Ultrafast optical recording reveals distinct capsaicin-induced ion dynamics along single nociceptive neurite terminals in vitro

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1 Introduction

The detection of painful stimuli takes place at the nociceptive peripheral sensory nerve terminals in the skin or the viscera. These miniature structures (0.5 to 5 μm in diameter, Fig. 1) are pivotal for the detection of noxious stimuli. Consequently, the comprehensive understanding of nociception and hence mechanisms of pathological pain can be achieved only by detailed exploration of nociceptive terminal function. However, not much is known about their properties because they are barely accessible by conventional experimental approaches. Here, we report the implementation of an ultrafast optical recording approach for studying cultured neurite terminals, which are readily accessible for assay manipulations. Using this approach, we were able to study capsaicin-induced calcium and sodium dynamics in the nociceptive processes, at a near-action potential time resolution. The approach was sensitive enough to detect differences in latency, time-to-peak, and amplitude of capsaicin-induced ion transients along the terminal neurites. Using this approach, we found that capsaicin evokes distinctive calcium signals along the neurite. At the terminal, the signal was insensitive to voltage-gated sodium channel blockers, and showed slower kinetics and smaller signal amplitudes, compared with signals that were measured further up the neurite. These latter signals were mainly abolished by sodium channel blockers. We propose this ultrafast optical recording approach as a model for studying peripheral terminal signaling, forming a basis for studying pain mechanisms in normal and pathological states. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.22.7.076010]

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Abstract. Pain signals are detected by terminals of nociceptive peripheral fibers situated among the keratinocytes and epithelial cells. Despite being key structures for pain-related stimuli detection and transmission, little is known about the functional organization of terminals. This is mainly due to their minute size, rendering them largely inaccessible by conventional experimental approaches. Here, we report the implementation of an ultrafast optical recording approach for studying cultured neurite terminals, which are readily accessible for assay manipulations. Using this approach, we were able to study capsaicin-induced calcium and sodium dynamics in the nociceptive processes, at a near-action potential time resolution. The approach was sensitive enough to detect differences in latency, time-to-peak, and amplitude of capsaicin-induced ion transients along the terminal neurites. Consequently, the basic questions in nociceptive physiology, such as where action potentials are generated and how they propagate along nonmyelinated tiny axons in normal and pathological conditions, still remain obscure.

Here, we introduce a new approach for high-temporal resolution optical recordings of ion dynamics from a single nociceptive neurite terminal in vitro. This approach allows us to monitor capsaicin-induced calcium (Ca²⁺) and sodium (Na⁺) dynamics at the terminal neurites and characterize their propagation along the distal neurites. Using this approach, we showed that capsaicin-induced calcium signals propagated in the terminals were not dependent on voltage-gated Na⁺ channels, whereas signals measured 15 μm away and onward, toward the cell body, were faster, stronger, and at least partially mediated by voltage-gated Na⁺ channels.
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Fig. 1 GCaMP6s-based Ca$^{2+}$ imaging of nociceptive-cultured DRG neurite terminals and distal processes. (a) Scheme depicting the experimental protocol (see Sec. 2). The cell bodies of DRG neurons were seeded in the middle compartment of the chamber and neurites were induced to grow to the lateral compartment. Upper inset, schematic representation of the neurite terminals and stimulating pipette. Lower inset, bright field image of a representative neurite terminal and puff pipette location. (b) Puff calibration experiment. Dispersion of a fluorophore (sulfarhodamine 101, 2 μM) following a 500-ms puff (see Sec. 2) is presented in relation to the representative neurite terminal [rendered from (a), lower inset], as a reduction profile of the fluorescence ($F - F_0$, indicated by arbitrary units—au, and color coded, left) with distance from the puff pipette located at coordinates of $x$, $y = 0$ (see Sec. 2). Coordinates of the neurite tip are indicated in parenthesis. Note that on the $x$-axis, the fluorescent intensity was reduced by about 50% when reaching a distance of 10 μm from the tip of the puff pipette. On the $y$-axis, intensity diminished almost completely after 15 μm. (c) Left, fluorescent image of the neurite’s terminal process expressing GCaMP6s. Arrows indicate locations of ROIs from which the recordings shown to the right were performed. Right, representative traces of changes in [Ca$^{2+}$]$_i$ following focal puff application of vehicle and 300-nM capsicain (far right) measured every 500 ms. (1) Vehicle, (2) 300 nM capsicain. Lower inset, bright field image of a representative neurite terminal and stimulating pipette. Upper inset, schematic representation of the neurite terminals and stimulating pipette. (Sigma) in DMEM (Biological Industries) was spread to cover the scratched region where the Teflon compartmentalized chamber is placed. Teflon compartmentalized chambers (Teflon dividers, Camp10 or Camp320, Tyler Research) were attached to the culture dish with silicon grease (Dow Corning), applied with a syringe grease applicator (CAMP-GLSS, Tyler Research). The central compartment was used for plating the cell soma, flanked by peripheral compartments, containing lanes in which neurites can extend [Fig. 1(a)].

