy at puff sites closely tracks the instantaneous Ca^{2+} flux through open IP₃R channels[33,34]. Thus, the integral under local fluorescence traces of puffs provides a relative measure of the cumulative amount of Ca^{2+} liberated by punctate release during a puff flurry. We utilized two approaches to calculate integrals; by directly estimating the area under local fluorescence traces from puff sites (Fig. 6A–E), and by using automated FLIKA analysis to sum the integrals under successive individual puffs (Fig. 6F–J).

A crucial requirement for the first approach is that the baseline is stable around zero, because any systematic deviations would accumulate large errors over time. After stripping

off the global Ca²⁺ elevation using our SPS algorithm we then corrected any remaining baseline deviations (typically $F/F_0 < 0.2$) by manually fitting a 20-point spline function using the baseline tool in Microcal Origin. Figs. 6A,B illustrate representative fluorescence ratio (F/F_0) traces from three puff sites after baseline correction, together with their respective cumulative integrals. As expected, the cumulative integrals show stepwise increases associated with each puff, and plateau when puff activity stops. The F/F_0 trace in Fig. 6C shows the sum of puff activity at 11 sites in a single cell, and the red trace in Fig. 6D plots the cumulative integral under that trace, alongside the global fluorescence signal from the entire cell (black trace, normalized to facilitate comparison). Similar data from five cells (different colors) are shown superimposed in Fig. 6E, with noisier traces on the left showing the sum of cumulative integrals from multiple sites in each cell and smoother traces on the right showing the corresponding increases in global Ca²⁺ signal in that cell. To facilitate comparison between punctate Ca²⁺ release and the global Ca²⁺ signal the amplitudes of all traces are scaled to the same peak height and are further normalized in time by aligning to the occurrence of the first puff (time zero) and stretching to align the time of half-maximal rise of the global signal (normalized time t=1).

Figs. 6F–J show corresponding data obtained by our second approach, calculating the integrals under each detected puff using an automated algorithm in FLIKA. Fig. 6F illustrates data from three puff sites in the same cell, with the height of the bars representing the integral of each successive puff. The cumulative sum of these integrals (Fig. 6G) follows a pattern closely similar to that derived by integrating under fluorescence traces (Fig. 6B), as do mean cumulative integrals derived from all detected puff in a cell (Fig. 6H,I) and from multiple cells (Fig. 6J).

Both methods of analysis show that the cumulative release of Ca^{2+} via punctate activity precedes the global rise of Ca^{2+} , being substantially complete (~85%) at the time the global signal is half-maximal (Figs. 6E,J), thereby reaffirming our proposal that the subsequent rise of global Ca^{2+} is primarily driven by release from a source other than puff activity[16].

3.9. The SD signal faithfully reports Ca²⁺ flux during puffs

Although determination of cumulative integrals under puff traces provides a direct measure of punctate Ca^{2+} release, this is a technically exacting procedure. We were thus interested to determine how well these measurements compared with more facile measurements of local and global SD signals.

Fig. S7A shows a scatter plot of measurements from 52 individual puffs, plotting the amplitude of the SD signal (y-axis) vs. the peak amplitude of the local F/F_0 signal (x-axis) for the each event. The data fit well to a linear relationship, indicating that the SD signal, on average, provides a good measure of peak puff amplitude, although there is appreciable scatter among paired measurements of individual puffs. Fig. S7B compares paired measurements of cumulative punctate Ca^{2+} release obtained by integrating the areas under the whole-cell SD trace and under the F/F_0 trace, summed from multiple sites as in Fig. 6D. The time course of Ca^{2+} liberation derived by these two approaches correspond closely, although with a consistent tendency for the puff integral to approach a plateau more rapidly than the SD integral. This relationship is further illustrated in fig. S7C showing a

scatter plot of measurements in five cells of SD integral vs. puff integral. The data closely approximate a linear relationship, but with a small deviation at higher values arising because the cumulative puff integral approaches a plateau value more rapidly.

