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Voltage Imaging with a NIR-Absorbing Phosphine Oxide Rhodamine Voltage Reporter

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ABSTRACT: The development of fluorescent dyes that emit and absorb light at wavelengths greater than 700 nm and that respond to biochemical and biophysical events in living systems remains an outstanding challenge for noninvasive optical imaging. Here, we report the design, synthesis, and application of near-infrared (NIR)-absorbing and -emitting optical voltmeter based on a sulfonated, phosphine-oxide (po) rhodamine for voltage imaging in intact retinas. We find that po-rhodamine based voltage reporters, or poRhoVRs, display NIR excitation and emission profiles at greater than 700 nm, show a range of voltage sensitivities (13 to 43% Δ F/F per 100 mV in HEK cells), and can be combined with existing optical sensors, like Ca²⁺-sensitive fluorescent proteins (GCaMP), and actuators, like light-activated opsins ChannelRhodopsin-2



(ChR2). Simultaneous voltage and Ca^{2+} imaging reveals differences in activity dynamics in rat hippocampal neurons, and pairing poRhoVR with blue-light based ChR2 affords all-optical electrophysiology. In *ex vivo* retinas isolated from a mouse model of retinal degeneration, poRhoVR, together with GCaMP-based Ca^{2+} imaging and traditional multielectrode array (MEA) recording, can provide a comprehensive physiological activity profile of neuronal activity, revealing differences in voltage and Ca^{2+} dynamics within hyperactive networks of the mouse retina. Taken together, these experiments establish that poRhoVR will open new horizons in optical interrogation of cellular and neuronal physiology in intact systems.

INTRODUCTION

Fluorescence microscopy has revolutionized the life sciences. Advances in both synthetic and genetically encoded fluorophores make fluorescence microscopy routine in laboratories across the globe. Coupled with progress in areas like super-resolution microscopy,¹ light sheet microscopy,² and tissue clearing,^{3–5} fluorescent dyes bring the high spatial and temporal resolution of optical microscopy to bear on samples ranging from single molecules to interacting cells.

However, noninvasive optical imaging in intact tissues and organisms remains challenging. Commonly used fluorophores, like fluorescein and rhodamine, possess excitation and emission profiles within the visible region of the spectrum and overlap extensively with endogenous chromophores, especially heme.^{6,7} Although alternative imaging modalities like magnetic resonance imaging, positron emission tomography, and photoacoustic imaging afford excellent penetration into thick tissue, these modalities require specialized instrumentation, are half-life limited, require radioactive reagents, and/or cannot access the micrometer spatial and millisecond time resolution routinely afforded by optical microscopy.⁸

In the context of neurobiology, an outstanding challenge is tracking neuronal voltage dynamics with submicron spatial resolution and submillisecond temporal resolution within intact tissues. Although two-photon (2P) microscopy provides access to structures in deeper tissue, traditional raster-scanning 2P microscopy cannot achieve the temporal resolution required to image large numbers of neurons simultaneously.⁹

Therefore, fluorescent dyes and voltage indicators that possess near-infrared (NIR, 700–1000 nm)^{10–14} excitation and emission profiles are of considerable interest. First, NIR photons scatter less than visible photons in thick tissue. Second, by avoiding endogenous chromophores like heme (<650 nm) and water (>900 nm),^{6,7,15} fluorophores that operate in the so-called NIR window gain profound advantages for tissue and *in vivo* imaging. Finally, NIR fluorophores can be more readily deployed for multiplex imaging alongside existing

Received: October 30, 2020 Published: January 27, 2021



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fluorescent labels, indicators, and actuators that use visible wavelengths of light, on account of the larger spectral separation between the two.

Despite the attraction of NIR dyes that both absorb and emit at wavelengths above 700 nm, traditionally employed NIR dyes, such as porphyrins and phthalocyanins,¹⁰ polymethines,¹⁰ BODIPY derivatives,^{16–18} and xanthenes with extended annulation^{19–21} tend to suffer from high molecular weights, low water solubility, a propensity to aggregate, and chemical or photochemical instability, complicating their use in biological contexts. A number of creative approaches have recently been employed to address these concerns.^{14,20,22–24} In particular, the compact, fused 3-ring system of xanthene-based dyes promotes stability and decreases molecular weight. Annulation of xanthenes can push their absorbance above 700 nm,¹⁹ but at the expense of decreased water solubility. Recent efforts replace oxygen at position 10 with carbon^{25–27} or heteroatoms such as Si,^{28–30} S,³¹ and P. Indeed, recent examples of phosphinate³² and phosphine-oxide based xanthene dyes^{35,34} demonstrate this unique scaffold can access absorption profiles at or above 700 nm while maintaining a compact, three-ring structure.

