Single-Molecule Imaging of L-Type Ca\textsuperscript{2+} Channels in Live Cells

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ABSTRACT  L-type Ca\textsuperscript{2+} channels are an important means by which a cell regulates the Ca\textsuperscript{2+} influx into the cytosol on electrical stimulation. Their structure and dynamics in the plasma membrane, including their molecular mobility and aggregation, is of key interest for the in-depth understanding of their function. Construction of a fluorescent variant by fusion of the yellow-fluorescent protein to the ion channel and expression in a human cell line allowed us to address its dynamic embedding in the membrane at the level of individual channels in vivo. We report on the observation of individual fluorescence-labeled human cardiac L-type Ca\textsuperscript{2+} channels using wide-field fluorescence microscopy in living cells. Our fluorescence and electrophysiological data indicate that L-type Ca\textsuperscript{2+} channels tend to form larger aggregates which are mobile in the plasma membrane.

INTRODUCTION

Single-molecule fluorescence imaging reveals great details about dynamical processes of biological interest in a multitude of in vitro systems (Weiss, 1999; Ishii and Yanagida, 2000; Edman et al., 1996; Dickson et al., 1997; Sase et al., 1997; Lu et al., 1998; Jia et al., 1999; van Oijen et al., 1999). This novel methodology, however, lacks a general utilization in vivo (Sako et al., 2000; Schütz et al., 2000), primarily because of the enhanced fluorescence background caused by the cellular autofluorescence and the lack of suitable fluorescence tags to label proteins in living cells. Fluorescent proteins provide a noninvasive and convenient means for the in vivo labeling of molecular components (Tsien, 1998). In the current study, we use a method which permits wide-field, single-molecule imaging of eYFP (enhanced yellow-fluorescent protein) fusion-proteins in living cells not masked by cellular autofluorescence.

The target of the current investigation is the cardiac L-type Ca\textsuperscript{2+} channel, one of the major subjects in molecular cardiology and research in Ca\textsuperscript{2+} signaling (Murphy et al., 1991; Fabiato and Fabiato, 1997; Rios and Brum, 1987; Gao et al., 1999). The channel protein consists of a pore-forming \( \alpha_{1C} \)-subunit and two auxiliary subunits (Hofmann et al., 1994; Catterall, 1995) (Fig. 1). Although a vast amount of functional information about this channel is available from electrophysiology (Catterall, 1995), our knowledge about its dynamic embedding in the cell membrane and its aggregation state in vivo is minute. Aggregation (Flucher and Franzini-Armstrong, 1996; Grabner et al., 1998; Flucher et al., 1993) has been previously anticipated from well ordered arrays of the Ca\textsuperscript{2+}-release channels on membranes of the sarcoplasmic reticulum (Saito et al., 1988), which are closely coupled to the L-type Ca\textsuperscript{2+} channels on the plasma membrane (Flucher and Franzini-Armstrong, 1996). In this study, we performed a detailed analysis of the fluorescence intensity in both position and time for functionally active individual eYFP-\( \alpha_{1C} \) fusion protein molecules in the plasma membrane of the HEK293 cell line. This allowed us to obtain information about their mobility and the state of molecular aggregation. Surprisingly, it seems that the aggregation of L-type Ca\textsuperscript{2+} channels is probably independent from its association with other structures of the excitation-contraction machinery (Gerster et al., 1999). The latter finding further confirms results obtained by electrophysiology (Kepplinger et al., 2000), and by in vitro studies of the purified channel when reconstituted in a phospholipid monolayer (Hinterdorfer et al., 1997).

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FIGURE 1  Diagram of the experimental setup (center). Signals of an entire cell localized in the object plane (left inset) are imaged on the CCD. The object plane is adjustable throughout the cell. Schematic structure of a L-type Ca\textsuperscript{2+} channel in the membrane.
MATERIALS AND METHODS

Cloning of eYFP-α1C,77

Cloning and electrophysiological characterization of eYFP-α1C,77 has been described in detail in Kepplinger et al. (2000). In brief, the 5′-terminal Hind III linker upstream Kozak sequence and 3′-terminal Bgl II linker were incorporated into the flanking regions of the peYFP DNA (Clontech Laboratories, Palo Alto, CA) open reading frame by polymerase chain reaction (PCR) using 5′-cttaacctggccacagcttgagc-3′ sense and 5′-ag- acctctggacgcttc-3′ antisense primers, respectively. The Hind III/Bgl II peYFP cassette was ligated with the BamH I/Not I HFCC77 cassette into the DNA3 vector (Invitrogen, Carlsbad, CA) at Hind III/Not I sites so that the eYFP-77pcDNA3 construct encoded the eYFP fused to α1C,77 via RSAT tetrapeptide. Integrity of the ORF was verified by sequencing.

