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Abstract. Voltage-sensitive fluorescence indicators enable tracking neuronal electrical signals simultaneously in multiple neurons or neuronal subcompartments difficult to access with patch electrodes. However, efficient widefield epifluorescence detection of rapid voltage fluorescence transients necessitates that imaged cells and structures lie sufficiently far from other labeled structures to avoid contamination from out of focal plane and scattered light. We overcame this limitation by exciting dye fluorescence with one-photon computer-generated holography shapes contoured to axons or dendrites of interest, enabling widefield detection of voltage fluorescence with high spatial specificity. By shaping light onto neighboring axons and dendrites, we observed that dendritic back-propagating action potentials were broader and slowly rising compared with axonal action potentials, differences not measured in the same structures illuminated with a large “pseudowidefield” (pWF) spot of the same excitation density. Shaped illumination trials showed reduced baseline fluorescence, higher baseline noise, and fractional fluorescence transient amplitudes two times greater than trials acquired with pWF illumination of the same regions.

Keywords: computer-generated holography; voltage imaging; action potential; axon; dendrite.

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1 Introduction

Understanding neuronal input–output transformations requires experimental characterization of how electrical signals generate and propagate in axonal and dendritic arbors. Although whole cell patch clamp techniques have enabled high temporal resolution and high signal-to-noise ratio (S/N) electrical recordings in somata and large dendrites, smaller structures remain difficult to access with patch electrodes. Voltage-sensitive dyes (VSDs) provide an alternative method to track membrane potential and have been effectively imaged with one-photon epifluorescence microscopy to characterize action potential propagation in small diameter axons and dendrites. Due to the close spatial mingling of neuronal substructures, improving lateral and axial confinement with confocal microscopy could enable discrimination of signals arising from adjacent or overlapping structures; however, the relatively low fractional sensitivity of voltage sensors (ΔF/F \( \sim 5\% \) to 20% per 100 mV in brain slices) necessitates excitation densities and collection efficiencies sufficient to overcome high fractional shot noise. Loss of photon flux through confocal pinhole and lens arrays stipulates extensive signal averaging or long integration time to increase the S/N. Among efforts to measure voltage transients with two-photon fluorescence, Acker et al. achieved high spatial specificity by implementing two-photon excitation in “single voxel” (nonscanning) mode to record voltage transients in single dendritic spines. However, low fractional voltage dye sensitivity, combined with fast sampling rates necessitated by rapid voltage transients, limits the number of points from which high S/N voltage-sensitive fluorescence changes can be collected in serial.

Here we propose an alternative method to obtain voltage signals from specific structures of interest while preserving the ability to record from several spatial locations in parallel. To this end, we implemented one-photon computer-generated holography (CGH) to target VSD excitation over laterally extended regions, imaging epifluorescence signals in widefield mode with a high frame-rate camera. Cells filled with voltage dye were imaged and based on structure shape and location, a phase profile was calculated and addressed to a spatial light modulator (SLM) such that illumination at the sample plane was confined to the dendrite or axon of interest. In contrast with full-field illumination, patterned excitation minimized signal degradation arising from nonspecific autofluorescence, spilled dye, or nearby labeled structures. Importantly, with CGH-shaped illumination we could discriminate kinetic differences between action potential–evoked fluorescence transients recorded from neighboring axons and dendrites, differences that could not be appreciated with widefield illumination. These results suggest that CGH can provide a seamless means to increase the spatial specificity of functional fluorescence measurements.

2 Methods

2.1 Slice Preparation, Cell Loading, and Recording

We prepared slices of somatosensory cortex from wild-type mice (C57BL/6J) in accordance with the guidelines of European Union and institutional guidelines of the care and use of...
laboratory animals (Council directive 86/609 European Economic Community). Male or female mice (3 to 4 weeks of age) were deeply anesthetized with isoflurane (5%) and killed through decapitation. The brain was rapidly removed and placed in ice-cold (≤5°C) or room temperature cutting solution containing the following (in mM): 110 choline chloride, 2.5 KCl, 7.0 MgCl₂, 0.5 CaCl₂, 25 NaHCO₃, 1.25 Na₂HPO₄, 20 glucose, and aerated with 95% O₂, 5% CO₂ to a final pH of 7.4. We dissected the cerebral cortex, blocking it to take coronal slices of somatosensory cortex. The cortical block was glued to an ice-cold stage on a Leica microslicer, and 300 μm slices were cut in ice-cold cutting solution. Cut sections were placed in an incubator at 35°C for 0.5 h in the cutting solution in a chamber containing ACSF (in mM): 125 NaCl, 2.5 KCl, 1.0 MgCl₂, 2.0 CaCl₂, 25 NaHCO₃, 1.25 Na₂HPO₄, 20 glucose, and 0.4 L-ascorbic acid. Slices were kept at room temperature for at least 0.5 h before dye loading and recording in a submersion chamber.

