

## Article

# Single-Molecule Tracking of Inositol Trisphosphate Receptors Reveals Different Motilities and Distributions

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**ABSTRACT** Puffs are local  $\text{Ca}^{2+}$  signals that arise by  $\text{Ca}^{2+}$  liberation from the endoplasmic reticulum through the concerted opening of tightly clustered inositol trisphosphate receptors/channels ( $\text{IP}_3\text{Rs}$ ). The locations of puff sites observed by  $\text{Ca}^{2+}$  imaging remain static over several minutes, whereas fluorescence recovery after photobleaching (FRAP) experiments employing overexpression of fluorescently tagged  $\text{IP}_3\text{Rs}$  have shown that the majority of  $\text{IP}_3\text{Rs}$  are freely motile. To address this discrepancy, we applied single-molecule imaging to locate and track type 1  $\text{IP}_3\text{R}$ s tagged with a photoswitchable fluorescent protein and expressed in COS-7 cells. We found that ~70% of the  $\text{IP}_3\text{R}$ 1 molecules were freely motile, undergoing random walk motility with an apparent diffusion coefficient of  $\sim 0.095 \mu\text{m}^2 \text{s}^{-1}$ , whereas the remaining molecules were essentially immotile. A fraction of the immotile  $\text{IP}_3\text{Rs}$  were organized in clusters, with dimensions (a few hundred nanometers across) comparable to those previously estimated for the  $\text{IP}_3\text{R}$  clusters underlying functional puff sites. No short-term (seconds) changes in overall motility or in clustering of immotile  $\text{IP}_3\text{Rs}$  were apparent following activation of  $\text{IP}_3/\text{Ca}^{2+}$  signaling. We conclude that stable clusters of small numbers of immotile  $\text{IP}_3\text{Rs}$  may underlie local  $\text{Ca}^{2+}$  release sites, whereas the more numerous motile  $\text{IP}_3\text{Rs}$  appear to be functionally silent.

## INTRODUCTION

Inositol trisphosphate receptors ( $\text{IP}_3\text{Rs}$ ) are  $\text{Ca}^{2+}$ -permeable channels in the membrane of the endoplasmic reticulum (ER) (1). They liberate  $\text{Ca}^{2+}$  sequestered in ER stores to generate cytosolic  $\text{Ca}^{2+}$  signals that control ubiquitous cellular functions as diverse as gene expression, secretion, and synaptic plasticity (2). The specific regulation of cellular functions is achieved through an exquisite spatiotemporal patterning of cytosolic  $\text{Ca}^{2+}$  signals (2) that arises from the properties of the  $\text{IP}_3\text{Rs}$  and their cellular organization. First, the gating of  $\text{IP}_3\text{R}$  channels is biphasically regulated by cytosolic  $\text{Ca}^{2+}$  itself. Small  $\text{Ca}^{2+}$  elevations (in the presence of  $\text{IP}_3$ ) induce channel openings via  $\text{Ca}^{2+}$ -induced  $\text{Ca}^{2+}$  release (CICR), whereas higher concentrations cause inactivation, thereby creating positive- and negative-feedback loops (3). Second, functional  $\text{IP}_3\text{Rs}$  are arranged in a clustered fashion on the ER membrane so that  $\text{Ca}^{2+}$  release through a single channel (known as a blip) can trigger the opening of neighboring channels within a cluster to generate local  $\text{Ca}^{2+}$  signals ( $\text{Ca}^{2+}$  puffs) (4). In turn, recruitment of multiple puff sites by successive cycles of  $\text{Ca}^{2+}$  diffusion and CICR can propagate saltatory  $\text{Ca}^{2+}$  waves throughout the cell (5,6). Thus, the localization of  $\text{IP}_3\text{Rs}$  is crucial for establishing and optimizing the hierarchical spatiotemporal patterning of cytosolic  $\text{Ca}^{2+}$  signals to ensure appropriate regulation of downstream signaling pathways (7).

$\text{Ca}^{2+}$  imaging studies in mammalian cell lines and in *Xenopus* oocytes have demonstrated that puffs arise repeatedly over many minutes at locations that remain essentially fixed within a cell, suggesting that the underlying clusters of  $\text{IP}_3\text{Rs}$  on the ER membrane are relatively stable entities (8–11).  $\text{Ca}^{2+}$  blips generated by lone  $\text{IP}_3\text{Rs}$  are similarly immotile (12,13). In marked contrast to the immotility of  $\text{IP}_3\text{Rs}$  as monitored by the  $\text{Ca}^{2+}$  signals they generate, studies of fluorescence recovery after photobleaching (FRAP) of GFP-tagged  $\text{IP}_3\text{R}$  proteins indicate that a substantial proportion of  $\text{IP}_3\text{Rs}$  are freely diffusible within the ER membrane (14–22). Moreover,  $\text{IP}_3\text{R}$  proteins have been reported to aggregate into clusters following sustained (minutes) activation of  $\text{IP}_3$  signaling and/or cytosolic  $\text{Ca}^{2+}$  elevation (14,19–23), and even to undergo clustering in response to  $\text{IP}_3$  within just a few seconds (24). There is also a marked disparity between immunostaining patterns that show  $\text{IP}_3\text{Rs}$  densely distributed throughout the entirety of the ER in mammalian cells (14,20,22) and observations that  $\text{Ca}^{2+}$  release arises at just a few discrete puff and blip sites (9–12,25–27).

To account for these apparent differences in the density, distribution, and motility of  $\text{IP}_3\text{Rs}$  as assessed by functional  $\text{Ca}^{2+}$  imaging versus protein localization, we previously hypothesized that a majority of  $\text{IP}_3\text{Rs}$  may be motile but are either functionally unresponsive or mediate  $\text{Ca}^{2+}$  liberation only during sustained global elevations of cytosolic  $[\text{Ca}^{2+}]$  (12,28). We further proposed that local  $\text{Ca}^{2+}$  signals arise from a much smaller subset of  $\text{IP}_3\text{Rs}$  that are anchored individually or in tight clusters by association with static

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cytoskeletal structures. Possibly as a consequence of this anchoring, these IP<sub>3</sub>Rs display a high sensitivity to IP<sub>3</sub> and are uniquely able to generate Ca<sup>2+</sup> blips and puffs (28).

Here, we sought to examine this hypothesis by using photoconvertible, genetically encoded proteins to track the motility of thousands of individual IP<sub>3</sub>R molecules with nanometer spatial resolution using single-particle-tracking photoactivation localization microscopy (sptPALM) (29). We expressed type 1 IP<sub>3</sub>Rs tagged with the photoconvertible fluorescent protein mEos2 in COS-7 cells. By using stochastic activation of mEos2 molecules at low density, we were able to track IP<sub>3</sub>Rs with high precision by fitting a two-dimensional Gaussian function to determine the centroid of the blurred spots (point-spread function) formed by the microscope in successive image frames. We found that the IP<sub>3</sub>Rs could be classified into two groups with relatively high or low/zero motility, and that the immotile IP<sub>3</sub>Rs appeared to be preferentially grouped within tight clusters. Further, we report that agonist activation of IP<sub>3</sub>/Ca<sup>2+</sup> signaling caused no appreciable short-term (seconds) changes in the motility or distribution of IP<sub>3</sub>Rs.

## MATERIALS AND METHODS

### Plasmid construction

The type 1 IP<sub>3</sub>R construct was a gift from Kevin Foskett. The IP<sub>3</sub>R1 was cloned into the EcoRI/NotI restriction sites of pcDNA3.1/Zeo(+), yielding pcDNA3.1-IP3R1. mEos2 was then tagged onto the N-terminus (20,30) of the IP<sub>3</sub>R1 by cloning into the multiple cloning site of pcDNA3.1-IP3R1 using the NheI/EcoRI restriction site. We named this final construct mEos2-IP3R1.