While numerous approaches to NIR-emitting fluorophores exist, there are relatively few examples of NIR-absorbing fluorescent reporters: fluorophores that sense and respond reversibly to changes in a specific physiological phenomena, for example, metal ions, reactive oxygen species, or transmembrane potential. The few examples of NIR-absorbing indicators employ polymethine, rather than xanthene, scaf-³⁷ Therefore, we were eager to adapt phosphorusfolds.4,35substituted xanthenes into a fluorescent voltage-sensing scaffold. Previous work in our lab suggests that xanthene dyes with a range of bridgehead $(O_{,38}^{,38} C_{,27}^{,27} S_{,39}^{,39})$ and terminal atoms $(O_{,38}^{27,38} or substituted N_{,39,40}^{39,40})$ can be employed as voltage-sensitive dyes via introduction of a lipophilic, conjugated molecular wire. We hypothesized that the installation of phenylenevinylene molecular wires into the context of a phosphorus-substituted xanthene dye would yield voltage-sensitive indicators with peak excitation above 700 nm. There are no examples of xanthene-based voltage-sensitive fluorophores with peak excitation above 700 nm, although there are a few examples of chemically synthesized⁴¹⁻⁴⁴ and genetically encoded 45-47 voltage indicators with emission maxima beyond 700 nm.

We now present the **p**hosphine-**o**xide **Rho**damine Voltage Reporters, or poRhoVRs, which all feature excitation and emission maxima above 700 nm, 13% to 43% Δ F/F per 100 mV voltage sensitivity, and compatibility with commonly used optical sensors and actuators, including Oregon Green BAPTA (OGB), GCaMP6, and ChannelRhodopsin-2 (ChR2). We use poRhoVR to provide the first direct voltage imaging of neuronal hyperactivity in retinas isolated from a mouse model of retinopathy.

RESULTS

Synthesis of Sulfonated Phosphorus Rhodamines. Although phosphorus rhodamines possess attractive optical properties, ^{32–34} NIR excitation and emission, photostability, ^{32–34} and good water solubility, their tendency to localize to intracellular membranes, including mitochondria³⁴ and lysosomes, ⁴⁸ precludes their straightforward deployment as cytosolic indicators or voltage sensors. Water solubility and exclusion from cellular uptake can be achieved through installation of anionic groups on the exocyclic rhodamine nitrogens;^{49,50} however, we recently found that sulfonation of the *meso* aromatic ring of rhodamine-type systems prevents cellular uptake of dyes.^{39,51} We hypothesized that a similar strategy could provide sulfonated phosphorus rhodamines for incorporation into voltage sensing scaffolds.

Phosphine-oxide RhoVRs were accessed in 6 steps from commercially available triphenylphosphonium salts (Scheme S1). Anilino phosphine oxide 4 was prepared in high yield over three steps.^{33,52} Phosphine oxide 4 was alkylated through a slightly modified procedure using ethyl iodide and potassium carbonate in dimethyformamide (DMF) to produce phosphine oxide 5 in 89% yield. The key sulfonated phosphorus rhodamine precursors with bromine substitution for installation of the phenylenevinylene molecular wire were prepared in one step by condensing bromosulfobenzaldehyes⁵¹ 6 or 7 with phosphine oxide 5 in the presence of urea⁵³ and glacial acetic acid (Scheme 1). Traditional methods for acid-catalyzed

Scheme 1. Synthesis of Sulfonated Phosphine Oxide Rhodamine Voltage Reporters (poRhoVRs)



construction of rhodamines or sulfonorhodamines, for example, condensation with methanesulfonic acid⁵⁴ or propionic acid without urea^{40,55} resulted in no reaction or were extremely low yielding.

The condensation between m-(6) or p-bromosulfobenzaldehyde (7) and phosphine oxide 5 produced two isomers of m- (8) and p- (9) bromosulfo-phosphine oxide rhodamines^{33,34} in which the P-methyl substituent and sulfonate exist in either a cis or trans relationship to one another (Figure 1b), consistent with previous reports and nomenclature³³ (Figure S1). For both 8 and 9, the two isomers were separable on a basic alumina column, eluting with 0-5% methanol in dichloromethane. After this step, no further purification was necessary for the major isomer of either 8 or 9. Preparative HPLC was required to isolate the minor isomer. Analysis of the NMR spectra of the purified major and minor isomers of 8 and 9 revealed differences in the chemical shifts of both the phosphorus atom and the P-methyl substituents (Table S1): for the P-methyl group, $\delta = 2.00$ ppm (¹H, major isomer) and $\delta = 1.96$ (¹H, minor isomer); for the phosphorus atom, $\delta =$ 20.3 ppm (³¹P, major) and 18.3 ppm (³¹P, minor) (Table S1 and Figure S1). Furthermore, X-ray crystal structures of the major isomer of both 8 and 9 reveal a cis relationship between



