Bright and photostable chemogenetic indicators for extended in vivo voltage imaging

Ahmed S. Abdelfattah1, Takashi Kawashima1,†, Amrita Singh1,2, Ondrej Novak1,3, Hui Liu1, Yichun Shuai1, Yi-Chieh Huang3, Luke Campagnola3, Stephanie C. Seeman5, Kaspar Podgorski1, Bei-Jung Lin4, Tsai-Wen Chen4, Glenn C. Turner1, Zhe Liu1, Yang-Ming University, Taipei 112, Taiwan. 5Allen Institute, Seattle, WA 98109, USA. 6Department of Neuroscience and Grossman Center for Brain Science, Columbia University, New York, NY 10027, USA.

Genetically encoded voltage indicators (GEVIs) enable monitoring of neuronal activity at high spatial and temporal resolution. However, the utility of existing GEVIs has been limited by the brightness and photostability of fluorescent proteins and rhodopsins. We engineered a GEVI, called Voltron, that uses bright and photostable synthetic dyes instead of protein-based fluorophores, thereby extending the number of neurons imaged simultaneously in vivo by a factor of 10 and enabling imaging for significantly longer durations relative to existing GEVIs. We used Voltron for in vivo voltage imaging in mice, zebrafish, and fruit flies. In the mouse cortex, Voltron allowed single-trial recording of spikes and subthreshold voltage signals from dozens of neurons simultaneously over a 15-minute period of continuous imaging. In larval zebrafish, Voltron enabled the precise correlation of spike timing with behavior.

Animal behavior is produced by patterns of neuronal activity that span a wide range of spatial and temporal scales. Understanding how neural circuits mediate behavior thus requires high-speed recording from ensembles of neurons for long periods of time. Although the activity of large numbers of neurons can now be routinely recorded using genetically encoded calcium indicators (GECIs) (1), the slow kinetics of calcium signals complicate the measurement of action potentials, and subthreshold voltage signals are missed entirely (2-3). Voltage imaging using genetically encoded voltage indicators (GEVIs) can overcome these challenges, enabling imaging of fast spikes and subthreshold dynamics in genetically defined neurons (4, 5). The high imaging speed and excitation intensity required for voltage imaging, combined with the smaller volume of the cellular membrane, place increased demands on voltage indicators relative to GECIs. Extant GEVIs rely on fluorescence from either microbial rhodopsins (6-8) or fluorescent proteins (9-13). These fluorophores lack the brightness and photostability to allow in vivo imaging from large fields of view over time scales of many behavioral events, precluding the millisecond-time scale analysis of neural circuits. Improved rhodopsine dyes such as the Janelia Fluor (JF) dyes can be used in complex biological experiments because of their high brightness and photostability (14), compatibility with self-labeling protein tags (15, 16), and ability to traverse the blood-brain barrier for in vivo delivery (17). We describe a “chemogenic,” or hybrid protein–small molecule, GEVI scaffold that we call Voltron, which irreversibly binds these synthetic fluorophore dyes. Voltron provides an increased photon yield that enables in vivo imaging of neuronal spiking and subthreshold voltage signals in model organisms with order-of-magnitude improvement in the number of neurons imaged simultaneously over substantially longer durations.

Our design for a chemogenetic voltage indicator combines a voltage-sensitive microbial rhodopsin domain (6, 7, 11) with a dye-capture protein domain (Fig. 1A) that irreversibly binds a synthetic fluorophore dye ligand (14, 15) (Fig. 1B), analogous to previously reported voltage indicators that use fluorescent proteins (10, 11, 18). Transmembrane voltage–dependent changes in the absorption spectrum (6, 19) of the rhodopsin domain of Voltron reversibly modulate the degree of fluorescence quenching of the nearby bound dye through Förster resonance energy transfer (FRET). We investigated the modularity of this approach, finding that three different rhodopsin domains—QuaAr1 (7), QuaAr2 (7), and Ace2N (11, 20)—modulated the fluorescence of the rhodamine dye Janelia Fluor 549 (JF549) after binding to either HaloTag (15) or SNAP-tag (21) dye-capture protein domains (figs. S1 to S8). Removing a small number of amino acid residues at the junction of the rhodopsin and self-labeling tag domains increased the amplitude of fluorescent voltage signals (fig. S1), presumably by decreasing average distance and thus increasing FRET efficiency between the dye and rhodopsin retinal cofactor. The configuration providing the best signal-to-noise ratio (SNR) for spikes was Ace2N fused to HaloTag with five amino acids removed at their junction (Fig. 1, A and B, and fig. S2), hereafter referred to as Voltron.

