

# Dual-color optical activation and suppression of neurons with high temporal precision

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
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

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## Abstract

A fundamental challenge in the optogenetic toolbox is that all opsins, regardless of their excitation spectra, are activated by blue light. We hypothesized that pairing a red-shifted channelrhodopsin with a blue light-sensitive anion channel of appropriately matching kinetics shall render neurons responsive to a red but not blue light. To achieve this, we used a semi-rational mutagenesis strategy to optimize the kinetics and light spectrum of a chloride channelrhodopsin. By pairing optimized variants of blue light-sensitive anion channel ZipACR with vfChrimson, a fast red-shifted channelrhodopsin, we created a system in which red light derives precise and faithful action potentials of high frequencies, while blue light, through shunting inhibition, nullifies the effect of the red-shifted ChR2. Additionally, by a simple switch between red and blue lights, one can effectively excite or inhibit the activity of the same neurons.

### eLife assessment

This study develops **valuable** tools for optogenetic control of neuronal activity. The basic characterization of the activation of a red-shifted channelrhodopsin paired with a blue-light sensitive anion channel engineered to obtain desired inhibitory current kinetics is **solid**. However, guidelines and feasibility for their practical use under simultaneous multi-color stimulation are **incomplete**, as are comparisons with current state-of-the-art tools.

Despite the short history of optogenetics, optical manipulation of neuronal activity has become an indispensable tool in the studies for functional analysis of brain circuits. The increasing needs for a more precise temporal and spatial manipulation of the neurons along with the untapped potentials of optogenetics have been driving forces in expanding the repertoire of optogenetics toolbox. Main development among these expansions is the discovery of opsins of diverse excitation spectra, with the current excitatory opsins showing light sensitivity ranging from near UV to near far-red spectra (Yizhar *et al.*, 2011 ; Deisseroth and Hegemann, 2017 ; Marshel *et al.*,

2019 [↗](#)). The main purpose for such developments is to independently as well as collectively control the activity of distinct populations of neurons. This aim, however, has not been satisfied, mainly because all opsins, in addition to their own excitation spectral peak, are activated by blue light, a problem known in the field as cross-over or bleed-through. Attempts to create blue light-insensitive opsins will be most likely unsuccessful due to the intrinsic blue absorption by the retinal molecule, the non-encoded chromophore, which captures light energy in channelrhodopsins (Waddell *et al.*, 1973 [↗](#); Wang *et al.*, 2012 [↗](#)).

There have been two stand out attempts to overcome the problem of cross-over. One uses a variant of blue-shifted ChR2 with fast kinetics in combination with a slow kinetics red-shifted ChR2 (R-ChR) (Klapoetke *et al.*, 2014 [↗](#)). The reasoning is that short pulses of blue light will activate only fast ChR2. Since it requires a tight regulation of the protein expression as well as the light intensity, this method has not been adopted for in-vivo application. Another approach is to expose axons expressing a R-ChR to prolonged photostimulation. Immediately following this, axons become unresponsive to light pulses and cease to fire. This method is limited to axonal regions as prolonged optical stimulation of the soma induces trains of action potentials (Hooks *et al.*, 2015 [↗](#)).

A potential solution to the problem of cross-over, the one that we propose here, is to co-express a blue-shifted anion channelrhodopsin (B-ACR) along with the R-ChR to ‘suppress’ its excitation by blue light. For this purpose, we chose vfChrimson—currently the fastest R-ChR—to gain more precise temporal control of red light excitation (Mager *et al.*, 2018 [↗](#)). This would, however, require a new variant of anion channelrhodopsin to optimize pairing. We used a semi-rational mutagenesis approach to modify the kinetic properties of ZipACR, an ultrafast blue light-activated chloride channel (Govorunova *et al.*, 2017 [↗](#)). We describe here two variants of ZipACR, I151T and I151V, with appropriate kinetics, light spectra, and photocurrent. We then paired the optimized anion channels with vfChrimson in a single expression unit. The new system, Zip-IvfChr, can drive time-locked high-frequency action potentials (APs) in response to red pulses (635 nm). In response to blue pulses (470 nm), on the other hand, Zip-IvfChr provides selective suppression of individual APs within trains of high frequency firing with high precision.

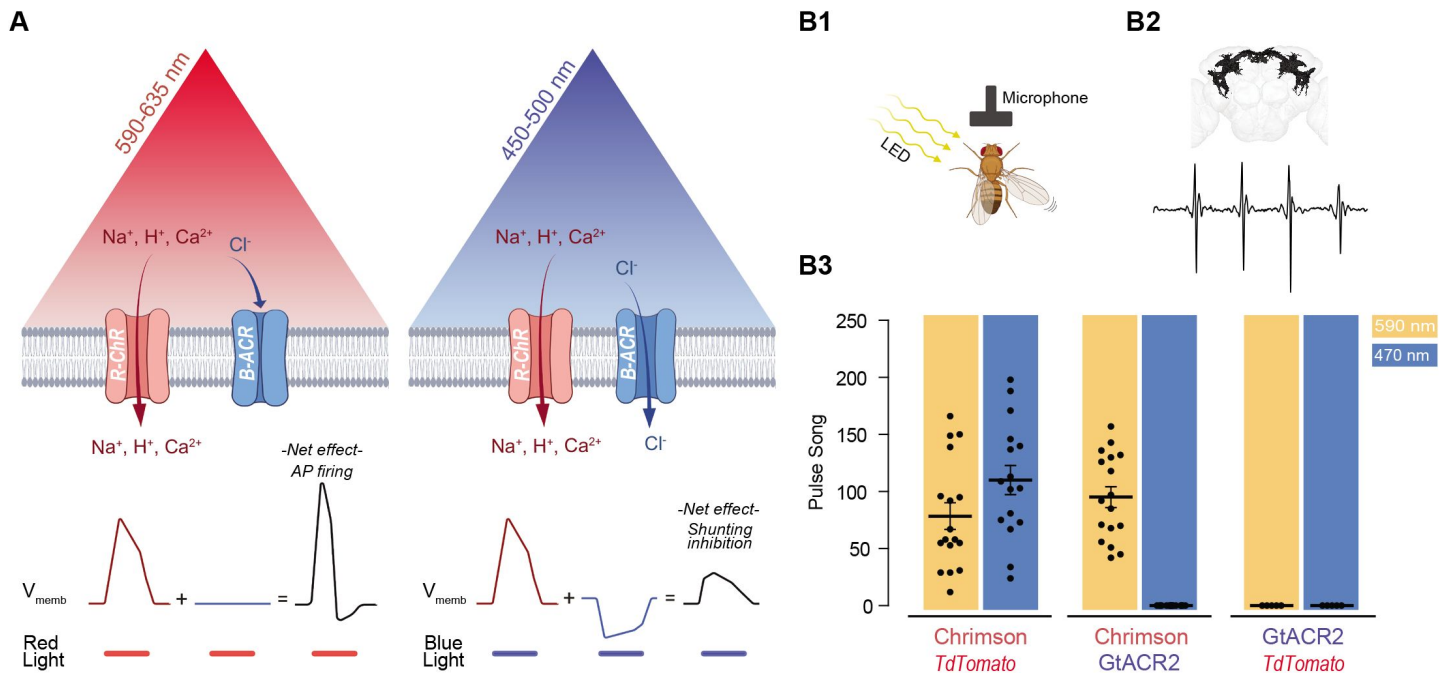
## Results

### The strategy to reduce blue light-mediated excitation by red-shifted channelrhodopsins

The channelrhodopsin Chrimson and its variants have the most red-shifted spectra (Klapoetke *et al.*, 2014 [↗](#)), and their response to 470 nm has a slower on-rate compared to the 590 nm light (Supplementary Fig. 1A-B [↗](#)). Despite this, 470 nm light is equally effective in inducing APs (Supplementary Fig. 1C [↗](#)). To overcome this limitation, we decided to take advantage of blue light-gated inhibitory chloride channels which effectively block action potential upon exposure to blue light (Govorunova *et al.*, 2015 [↗](#); Govorunova *et al.*, 2017 [↗](#)). By co-expressing a B-ACR along with a Chrimson variant, photostimulation of targeted neurons by red light only activates R-ChR, which causes photoactivation of these neurons. Blue light, however, activates R-ChR as well as B-ACR. This allows the chloride channel to induce shunting inhibition of the neurons which in turn prevents firing of the neurons by blue light (Fig. 1A [↗](#)).

### Validation of the co-expression strategy in *Drosophila*

As the first step for validating this approach, we took advantage of the well-studied male-specific P1 neurons in the fly central brain that elicit courtship singing upon activation (von Philipsborn *et al.*, 2011 [↗](#); Inagaki *et al.*, 2014 [↗](#); Ellenderson and von Philipsborn 2017 [↗](#); Mohammad *et al.*, 2017 [↗](#)) (Fig. 1B1-B2 [↗](#)). When Chrimson was co-expressed with an inactive control protein



**Figure 1.**

### Reducing blue-light mediated excitation by red-shifted channelrhodopsins-Proof of concept.

**(A)** Schematic representation of the approach. A red-shifted channelrhodopsin (R-ChR) is co-expressed with a blue-shifted anion channel (B-ACR). When red light is ON, there is an overall excitation of the cell due to the more dominant R-ChR response. When blue-light is ON, the shunting inhibitory effect of B-ACR reduces the excitation induced by R-ChR to blue light. **(B)** Validation of the approach in flies expressing Chromson, Chromson and GtACR2, or GtACR2 alone. **(B1)** Experimental setup to record the courtship song of solitary male flies. **(B2)** Top, Example of a reconstructed neuronal arborization of P1 neurons, Bottom, example of a courtship song induced by LED pulses. **(B3)** Courtship song production of the solitary male flies during 10-sec of constant illumination with 590 nm (amber) or 470 nm (blue) light. Each dot represents an individual (mean ± SEM)

(tdTomato), illumination at 470 nm and 590 nm wavelengths led to similar amounts of courtship song (respectively  $110.1 \pm 12.8$  and  $78.5 \pm 11.6$ ,  $n = 16-17$ ), demonstrating that blue light at the chosen intensity was efficiently activating Chrimson (**Fig 1B** [3](#) [4](#)). When Chrimson was co-expressed with the blue-sensitive anion channelrhodopsin GtACR2, 590 nm light illumination reliably elicited male courtship song ( $95.2 \pm 9.14$ ), however, 470 nm light illumination did not evoke any detectable pulse song ( $n = 17-18$ ). Neither 590 nm nor 470 nm light pulses induced courtship song in flies expressing only GtACR2 ( $n = 5$ ).

## Identifying a suitable pair of ultrafast light-gated anion channel and a red-shifted Chrimson variant

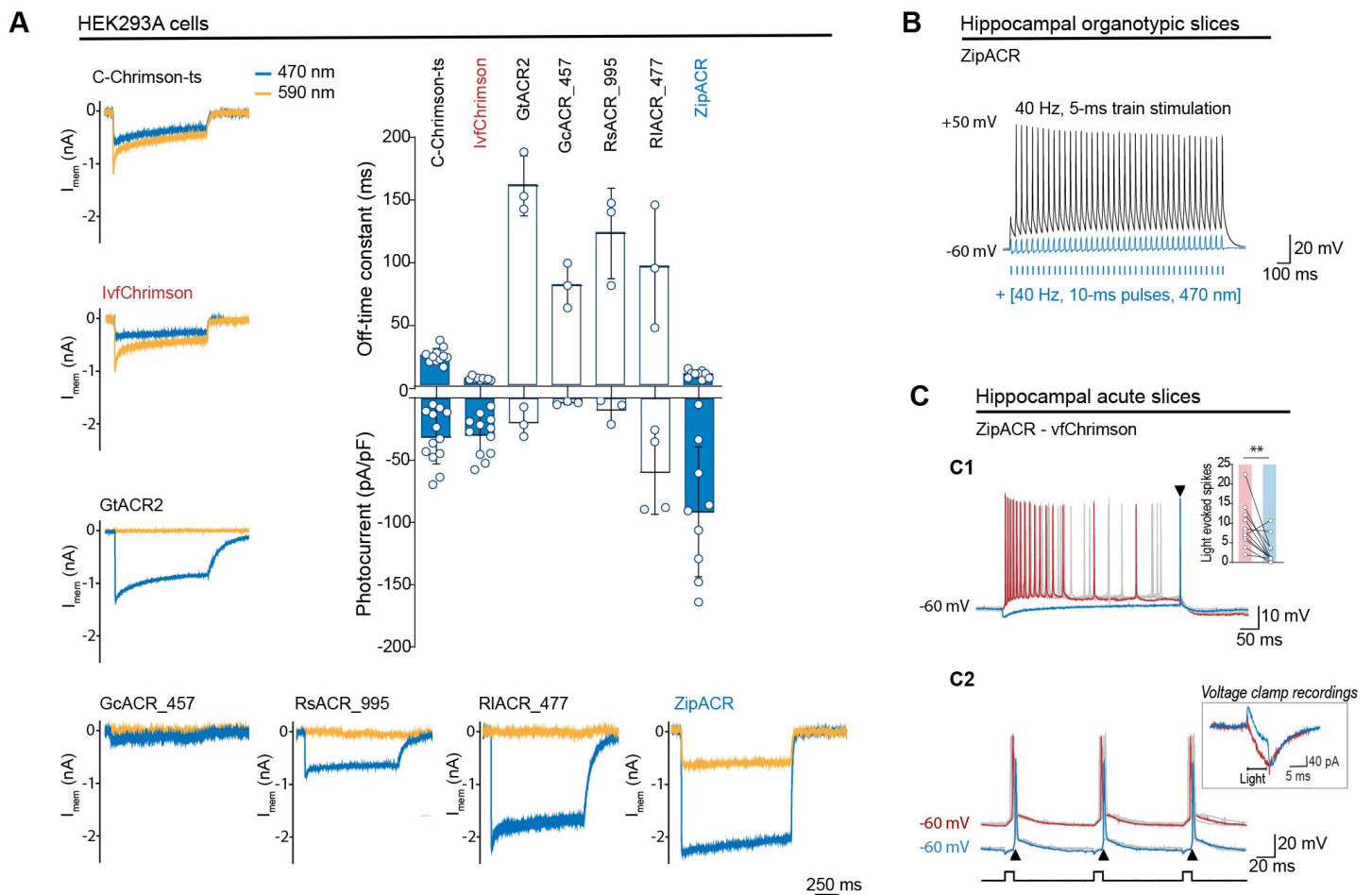
Although GtACR2 proved to be effective in protocols with pulses of long duration, its slow current decay (**Fig. 2A** [4](#), see GtACR2,  $161 \pm 24$  ms,  $n = 3$ ) limits its application to optical stimulations of lower frequencies (Govorunova *et al.*, 2015 [4](#); Govorunova *et al.*, 2017 [4](#)). For this reason, we searched for a pair of blue-shifted anion channelrhodopsin and red-shifted cation channelrhodopsin with fast and matching kinetics.

