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Spatiotemporal Control of Opioid Signaling and Behavior

Highlights

- A light-sensitive mu-opioid-like receptor (opto-MOR) was generated and characterized
- Opto-MOR initiates canonical MOR signaling both in vitro and in neurons
- Photoactivation of opto-MOR in selected GABAergic neurons induces reward or aversion
- Opto-MOR is a novel tool for in vivo optodynamic control of opioid signaling

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In Brief

Siuda et al. develop a photosensitive mu-opioid-like receptor (opto-MOR) that triggers cAMP inhibition and MAP kinase activation, couples to GIRK currents, and internalizes like the mu-opioid receptor. Photostimulation of opto-MOR within discrete GABAergic nuclei induces real-time preference or aversion.



Spatiotemporal Control of Opioid Signaling and Behavior

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SUMMARY

Optogenetics is now a widely accepted tool for spatiotemporal manipulation of neuronal activity. However, a majority of optogenetic approaches use binary on/off control schemes. Here, we extend the optogenetic toolset by developing a neuromodulatory approach using a rationale-based design to generate a Gi-coupled, optically sensitive, mu-opioid-like receptor, which we term opto-MOR. We demonstrate that opto-MOR engages canonical mu-opioid signaling through inhibition of adenylyl cyclase, activation of MAPK and G protein-gated inward rectifying potassium (GIRK) channels and internalizes with kinetics similar to that of the mu-opioid receptor. To assess *in vivo* utility, we expressed a Cre-dependent viral opto-MOR in RMTg/VTA GABAergic neurons, which led to a real-time place preference. In contrast, expression of opto-MOR in GABAergic neurons of the ventral pallidum hedonic cold spot led to real-time place aversion. This tool has generalizable application for spatiotemporal control of opioid signaling and, furthermore, can be used broadly for mimicking endogenous neuronal inhibition pathways.

INTRODUCTION

Opioid receptor-targeting drugs have been used as analgesics and recreationally abused for hundreds of years. Due to a lack of effective alternatives, they remain on the front lines for acute pain management, severe anti-tussive treatment, and other indications despite their high abuse potential. Mu (MOPR), kappa (KOPR), delta (DOPR), and nociceptin opioid peptide receptors (NOPR) persist at the forefront of basic science and drug discovery efforts on disorders ranging from gastrointestinal ailments to pain, addiction, and depression (Al-Hasani and Bruchas, 2011).

Among these receptor systems, the mu-opioid receptor has been the most intensely studied due to its long-established involvement in analgesia, euphoria, and reward in response to morphine-like analogs. However, despite years of study, opioid research had been limited by a fundamental challenge. Studying the specific effects of opioids with spatial, temporal, and cell-type-specific control is virtually untenable, especially in the context of the CNS. Systemic drugs bind MOPRs on heterogeneous cell populations across multiple brain regions. MOPR expression at both pre- and post-synaptic sites within overlapping discrete regions precludes determination of circuit level contributions. Therefore, approaches to selectively limit engagement of MOPR signaling to restricted cell populations with temporal control that closely mimics endogenous opioid kinetics is a first key step toward unraveling these issues and developing future therapeutic strategies where opioids are the most effective treatment regimen.

Recent developments in optogenetics and molecular biology provide an ideal strategy for addressing these questions. Class A G protein-coupled receptors (GPCRs), including the rat rhodopsin receptor and MOPR, have structural and functional similarities that can be exploited to create hybrid receptors with unique functional properties. More specifically, the light-sensitive external portion of rhodopsin receptors can be combined with internal signal transduction components of other GPCRs to produce so-called opto-XR chimeras capable of initiating and terminating receptor-specific signaling events with temporal precision enabled by pulses of light (Airan et al., 2009; Gunaydin et al., 2014; Kim et al., 2005; Masseck et al., 2014). Furthermore, packaging these receptors into Cre recombinase-dependent viruses using loxP flanked doubled inverted open (DIO or FLEX) reading frames allows for restricted expression in discrete cell types within isolated brain regions yielding spatial control of GPCR signaling *in vivo* (Atasoy et al., 2008; Zhang et al., 2010).