Figure 1. Characterization of sulfonated phosphine-oxide rhodamine dyes 8 and 9. (a) Plots of normalized absorbance (solid lines) and fluorescence intensity (dashed lines; excitation provided at 625 nm) of phosphine-oxide rhodamine 8 (thick blue lines) and 9 (black lines) in phosphate buffered saline (5 μ M dye, 1% DMSO). (b) Chemical structure of the *cis* isomers of 8 and 9. Thermal ellipsoid plots (50%) of parts c and d, 8 and 9, respectively. Hydrogen atoms, lattice solvent molecules, and resolved disordered fragments have been omitted for clarity.

the sulfonate and the methyl group on the phosphorus bridgehead atom (Figure 1c,d). Attempts to crystallize the minor isomer of 8 or 9 were unfruitful. In aqueous solution (phosphate buffered saline, 5 μ M dye), both bromo phosphine oxide rhodamine 8 – *cis* and 9 – *cis* had an absorption maximum at 704 nm, emission maxima around 728 nm (Figure 1a), and quantum yields near 20% (22% for 8 and 19% for 9).

Synthesis of Phosphorus Rhodamine Voltage Reporters. After determining that sulfonated po-rhodamines have reasonable quantum yields and desirable NIR excitation and emission maxima above 700 nm, we hypothesized that incorporation of phenylenevinylene molecular wires should render these NIR xanthenes voltage-sensitive. We installed molecular wires 10 (dimethyl aniline) or 11 (diethyl aniline and methoxy substituent) onto the phosphine-oxide rhodamine fluorophore using a Pd-catalyzed Heck coupling to generate a total of four new phosphine oxide Rhodamine VoltageReporters (poRhoVRs, 12–15, Scheme 1). For each reaction, the cross coupling was performed using the purified,

major isomer (cis) of either 8 or 9. However, in each case this did not prevent the formation of cis/trans isomers in the complete poRhoVR molecules: analytical HPLC/MS showed two peaks with the product mass corresponding to poRhoVR. Per previous reports,³³ separation of the isomers is extremely arduous but in the case of 13, extensive column chromatography on silica gel and preparative scale HPLC on C18functionalized silica afforded small quantities of major (9% yield) and even smaller quantities of minor isomers (1% yield). Comparison of the ¹H and ³¹P NMR data for the isomers of 13 closely match the chemical shifts observed for the cis and trans isomers of 8 and 9, suggesting that the major product is the cis isomer of 13 (Figure S1). Isomers of the other three poRhoVRs (12, 14, and 15) were not separated due to the difficulty of purification. ³¹P showed that for poRhoVRs 12, 14 and 15 major products isolated were *cis* (³¹P δ = 20 ppm), consistent with the results for poRhoVR 13 – *cis* (³¹P δ = 20.2 ppm).

All of the poRhoVR compounds possess excitation and emission profiles well-matched to the parent phosphine-oxide rhodamines: absorption was maximal at 700 nm, and emission peaked at around 728 nm (Table 1, Figure S3a-d).

Cellular Characterization of Phosphorus Rhodamine Voltage Reporters. All four poRhoVR dyes (12–15) localize to the cellular membrane and display differing cellular brightness when applied to HEK cells under identical experimental conditions (Figure S2a-d). Although all poRhoVRs localize to the membrane, poRhoVRs 13 and 15, with N,N-diethyl, methoxy-aniline substituents, are approximately 10-fold dimmer in cell membranes than poRhoVR 12 and 14 (N,N-dimethyl aniline) (Table 1, Figure S2e). The long wavelength excitation and emission of poRhoVRs enables multicolor imaging with organelle specific dyes. Live-cell imaging⁵⁶ of poRhoVR 14 (1 μ M, Figure 2a) in HEK cells costained with rhodamine 123^{57} (the methyl ester of rhodamine 110),⁵⁸ which localizes to the mitochondria on account of its overall positive charge (Figure 2b) and Hoechst 33342,⁵⁹ a bisbenzimidazole dye which binds to nucleic acid, labeling the nuclei of cells, (Figure 2c) demonstrate good membrane localization for poRhoVRs along with compatibility with simultaneous, multicolor imaging (Figure 2d).

The cellular fluorescence of poRhoVRs is voltage sensitive. Simultaneous fluorescence microscopy during sequential depolarizing and hyperpolarizing steps from a holding potential of -60 mV in HEK cells under whole-cell voltage-clamp conditions reveals linear fluorescence vs voltage relationships for poRhoVR dyes 12–14 (Figure 2e,f, 14; Figure S3 for all indicators). Voltage sensitivities range from 13% (poRhoVR 12) to 43% (poRhoVR 13) per 100 mV (Table 1). The sensitivity of poRhoVR 13, at 43%, is nearly 2-fold the

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Compound number	isomer ^a	\mathbb{R}^1	R ²	$\lambda_{\rm abs}/ \ {\rm nm}^{b}$	$\lambda_{\rm em}/ {\rm nm}^{b}$	Φ^b	rel. cell brightness ^c	$\Delta F/F^d$
8	meta	n/a	n/a	704	728	0.22	n/a	n/a
9	para	n/a	n/a	704	728	0.19	n/a	n/a
12	meta	Н	Me	705	731	0.01	15 ± 0.9	$13 \pm 0.3\%$
13	meta	OMe	Et	703	728	0.01	1.0 ± 0.1	$43 \pm 0.9\%$
14	para	Н	Me	704	723	0.07	12 ± 0.1	$17 \pm 0.5\%$
15	para	OMe	Et	704	726	0.09	1.3 ± 0.03	$31 \pm 0.5\%$

^{*a*}Relationship between fluorophore and molecular wire. ^{*b*}dPBS with 0.1% DMSO. ^{*c*}Relative cellular brightness, in HEK cells. ^{*d*}Per 100 mV, in HEK cells. Data are mean \pm SEM for at least 3 separate determinations.