For a fast R-ChR, we evaluated the recently published vfChrimson, the fastest red-shifted channelrhodopsin available at this time, which can induce APs up to the intrinsic limit of the neurons (Mager *et al.*, 2018 [4](#)). Previously, we found that Chrimson and its variants traffic poorly to the membrane and developed a strategy to improve the trafficking (C-Chrimson-ts, Buonaventura *et al.*, 2019). We adapted this modification to improve the trafficking of vfChrimson (referred as IvfChrimson). In our measurements in HEK293 cells, IvfChrimson has a channel off-rate consistent with previously reported (Mager *et al.*, 2018 [4](#)) (off-rate time constant of  $5.6 \pm 0.3$  ms,  $n = 10$  compared to C-Chrimson-ts  $21.5 \pm 1.2$ ,  $n = 10$ ; mean  $\pm$  SEM, **Fig. 2A** [4](#), **Supplementary Fig. 2** [4](#)). When comparing C-Chrimson-ts and IvfChrimson photocurrents, we observe 1.68 fold reduction of normalized photocurrent of IvfChrimson ( $0.086 \pm 0.055$  pA/pF/A.U.,  $n = 25$ ) compared to C-Chrimson-ts ( $0.145 \pm 0.126$  pA/pF/A.U.,  $n = 25$ ; mean  $\pm$  SD, **Supplementary Fig. 2B** [4](#)) when the peak photocurrent amplitude was adjusted to membrane capacitance and membrane fluorescence.

To counteract the IvfChrimson excitation, we estimated that desired properties of the blue light activated anion ChR should have the off-rate time constant of  $\sim 15-30$  ms, minimal response to light of wavelength  $> 590$  nm and a strong photocurrent. We identified ZipACR, RsACR\_995, GcACR\_457, RIACR\_477 and GtACR2 as possible templates for further development based on published data (Govorunova *et al.*, 2017 [4](#)). GtACR2, RsACR\_995 and RIACR\_477 are spectrally ideal with negligible response to 590 nm light (**Supplementary Fig. 3A** [4](#)), but their kinetics are slow (mean off-rate of  $161.0 \pm 24.1$ ,  $122.7 \pm 36.3$  and  $95.8 \pm 49.4$  ms, respectively,  $n = 3$ , **Fig. 2A** [4](#)) which could have undesired long lasting inhibitory effects after light termination. GcACR\_457 has very small photocurrents and cannot be characterized accurately, possibly due to poor membrane trafficking. ZipACR, although spectrally not ideal ( $\sim 515$  nm; Govorunova *et al.*, 2017 [4](#)) has a kinetic close to the desired range ( $9.5 \pm 3.0$  ms,  $n = 9$ ) and it has strong photocurrents (**Fig. 2A** [4](#)).

To test the inhibitory property of ZipACR, we injected 40 Hz somatic current in organotypic hippocampal neurons expressing the channel. Consistent with the original report, individual pulses of 470 nm light resulted in the time-locked suppression of APs (**Fig. 2B** [4](#)).

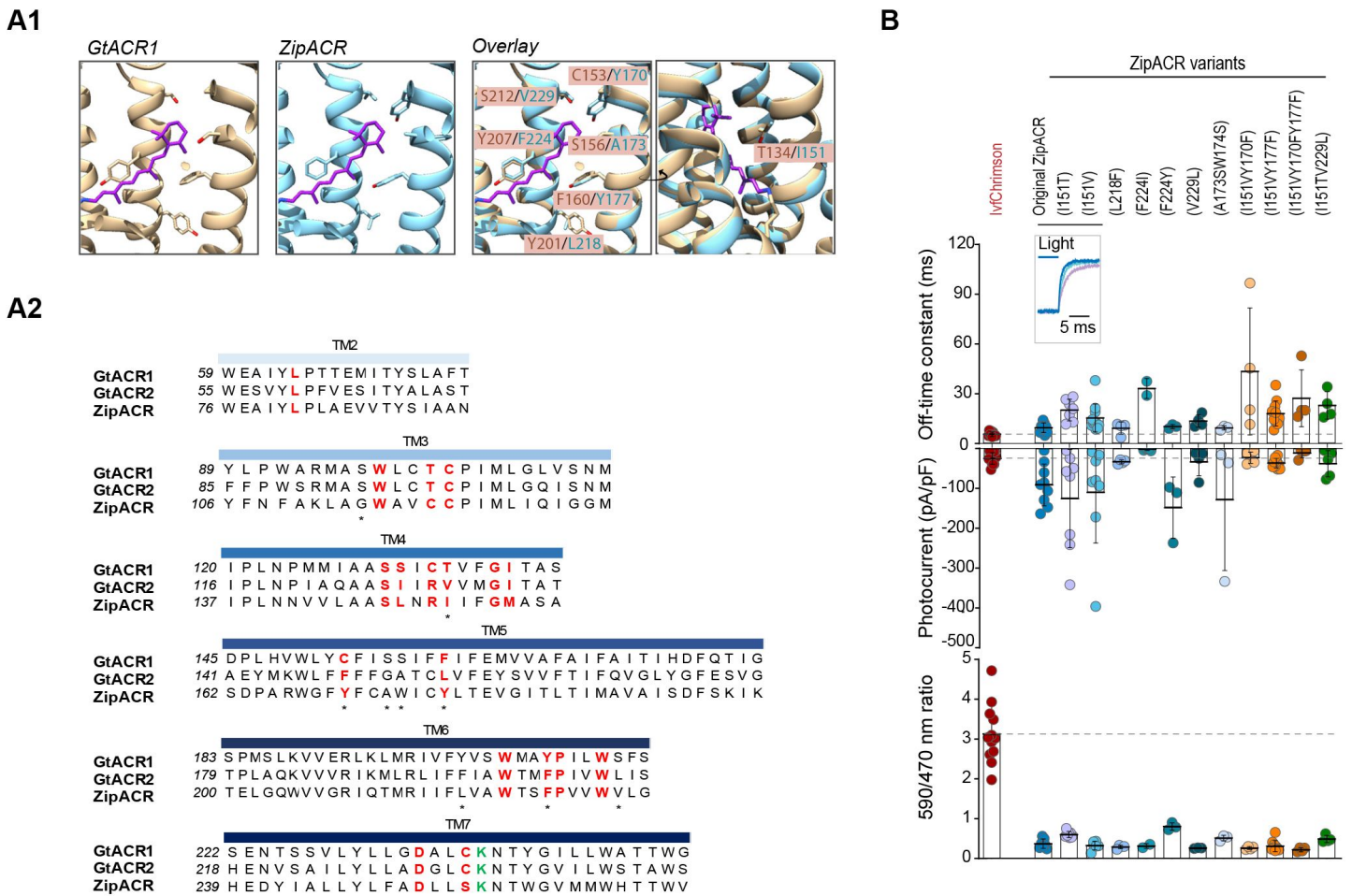
Next we examined whether ZipACR can block blue light-mediated APs in neurons expressing vfChrimson. Recordings from the granule cells in acute hippocampal slices co-expressing ZipACR and vfChrimson showed that, compared with 635 nm light, 470 nm light induced significantly fewer APs (**Fig. 2C** [1](#) [4](#)), in line with our prediction. However, we consistently observed the appearance of individual APs time-locked to the blue light offset (**Fig. 2C1-C2** [4](#)). We concluded



**Figure 2.**

### Comparison of channel properties of light-gated chloride channels and Chrimson variants.

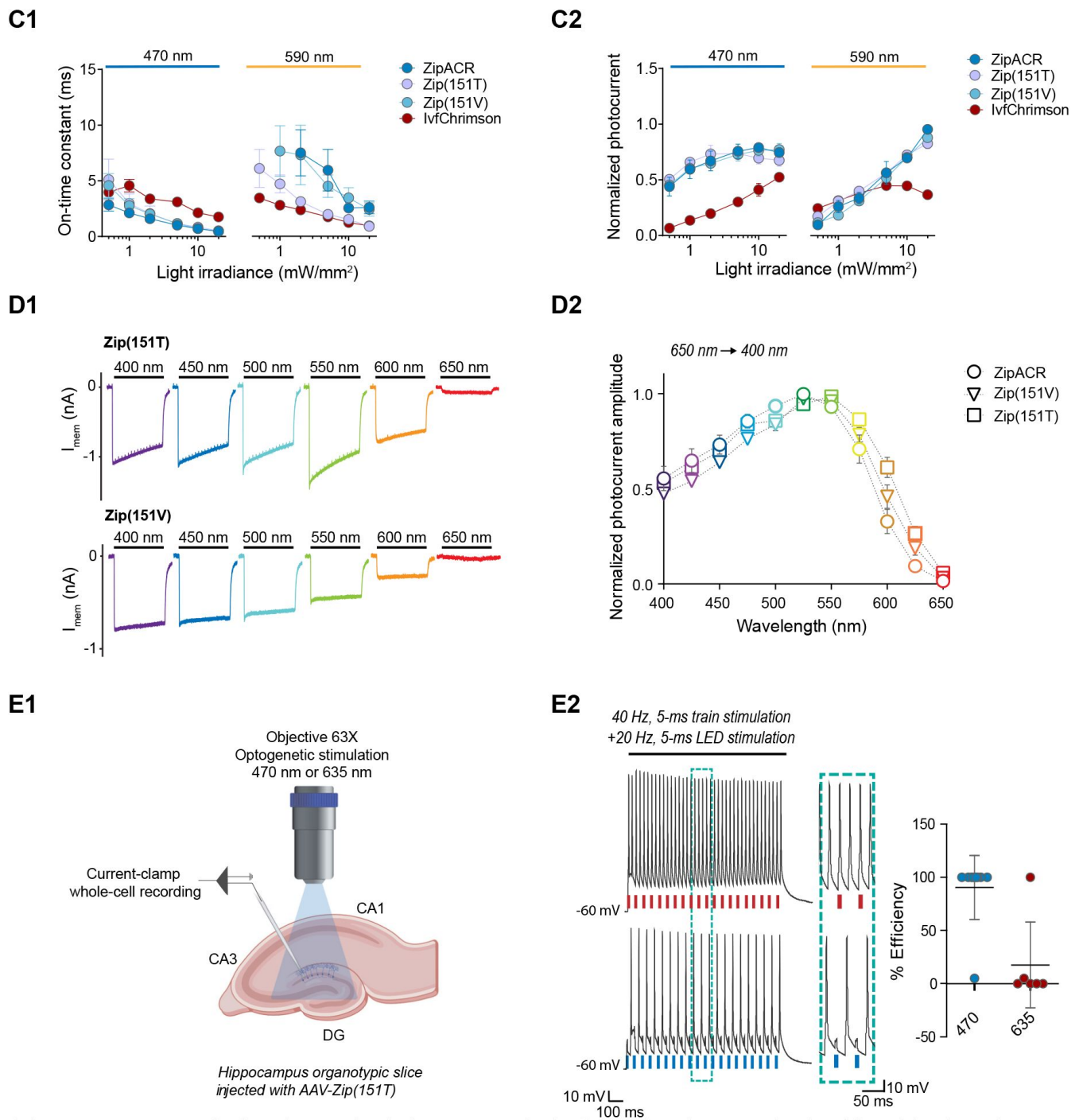
**(A)** Representative photocurrents of the Chrimson and ACRs variants in whole-cell patch-clamp recordings from HEK293 cells with 590 nm and 470 nm illumination ( $10\text{mW}/\text{mm}^2$ ). For Chrimson variants, a re-conditioning pulse at 405 nm illumination was applied 15 seconds before 470 nm LED stimulation. The plot shows the off-time constant (top) following 1 sec of 470 nm illumination, and the peak photocurrent (bottom) at -60 mV (mean  $\pm$  SD,  $n = 3-13$  cells). **(B)** Typical response of CA1 cells from hippocampal organotypic slices expressing ZipACR to 40 Hz, 5 ms pulses (top black trace) and to an illumination protocol using 40 Hz of 10 ms light pulses at 470 nm (bottom blue trace). **(C)** Recordings from dentate gyrus granule cells in acute slices expressing ZipACR and vfChrimson. **(C1)** Representative responses to 500 ms illumination at 470 nm and 635 nm (blue and red traces respectively). Note, 635 nm illumination evoked action potential firing while 470 nm illumination only evoked a single action potential time-locked to the end of the light pulse (arrowhead). Insert, plot showing the number of light-evoked spikes at 635 nm (red bar) and 470 nm (blue bar) illumination.  $n = 11$  cells, paired  $t$ -test  $**p < 0.01$ ; **(C2)** Example of light-induced firings by 635 nm and 470 nm illumination (red and blue traces respectively, 5 ms pulses at 10 Hz). The arrowheads show consistent APs time-locked to the end of the blue light pulse. Insert, example of a voltage-clamp recording of currents evoked by 5 ms pulses of 635 nm and 470 nm light (red and blue traces respectively). Light-induced currents with 470 nm illumination were initially outward but turned inward immediately following the light offset. Note the late inward current matches the red light induced current, reflecting a slower kinetics for vfChrimson compare to ZipACR.