2 Materials and Methods

2.1 Animals

All animal procedures were approved by the Ethics Committee of the Hebrew University. Five- to six-week-old male C57BL/6 mice were used.

2.2 Dorsal Root Ganglion Compartmental Cultures

The compartmented chamber dishes were prepared as described by Campenot et al. Briefly, poly-lysine- and laminin-coated dishes were scraped with a pin-rake (200-μm width of pins; Tyler Research) to create a series of parallel tracks on the laminin coating, limiting neurite growth to the areas left with laminin, in parallel lanes. A droplet of 0.4% methylcellulose (Sigma) in DMEM (Biological Industries) was spread to cover the scratched region where the Teflon compartmentalized chamber is placed. Teflon compartmentalized chambers (Teflon dividers, Camp10 or Camp320, Tyler Research) were attached to the culture dish with silicon grease (Dow Corning), applied with a syringe grease applicator (CAMP-GLSS, Tyler Research). The central compartment was used for plating the cell soma, flanked by peripheral compartments, containing lanes in which neurites can extend [Fig. 1(a)].

DRG culture preparation: Adult mice were deeply anesthetized (3% isoflurane), and DRG were isolated and dissociated in a manner similar to that described previously, with the following modifications: dissected ganglia were placed in ice-cold DMEM (Biological Industries), pelleted, and reconstituted for enzymatic digestion in solution containing 5 mg/ml collagenase and 1 mg/ml dispase II (Roche) for 45 min. Cells were triturated in the presence of 50 U DNase 1 (Sigma) and centrifuged through 10% bovine serum albumin (Sigma). The cell pellet was resuspended in 1 ml of Neurobasal (Gibco) medium containing B27 supplement (Invitrogen), penicillin and streptomycin (Biological Industries), 100 ng/ml 2.5 SS NGF (Promega), and 2 ng/ml GDNF (Sigma). Cells were plated on to poly-lysin- (500 μg/ml) and laminin- (5 mg/ml) coated 35-mm glass bottom tissue culture dishes [World Precision Instruments (WPI)] in the middle compartment of the compartmented chambers [Fig. 1(a), see also Ref. 20].

On the day of seeding the cells (day 0), both chambers contained the above-mentioned Neurobasal media with 100 ng/ml 2.5 SS NGF; thereafter, the concentrations of NGF were altered to promote the spouting neurites to grow toward the distal chambers as follows (cell bodies and peripheral compartments, respectively): day 1: 10 and 100 ng/ml; day 2: 1 and 10 ng/ml; and days 5, 8, and day 11: 0 and 1 ng/ml.

Cultures were maintained at 37°C in a humidified incubator containing 5% CO$_2$. The cultured DRG cell somata underwent viral coinfection on either of 8- to 10-days postseeding. Imaging experiments of neurite terminals were carried out on days 11 to 14 from seeding. Teflon dividers were removed prior to the experiment.

2.3 Viral Vectors

To measure the changes in intraterminal and intraneuritic Ca$^{2+}$ concentrations ([Ca$^{2+}$]$_i$), the adenov-associated virus serotypes 1 or 6, carrying an expression cassette for the genetically encoded Ca$^{2+}$ indicator (GCaMP6s) or the mRuby fluorescent protein indicator (under a CMV promoter; ELSC Virus Core Facility), were used. Cells were infected by the two viruses simultaneously.