3.10. Amount of Ca²⁺ liberated during a puff flurry

Measurements of the areas under Ca²⁺ fluorescence signals during a puff flurry allow us to estimate in absolute terms (moles) the cumulative amount of Ca²⁺ liberated into the cytosol by punctate release. Under our present recording conditions, we estimate that the Ca²⁺ flux through a single open IP₃R channel generates a mean local fluorescence signal of 0.135 F/F_0 (Fig. 5D–F, and see[20–22]). Further, the Ca²⁺ current passing through a single IP₃R under physiological conditions has been estimated to be about 0.15pA[35], corresponding to a Ca²⁺ flux of 7.5×10^{-19} moles s⁻¹. Thus, if a single channel were to remain open for 1 s, the integral under the fluorescence trace (0.135 $F/F_0 * s$) would be equivalent to a cumulative liberation of 7.5×10^{-19} moles of Ca²⁺ into the cytosol.

Taking as an example the top trace in Fig. 6F,G, the cumulative integral under the puffs at this individual site was about 1.4 $F/F_0 *$ s, corresponding to the liberation of about 7.8 × 10^{-18} moles of Ca²⁺. The sum of integrals from all sites detected by the FLIKA algorithm in the cell (Figure 6H,I) was 6.56 $F/F_0 *$ s, equivalent to liberation of 3.6×10^{-17} moles Ca²⁺. From similar measurements in eight cells we obtained a mean value for punctate Ca²⁺ liberation of $4.38 \pm 0.42 \times 10^{-17}$ moles (43.8 attomoles). This number reflects activity only at those puff sites detected within the TIRF footprint of the cell. Assuming a similar density of sites across the surface of the cell[36], and that the footprint of these highly flattened cells represents about one half of the total membrane area, we therefore estimate the total punctate release of Ca²⁺ into the cytosol to be roughly 90 attomoles.

We can further estimate the minimum number of functional IP₃R channels that must be present within the TIRF field by using the peak amplitude of the largest puff observed at each site as an indication of the number of channels simultaneously open at that site. Taking the cell in Fig. 6F–I as an example, the sum of amplitudes of the greatest puff observed at each of 97 sites was 40 F/F_0 , giving an estimate for the total number of channels of 296 (again, assuming a single-channel signal of 0.135 F/F_0). The puff flurry during which these channels were active lasted about 22s (Fig. 6I) and, as calculated above, resulted in a cumulative Ca²⁺ liberation of 3.6×10^{-17} moles. This would correspond to a mean of 2.2 channels remaining continuously open throughout the 22 s flurry, giving an estimated mean open probability among the 296 channels of P_{open} = 0.007. From the 8 cells in Fig. 6J the mean P_{open} = 0.0088 ± 0.001 during puff flurries with durations ranging between 13 and 47s.

3.11. Kinetics and relative magnitudes of punctate and diffuse modes of Ca²⁺ liberation

Having established that integration of fluorescence signals at puff sites provides a measure of punctate Ca^{2+} release, we could then address the relative contributions of punctate versus diffuse modes of Ca^{2+} release during global Ca^{2+} responses. Figs. 7A,B illustrate our procedure, using data from a representative cell. We first derived the kinetics of Ca^{2+} flux into the cytosol through IP₃Rs on the basis that the change in global fluorescence signal at

any given time reflects a balance between rates of Ca^{2+} release into and out of the cytosol. To obtain a rate constant for removal of cytosolic Ca^{2+} we fitted a single exponential to the decaying fluorescence signal of Cal520 following transient photorelease of Ca^{2+} from a caged precursor loaded into the cytosol of HEK cells[16]. As illustrated in Fig. 7A, we then differentiated the whole-cell fluorescence Ca^{2+} signal (black trace) and added to this the estimated rate of Ca^{2+} removal at each time point to derive the time course of total Ca^{2+} flux into the cytosol (cyan trace). By comparison, the blue trace shows puff integrals, reflecting punctate Ca^{2+} flux into the cytosol.