### Cell culture

COS-7 cells (ATCC Manassas, VA) were cultured according to the supplier's instructions in Dulbecco's modified Eagle's medium supplemented with 10% (v/v) fetal bovine serum (FBS). Cells were incubated at 37°C in a humidified incubator gassed with 95% air and 5% CO<sub>2</sub>, passaged every 3–4 days, and used for up to 20 passages. For expression of mEos2-IP3R1 in COS-7 cells, 5 × 10<sup>5</sup> cells were harvested and transfected using a Nucleofector (Kit V, program A-024; Lonza, Basel, Switzerland) or Neon Transfection System (Life Technologies, Carlsbad, CA). Transfected cells were plated at 5 × 10<sup>4</sup> cells/ml in Petri dishes with glass coverslips as the base (MatTek Ashland, MA). Cells were imaged 24–72 h later in HEPES-buffered saline (HBS; (in mM) NaCl 135, KCl 5, MgCl<sub>2</sub> 1.2, CaCl<sub>2</sub> 2.5, HEPES 5, glucose 10) at room temperature. For fixed-cell experiments, cells were rinsed in PBS, fixed for 15 min in 4% paraformaldehyde (PFA) at room temperature, and subsequently washed three times in PBS before imaging.

### Live-cell microscopy

Cells were imaged at room temperature (~21°C) using a custom-built total internal reflection (TIRF) microscope system based on an Olympus IX 70 microscope equipped with an Olympus 100× SAPO objective (NA 1.40). Activation and excitation lasers illuminated the sample in semi-TIRF mode, with the angle of light exiting the objective adjusted to create a light sheet restricted to the bottom few micrometers of the specimen. Images of emitted fluorescence were captured by an Evolve 512 electron-multiplied CCD camera (Roper Scientific Tucson, AZ) at a resolution of 256 × 256

pixels (1 pixel = 0.166 μm) at an exposure time of 150 ms (6.67 frames s<sup>-1</sup>). The camera was operated in quantitative mode, where one count = one detected photon. Cells expressing mEos2-IP3R1 were identified by their green fluorescence using 488 nm excitation and a longpass emission filter. A 561 nm laser was then used to preferentially excite the red, photo-switched form of mEos, which was imaged using a 568 nm sharp-edge longpass filter. Preexisting red fluorescence was bleached by exposing the cell to high-intensity 561 nm excitation and the intensity was then reduced (to ~100 W/cm<sup>2</sup>) to track single molecules. In some instances, spontaneous photoswitching of mEos provided a sufficient density of red fluorescent molecules. In other cases, continuous or pulsed activation by a 405 nm laser was used, with the intensity adjusted to give an appropriate density of molecules. Typically, the density of fluorescent spots in a given frame was ~1 per 30 μm<sup>2</sup>. Sequential image frames were captured using MetaMorph 7.5 (Molecular Dynamics Pittsburgh, PA) to accumulate continuous image stacks, which typically comprised 5000 frames (~12½ min).

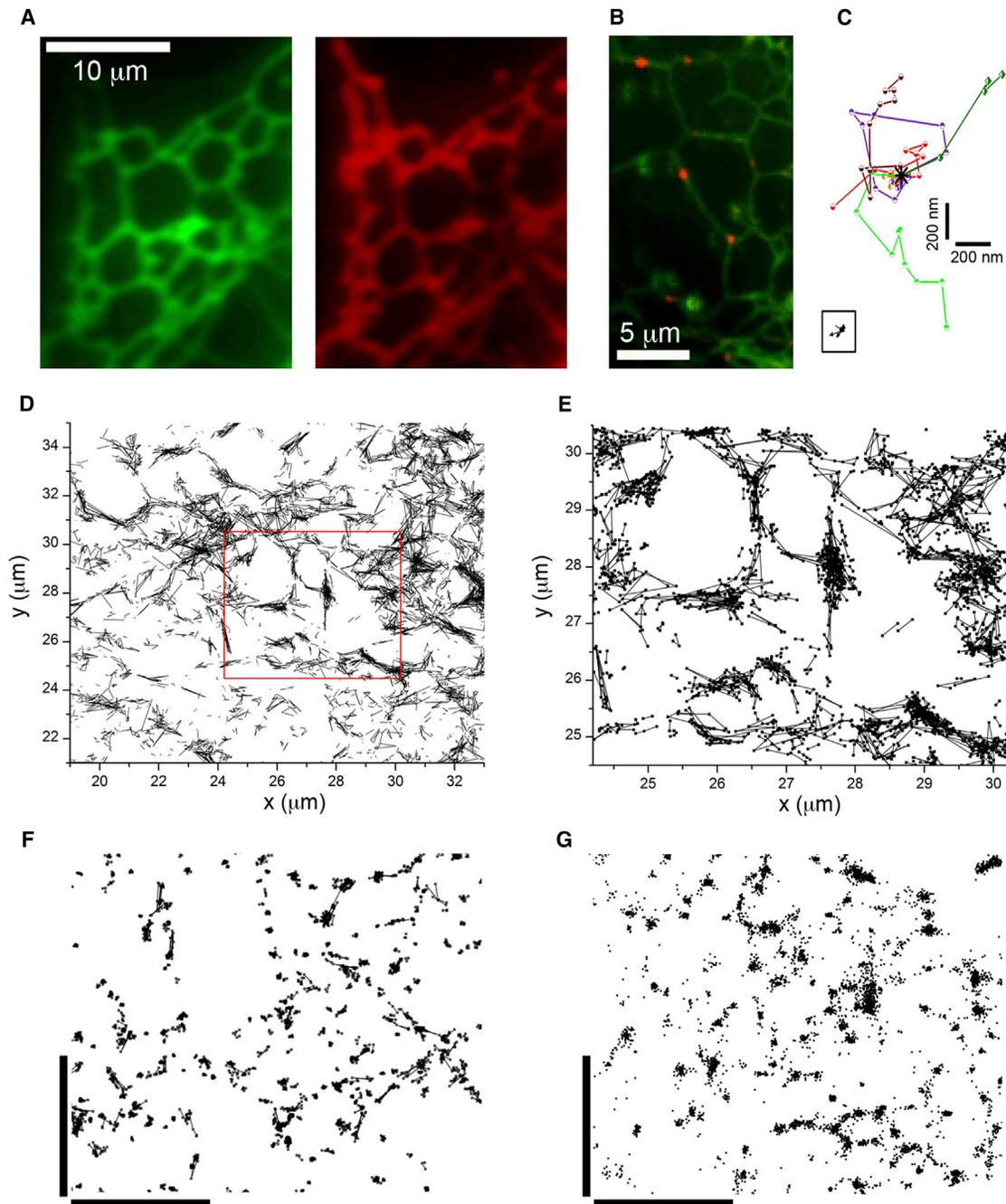
### Superresolution tracking and data analysis

Image stacks were imported into Insight3 (custom software provided by X. Zhuang, Harvard University) to track the motions of individual molecules. The program identified valid fluorescent spots (molecules) within each image frame on the basis of their intensity (>250 photoelectrons) and spatial width (200–600 nm), and localized them with subpixel precision by fitting a two-dimensional circular Gaussian function to the diffraction-limited spot. Molecular tracks were then assigned by linking a spot in one frame to the closest spot within a defined search radius in the following frame. The search radius was generally set to 4 pixels (664 nm) to minimize the possibility of false linkages between different adjacent molecules while being wide enough to encompass almost all Brownian displacements of a single molecule. Insight3 thus produced a data set of tracked molecules, listing sequential *x-y* coordinates during the track and the corresponding frame numbers (times) when they were observed.