**Figure 3.**

**Characterization of the optimized variants of the ultrafast anion channelrhodopsin ZipACR.**

(A) Structure-based design of ZipACR variants. (A1) The retinal binding pocket of GtACR1 and the putative retinal binding pocket of ZipACR homology model. (A2) The alignment of the transmembrane domains (TM) of GtACR1, GtACR2 and ZipACR. The residues surrounding the retinal binding pocket of GtACR1 and the corresponding residues from GtACR2 and ZipACR are highlighted in red. \*marks the residues targeted for mutation. The lysine that forms the Schiff base with retinal is indicated in green. The numbers on the left correspond to the number of the first residue in each line. (B) Basic properties of the ZipACR variants compared to Ivf-Chrimson. off-time constant (top) following 1 sec of 470 nm illumination, peak photocurrent (middle) and 590/470 ratio (bottom). The data have been obtained from whole-cell recordings in HEK293 cells. Each dot corresponds to one cell (mean  $\pm$  SD, n = 2-12).



**Figure 3.**

(C) On-time constant (C1) and normalized photocurrent (C2) of the selected variants Zip(151T) and Zip(151V) as compared to the original ZipACR and IvfChrimson, with 470 nm or 635 nm illumination of various light intensities (in mW/mm<sup>2</sup>). For the photocurrent, the values have been measured at steady-state. The data points are the mean value  $\pm$  SEM (n = 6-7). (D) The responses of the selected variants to various wavelengths of the same photon flux. (D1) The action spectra of ZipACR, Zip(151V) and Zip(151T) as measured from 650 nm to 400 nm (D2) For the action spectra, the maximum photocurrent amplitudes measured at each wavelength were normalized to the peak values obtained from the same cell across the spectrum. The data points are the mean value  $\pm$  SEM (n = 6-7). (E) Validation of the efficiency of the Zip(151T) variant in slices. (E1) The response of dentate gyrus granule cells expressing Zip(151T) variant at 470 nm and 635 nm light pulses delivered through a 63x objective. (E2) Representative traces of the firing induced by current injection at 40 Hz. Overlapping 470 nm but not 635 nm pulses block action potentials. The plot on the right shows the efficiency (in %) of Zip(151T) in blocking individual action potentials at 470 and 635 nm light pulses (mean  $\pm$  SD, n = 6-10 cells). The traces of the two outliers are presented in **supplementary Fig. 5A**.

that due to the fast closing rate, the photoinhibitory current generated by ZipACR decays too early to suppress the remaining excitatory response of vChrimson. Therefore, we decided to generate variants of ZipACR with slightly slower off-kinetics.

## Optimizing ZipACR off-kinetics

It is technically easier to generate mutations with slower kinetics and blue-shifted spectra than generating variants with faster on-and off-kinetics. Thus, we decided to use ZipACR as a template for further mutagenesis instead of further modifying GtACR2. To identify relevant mutation sites in ZipACR, we generated a homology model of ZipACR with the crystal structure of GtACR1 (Govorunova *et al.*, 2017) and identified the putative residues forming the retinal binding pocket in ZipACR (Fig. 3A). Multiple alignments were performed with the kinetically slower GtACR1 and GtACR2 to identify possible replacement of these residues that would result in slower kinetics and/or small blue-shift in ZipACR without disrupting the channel function (Fig. 3A2). We tested these variants in HEK293 cells for photocurrent ratio 590 nm / 470 nm LED stimulating light and measured the channel off-rate kinetics and photocurrent amplitude (Fig. 3B). Of these candidates, we identified I151V and I151T at the 4th transmembrane domain of ZipACR that resulted in a small increase in channel off-rate time constant ( $15.5 \pm 8.4$  ms for I151V and  $20.2 \pm 6.4$  ms for I151T, mean  $\pm$  SD, Fig. 3B) compared to the original ZipACR ( $9.5 \pm 2.9$  ms, mean  $\pm$  SD). We also generated double and triple mutants combining the I151V or I151T with Y170F, Y177F or V229L mutations but most of these double or triple mutants appeared to have smaller photocurrents and off-rate kinetics that could not be fitted well with single exponential back to zero baseline. Based on these results, we concluded the variants I151V (Zip(151V)) and I151T (Zip(151T)) are the best candidates for the pairing and proceeded for further characterizations.

Neither Zip(151V) nor Zip(151T) mutations changed the ion permeability of ZipACR as measured by channel reversal potential in potassium gluconate and cesium chloride based intracellular recording solutions (Supplementary Fig. 3B-C). In terms of light sensitivity, it is desirable that the ZipACR variant would respond to 470 nm light faster and stronger and to red-shifted light slower and weaker than IvfChrimson at the same intensity (Supplementary Fig. 4). With both Zip(151V) and Zip(151T), the channel on-rate kinetics have the desirable properties as described above to 470 nm and 590 nm light, although the Zip(151T) on-rate starts to approach the on-rate of IvfChrimson at light intensities  $>5$  mW/mm<sup>2</sup> at 590 nm (Fig. 3C1). With 470 nm light, IvfChrimson does not appear to reach saturating response even at 20 mW/mm<sup>2</sup> (Fig. 3C2). With 590 nm light, IvfChrimson reaches saturating response at 10 mW/mm<sup>2</sup> whereas none of the ZipACR variants appears to reach saturating response even at 20 mW/mm<sup>2</sup> (Fig. 3C2). Spectrally, Zip(151T) shows a wider red-shifted action spectra than ZipACR whereas Zip(151V) may have a slightly narrower blue-shifted action spectra than ZipACR (Fig. 3D).

Since Zip(151T) is a more red-shifted variant of the two (Fig. 3B,D), we asked whether red light pulses may induce photoinhibitory currents that suppress APs. We expressed Zip(151T) in the granule cells of the dentate gyrus in hippocampal slices using AAV vector and tested its inhibitory effect upon 470 nm and 635 nm light pulses. We injected somatic current at 40 Hz, while alternately delivering overlapping 5 ms pulses of light. Except in two instances, blue light pulses were consistently effective in blocking APs, while red light had no effect on APs induction (Fig. 3E,  $n = 10$  for 470 and  $n = 6$  for 635, the two outliers are presented in Supplementary Fig. 5A). It has been reported that at resting or hyperpolarized membrane potentials ZipACR may produce light-evoked APs (Kato *et al.*, 2018). We did not observe such a phenomenon under our experimental setup (Supplementary Fig. 5B).

## Validation of the co-expression strategy in brain slice

Since co-injection of two viruses results in variability in the ratio of expression in different neurons, we created constructs containing IvfChrimson and Zip(151T/V) (ZipT-IvfChr and ZipV-IvfChr) using the bicistronic 2A sequence (Fig. 4A). This ensures an equimolar expression of the



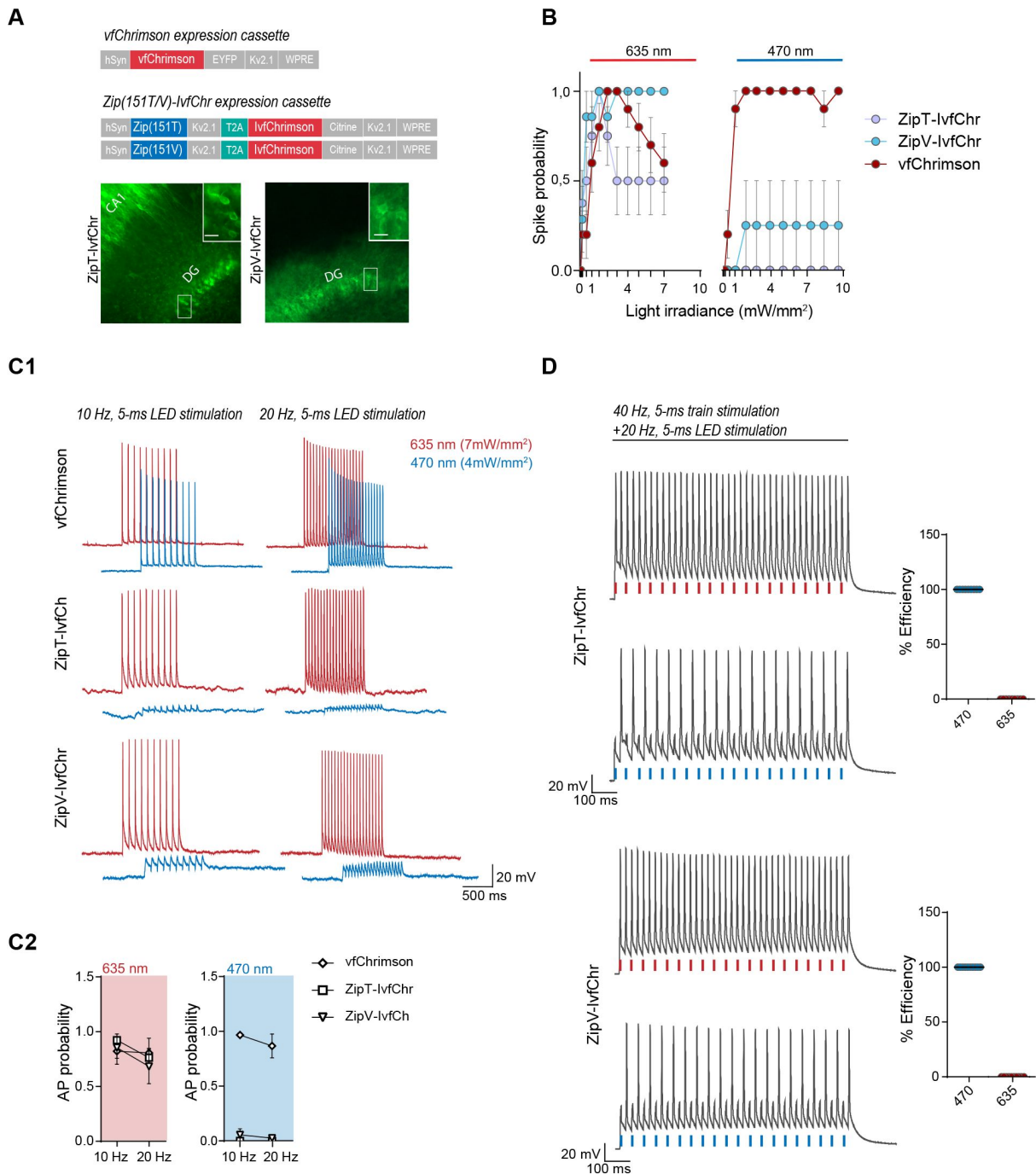
two channels. We also introduced a soma targeting Kv2.1 sequence into the constructs while eliminating the endoplasmic reticulum-exporting sequence of Ivf-Chrimson construct. Based on the photocurrent amplitudes of the constructs in response to 590 nm light (at -40mV holding potential with potassium gluconate-based intracellular solution to minimize Zip(151V) variant responses), the photocurrent amplitudes of IvfChrimson-only and ZipV-IvfChr constructs are comparable after adjustment to cell size and membrane fluorescence in HEK293 cells (**Supplementary Fig. 6** [↗](#)). We did not test the ZipT-IvfChr variant as the red-shifted response of Zip(151T) is difficult to eliminate to 590 nm light.

As the first step to test the performance of the system, we examined the power at which 470 nm and 635 nm lights induce APs in ZipT-IvfChr and ZipV-IvfChr, as well as in vfChrimson expressing neurons. We expressed AAV constructs in the granule cells of the dentate gyrus in the hippocampal slices (as described in **Fig. 3E1** [↗](#)). We recorded cells under the current-clamp mode and injected current through the patch pipette to maintain the membrane potentials at -60 mV. As expected, in vfChrimson expressing neurons 470 nm and 635 nm lights induced APs with light powers as low as  $0,5\text{mW}/\text{mm}^2$  ( $n = 10$ , **Fig. 4B** [↗](#)). In contrast, ZipT-IvfChr expressing neurons produced APs only to 635 nm light stimulation, whereas 470 nm light powers as high as  $10\text{mW}/\text{mm}^2$  failed to induce APs ( $n = 7-8$ ). As for ZipV-IvfChr, we occasionally observed blue light-induced APs ( $n = 4-7$ , **Fig. 4B** [↗](#)), which may be explained by its relatively faster decay time (**Fig. 3B** [↗](#)). In neurons expressing vfChrimson, we noticed a reduction in probability of AP upon successive optical stimulation specifically to 635 nm pulses (**Fig. 4B** [↗](#)). Consistent with this, we observed a reduction in response in HEK293 cells expressing IvfChrimson (**Supplementary Fig. 7A,B** [↗](#)). This phenomenon, likely caused by the desensitization of the channel, is consistent with previous reports (Lin *et al.*, 2013 [↗](#)).

To examine the efficiency of 635 nm and 470 nm lights in inducing high frequency APs, we delivered 10 Hz and 20 Hz light pulses of 5 ms duration (**Fig. 4C** [↗](#)). As expected, in vfChrimson expressing neurons both 470 nm and 635 nm lights induced APs ( $n = 8-9$ ). In neurons expressing either ZipT-IvfChr or ZipV-IvfChr, 635 nm pulses were equally effective in driving APs. On the other hand, 470 nm light only induced modest depolarization, which never reached the threshold for AP induction ( $n = 5-12$  for each opsin) (**Fig. 4C2** [↗](#)).

To test the temporal precision of the system in blocking AP, we stimulated the neurons by somatic current injection at 40 Hz, while alternately delivering perfectly overlapping 5 ms pulses of lights. Light pulses of 470 nm were consistently effective in blocking APs, while 635 nm light pulses had virtually no effect on APs induction as its amplitude, half-width and delayed time were indistinguishable from APs induced in the absence of light pulse (**Fig. 4D** [↗](#)).

Local mono-synaptic connection is a common feature in many brain regions. For this reason, independent dual optical activation requires that while optically stimulating pre-synaptic neurons, post-synaptic neurons remain responsive to the pre-synaptic inputs. To test whether our system meets this criterion, we asked how fast a neuron recovers from the inhibition induced by the ZipACR variants. We induced APs through somatic current injection at different time points from the offset of a light pulse. Blue light pulses blocked APs if the current was injected simultaneously. As expected from the off-rate of Zip(151T), within 5 ms from the light offset neurons expressing Zip(151T) or ZipT-IvfChr produced APs with high probability (**Supplementary Fig. 8** [↗](#)). The success rate reached its maximum within 10-15 ms. The recovery time for ZipV-IvfChr was immediate, with 100% success rate immediately following the light offset. This indicates that ZipV-Ivf-Chr could be used for dual optical activation without off target inhibition.