Cell culture

HEK293 cells were cultured in DMEM medium supplemented with streptomycin (100 μg/ml), penicillin (100 U/ml), and 10% bovine serum in a humidified atmosphere (95%) at 5% CO2 and 37°C. Cells were used for 12–14 passages and were transferred every 4 days. Transfection was performed using N-[2,3-dioleoyloxy]propyl]-N,N,N-trimethylammonium methysulfate (Amersham, Rosendaal, The Netherlands). Cells exhibiting confluence of ~30% were used for transfection with a total of, respectively, 0.25 (low transfection) and 2 μg (normal transfection) of cDNA (molar ratio of α1C,77-eYFP/β2A = 1/1.6 and α1C,77-eYFP/αβ = 1/1.4) in a 1 ml volume. The excess with β2A and αβ subunits assured a complete assembly of all α1C,77-eYFP units to a functional channel. This was concluded from the homogeneous behavior observed in the electrophysiological results in all patch-clamp trials. The transfection efficiency was in the range of 20–60%. For fluorescence measurement the cells were plated on #1 glass slides (Fisher, Zoetermeer, The Netherlands) in a bath of phosphate-buffered saline (150 mM NaCl, 10 mM Na2HPO4, pH 7.4).

Electrophysiology

Details on the electrophysiological experiments, including the single channel recording and analysis referred to later in this article, can be found in Kepplinger et al. (2000). Whole-cell patch-clamp recordings (Hamill et al., 1981) were obtained from HEK293 cells transfected with either α1C,77 or YFP-α1C,77 together with β2A and αβ subunits using an Axopatch 2B (Axon Instruments, Foster City, CA) or an EPC7 (Heka Elektronik, Lambrecht, Germany) amplifier. The pipette solution contained: 120 mM Cs methanesulfonate, 5 mM CaCl2, 2 mM MgCl2, 10 mM HEPES, 10 mM EGTA, 2 mM MgATP, pH (CsOH) 7.3. The bath solution consisted of: 154 mM N-methyl glucamine, 1 mM MgCl2, 5 mM D-glucose monohydrate, 10 mM HEPES, 5 mM 4-aminopyridine, 15 mM BaCl2, pH (HCl) 7.4. Soft glass pipettes (Microhematocrit tubes, No. 564, Fa. Assistant, Vienna, Austria) with a resistance of 1–4 MΩ were used for whole-cell recordings. Ba2+ currents were activated by repetitive (0.2 Hz) depolarizations from a holding potential of ~80 mV to test potentials (0.244 s) between −10 mV and +60 mV with an incremental increase of 5 mV or 20 mV. Current traces were filtered at 3 kHz, digitized at 8 kHz, and were neither capacity nor leak current corrected allowing to verify the quality of voltage-clamp. A liquid junction potential of 6 mV was not taken into account. This value should be subtracted from all voltages in whole-cell recordings (Neher, 1992). All experiments were performed at room temperature.

Single-molecule optical microscopy

The experimental arrangement for single-molecule imaging has been described in detail previously (Schmidt et al., 1995). Essentially, the samples were mounted onto an inverted microscope (Zeiss, Weesp, The Netherlands) equipped with a 100× objective (NA = 1.4, Zeiss), and illuminated for 5–10 ms at 514 nm from an Ar+ laser (Spectra Physics, Eindhoven, The Netherlands). The illumination intensity was set to 5 kW/cm2 in all experiments. The excitation polarization was selected by a Berek polarizer (New Focus, San Jose, CA). Use of appropriate filter combinations (DCLP530, HQ580/75, Chroma Technology, Brattleboro, VT, and OG530–3, Schott, Mainz, Germany) permitted the detection of individual eYFPs by a nitrogen-cooled charge coupled device (CCD)-camera system (Princeton Instruments, Vianen, The Netherlands). The total detection efficiency of the experiment was 0.048.

