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Nanodomain Ca^{2+} of Ca^{2+} channels detected by a tethered genetically encoded Ca^{2+} sensor

Lai Hock Tay¹, Ivy E. Dick¹, Wanjun Yang¹, Marco Mank³, Oliver Griesbeck³ & David T. Yue^{1,2}

Coupling of excitation to secretion, contraction and transcription often relies on Ca^{2+} computations within the nanodomain—a conceptual region extending tens of nanometers from the cytoplasmic mouth of Ca^{2+} channels. Theory predicts that nanodomain Ca^{2+} signals differ vastly from the slow submicromolar signals routinely observed in bulk cytoplasm. However, direct visualization of nanodomain Ca^{2+} far exceeds optical resolution of spatially distributed Ca^{2+} indicators. Here we couple an optical, genetically encoded Ca^{2+} indicator (TN-XL) to the carboxy tail of $Ca_V 2.2 Ca^{2+}$ channels, enabling near-field imaging of the nanodomain. Under total internal reflection fluorescence microscopy, we detect Ca^{2+} responses indicative of large-amplitude pulses. Single-channel electrophysiology reveals a corresponding Ca^{2+} influx of only 0.085 pA, and fluorescence resonance energy transfer measurements estimate TN-XL distance to the cytoplasmic mouth at ~55 Å. Altogether, these findings raise the possibility that Ca^{2+} exits the channel through the analogue of molecular portals, mirroring the crystallographic images of side windows in voltage-gated K channels.

¹ Department of Biomedical Engineering, Calcium Signals Laboratory, Ross Building, Room 713, 720 Rutland Avenue, Baltimore, Maryland 21205-2196, USA. ² Department of Neuroscience, Johns Hopkins University School of Medicine, Ross Building, Room 713, 720 Rutland Avenue, Baltimore, Maryland 21205-2196, USA. ³ AG Zelluläre Dynamik, Max-Planck-Institut für Neurobiologie, Am Klopferspitz 18, 82152 Martinsried, Germany. Correspondence and requests for materials should be addressed to D.T.Y (email: dyue@jhmi.edu).

a²⁺ signals extend pervasively across the intracellular expanse of most cells; yet, these signals exhibit remarkable specificity in activating appropriate Ca²⁺-modulated targets. Understanding how this selectivity arises remains a foremost question in Ca^{2+} biology^{1,2}. One strategy is to co-localize at molecular dimensions Ca²⁺ sources and targets. This co-localization is crucial to the local signalling of Ca²⁺ channels to nearby Ca²⁺-regulated ion channels and enzymes³⁻⁵ to neurotransmitter release⁶, as well as to excitation-contraction⁷ and excitation-transcription coupling^{8,9}. Intriguingly, co-localized sensors do not always respond preferentially to local Ca²⁺ sources, but can require the far weaker input of remote Ca²⁺ sources acting through a global Ca²⁺ selectivity paradigm¹⁰. Critical to signalling near Ca^{2+} sources is the nature of Ca^{2+} signals within a few tens of nanometers of a Ca^{2+} channel (nanodomain¹¹). Theoretical calculations¹²⁻¹⁶ postulate that nanodomain signals comprise Ca²⁺ pulses of equal and enormous amplitude (~100 µM), with each pulse synchronized to the millisecond stochastic openings of channels^{10,12}. Beyond the nanodomain, [Ca²⁺] rapidly dissipates with diffusion. Of note, predicted nanodomain [Ca²⁺] amplitudes vary considerably with assumed parameters¹⁷, and elegant experimental estimates of local Ca²⁺ signals^{18,19} only indirectly probe the nanodomain itself. Thus, the critical magnitude of nanodomain Ca²⁺ pulses (Ca_{spike}) has eluded explicit empirical comment. This uncertainty bears on key issues, including how spatial Ca^{2+} decoding occurs within the nanodomain^{10,20–22} and the Ca^{2+} channel number needed to trigger neurotransmitter vesicles^{23–25}.

Why have nanodomain Ca^{2+} signals remained elusive? Although fluorescent Ca^{2+} -sensitive chemical dyes have revealed much²⁶, visualization of nanodomain signals via freely diffusible dye molecules far exceeds the resolution limit of far-field microscopic approaches²⁷. In this regime, fluorescence from dye molecules outside the nanodomain will overshadow that from within²⁸. Even when imaged with total internal reflection fluorescence (TIRF) microscopy^{29–31} that illuminates a restricted TIRF volume within ~150 nm of the glass/cell membrane interface, freely diffusible dyes will still report Ca²⁺ from a region several-fold larger than a Ca²⁺ channel³².

One approach is to affix a Ca²⁺ indicator within the nanodomain, so as to support a form of near-field imaging³³. The Ca²⁺sensitive luminescent protein aequorin represents a traditional candidate³⁴. Recombinant aequorins have long been targeted to various subcellular compartments by genetic means³⁵⁻³⁷ but the limited-amplitude and consumptive nature of aequorin would challenge deployment in a channel nanodomain³⁴. Recently, promising results have been obtained using a biarsenical chemical fluorescent Ca²⁺ indicator (Calcium green FlAsH, CaGF) targeted to a tetracysteine tag implanted on Ca_V1.2 Ca²⁺ channels³⁸. CaGF features Ca²⁺ affinity and kinetics (K_d , ~55 µM; off rate, 2 ms⁻¹) well matched to the purported amplitudes and lifetimes of nanodomain Ca²⁺ pulses. Indeed, CaGF was potentially responsive to nanodomain Ca²⁺ fluctuations, but the observed fluorescence signals were highly heterogeneous, to an extent that precluded quantitative Ca²⁺ estimation. This heterogeneity was attributable to the diminutive and variable open probability of Ca_V1.2 channels (~0.1–0.3%), despite constitutive pharmacological manipulation by a channel opener. Indeed, the possibility of a large fraction of electrically silent Ca_V1.2 channels fits with their unusually large ratio of gating charge to ionic current³⁹. Also, although careful controls were performed for nonspecific labelling of other cysteine-rich sites in the cell by CaGF, such indiscriminate attachment would remain a generic concern for approaches involving chemical targeting of sensors. A third approach would be to employ genetically encoded Ca²⁺ indicators (GECIs) as near-field sensors. GECIs allow genetic fusion of sensor to a Ca²⁺ source, offering the greatest potential selectivity in positioning sensors. Among the most popular GECIs

are CFP/YFP-FRET (fluorescence resonance energy transfer)-based molecules such as cameleons⁴⁰, and GFP-intensity-based sensors like GCaMPs⁴¹. All these employ calmodulin (CaM) as their Ca²⁺ sensor, and convert binding of the sensor's own Ca²⁺/CaM with a target peptide into altered optical readouts. When targeted to plasmalemma, cameleons often lose Ca²⁺ sensitivity⁴², potentially via target peptide binding to elevated plasmalemmal CaM⁴³.

Accordingly, we here explore near-field imaging of nanodomain Ca²⁺ using TN–XL, a GECI⁴⁴ based on the Ca²⁺ sensor troponin C. TN-XL exploits Ca²⁺-driven conformational changes without a target peptide, thereby naturally resisting endogenous CaM interference⁴². Our specific approach is to fuse TN-XL to the carboxy terminus of the principle subunit of $Ca_V 2.2 Ca^{2+}$ channels, chosen for high open probability⁴⁵, relative absence of silent channels⁴⁶, and abundant plasmalemmal expression⁴⁷. To further improve the TN-XL signal-to-noise ratio at the surface membrane, we utilize TIRF microscopy to selectively illuminate only those Ca_V2.2/ TN-XL channels near the surface, thereby attenuating background signal from imperfectly targeted intracellular channels. Finally, to account for the high Ca²⁺ affinity and slow kinetics of GECIs⁴⁴ (off times, 100-900 ms for TN-XL) in the face of millisecond nanodomain Ca²⁺ pulses, we quantify the kinetics of TN-XL readouts, which our previous work suggests can distinguish between differing $Ca_{\rm spike}$ amplitudes⁴⁸. These strategies allow us to resolve nanodomain Ca^{2+} activity of $Ca_V 2.2$ channels undergoing native gating, without pharmacological manipulation.

Results

Functional Ca_V2. 2/TN-XL fusions at the surface membrane. Our strategy for probing nanodomain Ca^{2+} signals (Fig. 1) requires maintained Ca²⁺ responsiveness of genetically encoded sensors situated at the plasmalemma, which is a foremost challenge. Accordingly, we examined whether troponin-based TN-XL could respond to Ca²⁺ when targeted to this environment. As a baseline, Fig. 2a displays the properties of the well-studied, cytoplasmic form of TN-XL⁴⁴. Confocal images of an exemplar HEK293 cell, acquired under CFP and FRET imaging modes, reveal the expected pan-cytoplasmic distribution. An epifluorescence CFP image of another cell also demonstrates this generalized expression, and switching to TIRF mode indicates no preferential TN-XL accumulation within the plasmalemmal footprint of this same cell. To gauge TN-XL responsiveness, we measured, under TIRF imaging, the single-cell CFP (S_C, respective excitation and emission wavelengths at 442 and 485 nm) and FRET fluorescence signals (S_E excitation and emission at 442 and 545 nm), where the ratio of these signals $(R_{\rm E/C} = S_{\rm E}/S_{\rm C})$ furnishes the customary sensor readout of Ca^{2+} (Fig. 2b). In particular, the ratio $R_{F/C}$ was first determined in resting cells with cytoplasmic Ca²⁺ concentration ($\leq 0.1 \,\mu$ M) well below the K_d of the TN-XL²⁴, thus yielding R_{min} . To subsequently determine R_{max} , Ca_V2.2 channels were co-expressed as conduits of Ca²⁺ entry under whole-cell voltage clamp. To facilitate elevation of intracellular Ca²⁺, minimal internal Ca²⁺ buffering (1 mM EGTA) with elevated 10 mM extracellular Ca²⁺ was employed, such that repetitive step depolarizations (30 mV×500 ms pulses every 15 s) readily saturated $R_{F/C}$ at R_{max} . Thus, the ratio of R_{max} and R_{min} reports sensor dynamic range.