Further analysis of the data set was performed in MATLAB (The MathWorks, Natick, MA) using custom-written routines. For each track, we calculated the individual displacements between successive frames (single-lag displacements (SLDs)), as well as the mean location of the molecule averaged across its track. Data were filtered to exclude continuous tracks that persisted for fewer than four frames or longer than 40 frames. To further account for the blinking behavior of mEos, we applied a custom-written MATLAB algorithm to link together temporally (1–25 frames) and spatially (<167 nm) adjacent track segments of selected immotile molecules so that blinking molecules would not be overcounted.

### STORM imaging of native IP<sub>3</sub>Rs

We performed superresolution localization of native type 1 IP<sub>3</sub>Rs in COS-7 cells (Fig. 1 G) by employing STORM imaging (31) utilizing an anti-IP<sub>3</sub>R 1 (AB5882 (Millipore Billerica, MA), a rabbit polyclonal antibody raised to a synthetic peptide corresponding to amino acids 2732–2750 of rat IP<sub>3</sub>R1)) and a secondary antibody generated by dual labeling of donkey-anti-rabbit IgG (Jackson ImmunoResearch West Grove, PA) with Cy3 (mono-reactive dye pack; GE Healthcare Pittsburgh, PA) and Alexa Fluor 647 (Invitrogen Grand Island, NY). Cells were fixed and reduced by incubation for 15 min at room temperature in PBS containing 3% PFA and 0.1% glutaraldehyde (GA), followed by 7 min in a freshly prepared solution of 0.1% NaBH<sub>4</sub> in PBS (without additional Ca<sup>2+</sup>/Mg<sup>2+</sup>). The cells were rinsed quickly (×3) with PBS, permeabilized by incubation for 1 h in blocking buffer (3% (m/v) bovine serum albumin (BSA), 0.2% TritonX-100 (v/v) in PBS), and probed with the anti-IP<sub>3</sub>R 1 antibody (1:100 dilution in blocking buffer) for 1 h. The cells were quickly rinsed with wash buffer (0.2% (m/v) BSA, 0.05% TritonX-100 (v/v) in PBS), washed (×3) for 5 min with PBS, and fluorescently labeled by incubation with the Cy3-Alexa Fluor 647-conjugated secondary antibody (1:100 dilution in blocking buffer) for 1 h.



**FIGURE 1** Expression of mEos-tagged IP<sub>3</sub>R1 in COS-7 cells for single-molecule tracking. (A) Conventional fluorescence image showing the recticular distribution of tagged IP<sub>3</sub>R1 in the periphery of a COS-7 cell (*left*) and a pseudocolored image of the same region of the cell showing fluorescence of the dye ER tracker blue/white (*right*). (B) Overlay showing autofluorescence of the ER (*green*) with fluorescence of sparsely photoactivated mEos molecules in red. (C) An overlay of seven representative tracks (*different colors*) of mEos IP<sub>3</sub>R1s in live COS-7 cells, each localized over nine consecutive image frames (1.35 s) and plotted after aligning their starting positions (marked by *asterisk*). The inset trace (*boxed*) shows a representative track of an mEos IP<sub>3</sub>R1 in a fixed cell. (D) Overlay of >5000 mEos IP<sub>3</sub>R1 tracks in the periphery of a COS-7 cell. (E) Enlarged view of the region marked in red in (D). (F) Corresponding overlay of mEos IP<sub>3</sub>R1 localizations in a fixed COS-7 cell. (G) STORM superresolution image of native IP<sub>3</sub>R1 in the periphery of a control, nontransfected COS-7 cell. The plot depicts drift-corrected fluorophore localizations derived from a cell immunostained with a primary antibody raised against IP<sub>3</sub>R1 and a secondary antibody labeled with a Cy3-Alexa Fluor 647 dye pair. Bars in *F* and *G* = 2  $\mu$ m. To see this figure in color, go online.

Finally, the cells were rinsed with wash buffer and PBS (as previously), and fixed again by incubation with 3% PFA and 0.1% GA for 5 min. Imaging was carried out on a Nikon N-STORM system based on a Nikon Eclipse Ti microscope equipped with a Nikon 100 $\times$  APO TIRF objective (NA

1.49). In semi-TIRF mode, a 647 nm laser continually illuminated the cells, driving the Alexa Fluor 647 dye into a nonfluorescent state. Sparse single-molecule reactivation of individual Alexa Fluor 647 molecules was achieved by weak excitation of Cy3 using a 561 nm laser. The imaging

buffer was composed of 50 mM Tris, pH 8, 10 mM NaCl, 10% glucose, 0.14 mg/ml glucose oxidase (Sigma), 0.2 mg/ml bovine liver catalase (Sigma), and 0.14 M  $\beta$ -mercaptoethanol. Fifteen thousand frames were recorded from the center quad of an Andor iXon3 electron-multiplying charge-coupled device (EMCCD) camera with an exposure time of 15 ms per frame. Single-molecule fitting and drift correction were performed with the N-STORM software in NIS Elements (version 3.1.0.11901). Localization coordinates were exported into Microcal Origin Version 9.1 (OriginLab, Northampton, MA) for graphing.

## RESULTS

### Expression and imaging of fluorescently tagged IP<sub>3</sub>R molecules

To track single IP<sub>3</sub>Rs, we employed COS-7 cells, which flatten close to the coverglass, allowing the ER to be visualized in the cell periphery as an extended reticulum forming a nearly two-dimensional, fishnet-like sheet (Fig. 1, A and B). Cells were transfected 24–72 h before imaging with type 1 IP<sub>3</sub>Rs tagged with mEos2, a photoswitchable fluorescent protein that initially fluoresces green but can be switched to a red fluorescent form by weak illumination with near-UV (405 nm) light. Transfected cells were identified by their green fluorescence when excited by blue (488 nm) laser light. The pattern of green mEos2 fluorescence (Fig. 1 A, left) closely overlapped that of an ER marker (Fig. 1 A, right). After selecting a region of cell to image, we bleached out preexisting red Eos using strong 561 nm laser irradiation, and then visualized the fluorescence of individual red mEos molecules using a lower-power (561 nm) excitation and a 568 nm longpass emission filter (Fig. 1, B and C). The camera exposure time (150 ms) was chosen as a compromise between blurring of the single-molecule fluorescence spot by Brownian motion within a frame and allowing a measurable displacement between successive frames. [Movie S1](#) in the [Supporting Material](#) shows the image sequence from which Fig. 1 C was obtained.

We typically acquired image stacks of 5000 sequential frames (750 s) using the center quad of the camera chip (250 × 250 pixels; 167 nm/pixel at the specimen). Spontaneous photoswitching of mEos often provided a sufficient density of fluorescent molecules, and in other cases continuous or pulsed weak illumination with 405 nm light was used to induce photoswitching at a rate controlled by the intensity of the activating light.

### Tracking single IP<sub>3</sub>Rs

Molecules were identified, localized, and tracked by offline analysis of the image stacks using the Insight3 software package to generate a database of unique tracks with  $x, y, t$  coordinates for each position of a given molecule. Images were obtained from the cell periphery, where COS-7 cells flatten to a thickness of only ~800 nm. We thus analyzed the tracking data in only two ( $x, y$ ) spatial dimensions.

A molecule could typically be tracked for several consecutive frames (mean =  $6.34 \pm 0.22$  frames,  $n = >30,000$  tracks) before it reverted to a dark state. A majority of molecules in live cells showed apparently random displacements, with average track lengths of several hundred nanometers (*upper traces*, Fig. 1 C). A map obtained by superimposing all tracks from an image stack traced out the reticular pattern of the ER (Fig. 1, D and E). In contrast to this, localizations of individual IP<sub>3</sub>R molecules in cells after paraformaldehyde fixation were tightly constrained (*inset trace*, Fig. 1 C), although their distributions again mapped out a reticular pattern (Fig. 1 F).