Here, we present the generation and characterization of a new photosensitive mu-opioid-like chimeric receptor (we term opto-MOR). We show that opto-MOR suppresses cyclic AMP (cAMP) levels, activates MAPK signaling, and internalizes in a similar time course to native MOPR. Furthermore, it functionally couples

to GIRK channels in GABAergic neurons of the periaqueductal gray (PAG) and rostromedial tegmental region (RMTg), mimicking properties of native MOR (Blanchet and Lüscher, 2002; Ingram et al., 2008; Matsui et al., 2014; Vaughan et al., 2003). Finally, we demonstrate here that opto-MOR promotes classical MOR-mediated reward and aversion behaviors in distinct brain circuits. Together, these findings establish opto-MOR as a spatiotemporally precise MOR analog and support its utility for studying opioid circuitry and behavior.

RESULTS

Developing a Chimeric Photosensitive Receptor with Components of the Mu-Opioid Receptor

In order to effectively generate an optically sensitive MOR-based receptor chimera that would have high probability of structural similarity to mu-opioid receptors and retain native opioid Gi/o-protein signaling, we used the closest class A-related receptor as a backbone. Rat rhodopsin RO4 has been previously shown to couple to Gi/o signaling pathways *in vitro* and thus served as the ideal template for generation of an opto-MOR (Hille, 1994). Using the constraint-based multiple alignment tool (COBALT, NCBI; http://www.st-va.ncbi.nlm.nih.gov/tools/cobalt/re_cobalt.cgi), rat rhodopsin RO4 was aligned against rat MOR to identify and align transmembrane, intracellular, and extracellular domains (Figures 1A and S1A). To confer MOR-like coupling, we isolated the rhodopsin RO4 sequence, retained the critical photoisomerizing RO4 retinal binding site (Figures S1A and S1B), and inserted the rat MOR intracellular loops and C terminus (Figures S1A and S1C). A protein folding prediction was modeled by bioinformatics software (Roy et al., 2010) in order to accurately project how our various chimeras would result in a seven transmembrane protein with matched features (Figures 1A and S1D). This approach allowed for optimal conservation of Gi/o receptor dynamics, a high degree of photosensitivity, while simultaneously providing the critical intracellular communication components for engaging mu-opioid signaling pathways *in vitro* and *in vivo*.

Opto-MOR Is Photosensitive and Effectively Mimics Mu-Opioid Signaling Dynamics

In order to determine if opto-MOR is functional and traffics to the membrane, we first examined whether a YFP-tagged opto-MOR expresses at the cell membrane in a pattern similar to that of the wild-type MOR-GFP-tagged receptor. We found that both receptors express at the plasma membrane in a similar manner (Figures 1B and 1C) in unstimulated cells. Opioid receptor characterization has demonstrated that agonist stimulation of opioid receptors causes interactions with inhibitory G proteins ($G_{\alpha i}$) to induce inhibition of cAMP production (Childers and Snyder, 1978; Al-Hasani and Bruchas, 2011) (Figure 1A). In order to determine whether cells expressing opto-MOR couple to the same canonical MOR signaling pathways, we assessed their response to blue LED (465 nm, 1 mW, 20 s) stimulation in a battery of well-known *in vitro* opioid signaling assays in direct parallel with wild-type rat MOR using the high affinity, selective MOR agonist, [D-Ala², NMe-Phe⁴, Gly-o⁵]-enkephalin (DAMGO; 1 μ M). First, we show that opto-MOR maintains mem-

brane expression levels similar to that of MOR in unstimulated HEK293 cells (Figures 1B and 1C). Second, we found that photostimulation of opto-MOR and DAMGO activation of MOR caused a time-locked decrease in forskolin-induced intracellular cAMP levels with similar time constants of activation (Figures 1D–1F, S2A, S2B, S2D, and S2E). To verify the specificity of our constructs, we show that opto-MOR does not respond to the selective MOR full agonist DAMGO, and likewise wild-type MOR does not respond to photostimulation (Figures S2C and S2F). Additionally, we show that opto-MOR is maximally activated with 465 nm light and shows less efficacy at other wavelengths in cAMP inhibition (525, 630, and 660 nm) (Figure 1G). Finally, we found that opto-MOR is highly sensitive to light and requires very little light power for photoactivation (Figures 1H and S2G), while varying LED pulse lengths resulted in similar levels of cAMP inhibition (Figure 1I). Opto-MOR and MOR caused similar levels of cyclic AMP inhibition via photostimulation and agonist treatment, respectively, suggesting that opto-MOR couples to canonical mu-opioid signaling pathways, yet utilizes rapid time-locked photoswitching to engage $G_{\alpha i}$ -mediated inhibition of adenylyl cyclase signaling.