By contrast, for TN–XL fused to the farnesylation motif of Ras (TN–XL-Ras), sensors are sharply localized to the surface membrane in confocal and epifluorescence images (Fig. 2c), and there is marked enrichment within the TIRF footprint. Importantly, the Ca^{2+} dynamic range of TN–XL-Ras is nearly identical (Fig. 2d), despite residence within the plasmalemmal context. Yet more relevant are results for the Ca_V2.2/TN–XL fusions (Fig. 2e). Here constructs appear only partially localized to the surface membrane, consistent with typically imperfect trafficking of recombinant Ca^{2+} channels to plasmalemma⁴⁹. Still, there is substantial focalization



Figure 1 | Approach to resolving channel nanodomain Ca²⁺ signals.

(a) Conventional wide-field imaging using cytosolic chemical fluorescent dyes cannot resolve channel nanodomain Ca²⁺ signals. Blue shading denotes region of fluorescence excitation, which extends throughout the cell under wide-field imaging. (b), Design of the genetically encoded Ca²⁺ indicator TN-XL fused to the carboxy tail of the α_{1B} subunit of a Ca_V2.2 channel, yielding Ca_V2.2/TN-XL. For orientation, structure-function elements involved in calmodulin regulation are denoted on carboxy terminus²¹: EF, EF-hand region; IQ, IQ-domain for apoCaM binding. CFP denotes enhanced CFP. YFP denotes circularly permuted citrine. (c), Ca_V2.2/TN-XL constructs act as a 'near-field' sensor of nanodomain Ca²⁺. TIRF imaging evanescent wave illuminates only Ca_V2.2/TN-XL channels within -150 nm from the glass/cell membrane interface, as indicated by the blue-shaded region. This mode of excitation potentially excludes intracellular channels from consideration. When laser illumination angle Θ exceeds a critical angle Θ_C , TIRF illumination occurs.

of Ca_V2.2/TN-XL in the TIRF footprint, suggesting sufficient trafficking for nanodomain experiments (Fig. 1c). Critically, sensor dynamic range is spared when juxtaposed near Ca_V2.2 channels (Fig. 2f).

A second concern is the preservation of $Ca_V 2.2$ channel function upon fusion to TN-XL. Reassuringly, Ca^{2+} currents appeared nearly identical in Ca_V2.2/TN-XL (Fig. 3a, black) and Ca_V2.2 (grey) channels. In fact, fusing TN-XL to Ca_V2.2 enhances opening, as seen from hyperpolarization of tail-activation curves determined with 10-mM extracellular Ca^{2+} as charge carrier (Fig. 3b). Even at the single-channel level, Cav2.2/TN-XL fusions exhibited robust electrophysiological function, as illustrated by the exemplar single-channel current trace shown at high-gain magnification (Fig. 3c, noisy trace). We used 90 mM Ba^{2+} as a charge carrier to enhance resolution of elementary events, and evoked currents via ramp depolarization to readily resolve the open-channel conduction profile (downwardly convex grey relation). Multiple single-channel sweeps (Fig. 3d), displayed at lower-gain magnification, confirm the overall reproducibility and excellent resolution of elementary events under this protocol. Accordingly, numerous sweeps from multiple patches specify a highly reliable ensemble average current (Fig. 3c, red curve), closely similar to that of Ca_V2.2 channels without sensor fusion⁴⁵. Taking the ratio of the ensemble average current with the open-channel conduction profile (Fig. 3c, red and convex grey curves) yields a plot of open probability P_{Ω} versus voltage V (Fig. 3e, gray relation). This plot matches impressively with whole-cell activation (reproduced as open circles), after shifting along the voltage axis to account for contrasting surface-charge effects of single-channel versus whole-cell solutions. As with studies of Ca_V2.2 (ref. 45), the Ca_V2.2/TN–XL fusion exhibits a substantial P_{O} of 0.69 at 30 mV (Fig. 3e, red arrows), a feature advantageous for probing nanodomain Ca²⁺.

A final prerequisite is that $Ca_V 2.2/TN-XL$ fusions resist proteolysis. Accordingly, we utilized anti-GFP antibody to perform western immunoblots from cells expressing this construct (Fig. 3f, left lane). The other lanes correspond to cells expressing TN-XL-Ras (~71 kD), as well as untransfected cells (no signal). The absence of signal beneath the full-length ~320 kD band (Fig. 3f, left lane) substantiates maintained TN-XL fusion to channels (Fig. 3g).

Calibration of TN-XL fused to Ca_V2.2. channels. We next addressed more precise sensor calibration, to facilitate quantitative inferences about nanodomain Ca²⁺. Both kinetic and steadystate characterization would be ideal, given the millisecond kinetics of Ca²⁺ pulses within channel nanodomains, coupled with the comparatively slow response of free TN-XL (off rates of 1/150 to 1/850 1/ms⁴⁴). Accordingly, we devised a state-mechanism approximation of sensor performance, a 'forward transform' that maps rapid Ca²⁺ transient inputs onto slower sensor outputs, as previously established for an older variant of TN-XL (TN-L15), freely expressed in myocytes⁴⁸. Here we pursued an analogous approach, but attuned to Ca_V2.2/TN-XL constructs resident within the TIRF volume and driven by high-amplitude dynamic Ca²⁺ inputs pertinent to channel nanodomains. To produce such inputs, we activated ultra-large $Ca_V 2.2 Ca^{2+}$ currents (Fig. 4a) in HEK293 cells, featur-ing minimal exogenous intracellular Ca^{2+} buffering (1 mM EGTA); this configuration intentionally produced enormous spatial gradients of intracellular Ca²⁺ (Fig. 4b, inset), far exceeding that usually encountered. To estimate these gradients, we simultaneously monitored aggregate Ca²⁺ concentration (Fig. 4b, noisy dark trace), as deduced by imaging the entire cell for signals emanating from 10µM of the rapid chemical fluorescent Ca²⁺ indicator Fluo 4FF. The measured Ca^{2+} influx (Fig. 4a) and aggregate [Ca²⁺] signals were then used to explicitly constrain a radially symmetric Ca²⁺ diffusion mechanism, whose estimates of Ca²⁺ concentration, in differing cellular regions, are shown as red traces in Fig. 4b (Supplementary Fig. S1; Supplementary Methods). The estimated aggregate [Ca²⁺] concentration (Fig. 4b, lower red trace) closely approximates its measured analogue (dark noisy trace), and the estimated $[Ca^{2+}]$ at the surface membrane (upper red trace) peaks at ~80 μ M and later converges to the aggregate $[Ca^{2+}]$ (~20–25 µM). This estimate of submembranous $[Ca^{2+}]$ was reproducible in multiple cells



Figure 2 | Ca_V2.2/TN-XL fusion construct preserves function of sensor. (a) Behaviour of free TN-XL. Left to right: confocal (CFP filter), confocal (FRET filter), epi-fluorescence (CFP filter), and TIRF (CFP filter) images of HEK293 cells expressing cytoplasmic TN-XL. White scale bar, 10 μ m. Bar at far left pertains to all confocal images. Bar at middle right pertains to all epifluorescence and TIRF images. (b) TN-XL ratio ($R_{F/C} = S_F/S_C$) measured in resting cells (R_{min}) (grey bar) and in cells under high Ca²⁺ (R_{max}) (black bar). Data shown as mean±s.e.m. with number of cells in parentheses. (c,d), Behaviour of TN-XL-Ras (membrane targeted TN-XL). Format as in (a,b), respectively. (e,f), Behaviour of Ca_V2.2/TN-XL (N-type channel fused to TN-XL). Format as in (a,b), respectively.

exhibiting like Ca²⁺ influx, and was thereby taken as the relevant Ca^{2+} input in parallel experiments performed with $Ca_{V}2.2/TN-XL$ fusion constructs, under TIRF imaging. The green trace in Fig. 4c displays the corresponding sensor output from GECIs in the submembranous TIRF volume, averaged from multiple cells exhibiting Ca²⁺ influx matching that in Fig. 4a. The steady-state plateau of the response (Fig. 4c, grey dashed line at $R_{F/C} = 1.78$), at a near steady Ca²⁺ concentration of ~20–25 μ M, enabled us to calculate \sim 17.3µM as the half-response point of the steady-state response curve for Ca_V2.2/TN-XL constructs in the TIRF volume (Supplementary Methods). This differs from the 2.5 µM half-response point for free TN-XL⁴⁴, consistent with previous observations that sensor performance can change with cellular/molecular environment^{38,42}. Accordingly, steady-state response data for free TN-XL⁴⁴ were shifted along the $[Ca^{2+}]$ axis by ×16.3/2.5 to provide an appropriate steady-state profile for Ca_V2.2/TN-XL constructs (Fig. 4d, green symbols). In all, these steady-state data and the dynamic sensor response (Fig. 4c, green trace) to a specified submembranous Ca²⁺ input (Fig. 4b, upper red trace) furnished the constraints required to deduce a forward transform for Ca_V2.2/TN-XL.