Figure 2. Cellular characterization of poRhoVR indicators in HEK cells. Widefield, epifluorescence images of (a) poRhoVR 14 (1 μ M) in HEK cells. Cells are counter-stained with (b) rhodamine 123 (1 μ M) and (c) Hoechst 33342 (1 μ M) to visualize mitochondria and nuclei, respectively. (d) An overlay of poRhoVR 14, rhodamine 123, and Hoechst 33342. Scale bar for parts a–d is 20 μ m. (e) Plot of fractional change in fluorescence of poRhoVR 14 vs time for 40 ms hyper- and depolarizing voltage steps from a holding potential of –60 mV for a single HEK cell labeled with poRhoVR 14 (1 μ M). (f) Plot of Δ F/F vs membrane potential, summarizing data from n = 6 individual HEK cells. Error bars are ± SD. If error bars are not visible, the error is smaller than the marker.

sensitivity of our most red-shifted voltage indicator to date, the Si-rhodamine based BeRST $(24\% \Delta F/F \text{ per } 100 \text{ mV})^{39}$ and is comparable to the sensitivity of our previously reported, tetramethyl rhodamine based RhoVR (47% Δ F/F per 100 mV).⁴⁰ However, the low cellular brightness of poRhoVR 13 (more than 10x dimmer than poRhoVRs 12 and 14) means that indicators like poRhoVR 14, with intermediate voltage sensitivity (17% Δ F/F per 100 mV), but high cellular brightness (12x brighter, compared to 13) are also attractive for monitoring voltage dynamics in living systems. This is because, in a shot-noise-limited system, SNR scales with the square root of the number of collected photons, or the cellular brightness.^{60,61} Therefore, even indicators with only middling $\Delta F/F$ values can be quite effective detectors of rapid voltage changes.^{62,63} For poRhoVR indicators, it appears that the OMe-substituted compounds (13 and 15) show lower cellular brightness compared to the unsubstituted aniline molecular wires of 12 and 14 (Table 1). This effect appears to be specific to poRhoVR indicators, as rhodamine-based RhoVR indicators show the highest SNR values for OMe-substituted molecular wires.⁴⁰

We evaluated both poRhoVR 13 and 14 in cultured hippocampal neurons isolated from rat embryos. We selected 13 and 14 to investigate whether a bright, but less sensitive dye (14) could function as well as a dim but more sensitive dye (13). Both poRhoVR 13 (Figure S4a-c) and 14 (Figure 3a-c) localize well to cellular membranes and readily report on spontaneously firing action potentials in single-trial acquisitions, revealing a range of neuronal firing rates similar to previously reported ranges.⁶⁴ In some cases, poRhoVR 14 clearly visualizes subthreshold voltage dynamics, likely postsynaptic potentials (Figure 3c, cell 3, asterisks).

Consistent with its performance in HEK cells, poRhoVR 14 was approximately 8 to 12-fold brighter than 13 under



Figure 3. Voltage imaging in dissociated rat hippocampal neurons with poRhoVR 14. Transmitted light image of neurons loaded with (a) poRhoVR 14 (500 nM). (b) Epifluorescence image of neurons showing poRhoVR 14 staining. Scale bars are 20 μ m. (c) Plot of fractional change in poRhoVR 14 fluorescence (Δ F/F) vs time emanating from cells 1–4 in image b. Optical sampling rate is 500 Hz. Asterisks indicate subthreshold voltage changes.

identical experimental conditions (Figure S5a). In a complementary fashion, the Δ F/F value per spike/action potential for 13 was larger than for 14 (12% vs 6%, Figure S5b) but with lower overall signal-to-noise ratio (SNR, 20:1 vs 40:1, Figure S5c). Because of the higher brightness and SNR for detecting action potentials, we used poRhoVR 14 for subsequent experiments.