2.2 Dye Loading

We visualized recovered cortical slices with Olympus BX51WI’s native infrared differential-interference contrast (IR-DIC) path, further magnifying the image onto a scientific complementary metal oxide semiconductor (sCMOS, ORCA Flash 4, Hamamatsu) camera with a 2×/4× magnification changer (Fig. 1 “MC”; Luigs and Neumann, Ratingen, Germany) mounted above the tube lens. We loaded neurons with VSD with a glass patch electrode by filling the tip with dye-free internal solution (in mM: 130 K-glucurate, 7 KCl, 4 ATP-Mg, 0.3 GTP-Na, 10 phosphocreatine-Na, 10 HEPES; adjusted to pH 7.4 and 284 mol/kg), then back-filling with the dye-containing internal solution (JPW3028, 400 to 800 μM), and performed whole-cell somatic recordings from the selected neuron under DIC optics. The dye JPW3028 is a doubly positively charged analog of the aminostyryl-pyridinium series of lipophilic VSDs available from Invitrogen as D-6923. This electrochromic dye does not significantly increase the membrane capacitance of the labeled neuron, as evident from several control measurements showing that the waveform of the electrically recorded action potentials remains unaltered after intracellular dye loading. A dye-free solution in the tip is necessary because this highly lipophilic molecule would otherwise spill onto the slice before sealing and patching the cell of interest, as any dye bound to membrane outside the cell of interest increases the background fluorescence and thus degrades recording sensitivity. The cell was patched and stained at room temperature for 15 to 30 min by allowing passive diffusion of the dye into the cell. The dye diffuses slowly compared with other dyes of the same size and weight due to its high affinity for lipid membranes. After staining, we carefully detached the patch electrode from the neuron, forming an outside-out patch, after which the slice was incubated at room temperature for 1 to 2 h, allowing the dye to diffuse from the soma into the axon and neighboring basal dendrites.

2.3 Computer-Generated Holography

Dye-loaded cells were illuminated with a 450 mW frequency-doubled diode-pumped Nd:YVO₄ low-noise laser emitting at 532 nm (Fig. 1 “LS”; MLL-FN-532-450-5-LAB-TTL, Changchun New Industries Optoelectronics Tech. Co. Ltd., Changchun, China) with CGH to achieve desired spatial patterns of light at the objective focal plane. Specifically, the laser beam (Fig. 1 LS) is attenuated with a neutral density (ND) filter, a half-wave plate (∆/2) and polarizing beam splitter, reading and adjusting the power with a removable mirror (M2), and a power meter (PM, Newport 818-ST2). Lenses L₁₁₁₁ (f = 19 mm) and L₁₁₁₂ (f = 150 mm), and pinhole (PH, d = 15 μm) clean and expand the beam to fill the active area of the liquid crystal on silicon spatial light modulator (LCOS-SLM, Hamamatsu X10468-01). Lenses L₁ (f = 400 mm) converts the modulated wavefront into a spatial light pattern, which is demagnified into the sample plane by a telescope formed by L₁, L₂ (f = 200 mm) and the microscope objective (Fig. 1 “OBJ,” Olympus LUMFLN 60XW, NA = 1.1, f = 5 mm). The 532-nm fluorescence excitation light is reflected through the objective by a 560-nm long-pass dichroic (FP560-FDi01-25x36, Semrock, Rochester, New York). Fluorescence transmitted through the dichroic is long-pass filtered (FF513LP, FF01-593/LP-25, Semrock) and imaged onto a high speed scientific-CMOS camera (sCMOS, Hamamatsu ORCA Flash 4.0) or electron multiplying charge-coupled device (EMCCD, Andor 860 iXon3) by the native Olympus tube lens (LTB) and MC used to target patch clamp electrodes. In order to block the zero-order non phase-modulated component reflected from the SLM, we introduced a defocus in the beam by adjusting the distance between L₁₁₁₁ and L₁₁₁₂ in order to displace the zero-order focus by 30 to 40 nm after the Fourier plane of the first lens. For the defracted first order, the defocus was compensated with a spherical Fresnel lens at the SLM. Thus, with the zero-order displaced 30 to 40 mm from the effective Fourier plane of L₁, we could block the unwanted zero-order component with a point block [Fig. 1 (LB), “BL” tape on cover slip] without perturbing the propagation of the hologram (first-order beam). Phase holograms were calculated with an iterative Fourier transform algorithm (IFTA). The IFTA-generated phase profiles were computed and addressed to the SLM using “Wavefront Designer IV,” in house software written in C++ with Qt 4.4.0 and fftw 3.1.2.