Fig. 4e displays the corresponding state-diagram approximation of Ca_V2.2/TN–XL responsiveness to Ca²⁺, based on a mechanism for an older troponin-based GECI⁴⁸ (TN-L15). State UB₀ ('unbound') represents the Ca²⁺-free conformation of the sensor, characterized by a lower FRET-ratio output of R_{min} . On binding of a single Ca²⁺ to the N-lobe of troponin, the sensor adopts one of two alternate 'bound' conformations (B₁ or B₂), both featuring the same elevated FRET-ratio output of R_{mid} . The arrangement, thus far, is identical to that previously established for TN-L15 (ref. 48). The improved performance of TN–XL arises, via engineering the C-lobe of troponin, to dynamically bind and unbind two signalling Ca²⁺ ions, thereby driving a further conformational change of the sensor⁴⁴. To account for this feature, we allowed two additional Ca²⁺ ions to bind and induce a third bound state^{50,51} (B₃, dashed box), exhibiting the highest FRET-ratio output of R_{max} . Numerical simulations of this scheme (Supplementary Methods), coupled with error minimization via parameter variation, yielded impressive fits to the target constraints above (Fig. 4c,d, red curves), using the sensor parameter estimates in Table 1. Accordingly, these parameters and the scheme in Fig. 4e furnish a steady-state and dynamic representation of Ca_V2.2/TN–XL responsiveness to Ca²⁺, a 'forward transform' (Fig. 4f) that is potentially appropriate for making inferences about channel nanodomain Ca²⁺ fluctuations.

 $Ca_V 2.2/TN-XL$ sensors respond to nanodomain Ca^{2+} signals. This transform could only be utilized, however, if a substantial fraction of Ca_V2.2/TN-XL sensors respond to channel nanodomain Ca²⁺ signals, as specifically produced by individual channels fluxing Ca²⁺ into their own nanodomain. Thus far, we had only demonstrated that Ca_V2.2/TN-XL sensors in the TIRF volume respond well to a generalized increase in submembranous [Ca²⁺], but this would occur whether the majority of channel-sensor fusion constructs are properly trafficked to the surface membrane, or still plentiful in submembranous vesicles within the TIRF volume. To address this issue, we again undertook simultaneous recordings of whole-cell Ca²⁺ current and TIRF imaging of Ca_V2.2/TN-XL sensors, but here with high intracellular Ca2+ buffering present to restrict Ca²⁺ elevations to the nanodomain of active channels that flux Ca²⁺. Specifically, we dialysed cells with 10 mM EGTA, yielding a Ca²⁺ nanodomain radius of ~40 nm (ref. 13) (Fig. 5, top cartoon),



Figure 3 | Ca_V2.2/TN-XL fusion construct preserves function of channel. (a) Exemplar whole-cell currents of HEK293 cells expressing Ca_V2.2 channels (grey) or Ca_V2.2/TN-XL channels (black), using 10 mM Ca²⁺ as charge carrier, and 10 mM EGTA intracellular Ca²⁺ buffering. (b), Whole-cell tail-activation curves of Ca_V2.2 (grey) and Ca_V2.2/TN-XL (black), obtained from records as in (a). Data shown as mean±s.e.m. with number of cells in parentheses. (c), Single-channel analysis of Ca_V2.2/TN-XL. Top, On-cell ramp-voltage waveform delivered every 5-15 s. Bottom, exemplar single-channel sweep shown at high-gain magnification (100-200 per patch). 90 mM Ba²⁺ as charge carrier; 2 kHz low-pass filtering. Leak-subtracted traces were averaged, yielding red *I*-V curve. Unitary current relation (convex grey curve) was fitted to the open-channel current level using GHK equation¹⁰. (d), Multiple exemplar single-channel sweeps from another patch, shown at lower-gain magnification to impart a sense of the overall reproducibility and excellent resolution of elementary events. Format as in c. (e), Single-channel *P*_O-V relation (grey, with fit in black, average of *n* = 4 patches), determined by dividing *I*-V curve (c, red trace) by unitary current relation (c, grey, GHK fit). *P*_O-V curve was calibrated in voltage by aligning the single-channel activation curve with the whole-cell curve reproduced from (b) (open circles). From calibrated *P*_O-V curve, the *P*_o at 30 mV is 0.69. (f) Immunoblots of HEK293 cell lysates probed with anti-GFP antibody; (left) Ca_V2.2/TN-XL at ~320 kD; (middle) TN-XL-Ras at ~71kD; (right) control (untransfected) cells for reference. (g) Lack of lower molecular-weight species in leftmost lane of (f) excludes proteolysis of sensor from channel.

only twofold larger than electron micrographic estimates of Ca²⁺ channel diameters³². Additionally, we only investigated cells with modest Ca²⁺ currents (~1 nA with 10 mM Ca²⁺ as charge carrier) and little Ca²⁺-dependent inactivation²¹ (CDI), so as to exclude Ca²⁺ buffer depletion. Fig. 5a summarizes the results of an exemplar cell satisfying this criterion, with the displayed data evoked by a single voltage pulse. Clearly present is a transient decrease in CFP fluorescence (S_C, cyan trace), accompanied by a corresponding phasic elevation of the FRET ratio $R_{F/C}$ (green trace). Exponential fits (solid black curves) revealed dominant rise and fall times of 880 and 2,000 ms for $S_{\rm C}$, and 880 and 400 ms for $R_{\rm F/C}$. Both the return of TN-XL waveforms towards baseline, and the modest CDI of channels under elevated Ca²⁺ buffering²¹ (Fig. 5a compared with Fig. 4a), argue against unintended depletion of Ca2+ buffer. Accordingly, these results suggest responsiveness of Ca_V2.2/TN-XL sensors to genuine channel nanodomain Ca²⁺ signals. By contrast to the exemplar, about half the cells with currents of this magnitude and limited CDI exhibited no appreciable change of TN–XL readouts, presumably due to poor trafficking of active channels. Such cells were excluded from further analysis.

Forward transform estimates of nanodomain Ca^{2+} amplitudes. As a prelude to more rigorous assessment of $Ca_V2.2/TN-XL$ responses to nanodomain signals, we averaged signals from multiple responsive cells dialysed with 10 mM EGTA. The mean responses (Fig. 5b) were similar to those of the exemplar, with nearly identical characteristic time constants. This reproducibility encouraged us to quantitatively scrutinize these averaged waveforms. In particular, although the robust responses observed here under elevated Ca^{2+} buffering do indicate an appreciable fraction of active channels within the TIRF volume (Fig. 5c, active), the data do not exclude the possibility of a still substantial fraction of channels that



Figure 4 | Calibration of Ca_V2.2/TN-XL fusion construct. (a) High-amplitude Ca²⁺ influx into HEK293 cell expressing Ca_V2.2 channels, observed with 10 mM external Ca²⁺ and minimal internal Ca²⁺ buffering of 1mM EGTA. (**b**) Simultaneous $[Ca^{2+}]$ measurement (dark trace) deduced from 10 μ M Fluo-4FF Ca²⁺-indicator fluorescence from entirety of same cell (514 nm excitation, 545 nm emission). Ca²⁺ determination rendered ratiometric by inclusion of 2.5 μ M red Alexa 568 dye (514 nm excitation, 580 nm long-pass emission). Inset, cartoon of spatial Ca²⁺ gradients in this cell. Red traces, projected $[Ca^{2+}]$ waveforms from Ca²⁺ diffusion mechanism constrained by Ca²⁺ input in (**a**) and aggregate $[Ca^{2+}]$ (dark trace): lower red trace, aggregate $[Ca^{2+}]$; upper red trace, submembranous $[Ca^{2+}]$. (**c**), Average FRET-ratio ($R_{F/C}$) response of Ca_V2.2/TN-XL channels in TIRF volume (green trace); same protocol as in (**a**,**b**). Red trace, output of TN-XL forward transform in (**e**), when driven by submembranous $[Ca^{2+}]$ input (**b**, upper red trace). (**d**) Normalized steady-state FRET-ratio response ($R_{norm} = (R_{F/C} - R_{min})/(R_{max} - R_{min})$ of Ca_V2.2/TN-XL to $[Ca^{2+}]$ (experimental green symbols). Red trace, steady-state output of TN-XL forward transform in (**e**) when driven by differing fixed $[Ca^{2+}]$. (**e**) State-diagram representation of Ca_V2.2/TN-XL. Red circles, Ca²⁺ ions. UB, conformation with no calcium bound. B, various conformations with calcium bound. (**f**) System of states and corresponding fluorescence/FRET properties constitutes a forward transform that predicts TN-XL sensor outputs as a function of time-varying Ca²⁺ inputs.1)