2.4 Imaging

Image data were processed by Neuroplex software (RedShirt Imaging) and analyzed using pCLAMP 10.2 (Molecular Devices) and OriginPro 9.0.0 (OriginLab Corporation).

Number of repeats in this study ($n$’s) refers to the number of terminals examined in each experiment. In the Ca$^{2+}$ imaging assay, each analyzed terminal was from a culture made from a different animal. For the Na$^+$ imaging assay, $n$ reflects different terminals in different dishes, however, not all from different animals.

All analyses were performed on the processes in the same field of view.
We used a short imaging protocol of up to 5 s to avoid photo damage, which occurred during longer exposure, probably due to the recurrent excitation with high-intensity ultraviolet and blue excitation wavelengths used here.

### 2.4.1 Ca$^{2+}$ imaging

Fluorescent excitation was performed with a 75 W Xenon arc lamp (Lambda DG4; Shutter Instruments) and a GFP filter set (Ex 480, Em 535, dicroic Lp 510; Chroma). An inverted microscope (Eclipse Ti; Nikon) equipped with a x60 oil 1.4NA objective, Epi-Fl attachment, and perfect focus system was used. For bright field imaging and fluorescent still image acquisition, an Exi Aqua monochromator camera (QImaging) and Nikon elements AR software (Nikon) were used. Changes in intraterminal and intraneuritic fluorescence were acquired using a back-illuminated 80 × 80 pixel cooled CCD camera (NeuroCCD-SMQ; RedShirt Imaging), controlled by Neuroplex software (RedShirtImaging). In these conditions, pixel size was 1 × 1 μm. This spatial resolution was sufficient for analyzing the signal propagation along terminals and distal neurites, even though in some locations the diameter of the structure (0.5 to 5 μm) could have been less than the camera resolution.

Images were acquired at 1000 frames/second (fps).

### 2.4.2 Na$^+$ imaging

To measure changes in intraterminal and intraneuritic Na$^+$ concentrations ([Na$^+$]$_{in}$) in vitro, the cell-permeable Na$^+$ indicator SBFI-AM (Molecular Probes) was used. Five-millimolar stock was prepared in dimethyl sulfoxide with 10% Pluronic F-127. Cells were incubated for 1 h with a 5 μM final concentration of SBFI-AM in the culture media, followed by a 30-min rinse period. Images were acquired at 125 fps. Fluorescent excitation was performed using a CoolLed fluorescence excitation system, using a 380-nm LED and a Fura2 filter set (Ex: 380 nm, Em: 510 nm, dicroic Lp: 400 nm; Chroma). To compensate for time-dependent decrease in fluorescence, due to fluorophore bleaching, fitted linear curves derived from the first 500 ms of the acquired data were subtracted from the recorded data.

### 2.5 Solutions

All imaging experiments were performed in standard external solution (SES) composed of 145 mM NaCl, 5 mM KCl, 2 mM CaCl$_2$, 1 mM MgCl$_2$, 10 mM glucose, and 10 mM HEPES (pH = 7.4).

Capsaicin (300 nM) or vehicle (SES with 0.3 μl/ml ethanol) was focally puff applied (500 ms puff, 2 psi, PV820 Pneumatic PicoPump; WPI) by a pipette with 4 to 6 MΩ resistance, when filled with SES. The application regime was commanded by a Digidata 1440 A/D interface (Molecular Devices).

In some experiments, the Na$^+$ channel blocker, tetrodotoxin (TTX, 10 μM; Alomone Labs) together with a Na$^+$ 1.8 channel blocker, A803467 (10 μM; Tocris), were applied to bath for 10 min before the recordings.

### 2.6 Puff Calibration

We calibrated the dispersion profile of puffed substances by measuring changes in the sulfarhodamine (2 μM) fluorescent intensity profile with distance from the pipette tip [Fig. 1(b)] according to

\[
\frac{F_{x_1,y_1}}{F_{x_0,y_0}} \times C_{x_0,y_0} = C_{x_1,y_1},
\]

where $F$ is the fluorescence intensity at $x$, $y$ coordinates and $C$ is the concentration.