After the dye diffusion period, we imaged the neuron’s dendrites and axon at depths of 10 to 50 μm, weakly illuminating (excitation density ~1 nW/μm²) with a CGH-generated square shape (100 μm) leaving a light-free slot in the center ~30 μm wide [see Fig. 1 (LS)]. Positioning the soma in this light free slot enabled examination of axonal and dendritic structures, while sparing the dye-filled soma from unnecessary photodamage. Low-power density illumination was paired with long (500 ms) camera integration time and binning (2 × 2 or 4 × 4) to maximize collection and minimize light exposure and photodamage to stained axons and dendrites during preimaging acquisition. We then used fluorescence images of the axon and dendrites to define the spatial patterns of illumination for VSD signal acquisition trials, sculpted to the structure’s contours [Fig. 1 (LS)]. For VSD signal acquisition, we decreased the frame integration time to 156 μs (frame rate = 6.41 kHz) and modulated the laser power to achieve an excitation density in the range of 0.6 to 10 μW/μm² across the axon- and dendrite-shaped region. Laser light was gated onto the cell with a high-speed shutter [Fig. 1 (LS)] “Sh”; Uniblitz LS6, driver D880C, Rochester, New York] for 10 to 30 ms trials, with 1 to 2 min intertrial intervals. During each trial, we stimulated action potentials with brief current pulses (5 to 10 ms, 400 to 800 pA) applied through a dye-free pipette patched in whole-cell configuration. Electrical and optical waveforms were monitored.
for significant changes in width and fall-time indicative of photodamage, at which point the experiment was discontinued. CGH-sculpted light illumination trials were interleaved with “pseudowidefield” (pWF) trials illuminated with a large diameter (25 to 40 μm) spot with poor axial confinement [Fig. 1(d)], increasing laser power to maintain an excitation density equal to that of the sculpted light trials.

Current clamp signals were recorded at 20kHz with a MultiClamp 700B amplifier and 1440A digitizer (Molecular Devices, Sunnyvale, California), which also triggered camera acquisition and shutter opening. Micromanageπ piloted the sCMOS and Andor Solis drove the EMCCD. We analyzed fluorescence signals during experiments with the ImageJ time series plug-in and custom MATLAB scripts. Posthoc image visualization, processing, analysis, and statistics were performed in ImageJ and “VKAT: Voltage-imaging Kinetics Analysis Tool,” a wxpython-based GUI leveraging modules Numpy and Matplotlib.
2.4 Axial Propagation Simulations

In order to estimate and compare the z-confinement of CGH shaped and “pWF” configurations, we simulated the distribution of the holographic beam propagating along the optical axis around the objective focal plane as described in Ref. [26]. Briefly, we calculated the beam irradiance around the objective focal plane after the input phase hologram propagates through the telescope formed by L1, L2, and the objective using the angular spectrum approach of plane waves with a thin element approximation. Lutz et al. demonstrated that the simulations faithfully predict the experimentally measured propagation of shaped, CGH-generated spots. Although these simulations do not factor in depth-dependent brain tissue scattering, Zahid et al. have shown that holographic spots maintain axial confinement at depths of 30 μm in hippocampal slices. The axial confinement of the simulated beam propagation was quantified as full width at half maximum (FWHM) of intensity averaged over the “dendrite shaped” region of interest (ROI) in each axial plane [Fig. 1(c)].