For single-molecule detection cells were photobleached at 514 nm for typically 1 s at the intensity of 5 kW/cm2. Fluorescence images were taken consecutively with a delay between 50 and 500 ms with up to a possible 500 images in a sequence. An analysis program determined the lateral position of each signal with an accuracy of <30 nm by fitting to a two-dimensional (2-D) Gaussian surface. The extremely high positional

![FIGURE 2](image-url) Electrophysiological characterization of the eYFP-α1C,77 fusion protein. (A) Comparison of whole cell Ba2+ current of the wild-type L-type Ca2+ channel with the eYFP-α1C,77 fusion protein. The cells were depolarized to +20, +30, and +40 mV from a holding potential of ~80 mV. On average Iba = 1690 ± 180 pA (mean ± SE, N = 12) was observed on depolarization to +20 mV for the eYFP-α1C,77 channel. (B) Current-voltage characterization of the wild-type L-type Ca2+ channel compared with the eYFP-α1C,77 fusion channel. (C) Whole-cell Ba2+ current of the L-type Ca2+ channel with the fusion of the eYFP on the α1C subunit transfected with low concentration of plasmid in HEK cells. The cells were depolarized to −20, 0, +20 mV from a holding potential of ~80 mV. On average, Iba, was reduced to 200 ± 50 pA (mean ± SE, N = 6).
A Vogel-algorithm was used to correlate the images of molecules in subsequent observations from which the respective single-molecule trajectories were reconstructed (Schmidt et al., 1995). For each trajectory of length \( N \) with respective positions

\[
D_{st} = [4\Delta t (N - 1)]^{-1} \sum (\tilde{p}_i - \tilde{p}_{i-1})^2
\]

and time-lag \( \Delta t \), a mean diffusion constant (Sonnenleitner et al., 1999) was evaluated by:

**Fluorescence correlation microscopy**

Fluorescence correlation measurements were performed using a commercial system (ConfoCor, Zeiss). We used 514-nm excitation with standard optics, including a dichroic mirror (510 nm, Zeiss), a water-immersion objective (40\(\times\), 1.2 NA, Zeiss) and a band-pass filter (515–565 nm, Zeiss) to discriminate the fluorescence. The excitation intensity was <1 kW/cm\(^2\) to reduce photobleaching of eYFP. The emission light was filtered by a 45-\(\mu\)m diameter pinhole, and detected by an avalanche photodiode connected to a fast digital correlator. Correlation curves were selected which exhibited correlation times significantly longer than the mean photobleaching time of \(~50\) ms. The correlation curves, \( G(t) \), obtained for eYFP-\(\alpha_{1C,77}\) in HEK293 cells were fit by the combination of 2-D diffusion and photobleaching (Schwille et al., 1999):  

\[
G(t) = N^{-1} (1 + t/\tau_d)^{-1} \exp(-t/\tau_b),
\]

where \( \tau_d \) is the mean diffusion time, \( \tau_b \) is the mean photobleaching time, and \( N \) is the average number of fluorophores in the confocal volume. For diffusion analysis, only curves were accepted for which \( \tau_d > 2 \tau_b \). Calibration of the size of the focused beam was performed with tetramethylrhodamine in water (Widengren and Rigler, 1998), yielding a beam radius of 0.32 \(\mu\)m. For display, all curves are normalized by multiplication with the mean number of fluorophores, \( N \). The acquisition time was set to 10 s. Additional control experiments (not shown) were performed on purified eYFP in buffer solutions (including viscosity, \( pH \), and salt effects) which confirmed the submillisecond timescale dynamics of eYFP reported in literature (Widengren et al., 1999; Schwille et al., 2000).

**RESULTS**

**Single-molecule imaging**

Fig. 1 depicts a schematic diagram of the experiments as described in Methods. Cardiac L-type \( Ca^{2+} \) channels consisting of the fluorescent N-terminally labeled subunit, eYFP-\(\alpha_{1C,77}\), and the wild-type auxiliary subunits, \( \beta_2 \), were expressed in HEK293 cells. Fluorescence images were taken from \(~1-\mu\m\) thick slices through the middle of the cells. It was apparent from those images (Figs. 3A and 5A) that most channels were translocated to the cell membrane.