Agonist stimulation of all four opioid receptors has been shown to recruit various factors resulting in mitogen-activated protein kinase (MAPK) activation (Bruchas et al., 2011; Al-Hasani and Bruchas, 2011). MOR has been shown to elicit a rapid initial peak in the phosphorylation of extracellular signaling-regulated kinase (pERK) in neurons, astrocytes, and transfected cell cultures (Belcheva et al., 2005). Here, we examined whether opto-MOR and MOR produce similar kinetics and efficacy in engaging pERK signaling in HEK293 cells. In complementary experiments, we found a rapid and transient increase in pERK (~2–5 min) in response to blue LED photostimulation of opto-MOR and DAMGO application to MOR-expressing cells (Figures 1J–1L). pERK returned to basal levels 60–90 min after either photostimulation or DAMGO treatment. Furthermore, opto-MOR-mediated activation of ERK was mostly independent of LED pulse time (Figure S2H) and only mildly affected by light power (Figure S2I), suggesting that time-locked photoactivation of opto-MOR immediately engages the MAPK signaling cascade.

Opioid receptors are well known to be rapidly regulated by arrestin-clathrin-mediated internalization pathways. To assess whether opto-MOR exhibits similar activation-induced receptor regulation and engages canonical mu-opioid receptor internalization machinery, we performed side-by-side experiments whereby we compared the kinetics and efficacy of LED or agonist-stimulated receptor internalization on opto-MOR or MOR. Following photostimulation or DAMGO treatment, respectively, both opto-MOR and MOR internalized rapidly (within 15 min) with similar rates of internalization (τ_{in}), with a peak effect at 30 min after agonist or light stimulation (Figures 2A, 2B, S3A, and S3B). Opto-MOR receptors remained punctate in the cytosol at all post-stimulation times tested, lasting up to an hour. Opto-MOR internalization over this time course is consistent with previously reported wild-type MOR internalization by both widely used synthetic and endogenous ligands (Al-Hasani and Bruchas, 2011).

Agonists such as etorphine and DAMGO and endogenous opiate peptides such as Met-enkephalin and β -endorphin have