To estimate the proportion of the total IP₃-evoked Ca²⁺ flux contributed by punctate release we made a simplifying assumption that the initial, abrupt increase in total flux arose almost entirely from puffs, on the basis that puff activity is high during the first few seconds of the response when the global fluorescence signal shows only a small initial rise. In Fig. 7B we thus superimpose on the total Ca²⁺ flux trace (cyan) a smoothed trace of the puff integrals (blue) after scaling the latter to overlay the initial few seconds of the total Ca²⁺ flux increase. The difference between these traces (magenta trace) represents the additional Ca²⁺ flux that we attribute to diffuse mode release. Fig. 7C shows mean data derived in this way from seven cells, plotted on a normalized timescale. The integrals under the traces provide a proportional measure of the cumulative amounts of Ca²⁺ liberated by punctate and diffuse release modes during the rise to the peak of the IP₃-mediated global response. From Fig. 7C we estimate that punctate Ca²⁺ release contributes 39.7 ± 3.3% of the total, and diffuse release the remaining 60.3%.

4. **DISCUSSION**

4.1. Punctate and diffuse modes of IP₃-mediated Ca²⁺ liberation

We had previously shown that global IP₃-mediated Ca^{2+} signals involve two modes of Ca^{2+} liberation: 'punctate' release as a flurry of transient, local puffs during the initial rise of global Ca^{2+} , with the subsequent rise sustained by a distinct 'diffuse' release mode[16]. In those experiments we monitored puff activity via a statistical, population measure of Ca^{2+} fluctuations (SD signal), and now describe recordings of individual puffs that provide further insight into the mechanisms that terminate the flurry of puff activity.

In Fig. 2 we show that the decline in puff activity and its near cessation midway during the rise of a global Ca^{2+} signal are associated with both a decrease in amplitude of individual puffs and a marked fall-off in their frequency. The former effect, however, is minor. The decline in amplitude was restricted to a few sites in each cell that generated large amplitude puffs, where the largest puffs appeared early during the flurry and then progressively diminished in amplitude. However, an underswell of small events continued with undiminished amplitudes, so that the mean puff amplitude at these 'big' puff sites declined <50% from beginning to end of the puff flurry. At other, more numerous sites that showed only small puffs, their mean amplitude of puffs declined by only about 35% during the flurry. Moreover, this modest decrease in puff amplitudes was counterbalanced by an increase in mean puff duration, such that the average amount of Ca^{2+} liberated per puff remained substantially constant throughout the flurry.

In contrast, the mean frequencies of puffs averaged across sites throughout the TIRF footprint showed a progressive rise and then abrupt fall, following a time course that closely matched the punctate activity reported by the whole cell SD signal. A large majority (75%) of puffs arose before the global Ca^{2+} signal reached half-maximal and, correspondingly, cumulative liberation of Ca^{2+} liberation through puffs was >80% complete by this time.

4.2. Amounts of Ca²⁺ liberated by punctate and diffuse release

We estimate the amount of Ca^{2+} liberated by puffs into the cytosol of a cell during a typical i-IP₃-evoked global Ca^{2+} signal to be about 90 attomoles (9 × 10⁻¹⁷ moles). This value derived from summing the durations for which individual IP₃R channels were open. It is thus independent of any assumptions of indicator affinity or presence of cytosolic Ca^{2+} buffering, but does depend on other assumptions, notably estimates of the physiological Ca^{2+} flux through a single type 3 IP₃R channel obtained in a different cell type[35]. The volume of a HEK cell is about 3000 µm³ (3 × 10⁻¹² liters or 3 femtolitres). Assuming that the cytosol and nucleosol make up one half of that volume, diffusion of the Ca^{2+} liberated by puffs would result in a Ca^{2+} concentration rise of 60 µM. Under physiological conditions we further estimate that puffs contribute only about 40% of the total IP₃-mediated Ca^{2+} liberation, giving a total Ca^{2+} concentration rise of roughly 150 µM. The peak free Ca^{2+} concentration rise for IP₃-evoked global responses in naïve HEK cells is reported to be less than 1 µM[37], suggesting a minimal endogenous Ca^{2+} buffering power of about 150; compatible with the wide range (tens to >100) reported for various other cell types[38].