To test for the functional integrity of the eYFP-\(\alpha_{1C,77}\) fusion construct, an electrophysiological characterization was performed in parallel. We found (Fig. 2, A and B) that the N-terminal eYFP-fusion to the \(\alpha_{1C,77}\) subunit did not significantly affect \( Ca^{2+} \) channel function in both whole-cell and single-channel experiments (Keplinger et al., 2000). The channel inactivation was faster by \(~30\%\) and the activation curve was shifted by \(~10\ mV\) with respect to the wild-type \(\alpha_{1C,77}\) channel. These findings are in agree-
ment with work on homologous proteins (Grabner et al., 1998; Gerster et al., 1999).

The single-molecule fluorescence results are conceptualized in Fig. 3. Images obtained by fluorescence microscopy easily discriminate cells that contained fluorescence-labeled channels from untransfected cells (compare Fig. 3 A and 3 B). Diffraction-limited fluorescence signals (320 ± 80 nm full-width at half-maximum) are localized at the cell surface and are distinguishable from the background with high signal-to-background-noise ratio (~15, Fig. 3 C). These localized signals fit well to 2-D Gaussian surfaces yielding values for the integrated fluorescence and the lateral position on the membrane. Subsequent positional tracking follows those signals over time at a rate of up to 20 images/s. Such detailed analysis allows us to assign the signals to individual eYFP-α1C,77 subunits of the L-type Ca2+ channel. We find that the single fluorescent molecules have the following characteristics: (1) the mean amplitude of the smallest signal component of 34 ± 4 cnts/ms matches the mean amplitude found for individual eYFPs in buffer, 38 ± 4 cnts/ms, and mean signals found when eYFP is immobilized onto artificial- and cell-membranes, (36 ± 4 and 33 ± 4 cnts/ms, respectively), at identical illumination intensity and integration times (Harms et al., 2001). (2) All signals exhibited a stepwise photobleaching behavior (Fig. 3 D), a signature characteristic for a single-molecule event. (3) Fluorescence polarization imaging (Harms et al., 1999) (see Methods) shows that a fraction (<10%) of all signals exhibits fluorescence which is highly polarized. A high fluorescence polarization is a distinctive property for single quantum systems, such as a single fluorophore, if its rotation is slower than the detection time. The rotation of the fluorophore shown in Fig. 3 E results in a change of the direction of the transition dipole moment by 70° 20° in 110 ms. (4) Finally, fluorescence correlation experiments (Schwille et al., 1999; Widengren and Rigler, 1998) were performed from which the mean number of fluorescent entities was derived (~10 μm−2). This independent finding further corroborates our results from single-molecule imaging (Fig. 5 C). These arguments together demonstrate, for the first time, observation of individual eYFP-fusion proteins in a living cell.

To achieve this result, we applied a rigorous experimental optimization. For the single-molecule imaging experiments, the transfection procedure is adjusted such that the number of channels per resolution limited area of 0.08 μm2 was less than one. Fluorescence single-molecule measurements here require: (1) the reduction of the amount of plasmid by a factor of ~10 in comparison to transfection procedures normally applied in electrophysiology; (2) increasing the time between transfection and measurement to >4 days which additionally ensured a low cytosolic background; and (3) the selection of individual cells with low expression levels of the fusion protein (~104 copies per cell, corresponding to a surface density of ~30 μm−2). Additionally,
found that the photobleaching step also reduces the observable density of the eYFP-α$_{1C,77}$ to ∼10% of the initial level leading to ∼3 observable eYFP-α$_{1C,77}$ per $\mu$m$^2$. Together, the procedures permit reliable and reproducible single-molecule imaging and fluorescence correlation analysis of eYFP-α$_{1C}$ with a high discrimination from cellular autofluorescence.