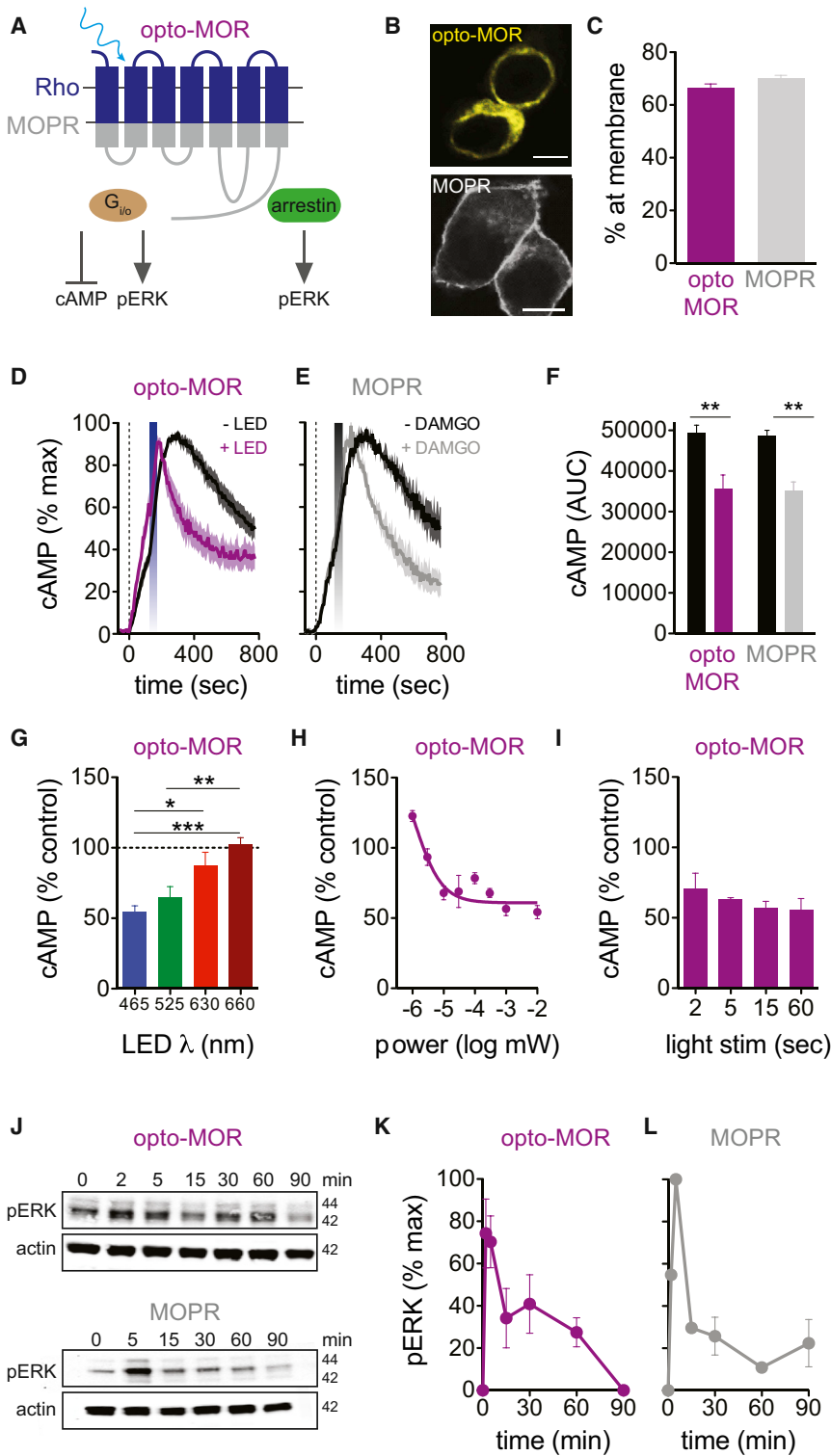


Figure 1. Opto-MOR and MOR Share Canonical Intracellular G Protein Signaling Mechanisms

(A) Schematic of opto-MOR showing activation of cAMP and pERK pathways. (B) Opto-MOR and MOPR are highly expressed at the membrane in HEK293 cells (scale bar, 10 μ m). (C) Opto-MOR (purple, n = 43 cells) and MOPR (n = 37 cells) share similar surface expression in HEK293 cells. (D) Optical stimulation (blue bar; 20 s, 1 mW) of opto-MOR reduces forskolin-induced (dashed line) cAMP in HEK cells (n = 9–10 experiments). (E) Application of DAMGO (gray bar; 1 μ M) reduces forskolin-induced (dashed line) cAMP in HEK293 cells expressing MOPR (n = 3 experiments). (F) Light and DAMGO significantly reduce forskolin-induced cAMP in opto-MOR (purple; n = 12) and MOPR (gray; n = 3). (G) cAMP inhibition by opto-MOR is most efficient at 465 nm (n = 6–8 experiments; *p < 0.05, **p < 0.01, ***p < 0.001 via one-way ANOVA with Bonferroni's multiple comparison test). (H) cAMP inhibition by opto-MOR is power dependent (n = 3–9 experiments). (I) cAMP inhibition by opto-MOR is not dependent on light pulse length (n = 2–10 experiments). (J) Representative immunoblots show similar kinetic increases in pERK in response to light (1 min, 1 mW) or DAMGO (1 μ M) in opto-MOR (purple) and MOPR (gray)-expressing cells. (K and L) Quantification of light-induced pERK in opto-MOR (n = 6) (K) and DAMGO-induced pERK in MOPR (n = 2) (L). Data are represented as mean \pm SEM. See also Figures S1 and S2.

addition to internalization, we photostimulated opto-MOR and induced inhibition of cAMP (Figures 2C and 2F). In parallel, we photostimulated opto-MOR expressing HEK293 cells at saturating power (1 mW, 20 s), waited varying degrees of time, and then photostimulated again (i.e., pre-pulsed). We found that at approximately 15 min following pre-pulse stimulation, there is a significant loss of opto-MOR function, with no difference observed from the unstimulated control cells (Figures 2D and 2F). However, longer time periods between the two light pulses show that within approximately 60 min there is restoration of opto-MOR function in inhibition of cAMP (Figures 2E and 2F). However, it is unclear whether this restoration is due to receptor recycling or de novo receptor synthesis. To address this, we repeated this experiment but pretreated cells with either brefeldin A, which blocks translocation of proteins from the ER to the Golgi and has been used in studies of MOPR recycling, or Dyngo-4a, an inhibitor of dynamin that prohibits receptor internalization and has

been shown to cause MOPR internalization 30 min after drug exposure, as receptor endocytosis and recycling to the membrane reaches steady state around 30 and 60 min, respectively. To test whether opto-MOR displays functional desensitization in

ment but pretreated cells with either brefeldin A, which blocks translocation of proteins from the ER to the Golgi and has been used in studies of MOPR recycling, or Dyngo-4a, an inhibitor of dynamin that prohibits receptor internalization and has