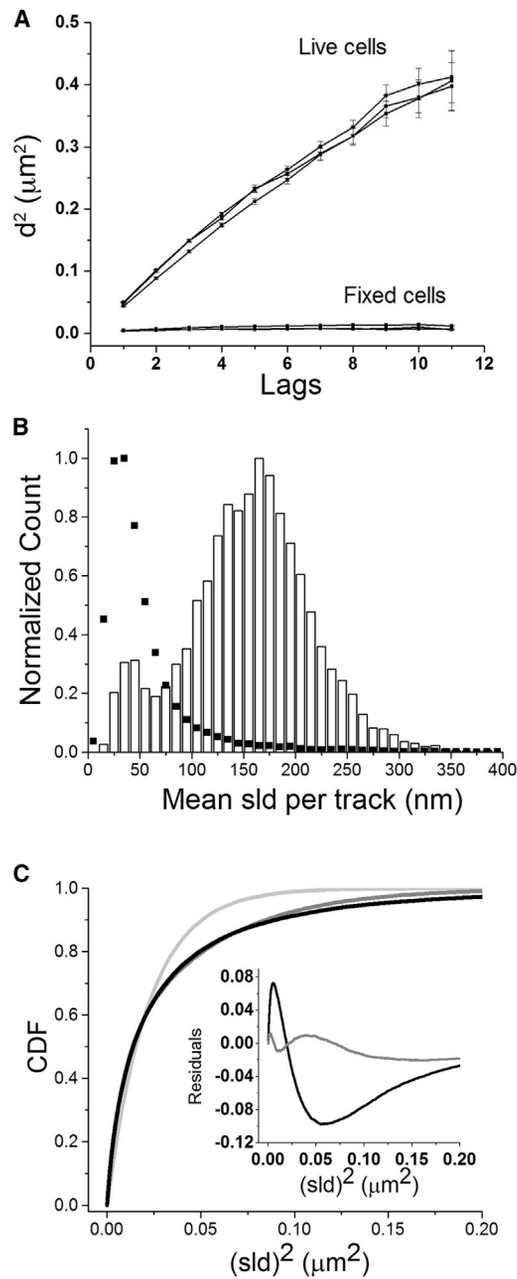
To confirm that the overexpressed, Eos-tagged IP<sub>3</sub>Rs showed a subcellular distribution similar to that of native IP<sub>3</sub>Rs, we further undertook superresolution STORM imaging of IP<sub>3</sub>Rs immunostained with a type 1-specific primary antibody and utilizing a secondary antibody optimized for single-molecule localization by dual labeling with Cy3 and Alexa Fluor 647 (32). The pattern of IP<sub>3</sub>R1 staining in fixed, control (nontransfected) COS 7 cells (Fig. 1 G) closely resembled that of expressed Eos-tagged IP<sub>3</sub>Rs in fixed cells (cf. Fig. 1 F).

To analyze the tracking data, we calculated the displacements  $d$  of molecules from any given point during their track to the point reached after a time  $t$  (expressed as the number of lags at a frame interval of 150 ms). The SLD is the displacement during a single lag (i.e., from one frame to the next), and the mean SLD is the average SLD across the entire length of a single-molecule track.

### Diffusional motility of IP<sub>3</sub>R

Fig. 2 A shows plots of the mean-squared displacements (MSDs: mean  $d^2$ ) of IP<sub>3</sub>R molecules in COS-7 cells as a function of time (lags). Square symbols show data from three representative experiments with live cells. Consistent with IP<sub>3</sub>R molecules undergoing random walk (Brownian) motion, the MSD increased approximately linearly with time (33). The slope of a linear fit to an MSD-versus-time plot provides a measure of the diffusion coefficient  $D$ . Assuming that diffusion in the flattened sheet of ER in COS-7 cells approximates a two-dimensional situation,  $D = \langle d^2 \rangle / 4t$ , where  $\langle d^2 \rangle$  is the MSD at time  $t$ . This gave an aggregate mean value of  $0.064 \pm 0.002 \mu\text{m}^2 \text{s}^{-1}$  for the live-cell data in Fig. 2 A. In contrast to this, data from fixed cells (*open symbols*, Fig. 2 A) showed little or no consistent increase in MSD with time, yielding an apparent diffusion coefficient of  $<0.0003 \mu\text{m}^2 \text{s}^{-1}$ .

Aggregate diffusion coefficients measured in live cells at cooler temperatures (10–12°C,  $n = 7$  cells) were only modestly lower than at 21°C ( $0.049 \pm 0.003$  and  $0.069 \pm 0.005 \mu\text{m}^2 \text{s}^{-1}$ , respectively;  $n = 14$ ). This slight temperature dependence ( $Q_{10} = 1.40$ ) is consistent with passive diffusion, whereas an active transport mechanism



**FIGURE 2** Diffusional characteristics of IP<sub>3</sub>R1 derived from single-molecule tracking. (A) Plot showing the square of the mean distance ( $d^2$ ) traveled by molecules from any given starting point as a function of lag time (1 lag = 1 frame interval of 150 ms). The data shown are from experiments in three live COS-7 cells (squares), with >2000 tracks in each case. Circles show corresponding data from recordings from three cells after fixation. (B) Distribution of mean SLDs from selected tracks longer than eight lags. Each count (y axis) represents a track with a given mean SLD. Open histogram bars show pooled data from six experiments in live COS-7 cells (3838 tracks). Black symbols show corresponding measurements from four experiments on fixed COS-7 cells (~25,000 tracks). Both data sets are normalized to the same peak height. (C) Plot showing the CDF of mean-squared SLDs ( $SLD^2$ ) from a representative experiment. The solid black curve shows the cumulative fraction of observations (y axis) for which the  $SLD^2$  exceeded a given value (x axis). Curves are single-exponential (light gray) and double-exponential (dark gray) fits to the data (black). The inset shows residual errors remaining after subtracting the single (black trace)- and double (grey trace)-component exponential fits from the data.

would be expected to display a much stronger temperature dependence ( $Q_{10} \sim 3$ ) (34)).

### Multiple components of IP<sub>3</sub>R diffusion

Diffusion coefficients derived as in Fig. 2 A for IP<sub>3</sub>R in live cells are a population aggregate from all tracked molecules. To determine whether multiple underlying components represent populations of IP<sub>3</sub>R in live cells with different diffusional motilities, we first analyzed the distribution of mean SLDs of individual molecules and then performed two-component fits to the cumulative density functions (CDFs) of molecular displacements.

Fig. 2 B shows the distribution of mean SLDs (histogram bars), which we pooled from 3838 tracks in six live COS-7 cells after selecting long (9–40 frames) tracks so as to obtain more precise estimates of the mean SLD of each track. The data show two clearly separable components with mean SLDs centered around 46 and 169 nm, corresponding to respective diffusion coefficients of 0.0035 and 0.0476  $\mu\text{m}^2 \text{s}^{-1}$ . For comparison, we overlay in Fig. 2 B the distribution of mean SLDs measured in fixed cells (solid symbols). This closely overlaps the slow component observed in live cells, indicating that the slow component represents a population of essentially immotile IP<sub>3</sub>Rs whose SLDs largely reflect jitter in the experimental localizations rather than true movements of the molecules.

We further analyzed the CDF of SLDs and multiple-lag displacements in live COS-7 cells, taking track lengths of 4–40 frames. Fig. 2 C shows representative data from one experiment, illustrating the CDF of SLDs. In the case of a single population of diffusing molecules, the CDF is expected to follow a monoexponential function (35,36) where the exponential fit parameter is a measure of MSD  $d^2$  and gives a measure of the effective diffusion coefficient  $D = d^2/4t$ .

The experimental data could not be fit adequately by a single exponential component, but were fit well (but not perfectly) by two exponential components, indicating two distinct diffusing populations with different diffusion coefficients and relative weights. Two-component exponential fits to the SLD data illustrated in Fig. 2 C corresponded to diffusion coefficients of 0.093 and 0.0075  $\mu\text{m}^2 \text{s}^{-1}$ , with respective weights of 69% and 31% for these motile and relatively immotile components.