We next examined the photostability, internalization rate, and toxicity of poRhoVR 14. The photostability of poRhoVR 14 is comparable to the photostability of the previously reported BeRST 1 (Figure S6).³⁹ We also measured the persistence of poRhoVR 14 on the cell membrane in HEK cells. We examined the degree of poRhoVR 14 internalization from the membrane by measuring the colocalization between calcein-AM⁶⁵ fluorescence, a cytosolic stain that becomes fluorescent and is retained in living cells, and poRhoVR 14 (Figure S7). Increased colocalization values indicates a higher degree of internalization of the membrane-associated po-RhoVR 14. We find a low degree of colocalization between poRhoVR 14 signal and cytosolic calcein-AM after an initial 20 min of loading (the Pearson's correlation coefficient is $0.22 \pm$ 0.2, mean \pm SEM for n = 3 coverslips) (Figure S7a). There is an increase in the degree of colocalization, to approximately 0.26, after additional observation for up to 2 h (Figure S7), but the changes are not statistically significant (P > 0.05, Dunnett's multiple comparisons test; see Figure S7). Finally, we examined the toxicity of poRhoVR 14 in HEK293T cells using calcein-AM. Comparison of the colocalization of Hoechst 33342 fluorescence, which stains the nuclei of both dead and live cells, and calcein-AM fluorescence reveals no difference in colocalization, and therefore toxicity, between vehicle-treated HEK293T cells (DMSO) and up to 5 μ M poRhoVR 14 (Figure S8).

poRhoVR 14 and Optogenetic Activator ChR2. poRhoVR 14 can be used in multicolor experiments not only to track static fluorescence associated with distinct organelles (Figure 2a–d) but also in concert with commonly employed optogenetic actuators and sensors. We expressed the

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Figure 4. All-optical electrophysiology using poRhoVR 14 and ChR2. (a) Transmitted light image of dissociated rat hippocampal neurons stained with poRhoVR 14 (500 nM). Scale bar is 20 μ m. (b) Epifluorescence image of neurons stained with poRhoVR 14. (c) Epifluorescence image of neuron displaying YFP marker of ChR2 expression. (d) Composite image depicting poRhoVR 14 labeling and ChR2-YFP expression. (e) Recording of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels a–d. (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f) Plot of Δ F/F from the cell bodies of neurons indicated in panels (f)



Figure 5. Simultaneous voltage and calcium imaging with poRhoVR 14. Epifluorescence image of neurons stained with both (a) poRhoVR 14 (1 μ M) and (b) Oregon Green BAPTA 1 AM (OGB, 1 μ M). (c) Plots of Δ F/F for voltage (poRhoVR 14, purple) and Ca²⁺ transients (OGB, green) in response to field stimulation driven at 5, 10, 20, and 30 Hz. (d) Epifluorescence image of a neuron transfected with GCaMP6s. (e) This same GCaMP6s (+) neuron is also stained with poRhoVR 14 and imaged simultaneously. Scale bar is 10 μ m. (f) Simultaneously recorded traces of voltage and calcium activity from neuron in panels d and e. Activity was evoked using field stimulation at a rate of 16 Hz. (g) The insets show that the onset and decay of voltage signals imaged with poRhoVR 14 precede that of the calcium signal visualized from the same cell with GCaMP6s.

blue light-activated opsin, ChannelRhodopsin-2 (ChR2)^{66,67} fused to yellow fluorescent protein (YFP) in a subset of hippocampal neurons isolated from rat (Figure 4a,b). Bath application of poRhoVR 14 to these same neurons results in membrane-localized fluorescence (Figure 4c) that is spectrally isolated from the YFP signal (Figure 4d).

The combined use of ChR and poRhoVR allows dissection of functional connectivity across a large number of neurons. Optical stimulation of cell 1 (475 nm, 5 ms, 1.92 mW/mm²) results in ChR2-evoked action potentials optically recorded at cell 1 (Figure 4e,f). Cells 4 and 5 appear monosynaptically coupled to ChR2-positive cell 1 with action potential latencies of 6.7 ms \pm 1.5 ms (SD n = 60 pairs of spikes) and than 10 fire act 5.6 ms \pm 1.3 ms (S.D., n = 20 out of 60 spikes), respectively. However, the data reveals differences in the relative strengths

However, the data reveals differences in the relative strengths of these connections with cell 1 triggering firing in cell 4 for 100% of action potentials, and only 33% for cell 5. Interestingly, bursts of spontaneous activity indicate strong recurrent connectivity between all neurons (Figure 4e,f, gray shaded areas), but firing initiated in the ChR2-expressing neuron (cell 1, Figure 4a–d) activates only a subset of these neurons (Figure 4e,f): notably, cell 4, nearly 50 μ m away from cell 1, and cell 5, over 120 μ m distant from ChR2-expressing cell 1.

The combination of poRhoVR and ChR2 enables interrogation of subthreshold potentials. In a few cases, during cyan light stimulation, we observe slower, subthreshold potentials in cells up to 400 μ m away from the ChR2-expressing neuron (Figure S9d,e, cells 2–4, asterisks), highlighting the ability of poRhoVR 14 to monitor subthreshold voltage changes (Figure 3a–c). The use of cyan light to stimulate ChR2 does not crossexcite poRhoVR 14, as indicated by the lack of stimulus artifact in the ChR2-negative cells in the same field of view (Figure 4d,e, Figure S9). Together, these experiments establish poRhoVR 14 as a powerful complement for all-optical electrophysiology utilizing NIR absorbing indicators.