2.5 Analysis

Axons and dendrites were discriminated visually, axons being thin and straight with collaterals issuing at obtuse angles, and dendrites thicker, spiny, and branching off in acute angle “Y”’s. We spatially averaged fluorescence signals from each imaged axon or dendrite over groups of 12 to 18 pixels. Acquiring frames at 6410 Hz, our sCMOS camera operated in “rolling shutter” mode, resulting in a systematic exposure delay of each horizontal line on the chip. Any temporal distortion potentially induced by the rolling shutter was canceled by spatially averaging axon and dendrite ROIs over pixels located across similar numbers of horizontal lines. All data shown and reported come from single-trial measurements. We applied a low-pass binomial filter (1 to 2 passes) to reduce high-frequency noise. In order to improve the temporal precision of spike kinetic measurements, the data collected at 6410 frames/s were reconstructed using cubic spline interpolation, and then resampled to 100 kHz before measuring spike width and rise time. Both for shaped and pWF trials, spike kinetic measurements were taken from ROIs in which both CGH excitation and widefield imaging were parfocal with the structure of interest. Axons and dendrites occupied different axial planes preventing simultaneous widefield epifluorescence imaging of both structures. Axonal and dendritic data were thus collected in separate, alternate trials. Because the \( \frac{dF}{F} \) signal from different parts of the neuron varies owing to factors other than voltage (e.g., partitioning of dye in inner and outer membranes), changes in voltage indicated by changes in fluorescence could not be calibrated on an absolute scale. We therefore limited comparison of signals emanating from axonal and dendritic compartments to kinetic parameters that do not depend on precise, absolute voltage scale calibration.

Since variable nonspecific fluorescence precludes signal amplitude comparisons between different ROIs, we performed paired comparisons of baseline fluorescence, noise, fractional change, and S/N in compartments illuminated with shaped and large spot pWF in alternate trials. To quantify spike-evoked fluorescence amplitude, we calculated the fractional change in fluorescence \( dF \), with respect to the baseline fluorescence for each signal-bearing pixel. Specifically

\[
\frac{dF}{F} = \frac{F - F_{BL}}{F_{BL} - F_{dark}},
\]

where \( F_{dark} \) is the average trace intensity 3 ms before the shutter opens and \( F_{BL} \) is the average “baseline” trace intensity during 3 ms after shutter opening but before spiking. For each trace, we also quantified the S/N:

\[
\frac{S}{N} = \frac{\mu}{\sigma},
\]

where

\[
\mu = \frac{dF}{F_{MAX}}
\]

and \( \sigma = \text{root-mean-square-deviation of samples acquired during “baseline” (BL, } t = 1:n) \):

\[
\sigma = \sqrt{\frac{1}{n} \sum_{i=1}^{n} \left( \frac{dF}{F} \right)^2 - \left( \frac{dF}{F} \right)_{BL}^2}.
\]

We quantified differences between the kinetics (spike width, rise time) and S/N characteristics between axonal- and dendritically generated waveforms, as well as CGH and “pWF” (large spot) configurations.

3 Results

We utilized CGH to confine VSD fluorescence excitation light to an axon or dendrite of interest [Figs. 1(c), 2(b) and 2(c)], comparing signals obtained in this fashion to those illuminated in “pWF,” that is with a large spot of light with poor axial confinement [Figs. 1(d) and 2(d)]. CGH-sculpted illumination generated an excitation spot that precisely covered the structure of interest and corresponded to approximately 1/20 of the area of the 25 to 40 μm-diameter circles used for pWF trials [Figs. 1(c) and 1(d)]. Axial propagation simulations of 532 nm CGH-generated shapes predict axial confinement between 7.0 and 9.6 μm for shaped illumination [Fig. 1(c)], and between 41 and 66 μm for pWF spots [Fig. 1(d)]. Both CGH and “pWF” excitation of dye in axons and dendrites generated fluorescence signals from which spike waveforms could be resolved in single trials [Figs. 1(e) and 1(f)].