Fluorescence parameters*	Value
Ca _V 2.2/TNXL R _{min}	1.10 (<i>n</i> =20 cells; Fig. 2f)
Ca _V 2.2/TNXL R _{mid}	1.53†
Ca _V 2.2/TNXL R _{max}	2.10 (n=15 cells; Fig. 2f)
Ca _V 2.2/TNXL α_0	2.00 (n=8 cells)
Ca _V 2.2/TNXL α_1	1.57‡
Kinetic Parameters†	
k ₀₁	174.4 ms ⁻¹ M ⁻¹
k ₁₀	0.00239 ms ⁻¹
k ₁₂	0.00174 ms ⁻¹
k ₂₁	$0.00309 \mathrm{ms}^{-1}$
k23	0.126×10 ⁶ ms ⁻¹ M ⁻²
k ₃₂	$1/733{\rm ms}^{-1}$

Table 1 | Cav2.2/TN-XL parameters for HEK293 cells under

are present in the plasmalemma but fail to open (Fig. 5c, silent), or resident within submembranous vesicles within the TIRF volume (Fig. 5c, intracellular). Such silent and/or intracellular channels would contribute static background fluorescence that could complicate quantitative interpretation. An indication of this scenario comes from exponential extrapolation of the S_C waveform decline (Fig. 5b, dashed curve), which asymptotes in the range of ~0.8. By contrast, under the assumption that all channels are active, simulations of the forward transform in Fig 4e,f indicate that, over a large range of plausible nanodomain Ca²⁺ pulse amplitudes (\geq 35µM), $S_{\rm C}$ waveforms would asymptote near Ca²⁺-saturating levels of ~0.5 (Supplementary Fig. S2; Supplementary Methods), as specified by the experimentally determined $1/\alpha_0$ value (defined in Table 1). Hence, the experimentally deduced asymptote of ~0.8 (Fig. 5b) suggests that ~40% of Ca_V2.2/TN–XL channels are active within the TIRF volume, whereas ~60% of channels are intracellular or silent. Cognizant of this configuration, we correct $S_{\rm C}$ and $R_{\rm F/C}$ waveforms to reflect only active channels, simply by subtracting ~60% of the baseline amplitude of CFP ($S_{\rm C}$) and FRET ($S_{\rm F}$) signals, and taking a ratio of these subtracted signals to obtain a corrected $R_{\rm F/C}$. Fig. 6a shows the correct signals ($\overline{S}_{\rm C}$, $\overline{R}_{\rm F/C}$), averaged over multiple cells. These waveforms were then suitable for making inferences about nanodomain Ca²⁺.

That nanodomain Ca²⁺ signals take the form of equi-amplitude Ca²⁺ pulses, synchronized to the millisecond opening of channels, is widely accepted^{10,12} (cartoon, Fig. 6b); however, the critical amplitude of such pulses (Ca_{spike}) has only been inferred via simulations of Ca²⁺ diffusion¹²⁻¹⁶. To estimate Ca_{spike} empirically, we analysed the corrected Ca_V2.2/TN-XL responses (Fig. 6a), using the forward transform in Fig. 4e,f. The experimental waveforms might initially appear difficult to conceptualize, as each TN-XL sees a different stochastic record of Ca2+ pulses driven by singlechannel openings. However, because the response of each TN-XL is slow (Table 1; compare Fig. 4b,c), relative to the millisecond duration of individual Ca²⁺ spikes, we can adopt a previously established kinetic simplification¹⁰ that permits the Ca²⁺-driven forward reaction rates in Fig. 4e to be treated, not as stochastic entities that fluctuate with each Ca²⁺ pulse, but as smoothly changing entities $k_{01} \cdot Ca_{\text{spike}} \cdot P_{O}(t)$ and $k_{23} \cdot Ca_{\text{spike}}^2 \cdot P_{O}(t)$. Importantly, these entities are to the first order equal for all active Ca_V2.2/TN-XL sensors, such that the collective output from multiple sensors, as displayed in Fig. 6a, would approximate that for any individual



Figure 5 | Ca_V2.2/TN-XL sensors respond to channel nanodomain Ca²⁺ signals. Top cartoon, Ca²⁺ nanodomains under elevated Ca²⁺ buffering, where Ca²⁺ elevation is restricted to nanodomains, each affiliated with individual channel. 40 nm nanodomain calculated with equation (1), f=1 (classical assumption), unitary current i = 0.085 pA, and conservatively requiring [Ca²⁺] ~60 nM (the approximate resting physiological Ca²⁺ concentration). (a) Simultaneous records of whole-cell current (top) and TIRF fluorescence signals (middle and bottom) from HEK293 cell expressing $Ca_V2.2/TN-XL$, using 10 mM external Ca²⁺ and 10 mM internal EGTA Ca²⁺ buffering. Shown are data from a single trial, evoked by the first step depolarization in the cell. Top, voltage and current waveforms, with moderately sized current chosen to minimize buffer consumption. Middle, corresponding CFP fluorescence signal after normalization to baseline level (S_C, noisy trace). Black curve, approximate exponential fits (initial phase: 200 ms (10%), 880 ms (90%); recovery phase: 2,000 ms (100%)). Vertical ticks, 0.1 increment. Bottom, corresponding FRET-ratio signal (R_{F/C}, noisy trace). Black curve, approximate exponential fits (initial phase: 200 ms (10%), 880 ms (90%); recovery phase: 400 msec (70%), 2,000 ms (30%)). Vertical ticks, 0.1 increment. (b) Average $Ca_{V2.2}/TN-XL$ responses to nanodomain Ca^{2+} signals, as isolated with 10 mM internal EGTA Ca^{2+} buffering. Protocol and format as in (a). n = 4 cells, with records taken from the first-step depolarization delivered in cells. CDI sometimes increased with repeated depolarizations, reflecting buffer depletion with attendant slowing of $R_{F/C}$ waveform decay. Cells exhibiting peak currents \leq 500 pA were not considered, as these cells demonstrated negligible $R_{F/C}$ transients, presumably reflecting insufficient trafficking of active channels. Solid-black-curve exponential fits were similar, with S_C exponential fit (initial phase: 200 ms (10%), 880 ms (90%); recovery phase: 500 ms (80%), 2,500 ms (20%)), and R_{F/C} fit (initial phase: 200 ms (10%), 880 ms (90%); recovery phase: 400 msec (70%), 2,000 ms (30%)). Dashed black curve, continuation of initial phase S_C exponential fit, which asymptotes at lower horizontal black line in middle. (c) Schematic of various classes of Ca_V2.2/TN-XL species in TIRF volume. Blue shading indicates region of excitation by evanscent wave in TIRF volume.

sensor on average. Thus, using our forward transform (Fig. 4e,f) to predict experimental waveforms becomes simple. Single-channel data (Fig. 3e) permit average whole-cell currents (Fig. 5b) to be rescaled as an open probability P_O waveform (Fig. 6a,b); this Po drives straightforward numerical integration of the differential equations corresponding to the transform in Fig. 4e, and this integration yields predictions of experimental waveforms, with the magnitude of Caspike, being the only unknown (Supplementary Methods). Whereas presumed Ca_{spike} values of 15 or 240 μ M yield predicted $R_{F/C}$ trajectories (Fig. 6a, grey smooth curves) that are patently inconsistent with the experimental curve, $Ca_{spike} = 60 \,\mu M$ produces an impressive fit to data (smooth red curves). Indeed, error analysis (Fig. 6c) confirms 60 µM as a best estimate of Ca_{spike}. As a conservative lower limit, we performed transform analysis on uncorrected waveforms (Fig. 5b), yielding a still-large estimated *Ca*_{spike} ~15 µM (Supplementary Fig. S3; Supplementary Methods). Fits to uncorrected data were notably inferior, further prompting us to favour Ca_{spike} estimates from the background-corrected data in Fig. 6a. For these data, the shading in Fig. 6c encompasses fits with errors similar to the minimum, yielding an approximate Ca_{spike} range of 45-80 µM. This estimate, combined with determination of the underlying single-channel current amplitude, can potentially furnish valuable insight into Ca²⁺ diffusion properties within the nanodomain itself (Fig. 7).

Discussion

The fusion of Ca_V2.2 channels to the genetically encoded Ca²⁺ sensor TN–XL here furnishes a new tool for detecting submembranous Ca²⁺ signals near these 'N-type' Ca²⁺ channels (Fig. 1b). Our particular application concerns the nanodomain Ca²⁺ transients accompanying individual channel openings and closings. These transients have eluded direct experimental comment, until the recent advent of TIRF microscopy to image single-channel activity^{30,31}, and the availability of targetable chemical-fluorescent Ca²⁺ indicators³⁸. Here we demonstrate an alternative approach exploiting TIRF/patch–clamp electrophysiology and our Ca_V2.2/TN–XL fusion. This tactic yields a first empirical estimate of the amplitude of nanodomain Ca²⁺ pulses (*Ca_{spike}*). This parameter figures crucially in the Ca²⁺ activation of co-localized channels and enzymes², evoked release of nearby neurotransmitter vesicles²⁵, and excitation–transcription coupling^{8,9}.

A limitation of the present study concerns the incomplete targeting of active Ca_V2.2/TN–XL channels to the surface membrane in the TIRF volume, necessitating approximate correction of static background fluorescence. The potential error in such correction renders our Ca_{spike} determination as a coarse approximate estimate. Nonetheless, the approach taken will prove valuable as a means to improve channel trafficking arise, and GECI technology progresses.