**Figure 4.**

**ZipACR variants co-expressed with the red Chr2 IvfChrimson prevent blue-light induced action potentials.**

(A) The expression cassette for vfChrimson and bicistronic Zip-IvfChr variants used in rAAV vector (Top). The selected ZipACR variants (Zip(151T) or Zip(151V)) are co-expressed with the red-sensitive IvfChrimson by using the 2A self-cleaving peptide (T2A). All the opsins are soma-targeted with a Kv2.1 peptide. (Bottom) Representatives showing the membrane enriched expression of ZipT-IvfChr and ZipV-IvfChr. Scale bar = 25μm. (B) Blue and red light-driven spike fidelity at various light intensities in DG cells (n = 4-10 cells for each opsin; 10 ms pulse width, single pulse). (C) Comparison of the high-frequency red and blue light-induced spiking (C1), and spike probability (C2, n = 6-13 for each opsin), of DG cells expressing vfChrimson or the Zip-IvfChr variants, with 10 and 20 Hz of light pulses (4-7mW/mm<sup>2</sup>). Holding potential = -60mV. (D) Optical inhibition of spiking in DG cells expressing the Zip-IvfChr variants. Representative traces of the firing induced by current injection at 40 Hz. Overlapping 470 nm but not 635 nm light pulses block action potentials. Holding potential = -60mV. The plots on the right show the efficiency (in %) of ZipT-IvfChr and ZipV-IvfChr in blocking individual action potentials at 470 and 635 nm (n = 9-10 for each opsin).

## Discussion

The main purpose of this work was to find a practical solution to the problem of cross-over where all opsins, regardless of their preferred excitation spectra, are activated by blue light. We describe a R-ChR2 expression system by which pulses of red light drive time-locked high frequency APs, while blue light remains ineffective. We created this system by pairing a B-ACR with the ultrafast red-shifted ChR2, IvfChrimson. This requires the B-ACR to have a significantly larger photocurrent compared to R-ChR2 as well as a faster channel opening and slightly slower channel closing. For this reason, we chose ZipACR as the template B-ACR for further modifications. Based on our homology modelling and multiple alignments with the kinetically slower B-ACRs, GtACR1 and GTACR2, we identified potential residue replacements within the putative retinal binding pocket which would result in slightly slower kinetics without impairing the channel function.

To our surprise, many mutations of ZipACR around all-trans retinal do not alter the channel properties or compromise channel conductance too strongly. This is contrary to our past experiences with channelrhodopsins where conductance is highly sensitive to mutations around the retinal binding pocket (data not shown). ZipACR double mutants appear to have more compromised conductance or altered properties. Among the tested positions, the I151 residue appears to have significant influences in kinetics and spectral properties. Indeed, we found that two variants, Zip(151T) and Zip(151V), sit in a sweet spot for our purpose. More specifically, the channels off-rate kinetics have the desirable property of being slightly but sufficiently slower than Ivf-Chrimson without compromising the opening rate, photocurrent and light spectrum. Thus, when paired with IvfChrimson, they prevent AP induction by blue light, without interfering with red light-induced high frequency neural spiking. On a similar note, although blue light pulses produced subthreshold depolarization, we did not observe charge integration that could have resulted in AP, even in frequencies as high as 20 Hz; this most likely reflects the reverse potential of the ZipACR. Of the two variants, we observed Zip(151T), and consequently ZipT-IvfChr, to be more consistent in blocking blue light-induced APs, which could be explained by its slightly slower closing rate. Although Zip(151T) is the more red-shifted of the two, within the tested range of light intensities and expression, only rarely did we observe a block of APs at 635 nm. However, we do not recommend the use of orange light pulses, as we observed a significant photocurrent in this wavelength.

Based on its impact on the kinetics and spectral properties, it will be of value to conduct saturating mutagenesis study on I151 residue that may produce variants of interesting properties or pairing that could outperform Zip(151V) and Zip(151T).

A recent study ([Vierock et al., 2020](#)) proposed a R-ChR2 expression system by relying on the pairing between GtACR2 and Chrimson, as we did in our early stage work with *Drosophila*. For physiological as well as experimental reasons, the priority we set was to create a system with fast closing kinetics. As channelrhodopsins are known to conduct calcium ions, long opening lifetime can evoke significant elevation of intracellular calcium ([Lin et al., 2009](#)) and activate second messenger pathways with complicating biological effects in experiments of long duration. The same concern applies to B-ACRs with slow closing kinetics as they could have additional inhibitory effects after the termination of blue light and alter the physiology to be studied. From an experimental viewpoint, a fast photostimulation system has the flexibility to perform a wide range of optical manipulations. This includes driving neuronal activity to their firing limits as well as suppressing their firing with single spike precision. Although not tested here, our results indicate that the fast closing kinetics of the B-ACR channels, particularly ZipV-IvfChr, should allow dual optical activation of monosynaptically connected neurons.

Before concluding, we must note that our approach, which runs based on chloride influx is unlikely to be suitable for axonal terminal activation, where the chloride concentration is high ([Mahn et al., 2016](#); [Mahn et al., 2018](#)). For this matter, we refer the readers to some published

methods by which independent dual channel photostimulation of the axons might be feasible (Klapoetke *et al.*, 2014 [↗](#); Hook *et al.*, 2015 [↗](#)). Thus, our system complements these studies as they are unsuitable for somatic photoactivation. This provides researchers with complementary methods to choose from based on their experimental needs.

## Methods

### Drosophila optogenetic activation experiments

Flies (*Drosophila melanogaster*) were kept on standard food medium (water, cornmeal, oatmeal, sucrose, yeast, agar, acetic acid, preservative methyl-4-hydroxybenzoate) at 25°C, 60% humidity, and a 12h/12h light-dark cycle. Male flies were collected after eclosion and aged for 7 days in isolation and protected from light on standard food, followed by 3 days on food containing 400 µM all-trans retinal (Sigma). The following genotypes were assayed: co-expression of Chrimson and GtACR2: w-; NP2631.GAL4/ UAS-CsChrimson.mVenus.attp40; tubP-FRT-stop-FRT-GAL80, fruFLP/ UAS-GtACR2.attp2; co-expression of Chrimson and td-Tomato: w-; NP2631.GAL4/ UAS-CsChrimson.mVenus.attp40; tubP-FRT-stop-FRT-GAL80, fruFLP/ UAS-myr-tdTomato. attp2; co-expression of GtACR2 and tdTomato: w-; NP2631. GAL4/ UAS-myr-tdTomato.attp2; tubP-FRT-stop-FRT-GAL80, fruFLP/ UAS-GtACR2.attp2.

For optogenetic activation experiments, we used a previously described multi-channel array of electret condenser microphones (CMP-5247TF-K, CUI Inc), amplified with a custom-made circuit board and digitized with a multifunction data acquisition device (NI USB-6259 MASS Term, National Instruments) (von Philipsborn *et al.*, 2014 [↗](#)). Single male flies were placed in recording chambers equipped with microphones at the bottom and light diodes (Amber 591 nm: Cree 5-mm C503B-AAS/AAN-015 or Blue 470 nm: Cree 5-mm C503B-BAS-CY0C0461) were placed 1 cm above the song recording chambers at a 60° angle, illuminating the whole chamber (10-sec of constant light). Tentative pulse song was detected by a MATLAB script and corrected manually, using a custom made user interface for visualization and annotation of song oscillograms (von Philipsborn *et al.*, 2014 [↗](#)).

### Cell culture, transfection and stable cell line generation

HEK293 cells (Invitrogen) were grown in DMEM (1g/L glucose) supplemented with 8% FBS and 1% Pen/Strep. For recordings, cells were plated at low density on glass coverslips and kept in DMEM media supplemented with 4% FBS and 1% Pen/Strep. Transfection was done with Xtremegene 9 (Sigma-Aldrich) according to the manufacturer's instruction. Recordings of the cells were performed between 38-50 hours after transfection.

For generation of recombinant lentivirus for stable HEK293 cell lines expressing IvfChrimson and ChrimsonR-tdTomato cells, HEK293 cells were co-transfected with psPAX2 and pMD2.G (courtesy of Trono laboratory) and the media was transferred to a new dish with untreated cells 2 and 3 days after transfection. The cells receiving the media were kept as stable cell lines of respective constructs.

### Structure alignment and modelling

Homology modelling of ZipACR structure was performed with Modeller within the UCSF Chimera to the published GtACR1 structure (PDB #6CSM). This was further aligned to the GtACR2 alignment from Govorunova *et al.*, (2017) [↗](#). Residues surrounding the retinal binding pocket of GtACR1 and the modelled ZipACR were identified and those residues that were different in ZipACR were changed to the corresponding residues of GtACR2 or GtACR1.

## Molecular biology

For GcACR\_457, RsACR\_995 and RlACR\_477, the cDNA were in the original pcDNA3.1 vector as EYFP fusion as available on Addgene (#103137, 103772, 103771). The GtACR2 construct was synthesized (IDT DNA) and modified with a N-terminal signalling peptide from the commercial pDisplay vector (Invitrogen) and a trafficking signal prior to tdTomato or citrine fusion and inserted in the pcDNA3.1+ vector (Invitrogen) for expression. For the ZipACR constructs, the original EYFP fusion from Addgene (#88844) was retained and the construct was either inserted into the pcDNA3.1+ vector or a second generation lentiviral vector pLenti with CMV promoter. No functional or fluorescent difference in the expressed protein were observed between expression from pcDNA3.1+ and pLenti vector in HEK293 cells and the results were combined for comparison and analysis.

Site-directed mutagenesis of the ZipACR, Chrimson and GtACR2 were performed with overlap extension PCR using Phusion DNA Polymerase (Life Technologies) and ligated into the pcDNA3.1+ or pLenti expression vector.

For ChrimsonR and vfChrimson, additional improvements for membrane trafficking was used as previously described (Buonaventura *et al.*, 2019). In brief, N-terminal leading sequence from ChR1/oChIEF/ReaChR was transplanted to the transmembrane A-G of ChrimsonR and vfChrimson. Trafficking sequence was added after the transmembrane G before fusion to tdTomato. The modified ChrimsonR/vfChrimson-tdTomato constructs were inserted into the 2nd generation lentiviral pLenti vector with CMV promoter and used to make stable ChrimsonR/vfChrimson-tdTomato HEK293 cell lines. The improved vf-Chrimson is named IvfChrimson.

For the comparison of IvfChrimson-citrine and Zip(151V)-Kv2.1-2A-IvfChrimson-citrine-Kv2.1 (ZipV-IvfChr) photocurrents, the two cDNAs were inserted into pcDNA3.1+ for expression. The IvfChrimson in both designs contains the modified N-terminal sequence from ChR1/oChIEF/ReaChR but the trafficking sequence was only used in the IvfChrimson-citrine design and not the Kv2.1 design.

## rAAV production

rAAV2-retro containing vfChrimson\_EYFP\_Kv2.1, Zip(151T)-Kv2.1\_IRES\_DsRed, Zip(151T)-Kv2.1-2A-IvfChrimson-citrine-Kv2.1 (ZipT-IvfChr), and Zip(151V)-Kv2.1-2A-IvfChrimson-citrine-Kv2.1 (ZipV-IvfChr) were produced according to the protocol at <https://www.salk.edu/science/core-facilities/viral-vector-core/resources/>. In brief, HEK293A cells (Life Technologies, Carlsbad, CA) were grown to 90% confluency and transfected with vectors containing opsin proteins, and the helper plasmids XX6-80 and rAAV2-retro (gift from A. Y. Karpova, Howard Hughes Medical Institute Janelia Farm Research Campus). rAAV2-retro were released and purified according to protocol at [https://www.takarabio.com/documents/User%20Manual/6232/6232\\_e.v1509Da.pdf](https://www.takarabio.com/documents/User%20Manual/6232/6232_e.v1509Da.pdf).

## Electrophysiological characterization and imaging of HEK293 cells

Whole-cell patch-clamping was performed with Axopatch 200A patch-clamp amplifier with Digidata 1322A data acquisition board and pCLAMP10 software. Cells were clamped at -60mV except in experiments with the ramp protocol and experiments comparing the 590 nm light-induced photocurrents between Ivf-Chrimson-citrine and ZipV-IvfChr expressing cells. In the experiment comparing the IvfChrimson-citrine photocurrent to the 2A configuration, cells were clamped at -40mV to reduce the influence of ZipACR response with IvfChrimson photocurrent. Series resistance for the recorded cells were mostly under 15 M $\Omega$  and series resistance were compensated up to 70% during recordings where possible.

The extracellular solution was composed of (in mM): 145 NaCl, 3 KCl, 2 CaCl<sub>2</sub>, 1 MgCl<sub>2</sub>, 20 D-glucose and 10 HEPES (pH 7.35). For most experiments, the intracellular solution contained (in mM): 110 CsCl, 15 TEACL, 10 HEPES, 5 Cs<sub>2</sub>EGTA, 1 MgCl<sub>2</sub> and 0.1mM CaCl<sub>2</sub> (pH 7.25). In

experiments comparing the 590 nm light-induced photocurrents between IvfChrimson-citrine and ZipV-IvfChr expressing cells and some ramp experiments, a K-gluconate based intracellular solution containing: 115 K-gluconate, 10 K2EGTA, 10 HEPES, 5 NaCl, 10 KCl, 2 MgATP and 0.1 Tris-GTP (pH 7.25), was used.

Channelrhodopsin-expressing cells were imaged and recorded on an Olympus IX73 microscope with a 40x air objective equipped with Photometrics Cascade II 512 EMCCD camera. Images were acquired using Micromanager 1.4 with no gain and no electron multiplication and exposure time of 250 ms. Excitation light was provided by a Thorlabs 4 channel LED (LED4D118 with 405, 470, 530 and 590 nm channels and only 470 nm and 530 nm light were used for imaging with LED driving currents of 150 and 80 mA, respectively). Citrine and EYFP expression were imaged with a GFP filter cube (Semrock, FF02-470/30 excitation, FF01-520/35 emission and FF495-Di03 dichroic) and tdTomato was imaged with a TRITC filter cube (Semrock, FF01-543/22 excitation, FF01-593/40 emission and FF562-Di03 dichroic).