4.3. Termination of puff activity

What might underlie the declines in puff amplitudes and frequencies during the rising phase of global Ca^{2+} responses? They cannot be attributed to a decay of i-IP₃ concentration, because photoreleased i-IP₃ is slowly degraded, and at low levels of i-IP₃ that evoke little rise in global Ca^{2+} puffs persist for minutes with stable amplitudes and frequency[29,39]. Based on our earlier findings that puff activity is reduced or abrogated by partial depletion of the ER Ca^{2+} content, but not by prior elevation of cytosolic free [Ca^{2+}][16], we propose that a declining concentration of Ca^{2+} in the ER lumen is the controlling factor. This then raises further questions. How may the luminal Ca^{2+} level affect the net flux through IP₃Rs at puff sites? Is the fall in luminal Ca^{2+} driven by puff activity or diffuse release?

A simple explanation for the overall decline in puff amplitudes during the rise of global Ca^{2+} could be that Ca^{2+} flux through each open IP₃R channel diminishes as a result of the reduced electrochemical driving force as Ca^{2+} levels fall in the ER lumen. Several observations, however, argue that this is not a major factor. The decrease in amplitude was restricted to large puffs that involve concerted opening of several IP₃Rs, while the mean amplitudes of small puffs at the same sites, as well as puffs at other sites that evoked only small puffs, remained steady. Similarly, measurements of the unitary Ca^{2+} signals (blips) generated by Ca^{2+} flux through openings of individual IP₃R channels[21,22], did not show any measurable change throughout the puff flurry. Moreover, although pre-treatment with CPA to partially deplete the ER Ca^{2+} content strongly attenuated puff activity evoked by photoreleased i-IP₃, the mean amplitude of remaining blips was comparable to that in control cells. We thus conclude that the reduced overall amplitude of puffs - driven by

the marked attenuation of 'large' puffs - does not arise in direct proportion to a reduced driving force for Ca^{2+} , which is expected to vary as a linear function of ER $[Ca^{2+}][40]$. Nevertheless, it is possible that a fractional decrease in single-channel Ca^{2+} flux that was too small for us to resolve might account for the decline of large puffs. Because puffs involve regenerative opening of channels via CICR[5,11,23,24] a small reduction in Ca^{2+} flux might result in an exaggerated, non-linear reduction in puff amplitude. A reduced single-channel Ca^{2+} flux would not, however, explain the fall off in frequency of either puffs or blips, as the probability of channel opening is a function of the cytosolic concentrations of Ca^{2+} and $IP_3[7,8]$.

Our simulations with synthetic blip data indicate that the near cessation of punctate Ca^{2+} release does not arise artifactually because of a detection failure as noise levels rise along with the increasing fluorescence of the Ca^{2+} signal; nor did we see any indication that puffs progressively decline during the flurry to a point where they would become undetectable. Our observations thus point to a mechanism by which the decline of puff activity during global responses is mediated primarily through modulation of the gating of IP₃Rs, rather than a diminution in the Ca^{2+} flux through open channels. In light of findings that puff activity is abrogated by lowered levels of Ca^{2+} in the ER lumen[16], and conversely that the frequency of puffs is accelerated by elevated ER Ca^{2+} levels[41], we infer that luminal $[Ca^{2+}]$ is the modulatory factor.

The issue of whether IP₃R channel activity is modulated, directly or indirectly, by $[Ca^{2+}]$ in the ER lumen is controversial[35,42,43]. Patch-clamp recordings of IP₃Rs in isolated patches of nuclear membrane showed evidence only for feed-through modulation by Ca²⁺ flux through an open channel acting on functional binding sites on the cytosolic face of the channel, without evidence for intrinsic functional Ca²⁺-modulatory sites on the luminal side of the channel[44]. However, putative accessory proteins would have been lost in the excised patch configuration, and nuclear IP₃Rs might not fully replicate the properties of the small numbers of 'licensed' IP₃Rs that generate puffs[15,20]. Regardless of specific mechanism, our observation of differential run-down of big and small puffs at the same 'big' puff sites suggests that regulation does not act the level of IP₃R clusters, but rather at the level of individual IP₃R channels within a cluster.