To obtain more precise values, we fitted two-component exponentials to CDFs calculated for all lags between one and 10, and averaged the values obtained within each experiment. Data from six experiments yielded mean diffusion coefficients of  $0.095 \pm 0.0057$  and  $0.0075 \pm 0.0005$   $\mu\text{m}^2 \text{s}^{-1}$ , with respective weights of  $69\% \pm 1.2\%$  and  $31\% \pm 1.2\%$  for these motile and relatively immotile components. The fraction of motile receptors was similar among all six experiments (range: 64–71%), although the expression level of tagged IP<sub>3</sub>Rs as assessed by the total

number of identified tracks varied ~3-fold (range: 2835–9136 tracks per experiment).

### Spatial segregation of motile and immotile IP<sub>3</sub>Rs

We were then interested in determining the respective spatial distributions throughout the ER of the populations of IP<sub>3</sub>R with different motilities. Based on the results in Fig. 2, B and C, we defined immotile molecules as those showing tracks with mean SLDs of <84 nm (i.e., similar to those seen for molecules in fixed cells and within our localization jitter). Correspondingly, we selected rapidly motile molecules as having mean SLDs of >200 nm.

Fig. 3 A shows a representative overlay of the locations (mean position) of immotile IP<sub>3</sub>Rs (*red dots*) and rapidly motile IP<sub>3</sub>Rs (*black dots*) in the periphery of a COS-7 cell. From visual inspection, it appears the immotile IP<sub>3</sub>Rs

do not closely colocalize with motile IP<sub>3</sub>Rs. To validate this subjective impression, we measured distances from every immotile IP<sub>3</sub>R to the closest neighboring motile IP<sub>3</sub>R, and distances from every motile IP<sub>3</sub>R to the closest neighboring motile IP<sub>3</sub>R. The distribution of immotile-to-motile nearest-neighbor distances showed a highly skewed distribution with a pronounced tail at long distances (*red*, Fig. 3 B). Although roughly two-thirds of immotile molecules were located within <500 nm of a motile IP<sub>3</sub>R, the remainder were located >500 nm and as far as 5 μm away. For comparison, the nearest-neighbor distances between pairs of motile IP<sub>3</sub>Rs showed a much tighter distribution (*black*, Fig. 3 B). Almost all motile IP<sub>3</sub>Rs were located within <1 μm of a motile neighbor, and only 13.1% were >500 nm away from a neighbor.

### Clusters of immotile IP<sub>3</sub>Rs

In light of the immotility of clusters of functional IP<sub>3</sub>Rs underlying Ca<sup>2+</sup> puffs (8,10–12), we sought to determine whether the population of immotile mEos-tagged IP<sub>3</sub>Rs might show a similarly clustered spatial organization, as distinct from a more diffuse distribution of motile IP<sub>3</sub>Rs.

This analysis was complicated by the fluorescence blinking behavior of mEos molecules (37). That is to say, after becoming photoactivated, a single mEos molecule could switch repeatedly between fluorescent and dark states within a few seconds before irreversibly bleaching. This would lead to artifactual identification of apparent tight clusters if a single immotile molecule were falsely identified as several distinct, closely localized molecules (38,39). To overcome this problem, we further processed the tracking data from immotile molecules (defined as mean SLD ≤ 84 nm) using a custom-written MATLAB algorithm to link together temporally (0.15–3.75 s) and spatially (<167 nm) adjacent track segments so that blinking molecules would not be overcounted.

After linking the tracks, we calculated the mean position of each molecule and measured the distance from every molecule to every neighboring molecule within an 8 × 8 pixel (1.3 × 1.3 μm) search box. In the case of a uniformly random spatial distribution, the number of neighboring molecules present at a given distance increases linearly with increasing distance (Fig. 4 A). The distribution of intermolecular distances of motile IP<sub>3</sub>Rs (Fig. 4 B) was closely similar to this linear relationship, as would be expected if their Brownian motion caused them to be randomly distributed throughout the ER. In marked contrast, the spacings between immotile IP<sub>3</sub>Rs (after linking the tracks) showed a considerable excess at short (<400 nm) distances (Fig. 4 C). This is further illustrated in Fig. 4 D, which shows the difference between the observed distribution of spacings between immotile IP<sub>3</sub>Rs and the distribution that would be expected if they were randomly distributed. Thus, a fraction of the immotile

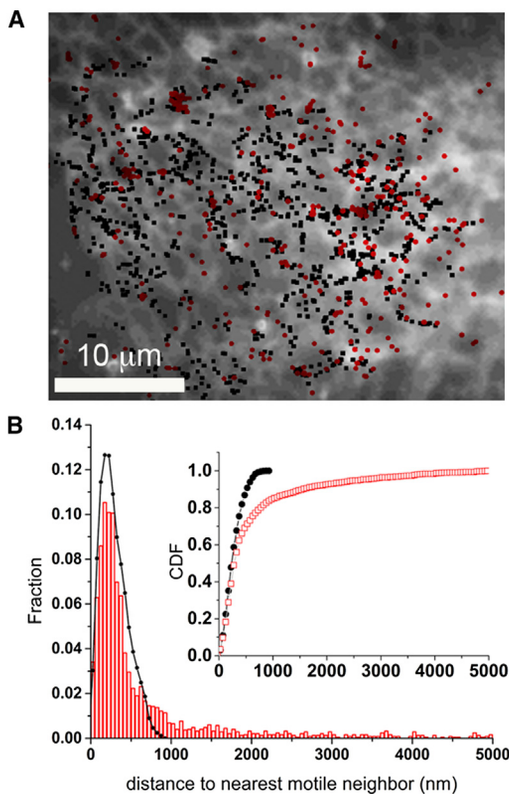


FIGURE 3 Spatial segregation of immotile and rapidly motile IP<sub>3</sub>Rs. (A) The background image is the prebleach fluorescence of a COS-7 cell expressing mEos-tagged IP<sub>3</sub>R1, showing the distribution of IP<sub>3</sub>Rs across the ER. Superimposed on this are the localizations (mean positions of tracks) of immotile (*red dots*) and rapidly motile (*black dots*) IP<sub>3</sub>R molecules. (B) The main graph plots the distributions of nearest-neighbor distances between each immotile molecule and its nearest motile molecule (*red bars*), and nearest-neighbor distances between pairs of motile molecules (*black circles*). The y axis is the fraction of molecules at a given distance from their nearest neighbor. The inset replots the same data as a CDF, i.e., the cumulative fraction of molecules with nearest-neighbor distances shorter than a given value. To see this figure in color, go online.