For most channelrhodopsin stimulation experiments 10 mW/mm<sup>2</sup> of 470 nm and 590 nm stimulation lights from Thorlabs 4 channel LED light source (LED4D118) were delivered to the specimen plane. The light intensity was controlled with a Thorlabs DC4104 4 channel controller electronically by the separate analogue outputs of a HEKA ITC-18 DAQ board under the control of WinWCP 5.5 software. In the spectral response experiment, a Sutter VF-5 system with a smart shutter in the neutral density mode was used to achieve cross spectrum stimulation at the same photon flux at 1.586 x 10<sup>15</sup> photons / s / mm<sup>2</sup> (25.1 mW/mm<sup>2</sup> at 400 nm and 15.4 mW/mm<sup>2</sup> at 650 nm) between 400-650 nm (25 nm increments). The VF-5 has the following filters and bandwidth settings: 451-398 nm with 15 nm bandwidth, 503-446 nm with 15 nm bandwidth, 564-498 nm with 14 nm band-width, 632-557 nm with 14 nm bandwidth, 703-627 nm with 13 nm bandwidth. A second Sutter shutter was placed in the light path before the light guide to control the light output. For experiments with ACRs, a 15-sec interval was used between stimulation episodes. For ChrimsonR/IvfChrimson stimulation, stimulation light pulses either have 15-sec intervals between stimulation episodes (no preconditioning) or a 15-sec interval -1-sec 405 nm light pulse – 15-sec interval stimulation pattern for preconditioning before the subsequent stimulation. The stimulation light was guided into the fluorescent excitation light path of the Olympus IX73 microscope as with imaging but reflected with a Semrock FF685-DiO2 dichroic mirror to the objective.

Light intensity (in mW/mm<sup>2</sup>) was measured with a Thorlabs slide based light detector (S170C) at the specimen plane with illumination area defined by the microscope aperture and manually calculated for photon flux or illumination intensity.

For comparison of the membrane expression of our Ivf-Chrimson-citrine and '2A', transfected cells were imaged on an Olympus BX51 microscope with a 40x Olympus water immersion objective, X-Cite 110LED and Hamamatsu Orca Flash4. The images were taken at 15% light intensity and 250ms exposure time with FF02-472/30 excitation filter, FF01-520/35 emission filter and FF495-Di03 dichroic mirror (Semrock, NY). A line profile was drawn across the cell and the membrane values were averaged and used for calculating the ratio with the mean cytosolic values.

## Organotypic slices and virus infection

All procedures involving animals were approved by the Danish Animal Experiment Inspectorate. Hippocampi were isolated from postnatal P5-6 wild-type Sprague-Dawley rats (Janvier). The pups were decapitated and the brain removed and kept in an icy low sodium ACSF solution composed (in mM) of 1 CaCl<sub>2</sub>, 4 KCl, 1 MgCl<sub>2</sub>, 26 NaHCO<sub>3</sub>, 10 D-Glucose, 234 Sucrose, and 0.1% Phenol Red Solution, in mQ H<sub>2</sub>O, and bubbled with a mixture of CO<sub>2</sub> (5%) and O<sub>2</sub> (95%). From that moment, all the steps were performed in a laminar flow tissue culture hood using sterile equipment and in total asepsie. The brain was poured into a petri dish filled with the low sodium ACSF solution for hippocampi dissection under microscope guidance (Stereomicroscope, Olympus x1 objective).

After extraction, the hippocampi were sliced at 400µm thickness using an automatic tissue chopper (McIL-WAIN) and moved to a dish with pre-heated culture medium containing MEM Eagle medium 78.8% (Gibco), 20% heat-inactivated horse serum (Gibco), 1 mM L-glutamine, 1 mM CaCl<sub>2</sub>, 2mM MgSO<sub>4</sub>, 170 nM insulin, 0.0012% ascorbic acid, 12.9 mM D-glucose, 5.2 mM NaHCO<sub>3</sub>, 300 mM HEPES (Sigma), pH = 7.2-7.3, osmolarity adjusted to 315–325. The slices with intact DG and CA regions were then transferred onto air-fluid interface-style Milli-cell culture inserts (Millipore) in 6-well culture plates (ThermoFisherScientific) with 800 µL of sterile medium added below each insert. The slices were kept in a sterile incubator at 37 degrees with 5% CO<sub>2</sub> (ThermoScientific, Steri-cycle i-160). The medium was replaced by a pre-warmed medium (37 degrees) three times a week.

After two to three days of culture, the slices were microinjected in DG (or in CA1 when specified) along the characteristic horseshoe pattern with a pulled glass pipette containing the following viruses: retro-vfChrimson (titer:  $7 \times 10^{12}$  particles/mL), retro-Zip(151T) (titer:  $9 \times 10^{12}$  particles/mL), retro-ZipT-IvfChr (titer:  $2 \times 10^{12}$  particles/mL), or retro-ZipV-IvfChr (titer:  $1.5 \times 10^{12}$  particles/mL). The injections were done under a microscope (using a Picospitzer III (Parker) connected to a Pulse Pal (10 ms pulse, every 0.5-sec).

### Whole-cell electrophysiology and light delivery in-vitro

Two to three weeks post-infection, the organotypic slices were transferred into the recording chamber and continuously perfused with Artificial Cerebrospinal Fluid containing (in mM): 119 NaCl, 2.5 KCl, 26 NaHCO<sub>3</sub>, 1 NaH<sub>2</sub>PO<sub>4</sub>, supplemented with 11 D-glucose, 4 CaCl<sub>2</sub>, 4 MgCl<sub>2</sub>. Additionally, APV (50mM), NBQX (100mM) and picrotoxin (100mM) were added to block excitatory and inhibitory fast transmission. The solution was adjusted to pH 7.4 (osm ~330) and bubbled with a mixture of CO<sub>2</sub> (5%) and O<sub>2</sub> (95%), at room temperature. The chamber was mounted on an upright microscope (Scientifica) linked to a digital camera (QImaging Exi Aqua) and the cells were visualized using 60X water-immersion objective (Olympus, LumiPlan). Acquisitions were performed in whole-cell configuration using Clampex 10.6 connected to a Multiclamp 700B amplifier via a Digidata 1550A digitizer (all from Molecular Devices). Voltage-clamp data were low-pass filtered at 200 Hz and digitized at 10 kHz and the whole-cell capacitance was compensated. Patch pipettes (2-4 MΩ of resistance) were filled with an internal solution containing (in mM): 153 K-gluconate, 10 HEPES, 4.5 NaCl, 9 KCl, 0.6 EGTA, 2 MgATP, and 0.3 NaGTP. The pH and osmolarity of the internal solution were close to physiological conditions (pH 7.4, osmolarity 297 mOsm). The access resistance of the cells in our sample was ~ 25 MOhm. For voltage-clamp experiment, the Dentate Gyrus granule cells (otherwise specified when done in CA1 pyramidal cells) were clamped at -60mV. For current-clamp experiments, the cells were maintained at -60mV (except for the protocol where the potential was brought from -70 to -40mV) by constant current injection (holding current ± 100pA).

The recorded neurons were illuminated with 470 nm and 635 nm (nominal wavelengths) light with a coolLED pE-4000 system connected to the Digidata via a single TTL. The light irradiance was controlled via the control pod and was adjusted to 4mW/ mm<sup>2</sup> for 470 nm and 7 mW/mm<sup>2</sup> for 635 nm (otherwise specified). Light irradiance was measured with a Thorlabs digital optical power meter (PM100D) at the specimen plane with illumination area defined by the microscope aperture. Between the recordings, wavelengths were switched from 470 nm to 635 nm and vice versa. At the end of the recordings, the location of the recorded cell was confirmed by inspection at 10x magnification. Data analysis was performed using Clampfit 10.6 (Molecular device).

After recordings, slices were kept in 4% formalin, and later mounted under coverslips with Fluoromount (Sigma-Aldrich). Pictures were taken with an Apotome.2 (Zeiss) and ZEN software at 10X and 40X magnifications.

## Acute slices

Experimental procedures were approved by the Animal Care and Use Committee of the University of Buenos Aires (CICUAL). Briefly, 4-week old C57 mice ( $n = 4$ ) were co-injected with AAV-8/2-hSyn1-ChrimsonVF\_EYFP\_Kv2.1 and AAV-8/2-hSyn1-ZipACR\_Kv2.1\_IRES\_DsRed2 within the dentate gyrus of the hippocampus (250nl of a 1:1 mixture, coordinates from Bregma: 1 mm mediolateral, -2 mm anteroposterior, -1.5 dorsoventral). After 3 weeks of expression, the animals were sacrificed and the brain removed and cut in a solution composed of (in mM): 234 sucrose, 11 glucose, 26 NaHCO<sub>3</sub>, 2.5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 10 MgSO<sub>4</sub>, and 0.5 CaCl<sub>2</sub> (equilibrated with 95% O<sub>2</sub>–5% CO<sub>2</sub>). The slices were maintained at room temperature before being transferred in a recording chamber mounted on a microscope (Nikon) connected to a Mightex Illumination system for 470 nm and 635 nm light delivery (at 10mW). For current clamp experiments the amount of current injected was corrected in the inter-sweep interval to keep it close to -60 mV. In voltage clamp experiments holding potential was -60 mV.

Recordings were done in a solution of composition (in mM): 126 NaCl, 26 NaHCO<sub>3</sub>, 2.5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 2 MgSO<sub>4</sub>, 2 CaCl<sub>2</sub> and 10 Glucose (pH 7.4).

Patch-pipettes (2-4 M $\Omega$  of resistance) were filled with a K-gluconate based internal solution (in mM, 130 K-gluconate, 5 KCl, 10 HEPES, 0.6 EGTA, 2.5 MgCl<sub>2</sub>·6H<sub>2</sub>O, 10 Phosphocreatine, 4 ATP-Mg, 0.4 GTP-Na<sub>3</sub>).

## Data Analysis

The data were analysed and plotted using GraphPad Prism V9 (GraphPad Software, La Jolla, CA, USA). All values are indicated as mean  $\pm$  standard deviation (SD) or mean  $\pm$  standard error of the mean (SEM), as specified. For statistical comparison between groups, data were tested for normal distribution using D'Agostino and Pearson, Shapiro-Wilk or KS normality test. Comparison of the mean between groups has been done with a one-way analysis of variance (ANOVA) followed by Dunnett's multiple comparisons test.

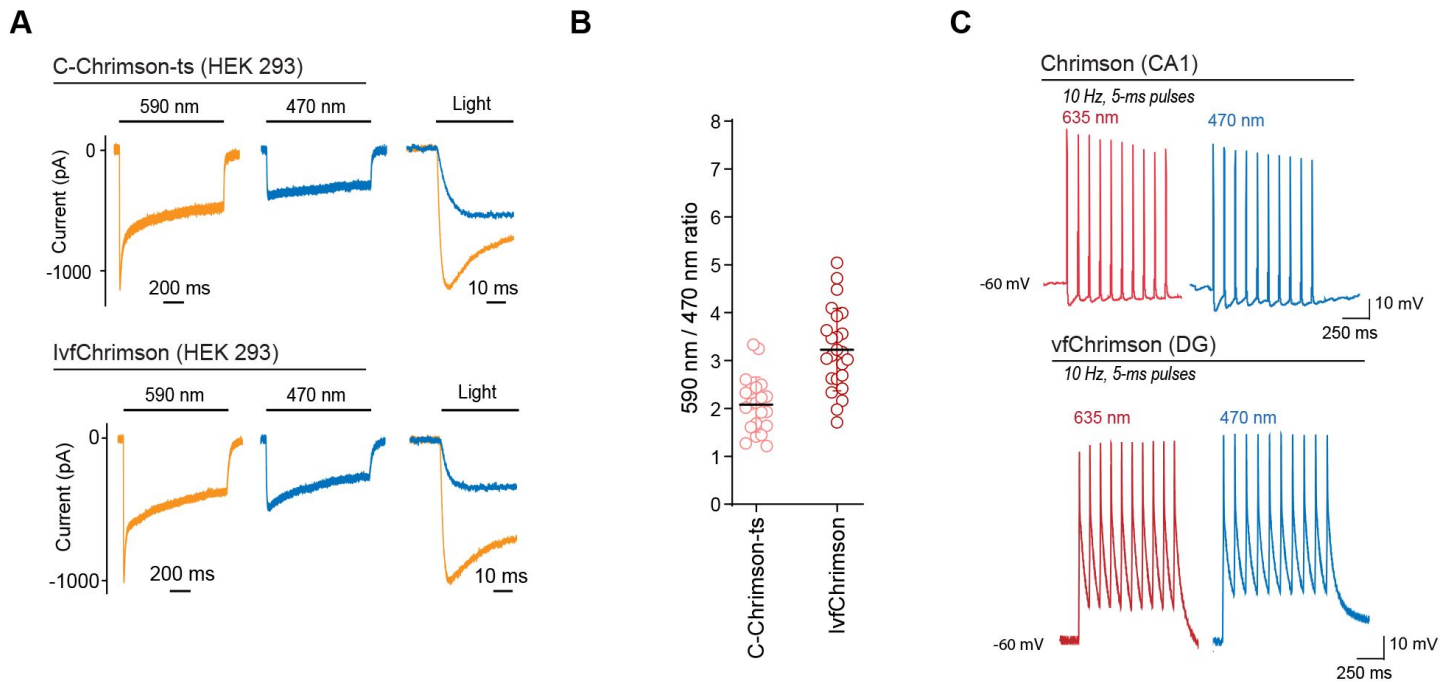
ZEN software (Zeiss) and FIJI (from ImageJ) were used for images processing.

Figure 1A, 1B1 [↗](#) and 3E1 [↗](#) have been created with BioRender.com.

## Acknowledgements

We thank the members of the Nabavi lab, in particular Islam Faress for suggestions, and Mariam Gamaleldin for the preparation of hippocampi organotypic slices. We also thank Dr. Sergio Almeida for his help on HEK293 cells culture at the early stage of the project, Peter Kerwin from the Philipsborn lab for his help on Drosophila work, and Anne-Katrine Vestergaard and Kathrine Meinecke Christensen for technical assistance. We are thankful to Dr. Thomas Mager and Dr. Ernst Bamberg for their generous gift of vfChrimson DNA vector. This study was supported by an ERC starting grant to S.N (22736), by the Danish Research Institute of Translational Neuroscience to S.N (19958), by PROMEMO (Center of Excellence for Proteins in Memory funded by the Danish National Research Foundation) to S.N (DNRF133), and by a Lundbeck NIH Brain Initiative grant to S. N and JY. L.