Two Color Imaging with poRhoVR 14 and Ca²⁺ Indicators. In addition to deployment alongside lightactivated actuators, poRhoVR can be used with optical indicators. Fluorescent sensors for Ca²⁺ are among the most widely used optical sensors. Despite some three decades since the initial reports of fluorescent indicators for this critical intracellular messanger,^{68,69} most Ca²⁺ indicators utilize excitation and emission profiles firmly centered in the blue/ green region of the visible spectrum (for example, Oregon Green BAPTA, OGB,⁷⁰ and the GCaMP family of genetically encoded indicators).⁷¹ Although promising new Ca²⁺ indicators, both synthetic^{72–75} and genetically encoded,^{76–79} possess red-shifted excitation and emission spectra, circularly permuted (cp) GFP-based indicators, like the GCaMP family,⁷¹ dominate the landscape of functional imaging.^{80,81} Therefore, fluorescent voltage indicators with orthogonal wavelengths are required.

We performed two-color, simultaneous voltage and Ca²⁺ imaging in the same cells using poRhoVR 14 and the synthetic Ca²⁺ indicator, OGB (Figure 5a–c). We treated hippocampal neurons with both poRhoVR 14 (500 nM) and OGB (1 μ M) simultaneously and imaged using an image-splitting device to project two emission wavelengths onto the same camera chip. Under these conditions, we observe clear membrane-associated fluorescence for poRhoVR 14 (Figure 5a) and cytosolic fluorescence for OGB (Figure 5b). We established that no cross-excitation exists under these conditions (Figure S10). Using field-stimulation electrodes, we evoked a series of 10 action potentials, across a range of frequencies, and simultaneously recorded voltage (Figure 5c, magenta traces) and Ca²⁺ (Figure 5c, green traces) dynamics.

Both poRhoVR 14 and OGB clearly resolve single action potentials when activity is evoked at rates of either 5 or 10 Hz. poRhoVR 14 clearly resolves action potentials at firing rates of 20 and 30 Hz (Figure 5c). OGB, despite its fast Ca^{2+} response kinetics (<5 ms to action potential peak)⁷¹ compared to GCaMP6f (~45 ms to peak)⁷¹ and other genetically encoded indicators,^{82–84} fails to accurately report individual action potential-evoked Ca^{2+} transients at firing frequencies higher

than 10 Hz (Figure 5c). Neurons in the brain and retina can fire action potentials at rates up to several hundred Hz, for example, in interneurons of the hippocampus,⁸⁵ Purkinje cells of the cerebellum,⁸⁶ and ganglion cells of the retina,⁸⁷ emphasizing the need for indicators with fast response kinetics.

 Ca^{2+} indicators are often characterized against varying numbers of action potentials arriving at a constant frequency. However, neural information is often encoded in the form of spike rates. Therefore, resolution of individual spikes and firing frequency is critical to an understanding of the underlying physiology of the system under observation. Even with very fast OGB, estimating spike frequency using Ca^{2+} imaging traces alone was unsuccessful. Neither peak $Ca^{2+} \Delta F/F$ (Figure S11a) nor integrated area under the curve (Figure S11b) were able to resolve differences at 5, 10, 20, or 30 Hz firing rates. In contrast, the optically recorded voltage transients revealed by poRhoVR 14 clearly discriminate between firing frequencies of 5, 10, 20, and 30 Hz (Figure S11c).

Simultaneous voltage and Ca²⁺ imaging in the same cells can also be achieved alongside genetically encoded indicators, like GCaMP6. We again stained neurons with poRhoVR 14 $(1 \ \mu M)$. This time, a subset of hippocampal neurons expressed GCaMP6s. Again, poRhoVR localizes to membranes (Figure 5d), while GCaMP6s fluorescence appears cytosolic (Figure 5e). Simultaneous voltage and Ca²⁺ imaging of spontaneous activity in hippocampal neurons reveals fast-spiking bursts resolved in voltage (Figure 5f-h, magenta trace), followed by slower, sustained increases in GCaMP6-associated fluorescence (Figure 5f-h, green trace). Notably, voltage imaging with poRhoVR 14 exhibits sufficiently high temporal resolution to distinguish individual action potentials in spike volleys (Figure 5g, 8 spikes; Figure 5h, 9 spikes), while Ca^{2+} imaging does not. Together, these experiments establish the utility of poRhoVR dyes for monitoring fast spiking in neurons alongside commonly used synthetic and genetically encoded Ca²⁺ indicators and emphasizes the care needed when interpreting Ca²⁺ imaging data.