We tested several Voltron-dye combinations in cultured rat neurons and acute mouse brain slices with high-speed imaging and simultaneous whole-cell patch clamp electrophysiology (Fig. 1C, figs. S6 and S9 to S13, and tables S1 and S2). Voltron could detect neuronal action potentials and subthreshold potential changes with a variety of JF dye ligands with emission maxima between 520 nm and 660 nm using fluorescence imaging with one-photon excitation (Fig. 1, C to E, and fig. S6) but was not compatible with two-photon imaging, as described previously for rhodopsin-containing GEVIs (22, 23). Voltron bound to JF525 (Voltron525) exhibited the highest sensitivity, giving a fluorescence change of ~23 ± 1% ΔF/F0 (ΔF/F0 for a voltage step from -70 mV to +30 mV (Fig. 1E and fig. S9)). Voltron525 showed similar sensitivity. Voltron525 responded to voltage steps with submilliseconds on and off time constants (table S3 and fig. S10). We compared the brightness and photostability of Voltron in neuronal cultures with those of two other fluorescent protein–based GEVIs: Ace2N-mNeon (17) and ASAP2f (13). Both Voltron525 and Voltron549 were brighter than Ace2N-mNeon (by a factor of 3 to 4) and ASAP2f (by a factor of 16 to 18) (Fig. 1P) in cell culture. This difference did not result from differences in expression; we compared the brightness of Voltron549 and Ace2N-mNeon at the single-molecule level and observed a similar brightness difference (factor of 3 to 4) (Fig. 1G). Voltron525 and Voltron549 were also more photostable in ensemble measurements (Fig. 1H, tables S4 and S5, and figs. S14 and S15) as well as in single-molecule assays, in which photobleaching times were longer for Voltron549 than those of Ace2N-mNeon by a factor of 8 (Fig. 1I). Overall, the improved brightness and photostability of Voltron increased the photon yield by at least a factor of 10 in neurons relative to existing GEVIs that rely on fluorescent proteins.

In vivo, Voltron could be reliably expressed and labeled with dye in mice, larval zebrafish, and adult fruit flies (Figs. 1 to 4 and figs. S16 to S19 and S21 to S45). Simultaneous in vivo electrophysiology and Voltron imaging in each of these organisms confirmed the detection of individual action potentials (Fig. 1, J and K, and figs. S17 to S19). For imaging in the mouse brain,
we used a variant of Voltron appended with a 63-amino acid sequence from the rat potassium channel Kv2.1 that restricts expression to the membrane of the cell body and proximal dendrites (24, 25) (Voltron-ST; fig. S20). The rapid kinetics of Voltron525-ST allowed clear observation of action potentials in fast-spiking parvalbumin (PV)-expressing interneurons in the CA1 region of the mouse hippocampus (Fig. 2, A to G, and fig. S21). We measured the orientation tuning of the spiking and subthreshold responses of cortical layer 2/3 pyramidal neurons in mouse primary visual cortex in response to the mouse observing directional movement of light and dark stripes, a benchmark for new indicators (1, 11) (Fig. 2, H to L, and figs. S22 to S24), and confirmed that spiking activity showed sharper orientation selectivity than did subthreshold voltage signals.