### 2.7 Data Analysis

To measure changes in [Ca$^{2+}$]$_{in}$ within a terminal at the highest possible resolution our conditions permit, the data were collected from 1 × 1 μm sized regions of interest (ROIs) [1 pixel; Fig. 1(c)]. To examine changes in Ca$^{2+}$ concentration propagating along the neurite and to improve the signal-to-noise ratio, the data were collected using Kernel smoother from 2 × 2 μm sized ROIs (2 × 2 pixels; Fig. 2), which were distributed every 5 μm along the neurite’s terminal and processes, dividing the terminals into $m$ segments. All the data were analyzed from the ROIs delimiting the examined segments, and the size of the ROI was neglected when the actual distance was calculated. Therefore, the distance is represented as an estimated location rather than an absolute distance.

The time of onset of the signal was measured as the time at the onset of a clear sharp deviation from fluorescent base level measured locally, while analyzing the trace from its positive peak toward time “0.” All the values were subtracted from the value at the terminal tip ($x = 0$) of the corresponding terminal.

Difference in time of onset between two adjacent segments, i.e., the delay in onset at one segment compared with the time of onset at the previous segment, were calculated according to

\[
t_m - t_{m-1},
\]

where $m$ is the $m$’th segment.

The fluorescence propagation velocity within 5 μm segments was calculated according to

\[
v = \frac{5}{t_m - t_{m-1}},
\]

where $v$ is the fluorescence propagation velocity and 5 is the distance between each ROI in micrometers (μm).

Maximum rate of rise of Ca$^{2+}$ transients was calculated as max(DF/DT) at different locations along the neurite terminal.

The values were normalized to the value of the terminal tip of the corresponding terminal and plotted as a function of distance from the terminal tip.

All averaged data are presented as the mean ± SEM. Assessment of statistical significance of differences between means was performed with repeated-measures of analysis of variance (ANOVA), with posthoc Bonferroni, as found appropriate.

### 3 Results and Discussion

To study processing of nociceptive information directly at its most relevant location, with sufficient time and space resolution, we introduced an approach for ultrafast optical recording of cultured nociceptive processes. To that end, we used compartmented chambers [Fig. 1(a), see Sec. 2] that have been previously used to study neurites of DRG neurons. In those chambers, DRG cell bodies are prompt to grow neurites, and terminal processes can be easily identified and accessed [Figs. 1(a) and 1(c)].