Figure 6 | Estimates of nanodomain Ca²⁺ amplitude from Ca_V2.2/TN-XL responses. (a) Average Ca_V2.2/TN-XL responses to nanodomain Ca²⁺ signals, corrected to reflect signal of active channels alone. Top, black trace plots average whole-current, rescaled to open probability P_O by normalizing peak current during 30-mV pulse to 0.69. Grey trace, exponential fit to P_O waveform. Middle, cyan trace plots average \overline{s}_{C} , corrected CFP signal. Red trace, forward transform prediction of experimental \overline{s}_{C} using $Ca_{spike} = 60 \,\mu$ M. Bottom, Green trace plots average $\overline{R}_{F/C}$, corrected FRET-ratio signal. Red trace, forward transform prediction of experimental $\overline{R}_{F/C}$ using $Ca_{spike} = 60 \,\mu$ M. Bottom, Grey traces, transform prediction with Ca_{spike} as labelled. (b), Nanodomain Ca²⁺ signals input to forward transform for prediction of Ca_V2.2/TN-XL responses in (a). Top, P_O waveform reproduced from (a). Middle, Schematic of stochastic Ca²⁺ pulses in nanodomain. Bottom, forward transform box with parameters in Table 1. (c) $Ca_{spike} = 60 \,\mu$ M). Cusp of plot denotes $Ca_{spike} = 60 \,\mu$ M as best fit. Shaded zone specifies approximate confidence region, where fits yield errors within 25% of minimum.

With this proviso, we consider other long-sought-after Ca²⁺ channel signalling properties that could, in principle, be accessed with our strategy. First, we estimate the gain factor A, defined as the ratio of Ca_{spike}^{2+} to unitary current *i* fluxing through individual Ca^{2+} channels¹². As such, the magnitude of *A* is crucial for local Ca²⁺ channel signalling to downstream nanodomain targets. Given knowledge of Ca_{spike}, we could estimate A itself, so long as the *i* were known under the conditions of Ca_{spike} determination (Fig. 6, with 10 mM Ca²⁺ as the charge carrier). The single-channel data thus far (Fig. 3c,d) were obtained with 90 mM Ba^{2+} to facilitate resolution of open probability; these data could not be utilized here. The relevant single-channel data for parameter A specification would entail several-fold smaller signals^{25,52}. Nonetheless, despite signals near the limits of detection, we acquired well-resolved unitary currents under these conditions (Fig. 7a). These data yield $i = 0.085 \pm 0.003$ pA (n = 5 patches) at 30 mV (as in Fig. 6), resembling values recorded in native Ca_V2.2 channels²⁵. Combining *i* with our estimates of Ca_{spike} (centre, ~60 μ M; range, 45–80 μ M), we obtain A, ~700 μ M/pA (Fig. 7b, shaded regions).

Second, if the distance from the cytoplasmic mouth of the channel to the sensor (r_{sensor}) were also known, near-field Ca²⁺ sensing could comment powerfully on Ca²⁺ diffusion within the nanodomain itself. We therefore gauge r_{sensor} from FRET measured between the CFP of our Ca_V2.2/TN–XL fusion, and YFP fused to the base of the channel amino terminus (Fig. 7c). To eliminate crosstalk, YFP within TN–XL itself is replaced with the inert analogue amber⁵³. FRET efficiency was thus determined⁵⁴ (Fig. 7d), yielding an estimated $r_{sensor} \sim 55$ Å (Methods).

With both parameter A and r_{sensor} in hand, we scrutinize nanodomain Ca²⁺ diffusion via classic point-source Ca²⁺ diffusion, as given by a generalized Neher-Stern equation^{13,16}

$$A = \frac{Ca_{\text{spike}}}{i} = \frac{(1/f)}{4 \cdot \pi \cdot F \cdot D_{\text{Ca}} \cdot r_{\text{sensor}}} \cdot \exp\left[\frac{-r_{\text{sensor}}}{\sqrt{D_{\text{Ca}}/(k_{\text{Bon}} \cdot B_{\text{T}})}}\right]$$
(1)

where r_{sensor} is the distance of the TN–XL sensor from the point source of Ca²⁺ influx; D_{Ca} is the diffusion coefficient of free Ca²; F is Faraday's constant; k_{Bon} is the on rate for Ca²⁺ binding to EGTA; and B_{T} is the internal EGTA concentration (10 mM). The parameter f, specifying the fraction of half-infinite space into which radial Ca²⁺ diffusion occurs, has always been set at unity (Ca²⁺ diffuses into a full half-infinite space), absent any evidence to the contrary. Intriguingly, our estimates of A and r_{sensor} square poorly with the traditional view (Fig. 7e, grey curve with f=1), but fit well with $f \sim 0.53$ ($0.53 \times 2\pi$ steradians, solid curve). This outcome may represent an early functional indication of fenestrated Ca²⁺ egress from the channel into cytoplasm (Fig. 7f), according with crystallographic structures of K channels⁵⁵.

Thus, nanodomain Ca^{2+} diffusion could focus Ca^{2+} through particular molecular geometries within the channel-signalling complex, raising the possibility of enhanced preferential signalling to target molecules near or within exit portals (Fig. 7f, grey ball). Indeed, although previous studies elegantly suggest that distinctive unitary current amplitudes *i* render $Ca_V 2.2$ channels favourable for triggering vesicle fusion²⁵, it is also plausible that differing *A* and *f* values also factor into such optimization. The latter possibilities represent an intriguing, but nearly unexplored, realm of Ca^{2+} biology. Although other interpretations are certainly viable, further empirical estimates of Ca_{spike} , parameter *A*, and r_{sensor} will likely aid in the dawning of this field of inquiry.

Methods

Molecular biology. Ca_V2. 2/TN–XL (Ca_V2.2 is of human origin (NM000718)) fusion construct was made by PCR, using fusion primers. Pfu polymerase (Strategene) was used for fidelity. To make Ca_V2.2/TN–XL, we first generated a Ca_V2.2 channel with the C-terminus truncated at the 2180th amino acid (without a stop codon), yielding Ca_V2.2A2180 stopless/XbaI in pcDNA3.1. This was made by cutting Ca_V2.2 with Xho I and Xba I, and replacing the resulting fragment with the PCR product amplified from Ca_V2.2 between the Xho I and 2180th amino-acid sites using forward primer 5'-CGCATCAGTTACAATGACA TG-3', and reverse primer 5'-CTGTCTAGAAGCACCAGATGTTGACAGCA-3'.



Figure 7 | Potential fenestrated Ca²⁺ egress within Ca_V2.2 channel nanodomain. (a) On-cell single-channel ramp experiments, with 10 mM Ca²⁺ as charge carrier to accord with whole-cell experiments; 700-Hz low-pass filtering. Voltages adjusted to align single-channel activation with whole-cell activation curves. Exemplars from single patch, with openings to GHK relation (grey curve). Unitary current *i* at 30 mV (green circles), 0.085 pA in 5 patches. (b) Gain factor *A*, slope of Ca_{spike} versus *i* relation. Shading, various estimates of *A* over confidence range of Ca_{spike} . Sloped dashed line, lower-limit estimate of *A* using uncorrected data. (c) Schematic, FRET construct to estimate r_{sensor} . Substituting circularly permuted amber for circularly permuted Citrine of TN-XL maintains sensor-channel configuration. (d) Estimating r_{sensor} via FRET. Each black symbol, individual HEK293 cell expressing construct in (c). Two outputs for each such cell obtained via 3³-FRET analysis⁵⁴ of fluorescence images: FRET efficiency and ratio of donor (CFP) to acceptor (YFP) molecules (D/A ratio), both plotted versus CFP intensity (\propto construct expression level). Data from cells exhibiting D/A ratio ~1 (as expected from c) are displayed and analysed (bottom subpanel, black line at unity). Invariant FRET efficiency with increasing CFP intensity (top subpanel) excludes spurious FRET, facilitating interpretation of FRET efficiency (-0.35, blue line, top subpanel, left axis) as indicative of r_{sensor} ~55Å (top subpanel, right axis). Grey symbols, control cells expressing YFP and CFP separately, yielding efficiencies ~0. (e) A versus sensor distance from channel mouth (r_{sensor}), calculated with Neher-Stern equation (equation (1)), using parameters^{13,60} D_{Ca} = 0.4 µm²ms⁻¹ and k_{Bon} = 10⁻³ µM⁻¹ms⁻¹ for EGTA. Grey curves, plots with *f* values as labelled. Mapping *A* from (b) with r_{sensor} from (d) yields *f* ~0.53 (black curve) and range (0.4-0.7, shading). (f) Schemat

Next, TN–XL was PCR-amplified with an Xba I site upstream of CFP (ATG removed) and a Nhe I/Spe I site downstream of the stop codon of circularly permuted Citrine. The forward and reverse primers used were 5'-CGGCCGCCA CCTCTAGAGTGAGC-3', and 5'-CAGACTAGTGCTAGTCCTCGATG TTGTGGC-3' respectively. Finally, the TN–XL PCR fragment was spliced into the Ca_V2.2Δ2180 stopless/XbaI construct at the Xba I site to give our Ca_V2.2/TN–XL construct.