**Fig. 1. Development of the chemigenetic voltage indicator Voltron.**

(A) Schematic of Voltron sequence: A rhodopsin (Ace2) is fused to a self-labeling tag domain (HaloTag) with additional sequences added to improve or localize membrane targeting: Golgi export trafficking sequence (TS), endoplasmic reticulum export sequence (ER), and somatic targeting sequence (ST). (B) Model of Voltron mechanism. (C) Left: Cultured rat hippocampal neuron expressing Voltron and labeled with JF525. Scale bar, 20 μm. Right: Single-trial recording of action potentials and subthreshold voltage signals from current injections in primary neuron culture using 400-Hz imaging (top, fluorescence) or electrophysiology (bottom, membrane potential). (D) Fluorescence emission spectra of different JF dyes overlaid with the absorbance spectrum of Ace2N. (E) Fluorescence change as a function of membrane voltage with different JF dye–Voltron conjugates. (F) Relative fluorescence of ASAP2f, Ace2N-mNeon, Voltron525, and Voltron549 in cultured neurons (n = 70, 68, 48, and 62 measurements, respectively, from five independent transfections for each construct). Illumination intensity was ~10 mW/mm² at imaging plane. ***P < 0.001 [one-way analysis of variance (ANOVA) followed by Bonferroni test on each pair]; NS, not significant. Fluorescence was normalized to ASAP2f mean intensity. (G) Relative single-molecule brightness of Ace2N-mNeon and Voltron549. ***P < 0.001 (two-tailed Student t test). (H) Bleaching curves for ASAP2f, Ace2N-mNeon, Voltron525, and Voltron549 in primary neuron culture. Illumination intensity was ~23 mW/mm² at imaging plane. Bleaching curves were normalized to mean cellular fluorescence from (F) or normalized to the zero-time value (inset). (I) Mean time to bleach of Ace2N-mNeon and Voltron549 during single-molecule imaging, 100-ms frames. ***P < 0.001 (two-tailed Student t test). (J and K) Simultaneous in vivo Voltron imaging (300 and 800 Hz, top) and electrophysiology (bottom) in larval zebrafish (extracellular) and adult Drosophila (whole-cell), respectively. Spike-triggered averages are shown at the right. Scale bars, 20 μm.
Fig. 2. Using Voltron to measure membrane voltage dynamics in hippocampal PV neurons and visual cortex pyramidal neurons of mice.

(A) Schematic of imaging of spontaneous activity in the CA1 region of the hippocampus of an awake mouse. (B) Image of hippocampal PV neuron expressing Voltron labeled with JF525. Scale bar, 20 µm. (C to E) Voltron525 raw ΔF/F₀ traces showing spontaneous spikes of a PV neuron (B) located at a depth of 60 µm in the hippocampal CA1 region of a fully awake mouse imaged at 3858 frames per second. Color-coded boxes indicate regions shown at expanded time scales. (D) is a zoom of the boxed region from (C); (E) is a zoom of the boxed region from (D). (F) Overlay of 177 spikes detected during a 15-s period (gray) and their average (black). (G) Spike shape of 11 PV neurons. (H) Schematic of imaging of primary visual cortex of an anesthetized mouse during display of drifting grating visual stimuli. (I) Example trace showing 500-Hz Voltron fluorescence during one trial of a sequence of visual stimuli. Arrows below represent the direction of movement of the drifting grating. (J to L) Top left: Images of a pyramidal cell at a depth of 148 µm, imaged three times over a period of 4 weeks at the indicated number of weeks after virus injection. Scale bar, 10 µm. Top right: Average of all spikes in session (black) and SD (gray). Middle: Raw ΔF/F₀ trace for five repetitions in each session, showing two orthogonal orientations (indicated with arrows below) from the neuron pictured at top left. Bottom: Orientation tuning to full-frame drifting gratings of the neuron pictured at top left, displayed as number of spikes during trials (blue), number of spikes during preceding intertrial intervals (gray), and subthreshold ΔF/F₀ (right y axis) after low-pass filtering traces using a 10-point median filter. For each orientation, subthreshold response is calculated by averaging the low-pass–filtered trace between 100 and 400 ms after trial onset, and baseline is calculated by averaging the low-pass–filtered trace from 80 ms preceding trial onset to 20 ms after trial onset. Data are displayed as response minus baseline. Error bars represent SEM (20 to 22 repetitions per session).
We extended the imaging period over several consecutive weeks by injecting additional JF525 HaloTag ligand before each imaging session (Fig. 2, J to L, and fig. S24).