Voltage and Ca²⁺ Imaging and Electrode Recording in a Mouse Model of Retina Degeneration. The NIR (>700 nm) excitation and emission spectra of poRhoVR dyes, along with their good voltage sensitivity, compatibility with commonly used optogenetic sensors and actuators, and ready uptake into cell membranes (Figure S12), make poRhoVR 14 a promising candidate for mapping voltage dynamics in intact neural tissue like retinas. The retina is a highly organized and accessible outpost of the central nervous system. Light responses initiated in rods and cones are synaptically transmitted to bipolar cells, which activate retinal ganglion cells (RGCs). RGCs generate the action potentials that carry visual information to the brain. In normally functioning retinas, the intrinsic light sensitivity of photoreceptors in rods and cones complicates optical imaging of both voltage and Ca²⁺ transients in RGCs, because visible light (or high intensity two photon excitation) used to excite the indicators triggers physiological responses.⁸⁸ We applied poRhoVR 14 to investigate membrane potential dynamics in retinas from a mouse model of retina degeneration.

In particular, retinas from rd1 mice are an attractive model system in which functional imaging can be applied to explore mechanisms occurring in inherited visual disorders, including the degenerative disorder retinitis pigmentosa (RP).⁸⁹ Lacking a functional β subunit of rod cGMP phosphodiesterase (β PDE),⁹⁰ rd1 mice suffer rapid loss of rod cells, followed



Figure 6. Simultaneous mapping of electrical and Ca^{2+} activity using poRhoVR, GCaMP6f and multielectrode arrays (MEA) in *ex vivo* retinas from *rd1* mice. Widefield fluorescence micrographs of retina stained with (a) poRhoVR 14; the retinal ganglion cells (RGCs) express (b) GCaMP6f. The black dots are MEA electrodes, labeled numerically, underneath the retina. Scale bar is 20 μ m. Recordings c–f depict MEA, GCaMP6f, and poRhoVR 14 signals vs time. Traces are as follows: raw MEA electrical signal (black), bleach corrected poRhoVR Δ F/F (arbitrary units) (magenta), and GCaMP6f Δ F/F (%) (green). Optical signals are from the regions of interest (ROIs) indicated in blue in panels a and b. Panels c and d depict the spontaneous activity in the retina prior to addition of synaptic blockers. Panels e and f show MEA, GCaMP6f, and poRhoVR 14 signals 15 min after the addition of synaptic blockers. Panels c and e correspond to signals associated with electrode 26 (e26), and panels d and f correspond to signals associated with electrode 25 (e25). Panels g and h show zoomed-in regions of e25, from the time period indicated by a gray bar in panel d and f, respectively. In panels g and h, the MEA signal (black) is inverted to facilitate comparison with optical voltage recordings with poRhoVR (magenta).

by a delayed loss of cone cells. As a result, these mice lack rods and cones and are therefore blind but still retain functional RGCs in the ganglion cell layer (GCL). Therefore, apart from the rare melanopsin-expressing, intrinsically sensitive RGC (<2% of the RGCs) the surviving RGCs in *rd1* retinas have no light response and continue to receive synaptic input from bipolar cells. RGCs in *rd1* mice develop hyperactivity, firing spontaneous bursts of spikes in darkness. This phenomenon has been observed only through electrophysiological recordings that sample only a fraction of cells, or by Ca²⁺ imaging with GCaMP expressed in RGCs, which is indirect and has low temporal resolution.^{87,91}

The precise mechanisms underlying hyperactivity are not understood, and voltage imaging could reveal where in the retinal circuitry this activity originates and how it propagates from neuron to neuron.

To explore the interplay between neuronal voltage and Ca²⁺ signaling in this model of retinopathy, we prepared *ex vivo*, flatmount retinas from *rd1* mice that express GCaMP6f in retinal ganglion cells (RGCs). Bath application of poRhoVR 14 (5 μ M, in oxygenated ACSF) results in diffuse poRhoVR 14 staining throughout the tissue, as assessed by widefield fluorescence microscopy (Figure 6a, Figure S14d,e). In retinas from *rd1* mice, we observed spontaneous oscillations of poRhoVR 14 fluorescence (Figure 6d,h, magenta traces). To confirm that these changes in poRhoVR 14 dynamics correspond to voltage changes in RGCs, we paired voltage

imaging with multielectrode array (MEA) recordings of extracellular potential.

The poRhoVR-stained retinas were mounted on a 64channel MEA to simultaneously record transmembrane potential (poRhoVR 14), extracellular potential (MEA), and Ca²⁺ transients (GCaMP6f) in many RGCs. Simultaneous transmembrane voltage imaging and MEA recording of extracellular potentials (black traces) confirm that poRhoVR 14 (magenta traces) accurately reports changes in transmembrane potential in RGCs of the retina (Figure 6c,f). The time courses of the MEA signals (black) and poRhoVR 14 signals (magenta) match well, establishing that poRhoVR senses voltage changes in mouse retina. The signs of the MEA and poRhoVR 14 signals are inversely related (Figure 6c-f): poRhoVR 14 measures transmembrane potential, while MEA records extracellular potentials. A zoomed-in view of poRhoVR 14 and MEA signals (Figure 6g,h), where the black MEA signals have been inverted to enable better comparisons, show excellent correspondence between the optically recorded transmembrane potential measured with poRhoVR 14 (magenta) and the MEA recording (black).