For the variant construct in Fig. 7c, the circularly permuted Citrine in TN-XL was replaced with circularly permuted amber⁵³, and YFP fused to the amino terminus of $Ca_V 2.2$ after removal of the first 81 amino acids²² (nearly the entire amino terminus). Complete sequence analysis of regions subjected to PCR was performed.

HEK293 cell culture and transfection. HEK293 cells were cultured on No-1 25-mm glass cover slips (Bellco glass) in 10 cm plates. These slips were coated overnight with 0.01% (wt/v; diluted 1:5 in 0.1 M borate buffer) poly-D-lysine (Sigma), and washed with ddH₂O on the day cells were seeded onto the slips. The poly-D-lysine coating facilitates proper whole-cell voltage control at the membrane/glass interface⁵⁶, and also increases electrostatic attraction between the glass surface and cells, thus favouring cell attachment and optimal TIRF imaging. Cells were transiently transfected, using calcium-phosphate precipitation⁵⁴, with Ca_V2.2/TN-XL, rat brain β_{2a} ⁵⁴, and rat brain α_{2b} ⁵⁴ (10µg each). Ca_V2.2 and β_{2a} -CFP were substituted for Ca_V2.2/TN-XL and β_{2a} , respectively, in some experiments. Cotransfection of channels with a number of molecules did not appreciably improve channel trafficking in our system: PI3K and Akt/PKB⁵⁷, ORL receptors⁵⁸, and

dominant-negative dynamin⁵⁹. TN–XL and TN–XL-Ras sensors were transfected using FuGENE6 (Roche).

Electrophysiology. Whole-cell and single-channel current records were obtained at room temperature 1–3 d post-transfection, using patch–clamp amplifiers Axopatch 200B and Axopatch 200A (Axon Instruments), respectively.

For whole-cell recordings, the cells were bathed in external solution containing (in mM): TEA-MeSO₃, 140; HEPES (pH 7.4 with TEA-OH), 10; and CaCl₂, 10; at 300 mOsm, adjusted with glucose. The internal solution contained (in mM): Cs-MeSO₃, 114–135; CsCl, 5; EGTA, 1 or 10; MgCl₂, 1; MgATP, 4; HEPES (pH 7.4 with CsOH), 10; and 295 mOsm with Cs-MeSO₃. In some experiments, 10 μ M Fluo 4FF and 2.5 μ M Alexa 568 (Invitrogen, Molecular Probes) were included for ratiometric Ca²⁺ determination. Electrodes were pulled from borosilicate glass capillaries (WPI MTW 150-F4) and had pipette resistances ranging from 1.5 to 2.5 MΩ before 75% series resistance compensation. Voltage pulses were applied at 90 s intervals. Currents were filtered at 5 kHz, and digitized at 25 kHz. Leak and capacitance transients subtracted by P/8 protocol. Data acquired and analysed with custom MATLAB scripts (Mathworks, Natick, MA, USA).

Single-channel recordings were all conducted in the on-cell configuration. The bath contained (in mM): K Glutamate, 132; KCl, 5; NaCl, 5; MgCl₂, 3; EGTA, 2; glucose, 10; and HEPES (pH 7.4 adjusted with KOH), 20 at 300 mOsm adjusted with glucose. This bath solution zeroed the membrane potential. The pipette solution contained, (in mM): BaCl₂, 90; TEA-MeSO₃, 20; HEPES (pH 7.4 adjusted with TEA-OH), 10 at 300 mOsm, adjusted with TEA-MeSO₃. In some experiments, 10 mM CaCl₂ was substituted for BaCl₂ and TEA-MeSO₃ correspondingly

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increased. To minimize electrical noise, $5-20\,M\Omega$ patch pipettes were pulled from ultra-thick-walled (2 mm outer diameter, 1.16 mm inner diameter) borosilicate glass (BF200-116-10, Sutter Instruments), and coated with Sylgard. Seal resistance was ~80 G\Omega. Voltage pulses applied at repetition intervals of 5-15 s; data sampled at 70 µsec intervals, and filtered at 2 kHz (Fig. 3c,d) or 700 Hz (Fig. 7a).

TIRF microscopy. On the day of experiments, the cover slips on which cells were cultured were mounted on a custom bath chamber for TIRF imaging. Fluorescence measurements of single cells were performed on a Nikon Eclipse TE2000U microscope, equipped with a Plan Apo 60× TIRF objective (NA 1.45 oil immersion). Nikon immersion oil, with nd = 1.515 (23 °C) was used. For TN-XL imaging, CFP excitation was delivered by a 16-mW Melles Griot 56ICS 442-nm diode laser. For Fluo 4FF/Alexa 568 imaging, excitation was delivered by a 40-mW Melles Griot IMA100 argon laser featuring a 514-nm line. For TN-XL imaging, the optical path diagrammed in Supplementary Fig. S4 was used, as described below. A high-resolution CoolSNAP HQ CCD camera (Photometrics), driven by IPLab (Scanalytics) scientific imaging software, was used for cell orientation and verification under TIRF microscopy. Fast quantitative data, averaged across single cells, were instead acquired via photomultiplier tubes (PMTs), as follows. Single-cell fluorescent signals was selected by the image-plane pinhole in a D-104C dual-channel photometer (PTI), and sent to a dichroic mirror (510dclp) that divides the fluorescence emission signals into CFP and FRET signals. These two signals passed through their respective emitters (HQ485 per 30 m and HQ545 per 40 m, Chroma), before being captured by two Hamamatsu R1527 PMTs. Fluorescent signals were filtered at 2 kHz and sampled at 25 kHz, and reflect the activity of several thousand Cav2.2/ TN-XL molecules. Shutter control, dark-current subtraction, and data acquisition were controlled by custom MATLAB programs. For Fluo 4FF/Alexa 568 imaging, the optical path was modified slightly. Single-cell fluorescence was isolated via the same pinhole and then sent to a dichroic mirror (545 dclp) that parsed signals into Fluo 4FF and Alexa 568 signals. These two signals passed through their respective emitters (545/40 nm bandpass and HQ580 nm lp, Chroma), before being captured by the same PMTs. The apparent K_d for Fluo 4FF/Alexa 568 ratios under our recording conditions was 20 µM, as determined in live HEK293 cells dialysed with Ca²⁺ dye and a series of calibrated [Ca²⁺] internal solutions.

Widefield FRET microscopy. 3³-FRET efficiency determination, along with estimation of the ratio of donor to acceptor molecules, was performed within individual live HEK293 cells imaged through a image-plane pinhole according to CFP, FRET and YFP filter cubes⁵⁴. Moreover, r_{sensor} in Fig. 7d was estimated using a Forster distance of 49 Å (ref. 54).

Immunoblots. HEK293 cells expressing Ca_V2.2/TN-XL or TN-XL-Ras, were washed and collected with PBS, centrifuged and resuspended in lysis buffer (20 mM Tris-HCl [pH 7.5], 300 mM sucrose, 1 mM DTT and protease inhibitor cocktail (Roche)). Lysis was completed with freeze-thaw cycles and sonication. Membrane proteins were separated by initial centrifugation (16,000g ×10 min at 4°C), followed by solubilizing the resulting pellet in 1% Triton X-100. This solution was then subjected to centrifugation (16,000g ×10 min at 4 °C), yielding membrane proteins in the supernatant. Membrane proteins were denatured (95 °C×1 min) on dilution with TGS buffer (Bio-Rad), and resolved by 5% SDS-PAGE with no CaCl2 added to the gel. Proteins were transferred to nitrocellulose membranes (Bio-Rad) by cooled transfer (Tris/Glycine transfer buffer/pH 8.3×19h). After blocking, membranes were sequentially incubated with rabbit polyclonal anti-GFP antibody (Abcam, 1:5,000 dilution, as directed by manufacturer) and HRP-conjugated secondary antibody (Sigma, 1:2400 dilution as recommended by manufacturer). Protein bands were visualized with enhanced chemiluminescence (Pierce ECL, Thermo Scientific).

Data analysis and modelling. Single-channel ramp data were analysed for $Ca_V 2.2/TN-XL$. unitary current and open probability using custom MATLAB programs¹⁰. For the analysis, all patches contained 1 or 2 active channels. TIRF fluorescent signals from $Ca_V 2.2/TN-XL$ were modelled using custom MATLAB scripts, using approaches detailed in Supplementary Methods

Generalization of the Neher-Stern equation (equation (1)), for f values other than unity, was accomplished by considering mass balance and substituting Q_1/f for Q_1 in equation 6 of the original Stern paper¹³.

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Author contributions

L.H.T. established the TIRF/patch-clamp apparatus, designed and performed research, analysed data, and co-wrote the paper. I.E.D. performed extensive electrophysiological experiments, FRET imaging, and data analysis. W.Y. performed western blots and molecular biology. M.M. and O.G. guided use of TN-XL. D.T.Y. conceived experiments and analyses, established and performed numerical simulations, analysed data, made figures, and co-wrote the paper.

Additional information

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Supplementary Information Nanodomain Ca^{2+} of Ca^{2+} channels detected by a tethered genetically encoded Ca^{2+} sensor

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Supplementary Figures

Supplementary Figure S1. Layout of radially symmetric Ca²⁺ diffusion mechanism. (a) Overall geometric features. (b) Finite-element approximation features. Rate constants k_2 , k_3 , ... k_{Nshell} refer to rightward and leftward movements between compartments.