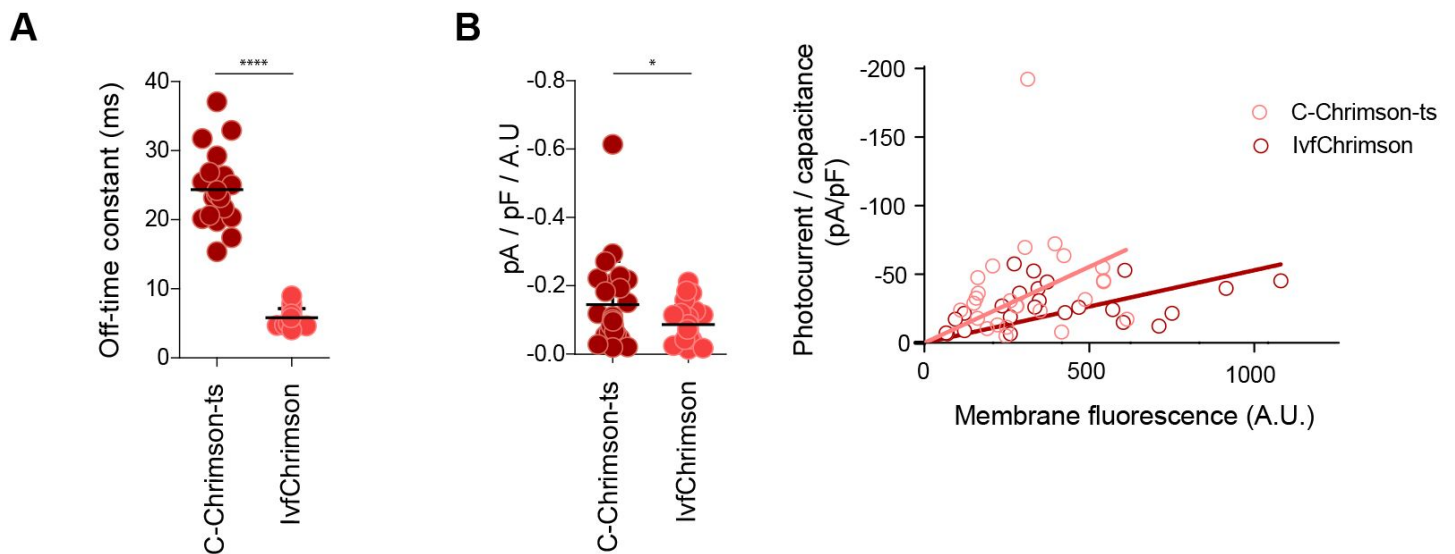




### Supplementary figure 1.

#### Red-shifted Chr2 variants are activated by 470 nm and 590 nm pulses of light.

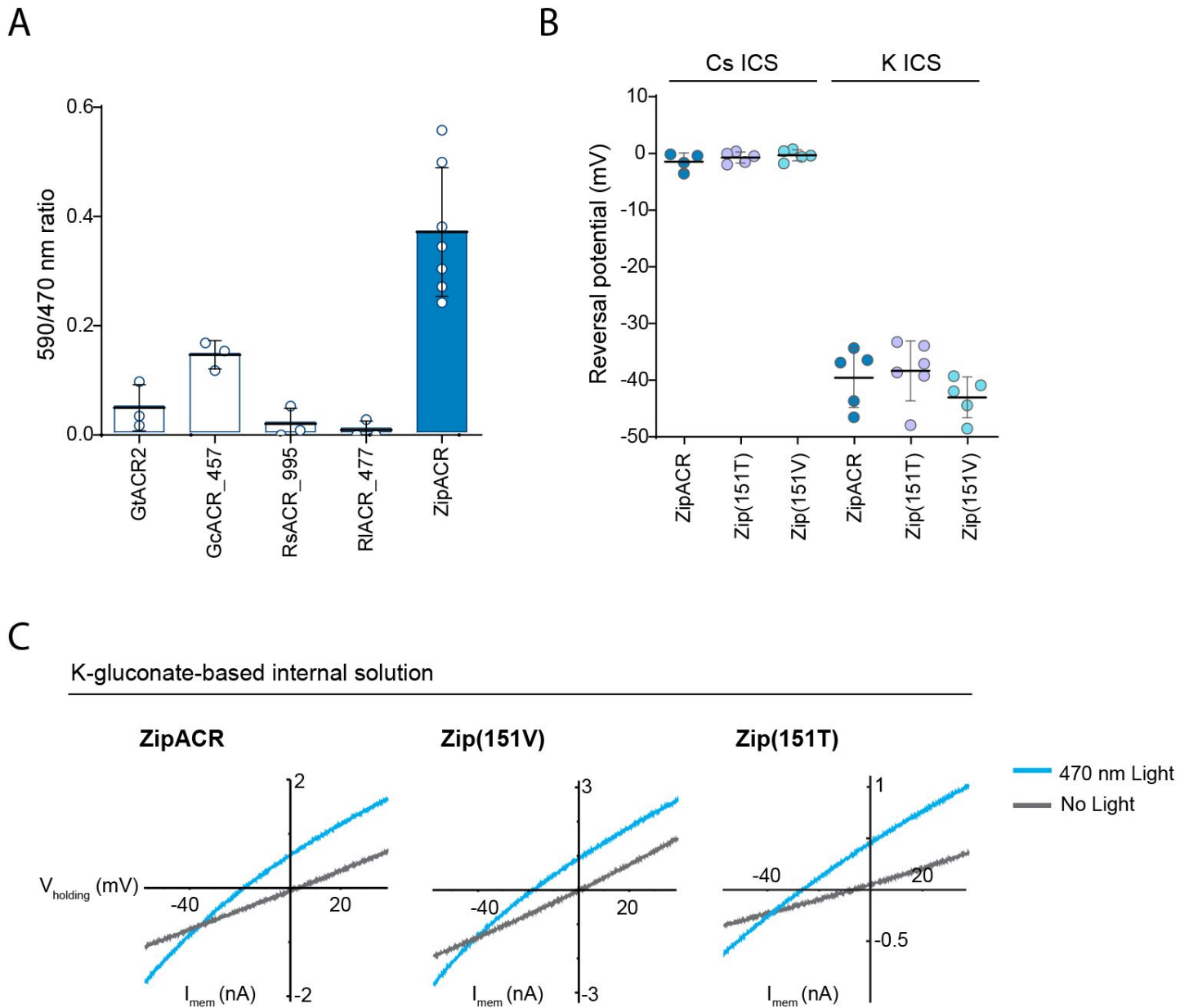
**(A)** Typical photocurrents (left and middle traces) and on-kinetics (right traces) of Chrimson-ts and IvfChrimson in HEK293 cells with 590 nm and 470 nm illumination at 10 mW/mm<sup>2</sup>. The traces have been recorded 15-s after a 405 nm reconditioning pulse. **(B)** Comparison of 590/470 nm ratio for Chrimson-ts and IvfChrimson, measured at peak. Each dot represents a cell (mean  $\pm$  SD). **(C)** Blue-light illumination of the Chrimson variants induces firing as efficient as red-light stimulation. This property is represented here with two examples of traces obtained from Chrimson expressing CA1 neurons and vfChrimson expressing DG neurons, in response to 10 Hz of 5 ms pulses at 635 nm and 470 nm.



### Supplementary figure 2.

#### Membrane trafficking of the improved ultrafast vf-Chrimson, IvfChrimson.

**(A)** Off-time constant of C-Chrimson-ts and IvfChrimson after 1-s of 635 nm illumination at 10mW/mm<sup>2</sup>. **(B)** Photocurrent amplitudes at the peak of C-Chrimson-ts and IvfChrimson, adjusted to membrane fluorescence (left), and comparison of current density and membrane fluorescence (right). The recordings have been obtained from HEK293 cells at -60mV. Each dot represents the values from a single cell. The horizontal black bars represent the mean  $\pm$  SEM for (A) and mean  $\pm$  SD for (B). Statistics for panels (A) \*\*\*\* $p < 0.0001$ ; unpaired t-test and (B) \* $p < 0.05$ ; unpaired t-test (Welch correction).

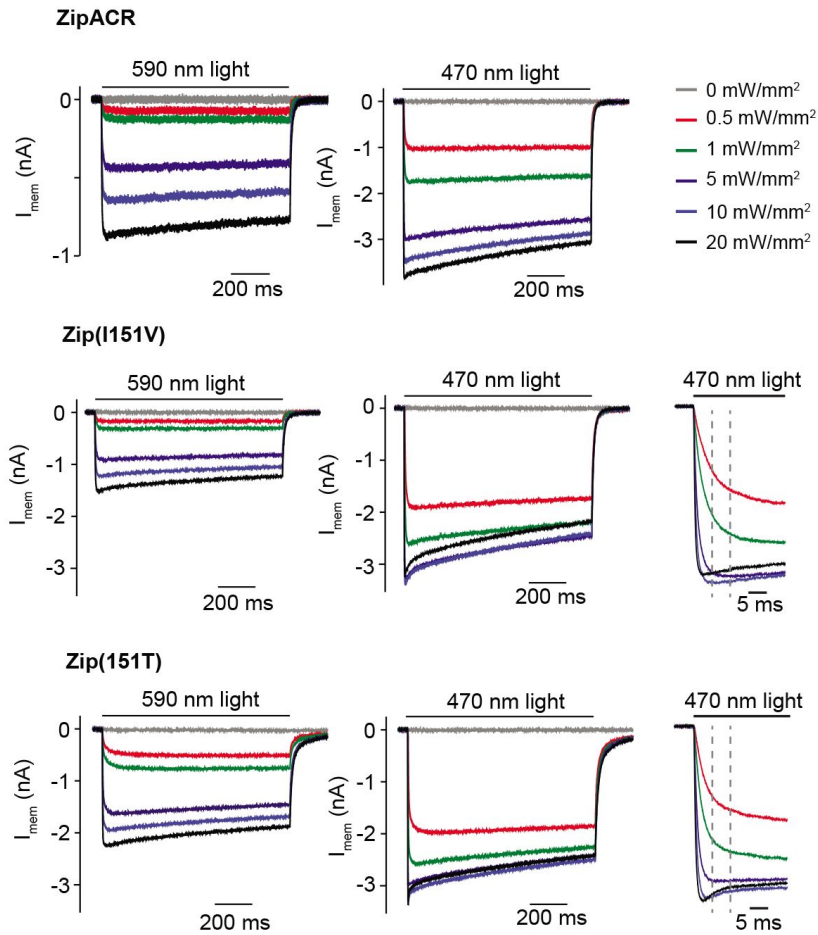


### Supplementary figure 3.

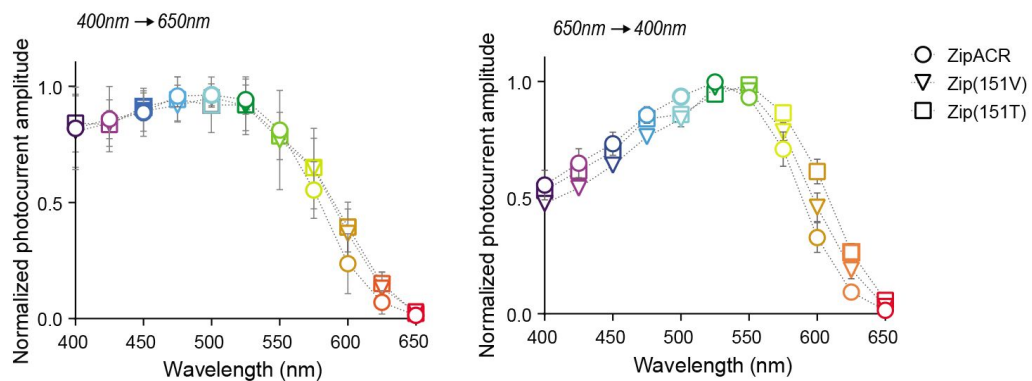
#### Channel properties of the ZipACR variants.

**(A)** 590/470 ratio measured from the ACR variants in HEK293 cells. Each circle corresponds to value obtained from one cell (mean  $\pm$  SD,  $n = 3-7$ ). **(B)** Reversal potential of the selected variants Zip(151T) and Zip(151V) compared to the original ZipACR with cesium (Cs) or potassium gluconate (K) intracellular solution. Each dot corresponds to the value obtained from one cell (mean  $\pm$  SD,  $n = 5-6$ ). **(C)** Typical I/V-relationship for the two variants and ZipACR. The recordings have been done with-or without 470 nm LED illumination.

A



B



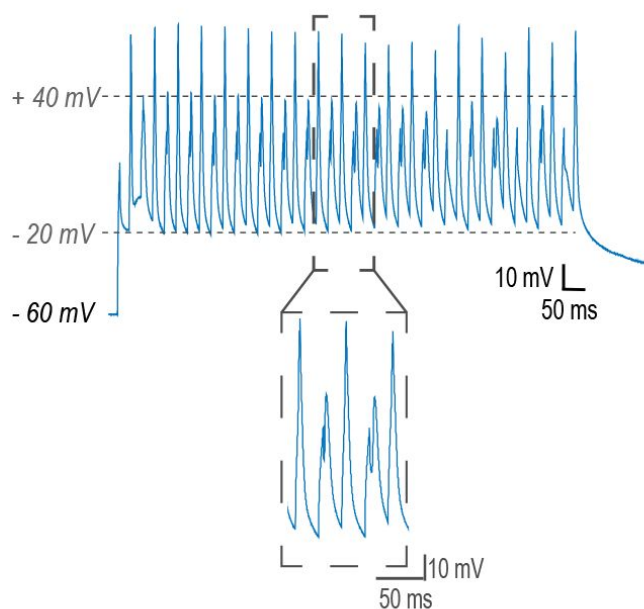
**Supplementary figure 4.**

**Photo-induced responses and action spectra of ZipACR variants.**

(A) Representative responses of the original ZipACR, Zip(151T) and Zip(151V) in HEK293 cells in response to different light intensities. (B) Normalized photocurrent amplitude in response to 400 nm-650 nm and to 650-400 nm light pulses (as presented in Fig. 3D2). The values have been measured on the photocurrents at the peak. The responses were normalized to the maximum response of each cell. The data points are the mean value  $\pm$  SEM (n = 6-7).