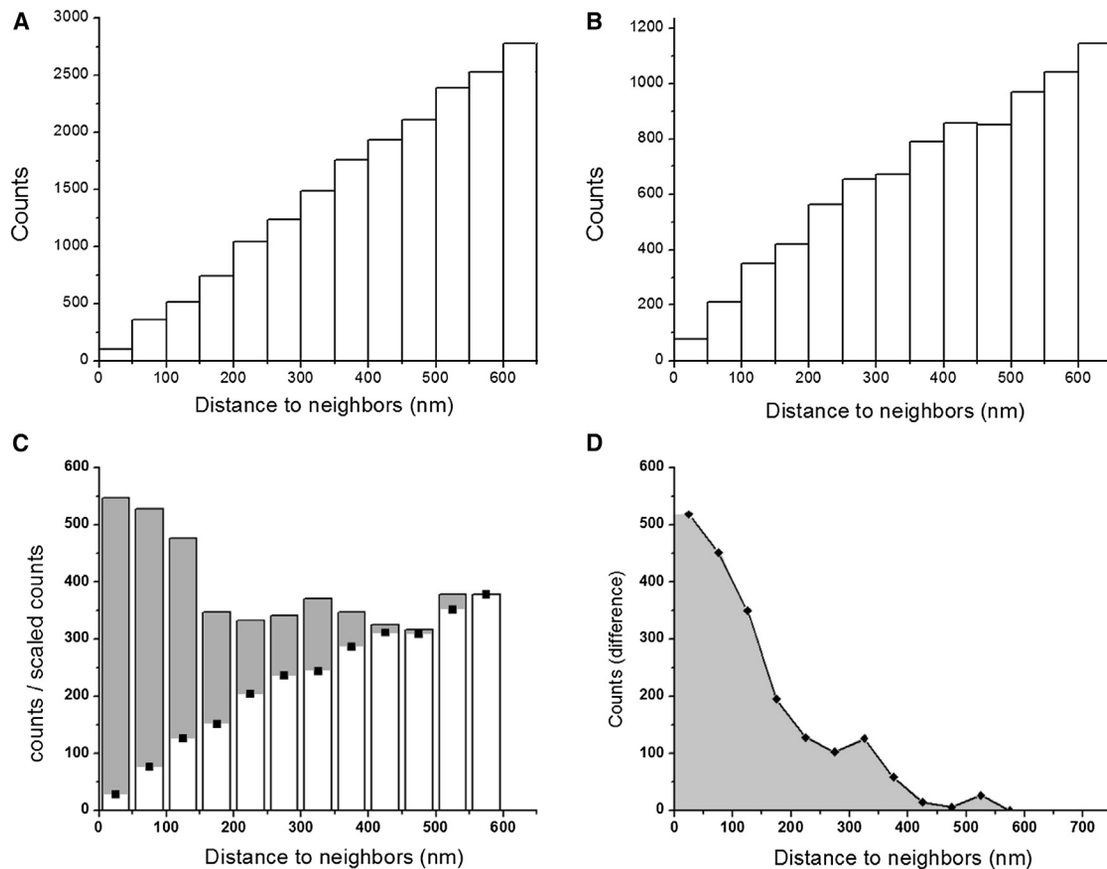


FIGURE 4 Cluster analysis of IP<sub>3</sub>R1 distribution. Panels show counts of instances in which two molecules were located within a given distance of one another. (A) Simulation showing the linear distribution of interparticle distances measured from a random spatial spread of 5000 particles. (B) Distribution of distances between motile IP<sub>3</sub>Rs (mean SLDs >200 nm) in live COS-7 cells. Molecule locations were defined as the mean position along each track, and data are from 9535 tracks in six experiments. (C) Histogram bars show the corresponding distribution of intermolecular distances between immotile IP<sub>3</sub>Rs. Black squares show the distribution of motile molecules, reproduced from (B) and scaled in amplitude to facilitate comparison. Gray shading indicates the excess numbers of immotile IP<sub>3</sub>Rs at close interneighbor distances above that expected for a random distribution. (D) Difference between the observed distribution of intermolecular distances of immotile IP<sub>3</sub>Rs and the scaled distribution of motile IP<sub>3</sub>Rs.

IP<sub>3</sub>Rs appear to be tightly grouped within <400 nm of neighboring, immotile IP<sub>3</sub>Rs.

### Nanoarchitecture of immotile IP<sub>3</sub>R clusters

We then identified putative clusters of immotile IP<sub>3</sub>R1 by filtering the data set to include only those molecules that met all of the following criteria: 1), IP<sub>3</sub>Rs with mean SLDs  $\leq 84$  nm; 2), molecules with  $\geq 2$  immotile neighbors within a 700 nm search radius; and 3), a mean distance of  $\leq 250$  nm to these surrounding neighbors. A representative spatial map of clusters identified in this way within the periphery of a COS-7 cell is shown in Fig. 5 A overlaid on the overall distribution of all IP<sub>3</sub>R tracks. Pooled over six cells, we identified 70 clusters from a total population of 3331 immotile molecules. To evaluate whether these putative clusters might arise simply from random coincidence, we used the population of rapidly motile IP<sub>3</sub>Rs as a control, on the premise that their locations (mean positions along their tracks) would be randomized by Brownian motion.

Using the same identification criteria and after normalizing for differences in molecule density by randomly sampling equivalent numbers of motile and immotile molecules from each record, we identified 25 apparent clusters, considering the same total number (3331) of rapidly motile molecules in the same cells.

Fig. 5 B shows magnified views of a representative cluster of immotile IP<sub>3</sub>Rs identified as above, plotting the mean localizations of each immotile molecule (*black dots*) and the individual localizations from which the mean positions were obtained (*open circles*). The distributions of IP<sub>3</sub>Rs within individual clusters typically were not circularly symmetrical, and instead clusters were elongated at random orientations, as might be expected from the location of IP<sub>3</sub>Rs along strands of the ER. Fig. 5 C shows an overlay of IP<sub>3</sub>R localizations obtained from 70 clusters that were formed by aligning their center positions (mean position of molecules within each cluster) and rotating them so that the long axis of each cluster aligned on the *x* axis. The mean distributions of molecules approximate Gaussian

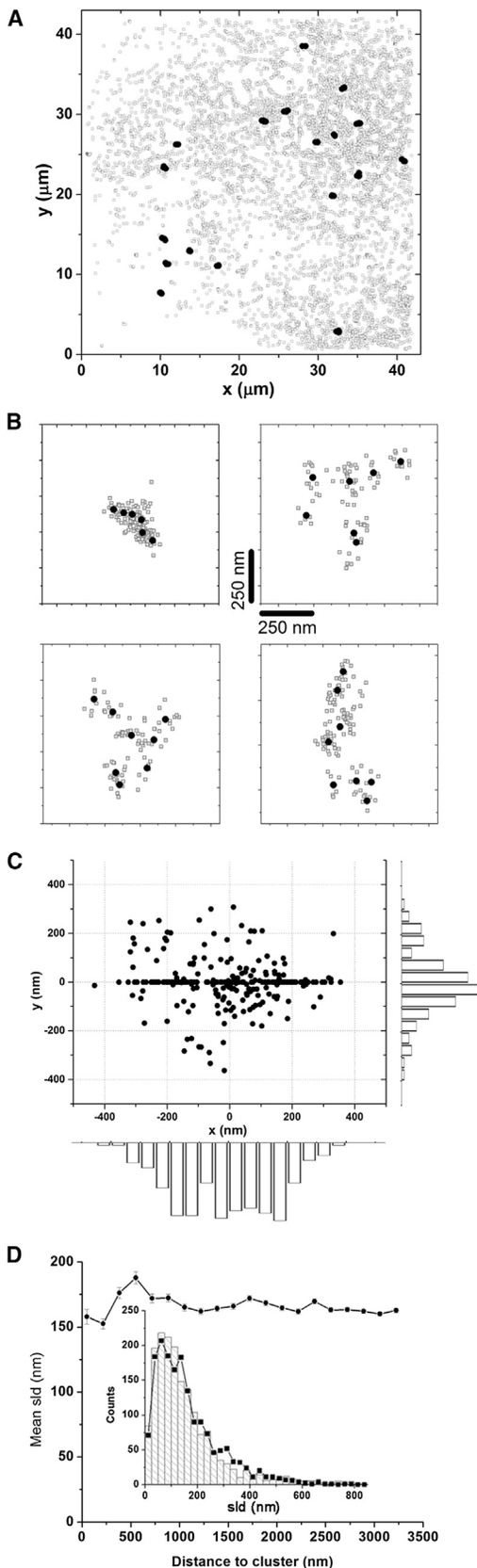


FIGURE 5 Clusters of immobile IP<sub>3</sub>Rs. (A) Gray circles denote the mean positions of all IP<sub>3</sub>R molecule tracks located in a live COS-7 cell. Black dots indicate the positions of clusters of immobile IP<sub>3</sub>Rs located by criteria

functions of distance from the cluster center, with widths (standard deviations (SDs)) of 195.2 nm along the long axis and 86.84 nm along the short axis.