Supplementary Figure S2. CFP-channel asymptotes as a function of Ca_{spike} . Simulations demonstrating that CFP channel asymptotes approach Ca²⁺-saturated value of ~0.5 for Ca_{spike} greater than or equal ~35 μ M (green-shaded region).



Supplementary Figure S3. Lower-limit estimate of Ca_{spike} by forward transform analysis of uncorrected average data from main text Fig. 5b. Format identical to main text Fig. 6, except for fit-versus-data residuals shown below S_C and $R_{F/C}$ traces in panel **a**. <residual_{norm}> metrics show mean residual over trace, normalized to maximum signal excursion of 0.1. (**a**) Average Ca_v2.2/TN-XL responses to nanodomain Ca²⁺ signals, uncorrected. Top, black trace, average whole-current from main text Fig. 5b, rescaled to open probability P_O by normalizing peak current during 30-mV pulse to 0.69 (see main text Fig. 3e, red arrows). Gray trace, exponential fit to P_O waveform. Middle, cyan trace, average S_C , uncorrected CFP signal. Red trace, forward transform prediction of experimental S_C , using $Ca_{spike} = 13 \ \mu$ M. Δ , Fit-versus-data residual below. Bottom, Green trace, average $R_{F/C}$, uncorrected FRET-ratio signal. Red trace, forward transform prediction of experimental $R_{F/C}$ with $Ca_{spike} = 13 \ \mu$ M. Δ , Fit-versus-data residual below. (**b**) Nanodomain Ca²⁺ signals input to forward transform for prediction of Ca_v2.2/TN-XL responses in **a**. (**c**) Ca_{spike} estimation criteria. Presumed Ca_{spike} values plotted versus fold-fitting error (sum of squared deviations between fit and data, normalized to sum with $Ca_{spike} = 13 \ \mu$ M. Cusp of plot denotes $Ca_{spike} = 13 \ \mu$ M as best fit.

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Supplementary Figure S4. Optical layout of TIRF/patch-clamp apparatus, for TN-XL imaging mode. See main text methods for precise optical element specification.

Supplementary Methods

Spatial Ca²⁺ gradients by radial Ca²⁺ diffusion

This section details enhanced methods regarding Figures 4a,b and Supplementary Figure S1. Supplementary Figure S1a illustrates a radially symmetric Ca²⁺ diffusion scheme used to estimate the spatial distribution of Ca^{2+} in Figures 4a and 4b. A known Ca^{2+} influx (Fig. 4a) injects Ca^{2+} at the periphery of the sphere with radius R_{cell} (dm), and the whole-cell pipet allows the gradual egress of Ca^{2+} from the cell center. Diffusion is characterized by the effective diffusion coefficient D_{Ca} (dm² ms⁻¹) appropriate for live-cell cytoplasm. This system is solved by finite-element approximation (Supplementary Figure. S1b), where the sphere is divided into multiple shells (N_{shell} in number, and indexed by $k = 1, 2, ..., N_{shell}$), each with thickness $\Delta R =$ $R_{\text{cell}} / N_{\text{shell}}$. The [Ca²⁺] within each cell (C_k (M), for the kth shell) is considered uniform. The surface area (inner boundary) and volume of the *k*th shell is then given by $A_k = 4\pi \cdot \Delta R^2 \cdot (k-1)^2$ (dm²), and $V_k = (4/3) \cdot \pi \cdot \Delta R^3 \cdot (k)^3 - (4/3) \cdot \pi \cdot \Delta R^3 \cdot (k-1)^3$ (dm³ = L). From the definition of a diffusion coefficient, the rate constant governing transitions between adjacent shell compartments (with indices k and k-1) is therefore given by $k_k = D_{Ca} \cdot A_k / \Delta R$ (moles M⁻¹ ms⁻¹, for $k \ge 2$). The rate of Ca²⁺ influx into shell N_{shell} is C_{input} (moles/s). The rate constant for Ca²⁺ egress into the whole-cell pipet is given by rate constant k_{exit} (moles M⁻¹ ms⁻¹). Accordingly, Ca²⁺ diffusion in this system is given by the following state-variable equation.

$$\frac{d}{dt}\begin{bmatrix} C_{1}(t) \\ C_{2}(t) \\ \vdots \\ C_{3}(t) \\ \vdots \\ C_{N_{shell}}(t) \end{bmatrix} = \begin{bmatrix} -(k_{exit} + k_{2})/V_{1} & k_{2}/V_{1} & 0 & 0 & 0 & 0 & 0 \\ k_{2}/V_{2} & -(k_{2} + k_{3})/V_{2} & k_{3}/V_{2} & 0 & 0 & 0 & 0 \\ 0 & k_{3}/V_{3} & -(k_{3} + k_{4})/V_{3} & k_{4}/V_{3} & 0 & \cdots & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & 0 & 0 & 0 & 0 & \cdots & k_{N_{shell}}/V_{N_{shell}} & -k_{N_{shell}}/V_{N_{shell}} \end{bmatrix} \begin{bmatrix} C_{1}(t) \\ C_{2}(t) \\ C_{3}(t) \\ \vdots \\ C_{N_{shell}}(t) \end{bmatrix}$$

+
$$\begin{bmatrix} 0 & 0 & \cdots & 0 & 1/V_{N_{shell}} \end{bmatrix}^T \begin{bmatrix} C_{input}(t) \end{bmatrix}$$
 (S1)

This equation could be numerically integrated by custom scripts in MATLAB (Natick, MA). Typically, $N_{\text{shell}} = 90$, and doubling N_{shell} to 180 made no discernible difference, substantiating a sufficiently fine grain in the finite element analysis.

To predict the aggregate $[Ca^{2+}]$ reported by collecting the ratio of Fluo 4FF and Alexa 568 fluorescence signals from an entire cell, we considered that the observed ratio would be given by the volume weighted average below:

$$R_{observed} = \frac{F_{Fluo/\min} + (F_{Fluo/\max} - F_{Fluo/\min}) \cdot \sum_{k=1}^{N_{shell}} \frac{V_k}{V_{cell}} \cdot \frac{C_k}{C_k + K_{d/Fluo}}}{F_{Alexa}}$$

$$= R_{Fluo/\min} + (R_{Fluo/\max} - R_{Fluo/\min}) \cdot \sum_{k=1}^{N_{shell}} \frac{V_k}{V_{cell}} \cdot \frac{C_k}{C_k + K_{d/Fluo}}$$
(S2)

where V_{cell} is the entire cell volume, and $K_{\text{d/Fluo}} = 20 \ \mu\text{M}$ in our cells (see main text Methods). We render this into the normalized ratio

$$R_{normalized/observed} = \frac{R_{observed} - R_{Fluo/min}}{R_{Fluo/max} - R_{Fluo/min}} = \sum_{k=1}^{N_{shell}} \frac{V_k}{V_{cell}} \cdot \frac{C_k}{C_k + K_{d/Fluo}}$$
(S3)

This normalized ratio is then used to calculate aggregate $[Ca^{2+}]$ ($C_{aggregate}$) via the equation

$$R_{normalized / observed} = \frac{C_{aggregate}}{C_{aggregate} + K_{d / Fluo}} \implies C_{aggregate} = K_{d / Fluo} \cdot \frac{R_{normalized / observed}}{1 - R_{normalized / observed}}$$
(S4)

Substituting $R_{\text{normalized/observed}}$ from Equation S3 into S4 yields our prediction of $C_{\text{aggregate}}$ based on the spatial gradient of $[\text{Ca}^{2+}]$

$$C_{aggregate} = K_{d/Fluo} \cdot \frac{\sum_{k=1}^{N} V_k \cdot \frac{C_k}{C_k + K_{d/Fluo}}}{V_{cell} - \sum_{k=1}^{N_{shell}} V_k \cdot \frac{C_k}{C_k + K_{d/Fluo}}}$$
(S5)

Importantly, the nonlinear Langmuir form of the C_k terms ($C_k / (C_k + K_{d/Fluo})$) ensures that $C_{aggregate}$ is not trivially equal to spatial average [Ca²⁺]; hence, $C_{aggregate}$ discriminates between various spatial gradients of [Ca²⁺]. Accordingly, fitting the predicted $C_{aggregate}$ (Equation S5, Fig. 4b, lower red trace)) to the experimental $C_{aggregate}$ (Fig. 4b, noisy dark trace), given a measured Ca²⁺ influx (Fig. 4a), yields a well constrained estimate of the three free parameters: R_{cell} , D_{Ca} , and k_{exit} in the radial diffusion mechanism. Using simplex nonlinear minimization of the sum of squared errors between predicted and measured $C_{aggregate}$ (Fig. 4b) yields the values $R_{cell} = 2.5 \times 10^{-4}$ dm, $k_{exit} = 2 \times 10^{-4}$ moles· M⁻¹ · ms⁻¹, and $D_{Ca} = 2.4 \times 10^{-12}$ dm² · ms⁻¹. Reassuringly, D_{Ca} is closely similar to that measured for native nerve and muscle^{61,62}. The estimate for the submembranous [Ca²⁺] in Figure 4b (upper red trace) plots C_{Nshell} for these parameters.