We selectively stimulated these neurite terminals with a calibrated focal puff application [Fig. 1(b)] of capsaicin, an agonist of the nocuous heat-sensitive transient receptor potential cation
Fig. 2 Propagation of capsaicin-induced Ca\textsuperscript{2+} transients along the neurite terminal. (a) Fluorescence image of representative nociceptive terminals expressing GCaMP6s with illustration of pipette location and with the arrows indicating the locations of ROIs from which the measurements shown in (b) and (c) were performed. (b, c) Representative traces of capsaicin-induced changes in [Ca\textsuperscript{2+}]\textsubscript{i} measured every 5 \(\mu\text{m}\) at distances shown in left, from the terminal tip before (b) and 10 min after (c) application of 10-\(\mu\text{M}\) TTX together with 10-\(\mu\text{M}\) A803467. Changes in fluorescent were measured at 1000 fps. Note the substantial increase in signal amplitude after 15 \(\mu\text{m}\), which was abolished by applying TTX and A803467. Insets: Expanded timescale of the traces presented in (b) and (c) emphasizing the signal onset. Arrows indicate calculated time of signal onset (see Sec. 2). Note that the timescale of the inset for (b) is further expanded to better illustrate the differences in time of signal onset along the neurite in the control conditions. (d) Time of onset of capsaicin-induced Ca\textsuperscript{2+} signals measured from terminal tip (point "0") in control conditions (SES, black squares) and after bath application of TTX together with A803467 (red circles), plotted versus distance from the terminal tip. Note much smaller changes in time of onset after 15 \(\mu\text{m}\) in control conditions. Application of TTX together with A803467 significantly increased the difference in time of onset after 15 \(\mu\text{m}\); ns, not significant; *\(p < 0.05\), **\(p < 0.01\), ***\(p < 0.001\); black letters and symbols, comparison between values measured at each location in control condition, \(n = 4\), one-way ANOVA with posthoc Bonferroni; orange letters and symbols, comparison between "SES" and "TTX + A803467," \(n = 2\); two-way ANOVA, with posthoc Bonferroni. (e) Peak amplitudes of capsaicin-induced changes in fluorescence, measured every 5 \(\mu\text{m}\) and plotted as a function of distance from terminal tip, in control conditions (SES, black) and after (red) application of TTX together with A803467. Note that in control conditions, signal amplitude significantly increased after 15 \(\mu\text{m}\). TTX together with A803467 did not affect the signal measured at the first 15 \(\mu\text{m}\), but significantly reduced the signal amplitude thereafter. Symbols and statistics as in (d). (f) Maximum rate of rise of Ca\textsuperscript{2+} transients, \(\max(dF/dt)\), at the different locations along the neurite terminal in control condition (SES, black), and TTX together with A803467 (red) plotted as a function of distance from the terminal tip. Note that in control conditions, the rise rate significantly increased at \(\sim 15\mu\text{m}\) and plateaued afterwards. Application of TTX together with A803467 prevented the increase in rise rate. Statistics as in (d).
channel subfamily V member 1 (TRPV1) channel, expressed by nociceptive neurons.25 We placed the pipette, containing 300 nM capsaicin, about 10 μm from the terminal tip [Figs. 1(a) and 1(b)] because at this distance, the application of vehicle did not induce changes in GCaMP6s fluorescence [Fig. 1(c); n = 20 terminals] or detectable movement of the processes. First, we calibrated the dispersion profile of the puffed substances [see Sec. 2; Fig. 1(b)]. Our data show that the predicted concentration at the terminal tip is ~50% of the initial value [Fig. 1(b)], i.e., in our conditions, the concentration of capsaicin at the terminal is close to the half maximal effective concentration (EC₅₀) of TRPV1 channels to capsaicin (~150 nM)⁵. This is an estimate of the peak concentration during a 500-ms puff, which dissipates in the medium when the puff ends. Additionally, considering that the minimal concentration of capsaicin to activate TRPV1 channels is ~30 nM,⁶ the prominent decrease in concentration along the y-axis [Fig. 1(b)] renders the concentration mostly ineffective once it reaches other parts of the fiber or adjacent terminals. Indeed, in all the performed recordings, we did not observe Ca²⁺ responses in neighboring terminal branches of the neurite.

We measured capsaicin-induced increase in [Ca²⁺]ᵢ at the neurite’s terminal process using the genetically encoded Ca²⁺ indicator GCaMP6slow [GCaMP6s, Fig. 1(c)]. We chose the slow variant of the GCaMP6 indicator due to its better signal-to-noise ratio and sensitivity (compared with other GCaMP6 variants²⁵), which enabled the detection of weak fluorescent emission signals from fine neurite processes. To study transducer channel- and action potential-mediated changes in terminal [Ca²⁺]ᵢ, we optically recorded changes in terminal GCaMP6s fluorescence intensity at a sampling rate of 1000 fps. As it has been shown that the half width of the action potential recorded from the distal neurite of nociceptive neurons is ~2.5 ms,⁸ this sampling rate should be sufficient to detect Ca²⁺ reflections of single action potentials. We achieved this using a 60x oil 1.4NA Plan Apo objective, in combination with a fast acquisition back-illuminated 80 × 80 pixel CCD camera (see Sec. 2). Puff application of capsaicin, but not vehicle [Fig. 1(c)], led to prominent increase in GCaMP6s fluorescence, detectable at a spatial resolution of 1 × 1 μm along the terminal tip [Fig. 1(c)].