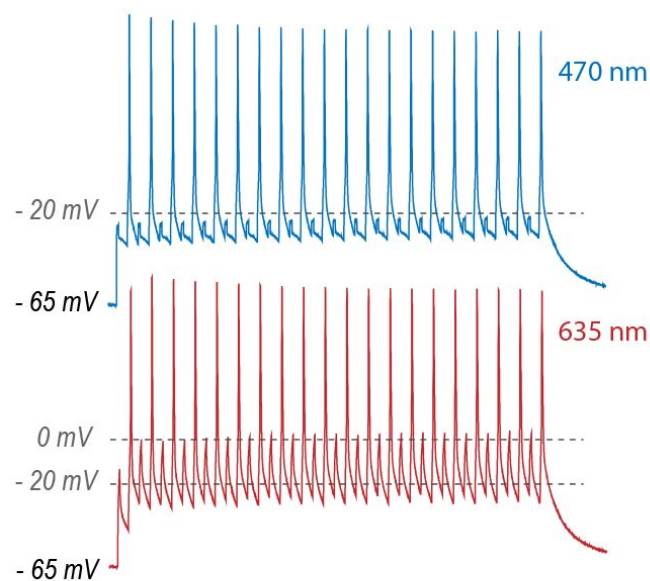
A1

40 Hz, 5-ms train stimulation  
+20 Hz, 5-ms 470 nm LED stimulation



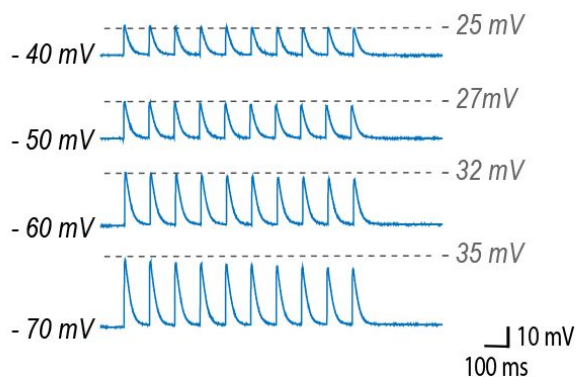
A2

40 Hz, 5-ms train stimulation  
+20 Hz, 5-ms 470 or 635 nm LED stimulation

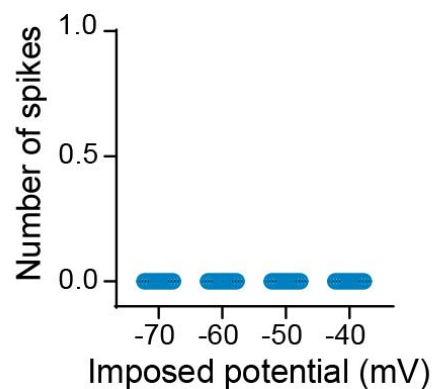


B1

10 Hz, 5-ms 470 nm LED stimulation



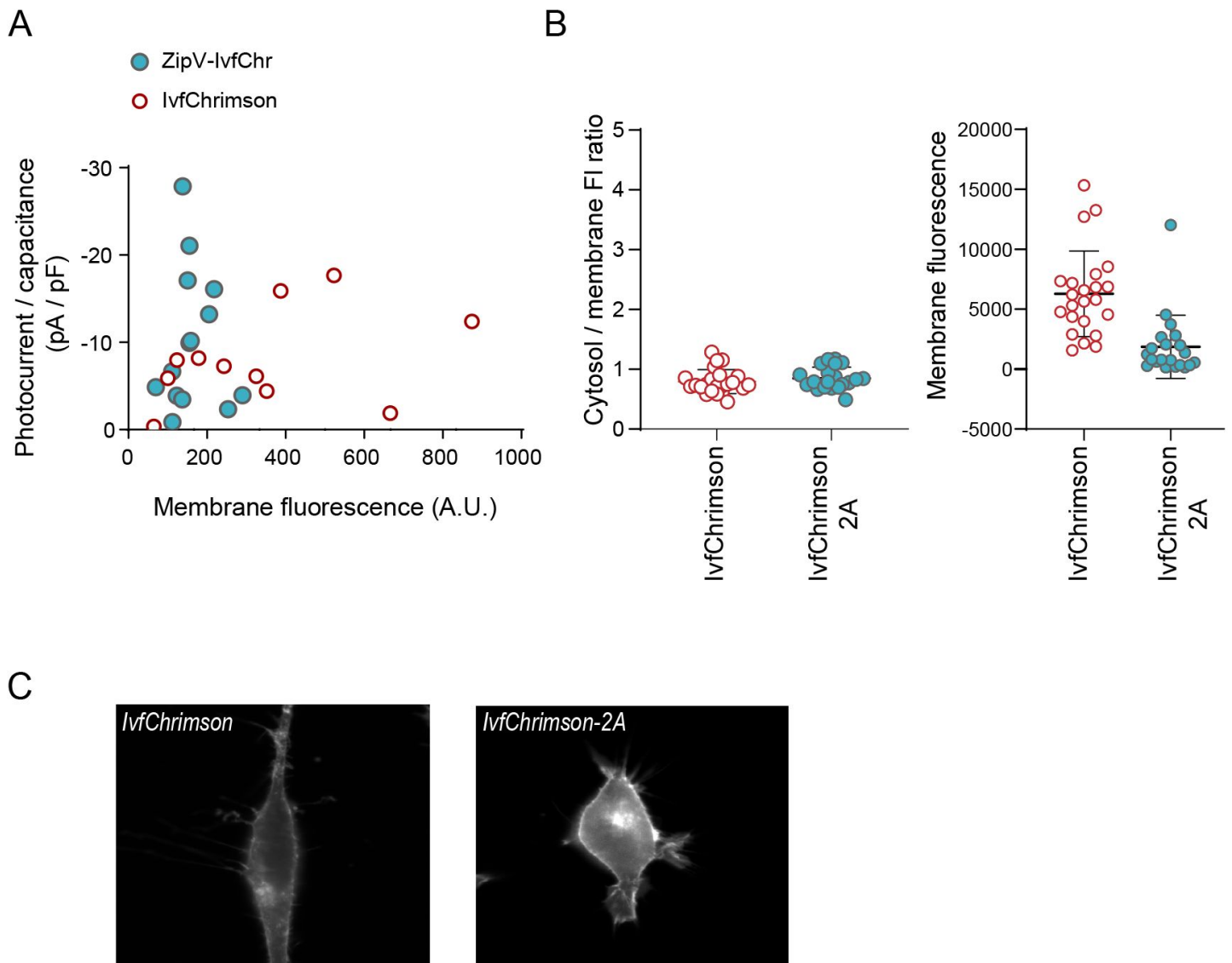
B2



Supplementary figure 5.

(A) The traces of the two outliers in Fig.3 E2.

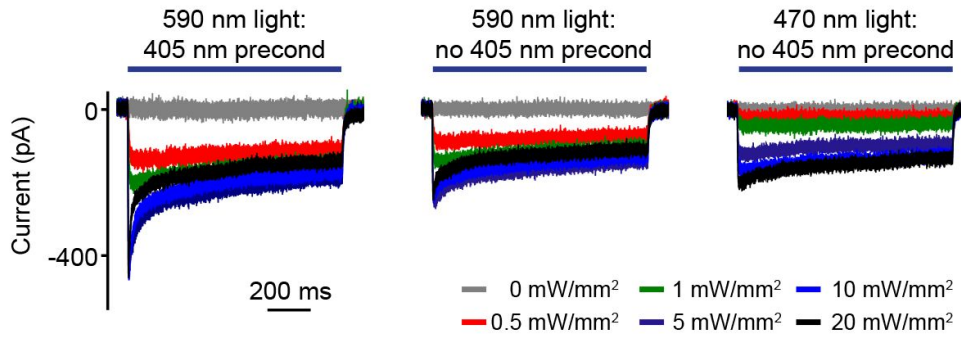
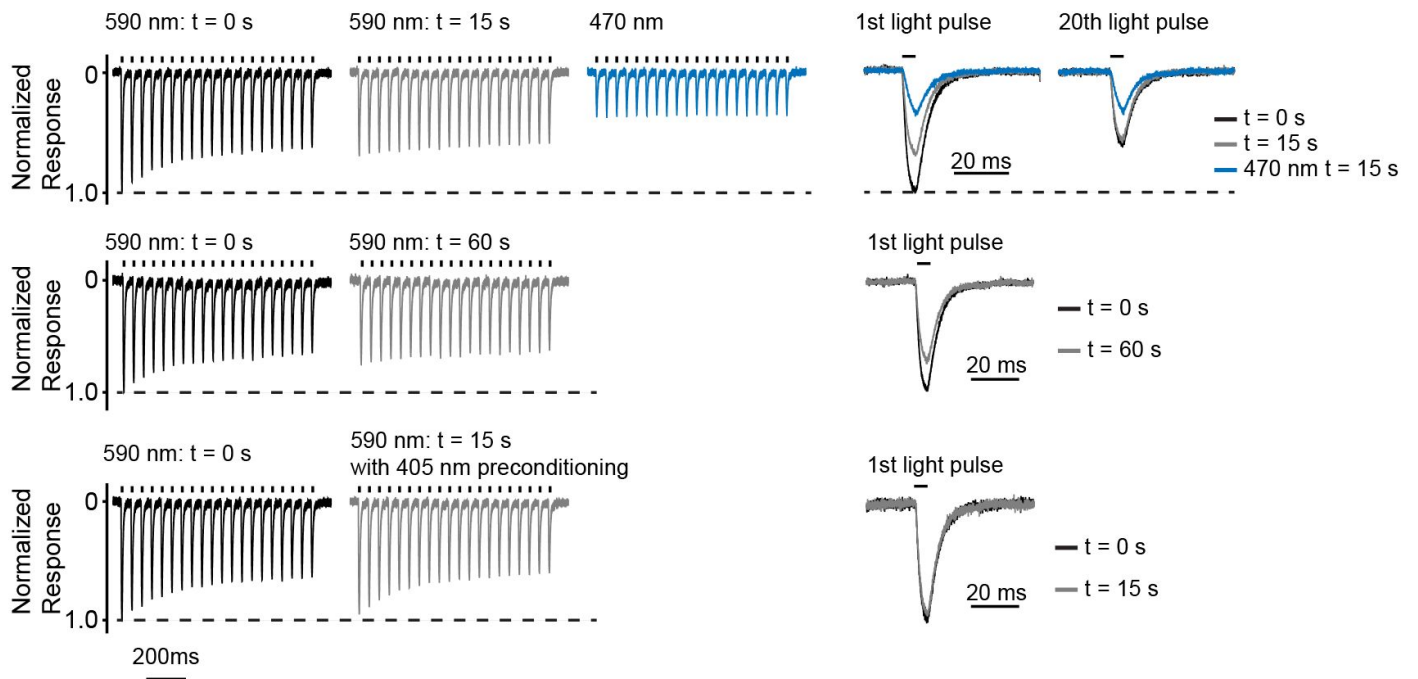
(A1), 470 nm illumination reduced the amplitude of the spikes, but did not abolish the firing. (A2), 470 nm illumination as well as 635 nm light blocked the firing. This cell expressed Zip(151T) channels which was significantly more red-shifted compared to the mean (635/470 nm ratio: 23% for this cell vs 6% for the averaged population). (B) For a range of imposed potentials 470 nm illumination of Zip(151T) does not trigger action potentials. (B1) Typical traces of the DG neurons at various imposed potentials, in response to 10 Hz of 470 nm light illumination. The dotted lines show the potential of the maximal response. (B2) Plot showing the responses in current-clamp of the 10 neurons recorded at -70, -60, -50 and -40 mV imposed potential. Each dot represents a single cell.



### Supplementary figure 6.

#### Improved membrane trafficking of vfChrimson.

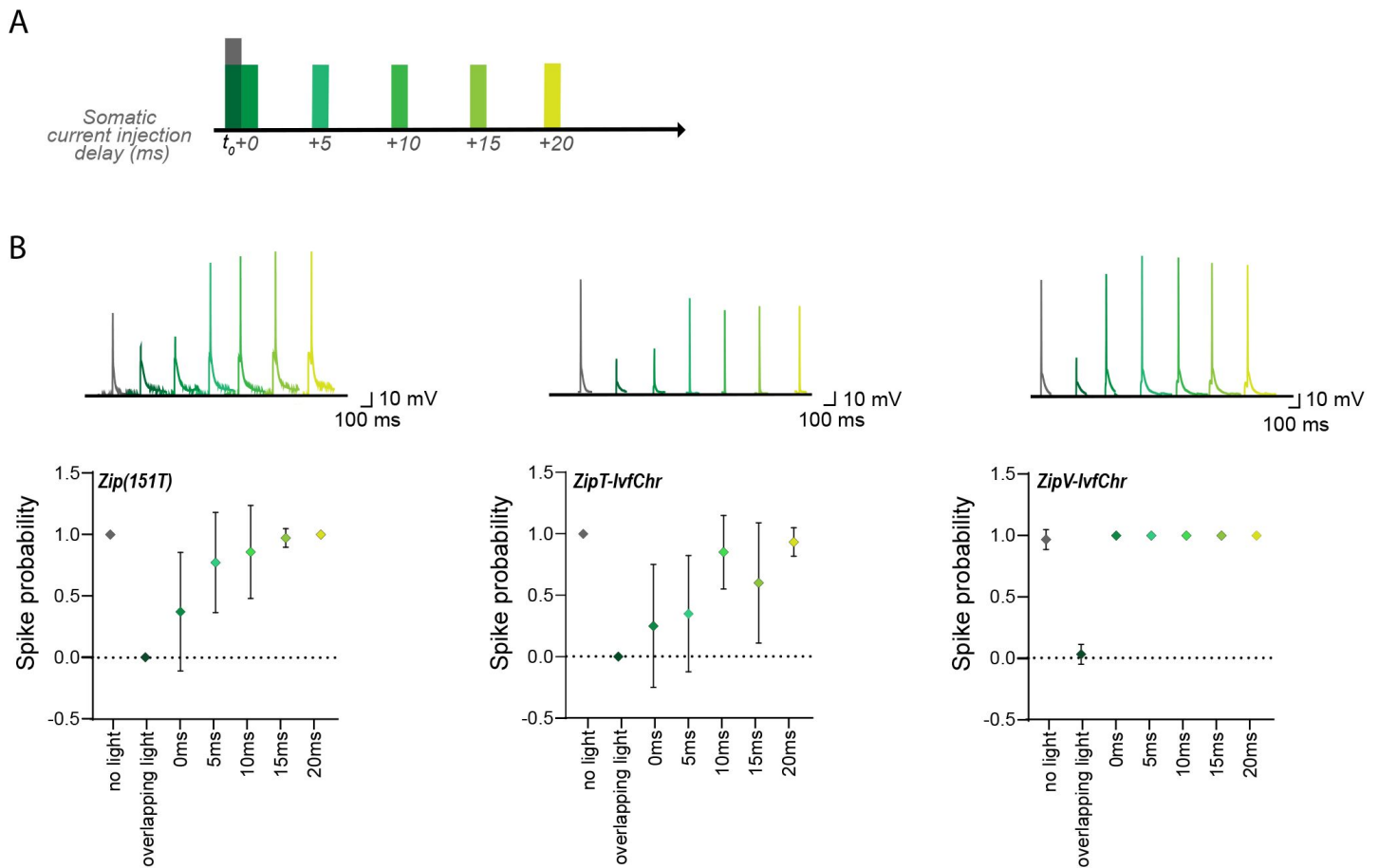
(A) Comparison of trafficking and photocurrent in HEK293 cells at 590 nm illumination for IvfChrimson and ZipV-IvfChr. (B) Comparison of the expression of IvfChrimson-citrine, and '2A'. A line profile was drawn across the cell and the membrane values have been averaged and used for calculating the ratio with the mean cytosolic values. Each dot represents the value from a single cell (n = 19-23, mean  $\pm$  SD). (C) Representative pictures of HEK293 cells expressing the IvfChrimson or '2A'.