Next, we attempted to image larger areas containing more neurons for longer times in vivo in mouse cortex (Fig. 3). By wide-field microscopy at illumination intensities between 3 and 20 mW/mm², we could clearly identify and distinguish action potentials from nearby neurons throughout 15 min of continuous imaging (SNR = 5.3 during the first minute, 4.4 during the final minute) (Fig. 3, B to E). We expanded the field of view to include dozens of cortical interneurons labeled with Voltron525-ST in a transgenic mouse line (NDNF-Cre) (27) while imaging at 400 Hz (Fig. 3, F and G, and figs. S25 to S42). Overall, we imaged a total of 449 neurons (12 fields of view in three mice), demonstrating routine voltage imaging of populations of neurons in superficial mouse cortex (Fig. 3G and figs. S25 to S42). This scale of in vivo voltage imaging enabled analysis of membrane potential correlations between many neuron pairs (fig. S26).


Fig. 3. Imaging of voltage activity in large fields of view of the mouse neocortex over long durations. (A) Schematic of the imaging setup. (B) Image of two neurons expressing Voltron525-ST in layer 1 of the visual cortex of an NDNF-Cre mouse. Scale bar, 10 μm. (C) ΔF/F₀ traces from neurons in (B), recorded over 15 min at 400 Hz. (D) Color-coded zooms of indicated regions of the traces in (C), with detected action potentials indicated by black dots above the fluorescence traces. (E) Average of all spikes in session (black) and SD (gray). (F) Left: Fluorescence image of a cranial window over primary visual cortex (V1) in an NDNF-Cre mouse showing Cre-dependent expression of Voltron525-ST (bright spots). Scale bar, 1 mm. Right: Zoomed image of left panel in the area indicated by the white rectangle, with neuron labels corresponding to fluorescence traces in (G). Scale bar, 100 μm. (G) Left: ΔF/F₀ traces during 3 min of recording at 400 Hz from neurons pictured in (F), in decreasing order of SNR. Right: Zooms of ΔF/F₀ traces from color-coded regions of (G) with detected action potentials represented as black dots above, illustrating representative traces with high (top), medium (middle), and low (bottom) SNR. Traces have been background-subtracted, which removes shared subthreshold membrane potential fluctuations (compare to fig. S25 without subtraction).
Fig. 4. Voltron reveals millisecond–time scale neural dynamics during swimming behavior in zebrafish.

(A) Schematic illustration of the setup. An immobilized zebrafish is placed under the light-sheet microscope, and the motor signals (inset) from its tail are recorded using a pair of electrodes. Visual stimuli (forward-drifting gratings) for eliciting swimming responses are presented below the fish. (B) Left: Anatomical location of the imaged brain region (midbrain nucleus; see fig. S44A). Center: Representative field of view of the imaged region expressing Voltron. Scale bar, 20 μm. Right: Position of neurons analyzed in (C). (C) Left: Periods of visual motion (pink) and swim signals (gray) are plotted above Voltron fluorescence traces (black) simultaneously recorded from 11 neurons shown in (B). Right: Zoom of swimming signals (top) and Voltron fluorescence traces from two representative neurons (bottom) are expanded from the dashed box in the left panel. Dots at the top of each trace represent action potentials recognized by the algorithm described in fig. S45, A and B. Inverted triangles and dotted gray lines indicate initiation of each swim bout. (D) Mean subthreshold signal (top), mean frequency of action potentials (middle), and raster plots of action potentials (bottom) near the initiation of swim bouts from three representative neurons: “Off” (green), “Onset” (red), and “Late” (blue). Shadows in the top and middle panels represent SEM across swim events. (E) Classification of recorded neurons by their mean subthreshold signals near the initiation of swim bouts; 179 neurons recorded from 43 fish were classified using non-negative matrix factorization and colored according to the weights for three factors: “Onset” (red), “Off” (green), and “Late” (blue). See supplementary materials for details of this classification. (F) Spatial organization of the same population of neurons as in (E). Neurons from multiple fish are superimposed on a single map according to the distance from the center of this midbrain nucleus. (G) Hypothetical model of neural activity modulation in this midbrain nucleus.
We used Voltron to image zebrafish larvae, which respond to visual input with fast, directed swim bouts that are tailored to the details of the stimulus (29). We sought to uncover how this sensory-to-motor transformation unfolds in neuronal populations at fine time scales that are inaccessible with calcium imaging. We verified that Voltron could detect action potentials and subthreshold voltage signals in live zebrafish after labeling with several different colors of dye ligands (figs. S37 and S40). We then used Voltron to monitor neural activity during swim bouts induced by visual motion (Fig. 4A). We recorded Voltron signals from 179 neurons across 43 fish in a motor-sensory nucleus in the tegmental area of the midbrain (Fig. 4B and fig. S44), yielding data on subthreshold membrane voltage modulation as well as automatically detected spike times (Fig. 4C and fig. S45). We found neuron populations with different temporal activity patterns, including neurons whose firing rate increased ~1 s before the fish started swimming (fig. S44, B and C, “Ramp”), neurons whose firing rate was suppressed each time the fish swam (Fig. 4D, “Off”), and neurons that fired each time the fish swam (Fig. 4D, “Onset” and “Late”). Of the latter types, some fired just before swimming (~20 ms before swim onset, “Onset”) and others fired just after swimming (~10 ms after swim onset, “Late”). There was a change in subthreshold voltage that preceded these firing rate changes by tens of milliseconds (Fig. 4D and fig. S44D). The neuron types were spatially intermingled within this midbrain nucleus (Fig. 4, E and F). The existence of neurons that fired before swimming and neurons that fired after swimming may indicate that this nucleus both partakes in the generation of swim bouts and is influenced by the motor output (Fig. 4G). Thus, Voltron allows for the dissection of population motor coding and sensorimotor integration circuits in ways that neither single-cell electrophysiology nor population calcium imaging can.