Voltage imaging reveals spontaneous membrane potential depolarizations that appear clustered in bursts throughout the imaging session (Figure 6c,d, magenta). These optically recorded oscillatory bursts are, to our knowledge, the first direct imaging of membrane potential dynamics in rd1 retinas, and are consistent with previous MEA recordings in rd1

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retinas.^{92,93} GCaMP6f recordings from the same areas revealed slower, Ca^{2+} transients that were delayed relative to increases in the transmembrane potential measured by poRhoVR 14 fluorescence (Figure 6c,d, green). This lag between Ca^{2+} and voltage is similar to our observations of simultaneous Ca^{2+} / voltage imaging in hippocampal neurons (Figure 5d,h) and provides the first direct and simultaneous observation of voltage and Ca^{2+} in the mouse retina. We observe no evidence of oxidative photobluing⁹⁴ of po-rhodamines under these imaging conditions (Figure S13).

Voltage imaging with poRhoVR 14 enables dissection of the temporal evolution of neuronal activity in *rd1* retina. Synaptic isolation of RGCs via blockade of all major forms of excitatory and inhibitory synaptic transmission results in evolution of activity from short, unsynchronized firing patterns (Figure 6c,d,g) to synchronous and sustained firing (Figure 6e,f,h), consistent with the idea that the driver of this behavior is presynaptic.⁹² Voltage imaging with poRhoVR 14 also allows investigation of the spatial differences in voltage dynamics with the retina. We recorded fluorescence from a region of interest (ROI) composed of a ring with a width approximately matching that of an RGC (~10 μ m) around the MEA electrode (Figure 6a,b). This enables a direct comparison to extracellular potentials recorded by the MEA (Figure 6c,d, black). For the two electrodes visible in the recording image (Figure 6a,b), the MEA recording is closely matched by the poRhoVR 14 dynamics (Figure 6c,d, black vs magenta). The use of poRhoVR 14 allows examination of the spatial relationships of voltage changes at locations distant from electrodes (Figure S14). Imaging of poRhoVR 14 signals recorded in ROIs in between electrodes reveal a unique activity profile that is not fully recapitulated by ROIs near the respective electrodes (Figure S14a,b), highlighting the ability of poRhoVR to map the spatial extent of voltage dynamics in retina. This methodology could prove a promising starting point for mapping the dendritic fields of RGCs by probing where voltage dynamics become convoluted with additional signals.

CONCLUSION

In summary, we present the design, synthesis, and application of phosphine-oxide rhodamines for voltage imaging. These new poRhoVRs have excitation and emission profiles above 700 nm and possess good voltage sensitivity. Their compatibility with other optical sensors and actuators makes them a powerful complement to existing approaches to dissect neuronal activity. We show that poRhoVR 14 can report on spontaneous action potentials in rat hippocampal neurons and enables all-optical electrophysiological manipulations with ChR2. Simultaneous voltage imaging with poRhoVR 14 alongside either fast synthetic Ca²⁺ indicators or genetically encoded Ca²⁺ indicators reveals the difficulty of relying solely on Ca²⁺ imaging to interpret underlying neuronal activity. Furthermore, poRhoVR 14 can be deployed in intact retinas alongside multielectrode arrays and Ca2+ imaging to record from many RGCs at once in a mouse model of retinal degeneration, providing a first direct visualization of voltage dynamics alongside simultaneous Ca²⁺ imaging in the retina.

In the future, we envision combining poRhoVR with chemical-genetic hybrid methods to enable cell type-specific labeling in intact tissues.^{95,96} Beyond the chemical sensing presented in this paper, innovations in microscopy will be needed to achieve the kilohertz framerates, with micrometer

resolution, across large, 3D volumes regions of the central nervous system that will be required for visualizing patterns of activity in arrays of neurons *in vivo*.⁹⁷ Promising advances microscopy,⁹ including light sheet and swept-field microscopy,⁹⁸ redesigned optical paths,⁹⁹ holographic imaging,^{100,101} and compressive sensing¹⁰² hint at possible avenues toward the use of poRhoVR for *in vivo* brain imaging.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c11382.

Supporting figures, spectra, procedures, and analysis (PDF)

Accession Codes

CCDC 2056525–2056526 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

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ACKNOWLEDGMENTS

Research in the Miller lab is supported by grants from the NIH (R01NS098088, R35GM119855) and Klingenstein-Simon Foundations (40746). E.W.M and R.H.K. acknowledge

support from NSF Neuronex (1707350). M.A.G. was supported in part by a training grant from the NIH (T32GM066698). Crystallographic data was acquired on an ApexII-Quazar, supported NIH grant S10RR027172. Research in the Kramer lab is supported by the NIH (R01EY024334 and R01NS100911). We thank Benjamin Smith for assistance with image analysis.

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