**A****B**

### Supplementary figure 7.

#### Excitation spectra of IvfChrimson in HEK293 cells.

(A) Representative responses of IvfChrimson to 1-sec of 470 nm and 590 nm with or without a preconditioning pulse at 405 nm. (B) Normalized responses in voltage-clamp to 590 nm pulses of light, at t = 0, t = 15 and t = 60 sec without 405 nm preconditioning. Due to desensitization IvfChrimson, responses do not recover at least within one minute. Prereconditioning pulse at 405 nm recovers the response.



### Supplementary figure 8.

#### Temporal recovery of neuronal excitability following light-induced inhibition by the ZipACR variants.

Immediate recovery of ZipV-IvfChr from the inhibition. **(A)** Experimental paradigm. DG granule cells expressing Zip(151T), ZipT-IvfCh or ZipV-IvfChr have been stimulated by a somatic current injection (5 ms pulse) overlapping or preceded by a 470 nm pulse (5 ms, 4mW/mm<sup>2</sup>) at 0, 5, 10, 15 or 20 ms intervals. **(B)** The traces in current-clamp (top) are representative of the responses to current injection for the three constructs. The spike probability (bottom) shows an immediate recovery of ZipV-IvfChr from the off-set of the 470 nm light (n = 7 per construct, mean  $\pm$  SD).



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## Editors

Reviewing Editor

### **Marcel Goldschen-Ohm**

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## Reviewer #1 (Public Review):

### Summary

In this manuscript, the authors generate an AAV-deliverable tool that generates action potentials in response to red light, but not blue light, when expressed in neurons. To do this, they screen some red light-excitatory/blue light-inhibitory opsin pairs to find ones that are spectrally and temporally matched. They first show that this works with Chrimson and GtACR2, however, they expand their search after finding that the tau-off (inactivation after light cessation) kinetics of these two opsins are not well-matched. They directly examine a small set of options based on a literature search and settle on a variant of red light-excitatory Chrimson and blue light-inhibitory ZipACR. To more closely match the kinetics of this pair, the authors create a structure homology model of the ZipACR retinal binding pocket and use this to guide the generation of a small mutagenesis panel, leading to a more optimized ZipACR mutant. They then show that a bicistronically expressed fusion arrangement of these opsins, plus some functional peptides, can drive action potentials up to 20hz with red light and does not do so with blue light, in hippocampal cells transduced by AAV. They also show function in vivo, in a mouse, using a physiological readout. They conclude that their new tool may be useful for complex experimental designs requiring multiple optical channels for write-in/read-out.

The major advantage claimed by the authors over existing tools is the temporal time-locking of their inhibitory opsin - this is driven by the contrast between the tau-off kinetics of their ZipACR variant compared to GtACR2, which is used by the leading competitor tool (BiPOLES).

### Big thoughts

While the authors were carefully thoughtful about the potential influence of temporal kinetics on the efficiency of a tool such as this one, there were no experiments conducted that made use of the unique properties of this molecular strategy. To understand why they embarked on this engineering program, I was required to put on my neuroscientist hat and contemplate this question myself:

First, experimental designs where I require multiple optical channels of control. This appears to be aligned with the author's thoughts, as they state, correctly, that opsins utilizing retinal as a light-sensing chromophore are universally activated by blue light (the so-called 'blue shoulder'). Therefore, their tool may be useful for stimulating multiple populations using a blue excitatory opsin in neuron A and their tool for red excitation of neuron B - or, in the author's own words, "A potential solution to the problem of cross-talk...". Yet, there are no data presented that showcases their new tool for this purpose (e.g. Vierock, Johannes, et al. "BiPOLES is an optogenetic tool developed for bidirectional dual-color control of neurons." *Nature Communications* 12.1 (2021): 4527. Figure 4f-I; 6). The same set-up could be imagined for green GECI (or equivalent) imaging of cells in the same volume that their tool is being used in - for instance, interleaving red stimulation light and blue imaging light, (perhaps) without the typical concern of imaging light bleed-through activating the opsin itself.

Second, for high-frequency temporal control over both excitation and inhibition in the same neuron. The red light turns the cell on, and blue light turns the cell off (see, for instance, Zhang, Feng, et al. "Multimodal fast optical interrogation of neural circuitry." *Nature* 446.7136 (2007): 633-639. Figure 2; Vierock as above, Figure 4a,b). Again, here the authors are long on theory ("The new system...can drive time-locked high-frequency action potentials in response

to red pulses") and short on data. While they do show that red light = excitation and blue light = inhibition, they neither show 1) all-optical on/off modulation of the same cell; nor 2) high-frequency inhibition or excitation (max stim rate of 20hz, which is the same as the BiPOLES paper used for their LC stimulation paradigm; Vierock, as above, Figure 7a-d).

Despite these major shortcomings, the further development and characterization of tandem opsins, such as this one, is of interest to the community. There is ongoing work by the BiPOLES team to create new iterations (e.g. Wahid, J., et al. "P-15 BiPOLES2 is a bidirectional optogenetic tool with a narrow activation spectrum and low red-light excitability." *Clinical Neurophysiology* 148 (2023): e16.). To make the case that the tool described in this manuscript is worth the effort that the authors are requesting the neuroscience community invest in trialing it in their own hands, they must revise the manuscript to show that their approach is both 1) different in some way when compared to BiPOLES (it is my understanding that they did not do this, as per the supplementary alignment of the BiPOLES sequence and the sequence of the BiPOLES-like construct that they did test) and 2) that the properties that the investigators specifically tailored their construct to have confer some sort of experimental advantage when compared to the existing standard.

There are a number of additional concerns and clarifications that will strengthen the manuscript that are communicated directly to the authors through this peer-review process.

## Reviewer #2 (Public Review):

### Summary:

One often wishes to combine activation of a neural population via red light with simultaneous modulation of a different population via blue light, or simultaneous imaging of a blue-excited fluorescent reporter. The problem is that all red-shifted opsins have an action spectrum with a long blue tail, leading to spurious opsin activation by blue light.

This valuable paper presents a clever solution to this problem, by pairing an engineered blue-shifted inhibitory chloride-conducting opsin with a red-shifted excitatory opsin. The combined effect is excitation by red light and shunting inhibition by blue light. The paper is very thorough, with convincing spectroscopic and patch clamp characterization of the tools, and tests in brain slices and in vivo. This tool is likely to be useful in the neuroscience community.

### Strengths:

The methods are solid, including the complete characterization of each tool separately, as well as the combination in vivo. The array of testing gives a strong degree of confidence that this tool will work as expected.

### Weaknesses:

There are two discussion points and one experimental point which would make the paper stronger.

1. In the Introduction or Discussion, the authors could better motivate the need for a red-shifted actuator that lacks blue crosstalk, by giving some specific examples of how the tool could be productively used, e.g. pairing with another blue-shifted excitatory opsin in a different population, or pairing with a GFP-based fluorescent indicator, e.g. GCaMP. The motivation for the current tool is not obvious to non-experts.
2. Simultaneous excitation and inhibition are not the same as non-excitation. The authors mentioned shunting briefly. Another possible issue is changes in osmotic balance. Activation of a Na<sup>+</sup> channel and a Cl<sup>-</sup> channel will lead to net import of NaCl into the cell, possibly changing osmotic pressure. Please discuss.

3. The authors showed that in ZipT-IvfChr, orange light drives excitation and blue light does not. But what about simultaneous blue and orange light? Can the blue light overwhelm the effect of the orange light? Since the stated goal is to open the blue part of the spectrum for other applications, one is now worried about "negative" crosstalk. Please discuss and, ideally, characterize this phenomenon.

3.1) Does the use of the new tool require careful balancing of the expression levels of the ZipT and the IvfChr? Does it require careful balancing of blue and orange light intensities?

3.2) Also, many opsins show complex and nonlinear responses to dual-wavelength illumination, so each component should be characterized individually under simultaneous blue + orange light.

3.3) I was expecting to see photocurrents at different holding potentials as a function of illumination wavelength for the co-expressed construct (i.e. to see at what wavelength it switches from being excitatory to inhibitory); and also to see I-V curves of the photocurrent at blue and orange wavelengths for the co-expressed constructs (i.e. to see the reversal potential under blue excitation). Overall, the patch clamp and spectroscopic characterization of the individual constructs was stronger than that of the combined constructs.

### Reviewer #3 (Public Review):

This study addresses the important topic of dual-color optogenetic control of neuronal activity, which is challenging due to significant optical crosstalk between channelrhodopsins of different absorption colors and ion selectivity. However, Mermet-Joret et al. demonstrate in flies that simple coexpression of a strong blue light-activated inhibitory opsin, such as the chloride-selective channelrhodopsin GtACR2, can suppress the blue light activity of a red-shifted excitatory opsin, such as Chrimson, and allow dual-color optogenetic control of the expressing neuron. The same concept was previously discussed by Vierock et al. and led to the generation of BiPOLES, which combines both channels in a single fusion protein. In the present manuscript, the authors introduce an alternative combination of channels with accelerated off-kinetics that are coexpressed by a bicistronic expression cassette. The goal is to better match the duration of illumination and optogenetic manipulation in order to reduce potential side effects induced by prolonged channel opening.

The major novelty of this work lies in the choice of the employed ion channels: the excitatory cation channel vf-Chrimson and the inhibitory anion channel ZipACR, alongside their subsequent modifications (Fig. 2 - 4). Both channels belong to the fastest known ChRs, but the choice of ZipACR raises questions. First, it has a peak absorption at 515 nm that is 40 nm further red-shifted than GtACR2 tested in Figure 1 and accordingly important optical crosstalk with the coexpressed Chrimson channel. Second, it was reported to have reduced chloride selectivity, first by Govorunova et al. in 2017 and later also by Kato et al. in 2018. Both of these aspects are also mentioned by the authors but were not resolved through molecular engineering. Instead site-directed mutagenesis primarily focused on membrane expression and photoreceptor kinetics of the employed channels. Nonetheless, improving the membrane targeting of the vf-Chrimson channel by exchange of the N-terminus finally provided sufficient red light activation at low light intensities to reliably activate expressing neurons and allowed in combination with the decelerated ZipACR mutants dual color optogenetic control with millisecond time resolution. At higher light intensities inactivation of Chrimson and the optical crosstalk of both channels seem to limit its performance.

The experimental results are well presented; but, certain questions persist:

1. The enhanced vf-Chrimson could potentially be a highlight of the manuscript, serving broader applications. Yet, gauging the overall improvements of ivf-Chrimson in comparison to other Chrimson variants remains intricate due to several reasons. First, photocurrents

from ivf-Chrimson seem smaller than those from C-Chrimson (Supplemental Figure 3), and a direct comparison with standard vf-Chrimson is absent. Second, while membrane expression of ivf-Chrimson appears enhanced in provided bright-field recordings, the quantitative analysis would necessitate confocal microscopy and a membrane marker (Supplemental Figure 2). Finally, other N-terminal modified Chrimson variants, like CsChrimson by Klapoetke et al. in 2014 and C1Chrimson by Oda et al. in 2018, have been generated. Comparing ivf-Chrimson to vf-CsChrimson or vf-C1Chrimson would be important to evaluate the benefits of the applied N-terminal modification.

2. The action spectra of ZipACR suggest peak absorption of ZipACR WT and its mutant at 525 - 550 nm (Fig. 3). This is even further red-shifted than previously reported by Govorunova et al. Further action spectra recordings differ for all constructs between recordings initiated with blue or red light (Supplementary Fig. 5). This discrepancy is unexpected and should be discussed. Additionally, the representative photocurrents of Zip(151V) in Fig. 3D1 do not align with the corresponding action spectrum in Fig. 3D2 as they show maximal photocurrents for 400 nm excitation.

3. The authors introduce two different bicistronic expression cassettes-ZipT-IvfChR and ZipV-IvfChR-without providing clear guidelines on their conditions of use. Although the authors assert that ZipT is slower and further red-shifted than ZipV, the differences in the data for both ACR mutants are small and the benefits of the different final constructs should be explained.

4. The ZipT/V-IvfChRs are designed as bicistronic constructs; yet, disparities in membrane trafficking and protein degradation between the two channels could lead to divergences in blue and red light photoresponses. For future applicants, understanding the extent of expression ratio variations across cells using the presented expression cassettes could be of significance and should be discussed.