We tested Voltron in adult Drosophila in vivo by expressing the protein in a pair of dopaminergic neurons, one in each brain hemisphere, which innervates a single compartment in the mushroom body. We detected strong spiking signals from axons and dendrites of these neurons with Voltron(4o) (Fig. 1K and fig. S18). The fluorescence signals matched action potentials detected using electrophysiology. In some neuronal cell types in Drosophila, calcium indicators located in the cell body have failed to exhibit fluorescence changes even under conditions where high spike rates are expected (29). However, spikes were clearly detectable when imaging from the soma of dopamine neurons with Voltron (fig. S1E). We could clearly distinguish spikes from the two neurons according to the amplitude of the spiking signals even when imaging from neuropil where their axons overlap extensively, likely because each bilaterally projecting cell contributes a denser innervation of the mushroom body in the ipsilateral hemisphere (fig. S1D).

Combining the molecular specificity of genetically encoded reagents with the superior photophysical properties of chemical dyes is an established path to improved imaging reagents (34). However, previous attempts to create hybrid protein–small molecule indicators by various approaches have not been successful for in vivo imaging (30). We engineered a modular sensor scaffold in which the targeting and sensor domains are genetically encoded and only the fluorophore and its protein-binding anchor are synthetic. The resulting chemigenetic indicator, Voltron, exhibits increased photon output, enabling in vivo voltage imaging of many more neurons over longer times—approximately 10^2 more neuron-minutes than other sensors. This improvement enables imaging experiments that can help to reveal how the precise electrical dynamics of neuronal populations orchestrate behavior over different time scales.

REFERENCES AND NOTES
Bright and photostable chemigenetic indicators for extended in vivo voltage imaging


Science 365 (6454), 699-704.
DOI: 10.1126/science.aav6416 originally published online August 1, 2019

Visualizing neuronal activity in vivo

Imaging the changes in fluorescence of voltage-sensitive reagents would enable monitoring of the activity of neurons in vivo. Abdelfattah et al. created such a voltage indicator by designing a protein that combines the voltage sensor domain from microbial rhodopsin with a domain that captures a dye molecule with exceptional brightness and photostability. When the protein was expressed in mice, flies, or zebrafish, they could monitor single action potentials in dozens of neurons simultaneously for many minutes.

Science; this issue p. 699

ARTICLE TOOLS
http://science.sciencemag.org/content/365/6454/699

SUPPLEMENTARY MATERIALS
http://science.sciencemag.org/content/suppl/2019/07/31/science.aav6416.DC1

REFERENCES
This article cites 61 articles, 7 of which you can access for free
http://science.sciencemag.org/content/365/6454/699#BIBL

PERMISSIONS
http://www.sciencemag.org/help/reprints-and-permissions

Use of this article is subject to the Terms of Service

Science (print ISSN 0036-8075; online ISSN 1095-9203) is published by the American Association for the Advancement of Science, 1200 New York Avenue NW, Washington, DC 20005. The title Science is a registered trademark of AAAS.

Copyright © 2019 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works