### Motilities of IP<sub>3</sub>Rs near cluster regions

Clustering of immotile IP<sub>3</sub>Rs might arise if these molecules are bound to a finite number of fixed anchoring sites, or it might result from a more general environmental influence, such as greatly reduced diffusion within a localized region of ER membrane that has formed a diffusion trap. To discriminate between these possibilities, we examined the motilities of IP<sub>3</sub>R tracks that passed through or close to (<3.5 μm) clusters, after excluding tracks of those molecules identified as forming the clusters themselves. Fig. 5 D plots SLDs along these tracks as a function of the mean distance of a molecule during that displacement from the centroid of the cluster. The mean data derived from 70 clusters showed little apparent change in SLDs in proximity to a cluster, and the distributions of SLDs within 2 pixels (334 nm) of cluster centroids (Fig. 5 D, inset and histogram) and between 8 and 10 pixels (1.33–1.67 μm) away (Fig. 5 D, inset, line, and symbol) overlapped closely. Thus, with the exception of those immotile IP<sub>3</sub>Rs that defined a cluster, other IP<sub>3</sub>Rs in the vicinity of a cluster did not show appreciably different diffusional motilities. A possible caveat to this conclusion is that we analyzed motility only as a single *x-y* plane, and it is possible that IP<sub>3</sub>Rs could diffuse freely above or beneath a cluster within the sheet of membrane on the opposite side of the ER lumen from the cluster. However, even if this were the case, we might still expect to detect an overall slowing if the fraction of molecules in the same sheet of membrane as the cluster showed a reduced motility in its vicinity.

### Activation of IP<sub>3</sub>/Ca<sup>2+</sup> signaling does not acutely alter IP<sub>3</sub>R motility or distribution

IP<sub>3</sub>Rs in the membrane of excised nuclei have been proposed to undergo clustering in response to IP<sub>3</sub> within just

as described in the text. (B) Four representative examples of individual IP<sub>3</sub>R clusters shown at high magnification. Black dots show the positions of individual molecules derived as the mean of all localizations along their tracks (gray circles). (C) The scatterplot is a superposition of IP<sub>3</sub>R molecules within 70 clusters after aligning the centroids of each cluster and rotating their long axes to lie along the *x* axis. Bar graphs indicate the numbers of molecules at different distances along the long and short axes to illustrate the mean density distribution of molecules within clusters. The plot along the short axis omits those molecules used to define the long axis of each cluster. (D) Diffusion of motile IP<sub>3</sub>Rs is not altered in the vicinity of clusters of immobile IP<sub>3</sub>Rs. The main graph plots mean SLDs (*n* > 79,000) of all tracks within 3.5 μm of identified clusters (*n* = 70) as a function of distance from the centroid of the cluster. Inset shows the distributions of SLDs recorded within 334 nm of cluster centroids (histogram bars) and at distances 1.33–1.67 μm away (black symbols and lines).

a few seconds (24). We therefore looked for possible changes in the diffusional motility and spatial distribution of IP<sub>3</sub>Rs following activation of endogenous IP<sub>3</sub> signaling in COS-7 cells. We tracked single-molecule displacements for 750 s at baseline and then for a further 750 s immediately after adding ATP (100 μM) to the bathing solution to activate endogenous IP<sub>3</sub>-linked purinergic receptors (40). In separate experiments with fluo-8-loaded cells, this concentration of ATP in Ca<sup>2+</sup>-free medium evoked robust intracellular Ca<sup>2+</sup> signals that persisted for ~1 min (Fig. 6 A). Plots of MSD versus lag from cells before and after addition of ATP were linear and showed no difference in slope (Fig. 6 B). Mean aggregate diffusion coefficients derived from six cells showed no significant difference (before ATP: 0.0956 ± 0.006 μm<sup>2</sup> s<sup>-1</sup>; after ATP: 0.0951 ± 0.007 μm<sup>2</sup> s<sup>-1</sup>; difference not statistically significant,  $p \gg 0.05$ ). To examine whether ATP treatment induced clustering of IP<sub>3</sub>Rs, we performed a near-neighbor distribution analysis of immotile IP<sub>3</sub>Rs, as in Fig. 4 C. The distributions of intermolecular spacings were closely similar before and after ATP application (Fig. 6 C). Moreover, we found no difference in the numbers of clusters of immotile IP<sub>3</sub>Rs identified using the same criteria as in Fig. 5 (18.7 clusters per cell in control, 18.3 clusters per cell after ATP;  $n = 6$  cells).

## DISCUSSION

Ca<sup>2+</sup> imaging experiments in numerous cell types have shown that local IP<sub>3</sub>-mediated Ca<sup>2+</sup> signals (Ca<sup>2+</sup> puffs) arise at just a few discrete sites within a cell, which are thought to represent tight clusters of several IP<sub>3</sub>R channels (4,9–11). For example, SH-SY5Y neuroblastoma cells, which express predominantly type 1 IP<sub>3</sub>R, were estimated to contain <20 discrete Ca<sup>2+</sup> release sites, totaling no more than 100 or 200 functional IP<sub>3</sub>Rs per cell (27). In comparison, estimates of the number of IP<sub>3</sub>R molecules within a cell range as high as several thousands (41) (E.J.A. Taylor and C.W. Taylor, personal communication), and immunostaining shows an extensive distribution throughout the ER (Fig. 1 G) (11,22). Moreover, several previous studies that applied fluorescence recovery after photobleaching (FRAP) techniques to examine the motility of overexpressed fluorescent protein-tagged IP<sub>3</sub>Rs in intact cells indicated that a substantial proportion of IP<sub>3</sub>Rs are freely diffusible within the ER membrane (14–22). This is markedly different from the functional puff sites that remain at essentially fixed locations over a period of minutes (13). Here, we employed single-molecule superresolution imaging to gain insights into how the subcellular localization and motility of individual IP<sub>3</sub>R proteins might underlie the distribution of functionally active channels and puff sites.

As with all experiments involving heterologous expression of a fluorescently tagged protein, there are caveats as to whether the expressed protein faithfully mimics the