Next, utilizing the high functional spatial resolution of our approach, which allows precise tracing of signal propagation along the terminal, we measured the amplitude, time of onset, and rise rate of capsaicin-induced Ca²⁺ signals along the terminal and terminal neurite [Figs. 2(b), 2(d)–2(f), black]. It is noteworthy that due to short acquisition time (see Sec. 2), we could not explore the mechanisms underlying the decay phase of the capsaicin-induced response at the neurite’s processes and, therefore, we focused our efforts on the rising phase of the response. We analyzed the difference in time of onset between two adjacent segments, consecutively for all adjacent segments (see Sec. 2), which reflects the time it took the signal to travel along a segment. We found that the signal traveled along m₁ for 58.5 ± 10 ms, m₂ for 62.2 ± 10 ms, and m₃ for 15.6 ± 5 ms (n = 4). After segment m₁, time of onset difference between two ROIs was 3.2 ± 2 ms [segments m₁ to m₃; Fig. 2(d), black, n = 4]. Using these data, we calculated the fluorescent propagation velocity of capsaicin-induced Ca²⁺ signals between segments (see Sec. 2) and demonstrated that the signal propagated along m₁ with a velocity of 97.2 ± 23 nm/ms, m₂ with 85.2 ± 11 nm/ms, and m₃ with 382 ± 95 nm/ms (n = 4). Signals propagated between segments m₁ to m₃ did not vary in their response peak amplitude or maximum rise rate [max(dF/dt); Figs. 2(b), 2(e), 2(f) black]. After ~15 μm (segment m₃), the fluorescence propagation velocity increased significantly and reached a peak of 870 ± 300 nm/ms at m₄ (~20 μm), and the speed of propagation between 5 μm segments remained similar along the rest of the measured terminal (p = 0.47, one-way ANOVA, n = 4). It is noteworthy that the values for propagation velocity that we have measured represent the propagation of fluorescent signals along the process and not the electrical conductance. The propagation of GCaMP6s fluorescence is dependent on electrical conductance; however, there are many additional factors that are involved, such as the Ca²⁺ channels³ and indicator kinetics (see also Ref. 28).

Remarkably, at 15 μm (segment m₁), a two-phase response appeared [Fig. 2(b)]. While the first phase was similar to the one observed at the terminal tip (p = 0.35 for rise rate; p = 0.58 for the amplitude, one-way ANOVA with posthoc Bonferroni), the second phase possessed a significantly larger peak amplitude and maximum rise rate [p < 0.05, one-way ANOVA, n = 6 terminals; Figs. 2(e), 2(f) black], reaching a plateau of about 3.5 times larger response peak amplitude and 7 times larger rise rate at the maximum. This large and fast component was completely abolished following application of a combination of TTX and A803467 [Figs. 2(c), 2(d)–2(f) red] to block TTX-s- and Na(v)1.8-mediated TTX-r Na⁺ channels,²⁹ which are expressed by nociceptive terminals.⁸ Moreover, blocking Na(v) channels delayed the time of onset [Fig. 2(d), red], reduced the fluorescence propagation velocity to 57 ± 11 nm/ms, and decreased the peak amplitude and maximal rise rate [Figs. 2(e) and 2(f), red] of capsaicin-evoked responses recorded along the neurite after ~15 μm, such that the values become similar to those recorded at the first 15 μm [Figs. 2(e) and 2(f)]. These data suggest that the increase in [Ca²⁺]ᵢ beyond the ~15 μm is partially dependent on activation of voltage-gated Na⁺ channels.

We further supported the latter notion by directly measuring capsaicin-induced Na⁺ fluxes in the neurite’s terminal process using the Na⁺ indicator SBFI-AM. Imaging of Na⁺ fluxes at different locations along the terminal revealed that application of capsaicin induced a small transient increase in [Na⁺]ᵢ in the neurite’s terminal tip and a larger step-like Na⁺ signals at a distance from the tip [Fig. 3(b), n = 3].