3.1 Targeted Fluorophore Excitation Enables Signal Discrimination in Neighboring Structures

We found that CGH-shaped voltage dye fluorescence excitation in neighboring axons and dendrites enabled discrimination of differing action potential kinetics not possible with large-field illumination. Specifically, we compared 10% to 90% rise time and FWHM of action potential-evoked fluorescence transients recorded in axons and dendrites illuminated with CGH-sculpted shapes in alternate trials (Fig. 2). Corresponding to previously reported electrical [13] and optical measurements [12] action potential rise time and FWHM were shorter \( p \)-values \( \leq 0.05 \) Student’s t-test; Figs. 2(a), 2(b), and 2(c)] in axons (FWHM: 1.62 ms ± 0.24 standard error of the mean, S.E.M.; rise time: 221.1 μs ± 95.6 S.E.M., \( n = 3 \) trials from two cells) than in dendrites (FWHM: 2.37 ms ± 0.30 S.E.M.; rise time: 1154.6 μs ± 194.0 S.E.M.; \( n = 4 \) trials from two cells). Displayed action potential-evoked fluorescence transients...
emanating for the dendrite [Fig. 2(b)] and axon [Fig. 2(c)] were collected in separate trials since these structures occupied two different planes of focus. Figure 2(e) redisplays these two traces, amplitude normalized and peak aligned to show the difference in spike kinetics undetectable with pWF illumination [Fig. 2(a)]. Trials interleaved in which neighboring axons and dendrites were simultaneously illuminated with large spots did not show differences [Fig. 2(f), 2(g), and 2(h), p-values > 0.3 Student’s t-test] in spike kinetics in the same axonal (FWHM: 1.87 ms ± 0.09 S.E.M.; rise time: 618.5 µs ± 68.2 S.E.M.; n = 4 trials in two cells) and dendritic ROIs (FWHM: 1.80 ms ± 0.15 S.E.M.; rise time: 600.2 µs ± 170.2 S.E.M.; n = 3 trials in two cells), even though these structures occupied different planes of focus.

3.2 Targeted Excitation Increases Fractional Spike-Evoked Transient Amplitude but not Signal-To-Noise

Paired comparisons of CGH shaped (n = 4) or “pWF” (n = 4) trials revealed significantly decreased background fluorescence (F_{Bkg} = F_{BL} – F_{dark}, averaged over time postshutter opening and prespike), increased RMS noise and increased peak action potential evoked dF/F [dF/F₉₅ summarized by Fig. 3(table)]. Although inhomogeneous partitioning of the lipophilic voltage dye precludes comparison of the signal amplitude between axonal and dendritic compartments, here we compare dF/F₉₅ emanating from the same ROIs under sculpted and pWF illumination.
illumin
tion conditions. Since both the signal ($\mu = dF/F_{MAX}$) and RMS noise increased with CGH-shaped excitation, we observed no change in S/N.

4 Discussion

We demonstrate that the CGH light targeting to neuronal structures enables functional fluorescence transient spatial specificity unachievable with widefield epifluorescence. Importantly, differences in action potential kinetics between adjacent axons and dendrites could be discerned with targeted CGH illumination, but not with pWF illumination. This gain in spatial specificity, despite reducing baseline fluorescence and total photon flux, did not decrease signal-to-noise ratios (Fig. 3, S/N). Specifically, the increased baseline noise (RMS$_{BL}$) was compensated by increased transient $dF/F$, presumably due to the higher fraction of photons emanating from external membrane-embedded fluorophores undergoing synchronous electrophoretic shifts during action potential propagation. In the pWF case, increased baseline fluorescence drives an expected decrease in RMS noise, as shot-noise decreases in proportion to lateral area, with temporal focusing the lateral area and axial confinement are decoupled for laterally extended axons and dendrites [Fig. 1(c) and 1(d)].

In addition, temporally focused shapes show high robustness to scattering, further recommending this approach for extended shape fluorescence excitation in depth.

One undisputable disadvantage of CGH is the inhomogeneous light distribution within generated light patterns. These intensity "speckles" vary by 15% to 20% for one-photon excitation, and up to ~50% for two-photon, due to the quadratic dependence of the signal on the excitation density. Speckles arise primarily from the approximation in the IFTA and cross talk between adjacent pixels of the LCOS-SLM. Techniques such as rotating diffusers and phase mask shift-averaging can average over speckles and smooth the spatial distribution, but at the cost of light efficiency, axial or temporal resolution. Speckle-free illumination can also be accomplished with generalized phase contrast and interferometric method for light shaping. Overcoming the problem of speckle, especially in the two-photon case, will be important since large intensity fluctuations can exceed photodamage thresholds.

In conjunction with the rapidly expanding toolbox of organic and protein-based, genetically targetable voltage calcium, and neurotransmitter reporters, holographic light shaping can enable high S/N, parallel detection of neuronal-evoked fluorescence transients from neighboring cells or subcellular compartments. Here, we have demonstrated that light targeting increases spatial specificity even for cases in which the dye itself is confined to the membrane of a single cell. Targeted light spatial specificity gains could prove more critical with dye bulk loading or with dense expression of genetically targeted voltage-sensitive fluorescent proteins.

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References


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