Recent studies identify two different, putative mechanisms of luminal Ca^{2+} regulation of IP₃Rs via interacting Ca^{2+} -sensitive proteins. Vais et al[45] describe regulation of IP₃R gating mediated by a Ca^{2+} -binding luminal protein – likely annexin A1 (ANXA1); and Emrich et al[46] report that the STIM proteins that sense ER luminal Ca^{2+} levels to regulate plasmalemmal Orai channels[47] also modulate IP₃Rs by an unknown mechanism independent of Orai activation. Notably, both mechanisms operate such that IP₃Rs are inhibited under resting conditions when the ER Ca^{2+} is replete. The presence of ANXA1 on the luminal side of excised nuclear membranes was without effect on IP₃R gating when the luminal [Ca^{2+}] was low, but the open probability of the IP₃R channel dropped abruptly as luminal [Ca^{2+}] was raised above about 50µM[45]. Similarly, Emrich et al[46] propose that unactivated STIMs inhibit IP₃R channels, and cells in which STIM1 and STIM2 were knocked out showed greater numbers of puffs and puff sites in response to photoreleased i-IP₃. It is thus difficult to reconcile either of these proposals with our findings that puff

activity becomes depressed as ER Ca^{2+} levels fall during global signals. The mechanism underlying puff termination thus remains to be elucidated. On the other hand, relief from constitutive inhibition of IP₃Rs by ANXA1 or STIM as ER Ca^{2+} levels decline might contribute to a regenerative activation of the diffuse mode of Ca^{2+} liberation.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

We thank Dr. George D. Dickinson for creating the scaled peak subtraction (SPS) and puff simulator plugins for FLIKA used in this manuscript.

Funding Sources

This work was supported by National Institutes of Health grant R37 GM048071 (IP).

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Fig. 1: Resolution of Ca²⁺ puffs during global IP₃-evoked Ca²⁺ responses.

(A-D) Records are from a single WT HEK cell loaded with Cal520 and stimulated by photorelease of i-IP₃ to evoke a global Ca^{2+} elevation. (A) Upper panels show TIRF fluorescence ratio (F/F_0) images of the cell at different times (marked by Roman numerals in B) during the rising phase of the global Ca^{2+} signal. Each panel is a single frame snapshot (8 ms exposure). Grey scale depicts fluorescence ratio intensities as indicated by the bar on the right. Middle panels show corresponding SD images at the same times, with grey scale depicting the temporal standard deviation of fluorescence intensities in arbitrary units. *Lower panels* show corresponding fluorescence ratio (F/F_0) images after processing to strip off the global fluorescence rise, using an algorithm (SPS, scaled peak subtraction) to subtract a scaled version of an averaged image captured at the peak of the response. (B) Traces show measurements from a region of interest (ROI) encompassing the entire cell of the fluorescence ratio signal (F/F_0 ; smooth black trace) and the SD signal (noisy red trace). (C) Representative traces of SD signals from two small ROIs (4×4 pixel; $\sim 2 \times 2 \mu m$) centered on puff sites. (D) Fluorescence ratio traces (F/F_0) after SPS processing from the same two ROIs. (E–H) Representative images and traces corresponding to those in A–D, but obtained in a single WT HEK cell that was additionally loaded with EGTA by incubation for 60 min with 15 µM EGTA/AM.



Fig. 2: Frequencies, amplitudes, durations and integrals of puffs during slow- and fast-rising global Ca²⁺ responses.