It is noteworthy to mention that TRPV1 channels are also permeable to Na⁺.³⁰ Thus, the signal at the terminal tip may reflect Na⁺ influx via active TRPV1 channels. The fast and step-like nature of the signals recorded away from the neurite may underlie burst-like firing of the action potentials.³¹ From a methodological point of view, this in vivo approach, which is based on a distinctive compartment containing only neurite terminals, is currently the only approach allowing direct study of terminal Na⁺ dynamics. It is advantageous over an in vitro assay as the usage of AM-based dyes in vivo will likely produce a strong background noise due to AM dye loading of not only the terminals but also nonneuronal cells in the imaged area. The lack of genetically encoded Na⁺ indicators for specific measurements of Na⁺ flux in distinct cell types render measurements from small structures in vivo or ex vitro extremely challenging. Importantly, low initial fluorescence intensity of SBFI-AM in the neurite will likely produce a strong background noise due to AM dye loading of not only the terminals but also nonneuronal cells in the imaged area. The lack of genetically encoded Na⁺ indicators for specific measurements of Na⁺ flux in distinct cell types render measurements from small structures in vivo or ex vitro extremely challenging.
Altogether our data suggest that the nociceptive neurite terminals and processes, in terms of \( \text{Ca}^{2+} \) dynamics and its dependence on \( \text{Na}^+ \) channel blockers, are composed of two functionally separated zones. Our results imply that signal generation and propagation in the first 15 \( \mu \text{m} \) is not mediated by voltage-gated \( \text{Na}^+ \) channel-induced depolarization. This \( \text{Ca}^{2+} \) signaling could be mediated by \( \text{Ca}^{2+} \)-permeable TRPV1 channels, low threshold voltage-gated \( \text{Ca}^{2+} \) channels, \( \text{Ca}^{2+} \)-induced \( \text{Ca}^{2+} \) release, and capsaicin-mediated \( \text{Ca}^{2+} \) release from TRPV1-expressing internal stores.\(^{32}\) Given the focality of our application and the susceptibility of the late response to \( \text{Na}^+ \) channel blockers, TRPV1-mediated \( \text{Ca}^{2+} \) release from internal stores has a lesser contribution to the capsaicin-induced \( \text{Ca}^{2+} \) response, which we observed after 15 \( \mu \text{m} \). Hence, as suggested by our data, signal propagation beyond the first 15 \( \mu \text{m} \) is largely driven by voltage-gated \( \text{Na}^+ \) channels leading to \( \text{Ca}^{2+} \) elevation possibly via voltage-gated \( \text{Ca}^{2+} \) channels and \( \text{Ca}^{2+} \)-induced \( \text{Ca}^{2+} \) release. Interestingly, the maximal propagation velocity, we have measured from terminal neurites of nociceptive processes, was substantially smaller than those measured using single fiber recordings from mice sciatic-tibial nerve \textit{in vitro}.\(^{3} \) This discrepancy could result from the differences between electrical conductance and fluorescent propagation, as stated above. Indeed, the velocities of fluorescence propagation that we and others\(^{3, 20, 28} \) have measured in cultured neurites are slower than that measured for propagation of electrical signals.\(^{6} \)

In this study, we have analyzed the activity of individual terminal neurites, which are easily identified and traced. It is difficult, however, to trace and record individual neurite activity along its full length because proximal neurites in culture create dense bundles of neurites.\(^{20, 24} \) Utilization of our approach on individually labeled neurons, via micropipette injection of a spectrally different dye, may overcome this difficulty and provide essential information about the calcium dynamics along different compartments of peripheral processes.

In conclusion, we have introduced an approach for studying nociceptive terminal processes by bringing together and utilizing fast and sensitive optical recordings of ion dynamics from easily accessible cultured neurite terminals. Altogether, the data shown here demonstrate that our methodology gains in sensitivity and temporal acquisition capabilities, compared with standard imaging techniques, allowing fast optical recordings of weak signals from minute nociceptive terminal processes. Moreover, our approach gains in functional spatial resolution, compared with single electrode electrophysiological approaches, permitting characterization of signal propagations along the terminal and terminal neurites by sampling the signals in action potential relevant time resolution. Thus, the ultrafast and high-resolution optical recording technique described here provides the only tool for a detailed study of a nociceptive terminal functional molecular network, which underlies noxious stimuli detection and transmission in normal and pathological conditions. This could advance the very much needed knowledge for understanding pain physiology and pathophysiology. Moreover, this platform can be utilized to enable studying the terminals of other neuronal types, such as other primary sensory neuron terminals.

**Disclosures**

No conflicts of interest, financial or otherwise, are declared by the authors.

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