As an independent confirmation of our findings, the influence of the variation in the transfection procedures on the channel activity was investigated (Fig. 2 C). The electrophysiological experiments showed that under described conditions, the average peak Ba$^{2+}$ inward current was $200 \pm 50$ pA (mean ± SE, $N = 6$) at a depolarization to $+20$ mV, which is eight times less than the current recorded with the transfection procedures as used in Fig. 2, A and B ($1690 \pm 180$ pA, $N = 12$). Given a single-channel conductivity of $\sigma = 29$ pS, the reverse potential of 63 mV (Fig. 2 B), and an open probability of $p_0 = 0.03$ (Kepplinger et al., 2001), the average density of channels in a typical cell of diameter 10 $\mu$m using the low transfection protocol is $n \sim 17$ $\mu$m$^{-2}$ in registry with our estimations from the fluorescence. We have further tried to obtain single-channel recordings from cells which were transfected with the reduced amount of channel plasmid to determine a possible influence on the channel behavior. However, we were unable to obtain any single-channel recording in $N = 12$ trials.

**Ca$^{2+}$ channel clustering**

Achievement of single-molecule sensitivity in fluorescence permits for a detailed analysis of local stoichiometries yielding information on channel aggregation. Fig. 4 A shows the fluorescence intensity distribution calculated from 271 individual fluorescence signals on the surface of an HEK293 cell. The distribution is far from unimodal, consisting of multiple central intensity values. This observation was confirmed to be characteristic for eYFP-α$_{1C}$ expressed in HEK293 cells by comparison to the intensity distribution obtained for purified eYFP when anchored to a membrane (Harms et al., 2001). The intensity distribution of that control measurement was found to be close to a single Gaussian with a width mostly accounted for by the shot-noise and the instrument read-out noise of the CCD-camera. It should be noted, that the long timescale (>100 ms) “blinking” behavior of the autofluorescent proteins (Dickson et al., 1997; Schwille et al., 2000) could not be distinguished from diffusion in our experiments. The fast photophysical dynamics of the autofluorescent proteins observed on submillisecond timescales (Widengren et al., 1999; Schwille et al., 2000; Garcia-Parajo et al., 2000), are averaged out on the 5-ms illumination time used here, giving rise to a slightly larger widths of the intensity distributions compared with that of conventional fluorescence labels (Schmidt et al., 1996a).
For quantitative interpretation of the multimodal intensity distribution shown in Fig. 4 A, algorithms were used which have been described previously (Schmidt et al., 1996b). In brief, the lower part of the distributions (up to 1000 cnts) were first fit to four Gaussians for which the positions of the maxima were found to be equidistant with a spacing of \(168 \pm 20 \text{ cnts/5ms}\) (Fig. 4 A, inset). It was also found that the squared widths of those Gaussians scaled linearly with the aggregation number as being predicted for statistically independent distributions. The spacing matches that of the mean intensity of an individual eYFP-\(\alpha_{1C,77}\) (34 \(\pm\) 4 cnts/ms, Fig. 3 D), providing strong direct evidence for a local clustering of L-type \(\text{Ca}^{2+}\) channels. Signal amplitudes \(>1000 \text{ cnts/5 ms}\) were not taken into account for such detailed analysis. For those signals, the increased width of the Gaussian deems a reliable assignment unjustified (Schmidt et al., 1996b).

Subsequent to this global analysis, a more detailed analysis was applied for classification of each individual signal to a local stoichiometry (Schmidt et al., 1996b), \(N (N\) ranging from 1 to 4, and \(\cong 5\) of detected eYFP-\(\alpha_{1C,77}\); the intensity of the signal was compared with that obtained from individual eYFP protein molecules when anchored to a membrane (Fig. 4 A, red curve, and Harms et al., 2001). A priori knowledge of the monomer signal-distribution was used as a solid basis for our stoichiometry assignment. In total, 7 cells and 3417 signals were analyzed, yielding a probability of 0.13, 0.21, 0.24, 0.19, 0.23 for one, two, three, four, and more colocalized eYFP-\(\alpha_{1C,77}\), respectively (Fig. 4 C). The distribution in Fig. 4 C is characterized by an average number of 3.1 \(\pm\) 0.3 colocalized and detected eYFP-\(\alpha_{1C,77}\) molecules. For determination of the overall size of the aggregates, the initial photobleaching step must be taken into account. Including photobleaching the observed distribution is given by a binomial characterized by the mean cluster size \(M\) and the photobleaching probability, \(1-P\). A least-squares fit yielded \(M = 40 \pm 15\) and \(P = 0.07 \pm 0.03\).