(A-G) Records are from 4 EGTA-loaded HEK cells that exhibited relatively slow-rising global Ca²⁺ signals in response to photoreleased i-IP₃. (A) Overlaid traces (each cell a different color) show whole-cell Cal520 F/F₀ signals. To compensate for minor differences in time course between the cells (inset), data in the main graph, and in other panels are depicted on a normalized timescale, with t=0 corresponding to the time of the photolysis flash and t=1 the time when the global Ca^{2+} signal rose to 50% of its peak in each cell. (B) Superimposed traces show corresponding whole-cell SD signals. (C) Overlaid traces of representative puff activity (Cal520 F/F₀ after SPS processing) at 12 sites in one cell. (D) Frequencies of Ca²⁺ puffs (total numbers of events from four cells per normalized 0.1 time interval) during the rise of the global signal. (E-G) Plots show measurements of individual puffs (grey) and mean values per binned time interval (black, ± 1 SEM) during the rise of the global signal. (E) Puff amplitudes, F/F_0 , measured as rise from the immediately preceding baseline to peak. (F) Puff durations, measured as time the fluorescence remained above 20% of the peak value. (G) Puff integrals, measured as the area under each Ca²⁺ puff in units of F/F₀*s. (H–N) Corresponding data from 4 cells that exhibited faster-rising global Ca²⁺ signals. Data are from 788 puffs in (D–G) and 793 puffs in (K–N). For clarity of presentation the y-axes are scaled so that a few (< 5%) individual data points are truncated.

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Fig. 3: Differing evolution of puff amplitudes at sites showing large, or only small events. (A) Representative examples of i-IP₃-evoked Ca²⁺ puffs in EGTA-loaded HEK cells, shown on a normalized timescale as in Fig. 2. Traces show SPS-corrected Cal520 fluorescence F/F_0 from 3×3 pixel ($1.5 \times 1.5 \mu m$) ROIs. The upper three traces are from sites that generated large puffs, and the lower three traces from sites where only small puffs occurred. (B) Plot of individual puff amplitudes as a function of normalized time from a subset of 158 'big' sites where at least one puff was observed with an amplitude $> 0.5 F/F_0$. (C) Scatter plot of puff amplitudes from 133 'small' puff sites where no puff was greater than 0.5 F/F_0 . (D) Mean puff amplitudes at big and small puff sites, binned at normalized time intervals throughout the global response. Filled red squares are means of all puffs at big sites; open red squares are means of puffs at big sites. Error bars are ± 1 SEM. (E,F) Frequencies of puffs (numbers of events per normalized time bin) throughout the global response plotted, respectively, for those sites exhibiting large or only small events. Data were derived from the cells illustrated in Fig. 2 and fig. S4.



Fig. 4: Dwell-state amplitude measurements.

(A) Examples of puffs and single-channel events illustrating different dwell-state levels (red arrows) of Cal520 Ca²⁺ fluorescence signal during individual puffs. (B) Superimposed traces show global Ca²⁺ signals (F/F_0) evoked in 5 HEK cells by photorelease of i-IP₃ when marked by the arrow. (C) Corresponding whole-cell SD signals. (D) Pooled data plotting all dwell-state amplitudes measured from 43 puff sites. (E) Frequency of occurrence of all dwell-state events binned at 2 s intervals throughout the global Ca²⁺ responses. (F) Pooled data plotting dwell-state levels measured from a subset of 23 sites where at least one puff was observed with an amplitude > 0.5 F/F_0 . (G) Pooled data plotting dwell-state amplitudes at those sites exhibiting large events (filled red circles) and amplitudes at these big sites excluding events > 0.5 F/F_0 (open red circles); and mean dwell state amplitudes at those sites exhibiting only small events (blue triangles). Error bars are ± 1 SEM. (I,J) Frequencies of occurrence of dwell-state events plotted, respectively, for sites exhibiting large or only small events.



Fig. 5: Amplitudes of single-channel Ca^{2+} signals when the ER Ca^{2+} content is replete or partially depleted.