Ca_V2.2/TN-XL forward transform calibration

This section details enhanced methodology for Figures 4c-f. To determine $K_{d/TN-XL}$ for Ca_V2.2/TN-XL in the TIRF volume, we considered that the steady-state relation between the normalized FRET ratio output of TN-XL (R_{norm} in Fig. 4d) and [Ca²⁺] is well described⁴⁴ by the

Hill function $R_{\text{norm}} = 1 / (1 + (K_{d/\text{TN-XL}} / [\text{Ca}^{2+}])^{1.7})$. The steady FRET ratio $(R_{\text{F/C}})$ signal of 1.8 (Fig. 4c, dashed gray line) corresponds to an R_{norm} of 0.7, as given by $(1.8 - R_{\text{min}}) / (R_{\text{max}} - R_{\text{min}}) = (1.8 - 1.1) / (2.1 - 1.1)$ and Table 1. This steady R_{norm} value is achieved at a [Ca²⁺] of 26.9 μ M (Fig. 4b, upper red trace). The Hill function then yields that $K_{d/\text{TN-XL}} = 17.3 \ \mu\text{M} = 26.9 \ \mu\text{M} \cdot ((1 - 0.68) / 0.68)^{1/1.7}$ for Ca_V2.2/TN-XL in the TIRF volume.

The probabilities of occupying the various states of the Ca_v2.2/TN-XL scheme in Figure 4e (define these probabilities as UB_0 , B_1 , B_2 , and B_3) can be deduced in the usual fashion for ionchannel kinetics⁶³, with the rate constant for UB₀ to B₁ transitions given by the potentially timevarying rate constant k_{01} . $Ca_{sensor}(t)$, and the rate constant for B₂ to B₃ transitions by a corresponding k_{23} . $Ca_{sensor}(t)^2$ term. $Ca_{sensor}(t)$ would be the Ca²⁺ concentration in the channel nanodomain, at the sensor binding site. To convert state occupancy into fluorescence output, let C_0 and C_2 be the CFP fluorescence detected in our system for a single Ca_v2.2/TN-XL molecule in states UB₀ and B₃, respectively. Let C_1 be corresponding CFP fluorescence for states B₁ and B₂. Let F_0 , F_1 , and F_2 be similarly defined for fluorescence detected through the FRET channel. The corresponding fluorescence signals for N_{sensor} number of independent and identical sensors would be given by multiplying the above terms by N_{sensor} . To render measurements ratiometric, we will utilize various ratios of these terms, defined as follows.

$$R_{\min} = \frac{F_0}{C_0}, \ R_{\min} = \frac{F_1}{C_1} = \frac{F_2}{C_2}, \ R_{\max} = \frac{F_3}{C_3}$$
 (S6)

and

$$\alpha_0 = \frac{C_0}{C_3}, \ \alpha_1 = \frac{C_1}{C_3} = \frac{C_2}{C_3}$$
 (S7)

Accordingly, the FRET channel fluorescence of a single Cav2.2/TN-XL sensor would be

$$F(t) = F_0 \cdot UB_0(t) + F_1 \cdot (B_1(t) + B_2(t)) + F_3 \cdot B_3(t)$$
(S8)

The CFP channel fluorescence of a single Cav2.2/TN-XL sensor would be

$$C(t) = C_0 \cdot UB_0(t) + C_1 \cdot (B_1(t) + B_2(t)) + C_3 \cdot B_3(t)$$
(S9)

Finally, the ratio of FRET and CFP signals, for both a single or ensemble of $Ca_V 2.2/TN-XL$ sensors would be

$$R_{\rm F/C}(t) = \frac{F(t) = F_0 \cdot UB_0(t) + F_1 \cdot (B_1(t) + B_2(t)) + F_3 \cdot B_3(t)}{C_0 \cdot UB_0(t) + C_1 \cdot (B_1(t) + B_2(t)) + C_3 \cdot B_3(t)}$$

$$= R_{\rm min} \cdot \alpha_0 \cdot \frac{UB_0(t)}{D(t)} + R_{\rm mid} \cdot \alpha_1 \cdot \frac{B_1(t) + B_2(t)}{D(t)} + R_{\rm max} \cdot 1 \cdot \frac{B_3(t)}{D(t)}$$
(S10)

where $D(t) = \alpha_0 \cdot UB_0(t) + \alpha_1 \cdot (B_1(t) + B_2(t)) + 1 \cdot B_3(t)$.

To simulate the constraints in Figs. 4c and 4d, R_{mid} , R_{max} , α_0 , and α_1 were fixed by experiments as detailed in Table 1. R_{mid} and rate constants k_{01} , k_{10} , k_{12} , k_{21} , k_{23} , and k_{32} were then varied by numerical simplex optimization to minimize the sum of squared deviations between data (Figs. 4c and 4d, green) and predictions (red). The optimal parameter set is summarized in Table 1. For the simulation in Figure 4c, $Ca_{sensor}(t)$ was set equal to the submembranous [Ca²⁺] deduced in Figure 4b (upper red trace), and the differential equations corresponding to the forward transform in Figure 4e integrated numerically by custom-written scripts in MATLAB (Natick, MA). For the simulation in Figure 4d, Ca_{sensor} was set at a number of fixed values, and the same differential equations numerically integrated until steady state was reached.

Treating silent/intracellular Cav2.2/TN-XL

This section details in-depth methods relating to Figure 6a, and to Supplementary Figure S2. Here, we assess how the asymptote of $S_{\rm C}$ signals (as in Fig. 5b, middle, dashed curve) would change with $Ca_{\rm spike}$ (amplitude of Ca²⁺ pulses in nanodomain, defined in Fig. 6b), under the assumption that all channels are active (Fig. 5c). To do so, we utilized the forward transform in Figures 4e and 4f to predict Ca_V2.2/TN-XL sensor outputs in the presence of high levels of Ca²⁺ buffering to isolate nanodomains. Under these conditions, these predictions can be accomplished by setting the rate constant for transition from UB₀ to B₁ as $k_{01} \cdot Ca_{\rm spike} \cdot P_0(t)$, and from B₂ to B₃ as $k_{23} \cdot Ca_{\rm spike}^2 \cdot P_0(t)$, as described in the main text. These approximations hold so long as either Ca²⁺ binding or unbinding steps to TN-XL are slow relative to the millisecond kinetics of Ca²⁺ pulses within the nanodomain (true in Table 1), according to the 'slow CaM approximation' detailed in section 2A of the Supplemental Data of our prior work¹⁰. In conjunction with parameters in Table 1, the differential equations corresponding to the forward transform in Figure 4e can then be integrated numerically as described in the section immediately above, using the experimentally determined $P_O(t)$ waveform (Fig. 6b, top). We simulated S_C trajectories (as in Fig. 5a, middle) by calculating $C(t) / C_O(t)$ from Equation S9, and performing exponential extrapolation of asymptotic values for different values of Ca_{spike} . Supplementary Figure S2 plots these asymptotes as a function of Ca_{spike} , demonstrating that asymptotic values approximate the Ca^{2+} -saturated level of $0.5 (= 1/\alpha_O)$, with α_O determined experimentally in Table 1) for all Ca_{spike} $\geq 35 \ \mu$ M. Thus, estimation of S_C asymptotes via exponential fitting was performed within individual cells, and static background fluorescence determined through knowledge that active sensors would asymptote at 0.5. Background-fluorescence and minor-bleach corrections were then undertaken on CFP and FRET channel signals, yielding corrected \overline{S}_C , \overline{S}_F , and thereby $\overline{R}_{F/C}$ signals. These were averaged across cells, yielding the signals shown in Figure 6a.

Predicting Ca_v2.2/TN-XL responses to nanodomain Ca²⁺

This section details enhanced methods for Figures 6a,b. Predictions of Ca_v2.2/TN-XL sensor outputs (Fig. 6a, in the presence of high levels of Ca²⁺ buffering to isolate nanodomains) can be accomplished as described above in the section immediately above. The one new feature is that $\overline{R}_{F/C}$ predictions were performed via Equation S10, using parameters as stated in Table 1.

Ca_v2.2/TN-XL responses from entire TIRF volume

This section describes on the lower-limit fit relating to Figure 6a, and expands on Supplementary Figure S3. To furnish a conservative lower-limit estimate of Ca_{spike} , we applied forward transform analysis to wholly uncorrected data from Figure 5b. The analysis was performed as described for Figure 6, with results shown in the same format (Supplementary Figure S3). The best fit of the forward transform was obtained with $Ca_{spike} = 13 \mu M$, yielding the red trajectories in Supplementary Figure S3a, and the nadir of the error analysis in Supplementary Figure S3c. We note that the forward transform to uncorrected data furnishes a comparatively poor fit, as follows. The residuals between forward transform fits and data are displayed underneath the S_C and R_{F/C} traces in Supplementary Figure S3a. From these, visual inspection reveals systematic upward deviations, even for the best-fitting Ca_{spike} value of 13 μ M. These deviations are also evident from mean of residuals, averaged across traces, and normalized by maximum waveform excursions of 0.1. These <residual_{norm}> values are 0.104 for the S_C signal, and 0.06 for the R_{F/C}

signal. For forward transform analysis of corrected data in Figure 6, the comparable metrics were 200- and 3-fold smaller, respectively. This contrast in the quality of forward transform fits further suggests that correction of static background fluorescence, as in Figure 6a, is warranted.

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