As the individual classification algorithm applied here is statistical (Schmidt et al., 1996b), it was occasionally confirmed by a direct method: for this the fluorescence intensity analysis was applied for classification of each individual signal to a local stoichiometry (Schmidt et al., 1996b), \(N (N\) ranging from 1 to 4, and \(\cong 5\) of detected eYFP-\(\alpha_{1C,77}\); the intensity of the signal was compared with that obtained from individual eYFP protein molecules when anchored to a membrane (Fig. 4 A, red curve, and Harms et al., 2001). A priori knowledge of the monomer signal-distribution was used as a solid basis for our stoichiometry assignment. In total, 7 cells and 3417 signals were analyzed, yielding a probability of 0.13, 0.21, 0.24, 0.19, 0.23 for one, two, three, four, and more colocalized eYFP-\(\alpha_{1C,77}\), respectively (Fig. 4 C). The distribution in Fig. 4 C is characterized by an average number of 3.1 \(\pm\) 0.3 colocalized and detected eYFP-\(\alpha_{1C,77}\) molecules. For determination of the overall size of the aggregates, the initial photobleaching step must be taken into account. Including photobleaching the observed distribution is given by a binomial characterized by the mean cluster size \(M\) and the photobleaching probability, \(1-P\). A least-squares fit yielded \(M = 40 \pm 15\) and \(P = 0.07 \pm 0.03\).

As the individual classification algorithm applied here is statistical (Schmidt et al., 1996b), it was occasionally confirmed by a direct method: for this the fluorescence intensity of individual signals was monitored over time. Some of the signals showed a one-step photobleaching behavior indicative for an individual eYFP-\(\alpha_{1C,77}\) (Fig. 3 D), whereas others exhibited a multistep bleaching behavior (Fig. 4 B). Taken together with the equidistant fluorescence intensity distribution (Fig. 4 A, inset) from which the fluorescence levels for aggregates was predicted (horizontal lines in Fig. 4 B) those events were taken as a signature of multiple eYFP-\(\alpha_{1C,77}\) bleaching.

\(\text{Ca}^{2+}\) channel mobility

To complete our findings, the dynamic behavior of the channels in the plasma membrane was directly visualized. Repetitive imaging at a rate between 2 and 20 images/s was used to obtain a detailed picture of molecular movement in a biomembrane with a lateral resolution of \(<50\ \text{nm}\) (Schmidt et al., 1996a). Analysis of image sequences of a typical length of 3–5 observations, limited by photobleaching, is used to construct the trajectories of individual signals on the cell membrane. All trajectories are confined to the cell perimeter (Fig. 5 A), which further confirms localization of the proteins in the plasma membrane. The lateral diffusion constant, \(D_{\text{lat}}\), for each individual signal is calculated from the ratio of the mean-square displacement (msd) with the time-lag \(t\), \(D_{\text{lat}} = \text{msd}/4t\), assuming normal diffusion behavior. That analysis yielded the histogram presented in Fig. 5 B. A Gamma distribution (Sonnenleiter et al., 1999) with a mean of \(D_{\text{lat}} = 0.15 \pm 0.05 \text{ \mu m}^2/\text{s}\) is used to describe the width of the histogram. That value falls well within the range of diffusion constants found for membrane proteins (Edidin, 1987). The diffusion behavior is further examined by a complementary analysis in which the time dependence of the mean-square displacement, averaged over all trajectories, was calculated (see Fig. 5 B, inset). We find that the msd is close to linear in time, at least up to the length scale of \(\sqrt{0.5 \ \text{\mu m}^2} = 0.7\ \text{\mu m}\). Hence, in that analysis any deviation from normal diffusion, which is expected for membrane proteins (Sonnenleiter et al., 1999; Edidin, 1987; Simons and Ikonen, 1997), may occur on length scales \(\cong 1\ \text{\mu m}\). Indeed, preliminary analysis according to anomalous subdiffusion behavior (Qian et al., 1991; Saxton, 1989) (msd \(\propto t^\alpha\)) yielded an exponent of \(\alpha = 0.79 \pm 0.06\).