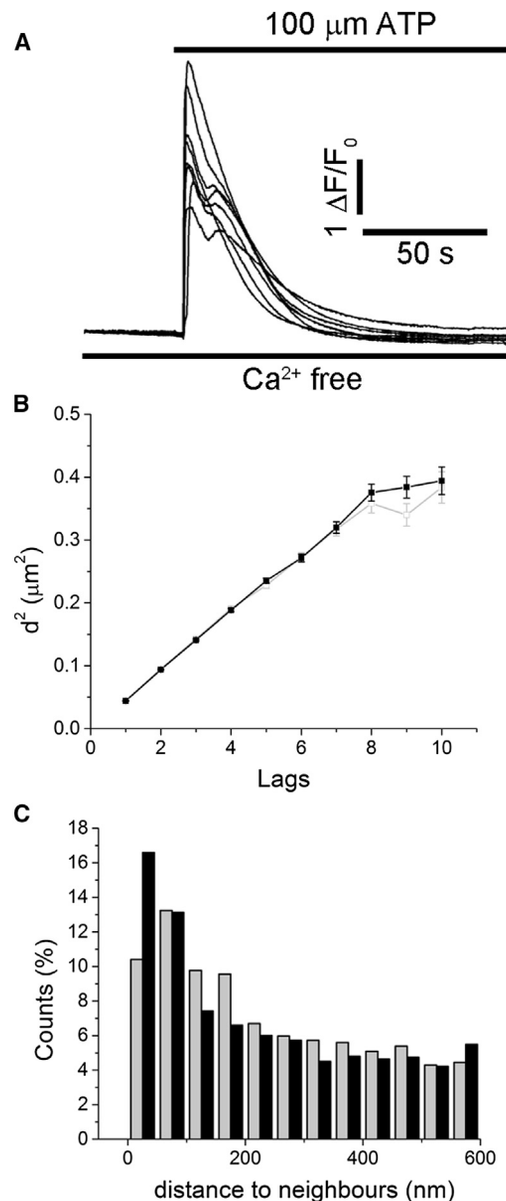


FIGURE 6 Activation of IP<sub>3</sub>/Ca<sup>2+</sup> signaling has little effect on the motility and spatial distribution of IP<sub>3</sub>Rs. (A) Superimposed traces show Ca<sup>2+</sup>-dependent changes in fluo-8 fluorescence from six COS-7 cells in response to bath application of 100 μM ATP in Ca<sup>2+</sup>-free extracellular solution. (B) The plot shows the MSD of molecules from their origin as a function of increasing lag. The data shown are from a single cell and are representative of results from six cells. Open gray symbols are measurements before application of ATP, and black symbols are measurements after ATP. (C) Distributions of mean interneighbor distances of immotile IP<sub>3</sub>Rs (mean SLD < 84 nm) before (gray bars) and after (black bars) addition of ATP. Data were pooled from six cells.

behavior of the native protein, and in particular whether overexpression might overwhelm the cellular processes that determine the normal localization and motility of the native protein. However, we found that superresolution snapshots of fixed COS-7 cells showed similar distributions of immunostained native IP<sub>3</sub>R1s and Eos-tagged,

overexpressed IP<sub>3</sub>R1s, and that the fraction of motile receptors remained approximately constant over a 3-fold range of expression levels. Moreover, GFP-tagged IP<sub>3</sub>R1s expressed using a construct closely similar to ours have been reported to be functionally active (30), and two reports (16,42) described little difference in FRAP measurements of IP<sub>3</sub>R motility across a wide range of expression levels, providing reassurance that overexpression would not be a major concern.

From single-molecule tracking, we estimate that ~70% of the fluorescently tagged IP<sub>3</sub>R1s we expressed in COS-7 cells were motile, with an apparent two-dimensional diffusion coefficient of  $0.095 \pm 0.005 \mu\text{m}^2 \text{s}^{-1}$ . This diffusion coefficient is well within the extensive range ( $0.01$ – $0.3 \mu\text{m}^2 \text{s}^{-1}$ ) reported from FRAP experiments in various cell types (see Table S2 of Pantazaka and Taylor (42)), but is ~5-fold greater than that derived from previous FRAP measurements of the mobile fraction of EGFP-tagged IP<sub>3</sub>R1 expressed in COS-7 cells (42). The reason for this discrepancy is not clear, but it may involve both methodological differences and fundamental differences between measurements of single-particle and collective (population) diffusion coefficients in the presence of binding traps (43).

We found that ~30% of the expressed IP<sub>3</sub>R1s were essentially immotile, with mean SLDs similar to those measured in fixed cells and thus attributable to localization jitter. The spatial distribution of these immotile IP<sub>3</sub>R1s showed an excess of closely neighboring molecules over that predicted for a random distribution, and we identified clusters of several molecules spaced within a few hundred nanometers of one another. Representative examples of localizations within individual clusters are shown in Fig. 5 B, although these should not be interpreted as necessarily depicting the true numbers of IP<sub>3</sub>R1 proteins present in a cluster. Native, unlabeled IP<sub>3</sub>R1 obviously would not be observed, and it is possible that not all of the mEos-tagged molecules were activated during the imaging sequence. Conversely, the tetrameric nature of IP<sub>3</sub>R1 and the fact that an mEos molecule may undergo multiple reactivations (38) would tend to result in an overcount, even after correcting for photoblinking artifacts. Nevertheless, the composite overlay Fig. 5 C should provide a representative view of the average dimensions of stationary IP<sub>3</sub>R1 clusters at sub-micron resolution.

In principle, clustering of immotile IP<sub>3</sub>R1s might result from their binding to a limited number of anchoring sites (e.g., cytoskeleton-associated proteins) or from an environmental effect that greatly slows down diffusional motility, such as a local change in ER membrane lipid composition. We favor the first possibility because tracks of IP<sub>3</sub>R1s that passed through or nearby identified clusters showed no evidence of a slowing of diffusional motility. A related question concerns the kinetics of IP<sub>3</sub>R1 binding to anchoring sites. The very fact that we observe clusters of immotile, fluorescently tagged, exogenous IP<sub>3</sub>R1s suggests that this binding must be

reversible over a timescale of hours; otherwise, the sites would be permanently occupied by native (nonfluorescent) IP<sub>3</sub>R1s and hence would be inaccessible. However, our ability to discern an interchange of IP<sub>3</sub>R1s at cluster sites is severely limited by the short time (a few seconds) in which the Eos reporter on a given molecule remains fluorescent.

It has been proposed that IP<sub>3</sub>R1 clusters are dynamic and that IP<sub>3</sub> regulates the assembly and function of Ca<sup>2+</sup> puff sites under physiological conditions (24). We previously argued against this hypothesis on the basis that the low density and restricted motility of functional IP<sub>3</sub>R1s (i.e., those that give rise to Ca<sup>2+</sup> signals) are incompatible with dynamic clustering on a timescale that is fast enough to account for the short latencies (a few hundred microseconds) of puffs evoked by strong flash photorelease of IP<sub>3</sub> (12). On the other hand, the results presented here show a high density of freely motile IP<sub>3</sub>R1 proteins, suggesting that aggregation of IP<sub>3</sub>R1s into clusters could occur within hundreds of milliseconds if IP<sub>3</sub> were to create a diffusion trap by substantially increasing the affinity between IP<sub>3</sub>R1s and putative anchoring sites. Nevertheless, we do not believe this to be the case in light of our observations of clusters of immotile IP<sub>3</sub>R1s in quiescent cells, as well as our failure to observe any changes in diffusional motility or spatial clustering after stimulating cells with ATP to activate the endogenous IP<sub>3</sub>/Ca<sup>2+</sup> signaling pathway.

Given that only a tiny fraction of the IP<sub>3</sub>R1s in a cell (<5% based on the estimates in SHSY-5Y cells given above) appear to underlie the generation of calcium puffs, a search for those IP<sub>3</sub>R1s presents a needle-in-a-haystack problem. Although thus far we have been unable to directly correlate the clusters of small numbers of immotile IP<sub>3</sub>R1s we describe here with active calcium release sites, we believe they are promising candidates for the clusters underlying puff sites. In particular, the IP<sub>3</sub>R1 clusters have dimensions (extending over roughly 300–400 nm) that are compatible with estimates of the size of clusters of functional IP<sub>3</sub>R1s underlying puff sites made via functional Ca<sup>2+</sup> imaging, based on simulations of the spatial spread of the Ca<sup>2+</sup> fluorescence signal during puffs (44) and on movements of the centroid of the Ca<sup>2+</sup> signal during puffs (13). We thus propose the parsimonious explanation that a small fraction of IP<sub>3</sub>R1 channels are stably aggregated under resting conditions. Unlike the greater numbers of motile IP<sub>3</sub>R1s, these clustered IP<sub>3</sub>R1s are functional. In response to increased [IP<sub>3</sub>], their openings are concerted via CICR to generate puffs following a brief latency that is determined largely by stochastic Ca<sup>2+</sup> activation (45) rather than diffusional aggregation. The functional role of the substantial majority of freely motile IP<sub>3</sub>R1s remains a mystery.

## SUPPORTING MATERIAL

One movie is available at [http://www.biophysj.org/biophysj/supplemental/S0006-3495\(14\)00713-9](http://www.biophysj.org/biophysj/supplemental/S0006-3495(14)00713-9).

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