(A-C) Representative data are from a HEK cell not loaded with EGTA that was exposed to CPA (50 μ M) for 1 min and then maintained in Ca²⁺-free medium for 30 min before imaging. Photorelease of i-IP₃ evoked a substantial global rise in Ca^{2+} and, although puff activity was strongly suppressed, some small, local Ca^{2+} transients were still evident. (A) Maximum intensity projections (1250 frame, 10s) from the SD image stack before the photolysis flash (top), and during the rising phase of the global signal (bottom). (B) Traces show the global F/F_0 signal (black) and SD signal (red) derived from a ROI encompassing the whole cell. Gray box indicates the time period used to generate the lower panel in A. (C) Traces show Ca^{2+} activity (F/F_0) at four of the hotspots identified in the lower panel of A, displayed on an expanded timescale beginning at the onset of the SD signal. (D-F) Plots show distribution histograms of dwell-state amplitudes recorded in different situations. The black curves are Gaussian distributions with the same mean (0.135 F/F_0) and standard deviation (0.069 F/F_0) but scaled individually in amplitude to match the data in each panel. (D) Distribution of step-level amplitudes in three cells that had been pre-treated with CPA to partially deplete the ER Ca²⁺ store content. The Gaussian fit was determined from these data. (E) Distribution of step-level amplitudes from naïve EGTA-loaded cells (the same data set as in Fig. 4E) measured during the initial 8 s following photorelease of i-IP₃. (F) Distribution of step-level amplitudes from the same sites as in E at times between 16 and 32 s following photorelease of i-IP3 when the event frequency had markedly declined.



Fig. 6: Estimating cumulative Ca²⁺ liberation via punctate release by integrating under puff traces.

Estimates of the cumulative amount of Ca²⁺ released by puffs during global responses were made by integrating under fluorescence traces from selected puff sites (A-E) and by summing the integrals under individual puffs as derived by FLIKA analysis (F–J). (A) Traces show representative puffs evoked by photorelease of i-IP₃ at three sites in the same cell. The F/F₀ signal was flattened by first applying the SPS algorithm and then correcting residual baseline fluctuations by manually fitting a 20-point spline function using the 'baseline' function in Microcal Origin. (B) Respective cumulative integrals under the three traces in A. (C) Summation of F/F_0 records from 11 sites in a different cell. (D) Red trace shows the cumulative integral under the trace in C. Black trace shows the global rise in F/F_0 signal from the whole cell. To facilitate comparison, both traces are scaled to the same peak amplitude (100%). (E) Superimposed scaled traces plotted on a normalized time scale from five cells (different colors) showing cumulative integrals of puff activity (noisier traces to the left) and global F/F₀ signals (right). (F–J) Corresponding data derived from different cells using automated measurement by FLIKA to calculate integrals under individual puffs. (F) Bars indicate the integrals of Ca^{2+} puffs at three different sites in the same cell during a global cytosolic Ca^{2+} response. (G) Respective cumulative sum of the traces in F. (H) Summation of puff integral trace measurements from all detected Ca²⁺ puffs within a single cell. (I) Red trace shows the cumulative sum of the integral trace in H, and the black trace shows the global Ca^{2+} signal, both scaled to a peak amplitude of 100%. (J) Superimposed traces plotted on a normalized time scale from eight cells (different colors) showing cumulative integrals of puff activity (traces to the left) and global F/F_0 signals (to the right).



Fig. 7: Kinetics and relative contributions of punctate and diffuse modes of Ca²⁺ release during IP₃-mediated global Ca²⁺ signals.

(A) Representative records from a single EGTA-loaded HEK cell. Traces (scaled to the same peak height) show the whole-cell Cal520 fluorescence F/F_0 (black), summed puff integral measurements from all detected Ca²⁺ puffs at the times of their occurrence (blue), and the estimated rate of Ca²⁺ efflux into the cytosol (cyan). The rate of Ca²⁺ efflux was derived by differentiating the Cal520 F/F_0 trace and adding to it an estimated rate (0.2 s^{-1}) of Ca²⁺ removal from the cytosol. (B) Traces showing Ca²⁺ fluorescence (black) and estimated Ca²⁺ efflux (cyan) are replicated from A. The blue trace depicts the summated puff integral trace from A, after strong smoothing (200 point, 1.6s, adjacent averaging) and scaling in amplitude to match the initial portion of the Ca²⁺ efflux trace. The magenta trace shows the difference between the cyan (total Ca²⁺ efflux) and blue (punctate release) traces, giving an estimate of the magnitude and time course of Ca²⁺ liberation by diffuse mode Ca²⁺ release. (C) Mean traces from seven cells showing normalized global Ca²⁺ fluorescence(black), estimated total Ca²⁺ efflux (cyan), punctate release (blue), and diffuse release (magenta).