Fluorescence correlation spectroscopy (FCS) and fluorescence recovery after photobleaching experiments were used on the apical membrane of identical cells to confirm the single-molecule imaging results. In FCS an average of three eYFP-\(\alpha_{1C,77}\) were present in the detection area of 0.32 \(\text{\mu m}^2\), which corroborates the density found from the imaging experiments. The diffusion time obtained from the correlation experiments is \(200 \pm 100\ \text{ms}\) (Fig. 5 C), yielding a lateral diffusion constant of \(D_{\text{lat}} = 0.11 \pm 0.07 \text{ \mu m}^2/\text{s}\). This value agrees with our findings of single-molecule microscopy (Fig. 5 B). It is noteworthy that on axially scanning through the cell, such long correlation times were only observable at the cell membrane (data not shown). The diffusion constant recorded at the membrane is a factor of 100 smaller than that obtained for cytosolic diffusion of nonfused eYFP, \(D_{\text{lat}} = 20 \pm 5\ \text{\mu m}^2/\text{s}\), and that for purified eYFP in solution, \(D_{\text{lat}} = 21 \pm 5\ \text{\mu m}^2/\text{s}\) (Fig. 5 C, inset). Similar findings for membrane-bound and free molecules have been reported previously (Schwille et al., 1999; Swaminathan et al., 1997). Hence, we are confident to conclude that the L-type \(\text{Ca}^{2+}\) channels observed were localized at the plasma membrane. Both the single-molecule imaging and FCS measurements further agree with fluorescence recovery after photobleaching measurements (data
not shown), yielding a diffusion constant of $D_{\text{lat}} = 0.13 \pm 0.05 \ \mu m^2/s$.

**DISCUSSION**

To draw conclusions from the distribution of observed colocalized eYFP-\(\alpha_{1C,77}\) molecules as displayed in Fig. 4 C, it must be compared with a theoretical model. For purely random colocalization within the resolution-limited area of the experiment (0.08 \(\mu m^2\)) a Poissonian distribution is predicted. The latter is characterized by the average number of detected eYFP-\(\alpha_{1C,77}\) per resolution-limited area at the cell surface which was estimated from the number of identified signals in the experiments (see e.g., Fig. 3 B). The average density per resolution-limited area was \(\sim 0.24 \) eYFP-\(\alpha_{1C,77}\) leading to the Poisson distribution indicated in Fig. 4 C (blue open bars). The predicted and the experimentally obtained distributions are significantly different. Hence, from our experiments we conclude that the L-type Ca\(^{2+}\) channels form aggregates.

The conclusion is evidenced further by the corresponding electrophysiology experiments. Grouping of functional eYFP-\(\alpha_{1C}\) channels rather than an even distribution in the plasma membrane sharply decreases the chance to detect channel activity in membrane patches. Clustering of the channel with a mean cluster size \(M = 40\) will, at low plasmid transfection (Fig. 2 C), lead to the formation of <1 electrophysiological active patch per \(\mu m^2\). This might explain our unsuccessful trials of patch-clamp recordings (1.7 \(\mu m^2\) was the typical size of a membrane patch) at low plasmid concentration. At higher plasmid concentrations, clustering of functional L-type Ca\(^{2+}\) channels has been reported in patch-clamp recordings (Keplinger et al., 2000).

It seems surprising that clustering of the L-type Ca\(^{2+}\) channel occurs in the expression system used, taking into account that HEK293 cells lack the excitation-contraction machinery of muscle cells which is believed to also govern the organization of L-type Ca\(^{2+}\) channels (Hofmann et al., 1994; Flucher et al., 1993; Flucher and Franzini-Armstrong, 1996). It appears from our experiments that interaction of the channels with the ordered arrays of the Ca\(^{2+}\)-release channels inside the muscle cell (Flucher et al., 1993) seems to not be essentially needed for aggregation of L-type Ca\(^{2+}\) channels. This result suggests that clustering is driven by self-aggregation of the protein.

In summary, we have demonstrated the viability of single-molecule fluorescence in vivo studies for quantitative elucidation of the dynamical embedding of L-type Ca\(^{2+}\) channels in the plasma membrane as characterized by their lateral and rotational mobility and state of aggregation. With the swift rate of advances occurring in molecular biology along with the complex interactions of proteins taking place, we expect single-molecule microscopy to present a new perspective for research on molecular interactions, protein dynamics, and signaling pathways in living cells.

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