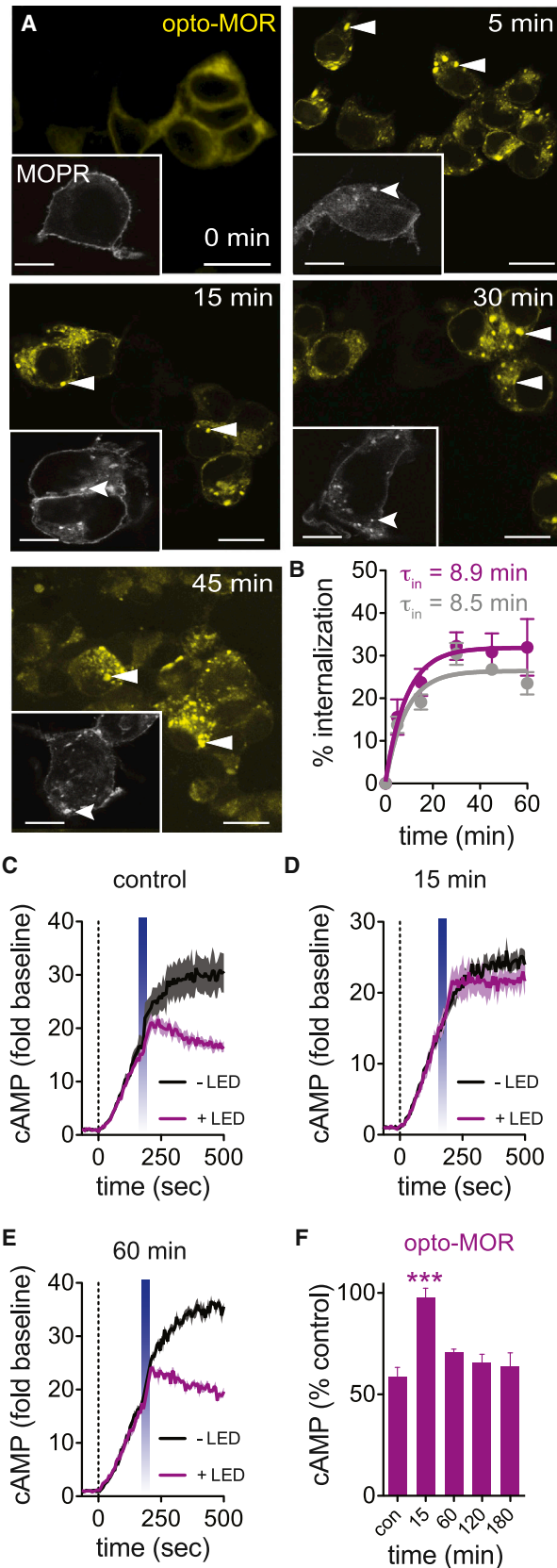


Figure 2. Opto-MOR and MOPR Internalization and Recovery from Desensitization

(A) Representative images show internalization of opto-MOR (colorized yellow; scale bar, 50 μ m) and MOPR (inset, gray; scale bar, 10 μ m) expressed in HEK293 cells in response to light and DAMGO. 0, 5, 15, 30, and 45 min time points are represented. Arrowheads show examples of internalized receptor. (B) Quantification of receptor internalization in opto-MOR (purple; τ_{in} = 8.9 min; n = 16–43 cells per time point over 2 experimental replicates) and MOPR (gray; τ_{in} = 8.5 min; n = 24–38 cells per time point over 3 experimental replicates) in response to light and DAMGO-induced activation, respectively. (C) Opto-MOR inhibits forskolin-induced (dashed line) cAMP following light stimulation (n = 4 traces). (D) A second light pulse 15 min following the first shows a loss of opto-MOR activity (n = 3–4 traces). (E) cAMP inhibitory activity returns to baseline levels 60 min following an initial light pulse (n = 3 traces). (F) Time course of recovery from desensitization (n = 3–11 replicates; ***p < 0.001 via one way ANOVA followed by Dunnett’s multiple comparison test to control). Data are represented as mean \pm SEM. See also Figure S2.

recently been shown to prevent intracellular GPCR cAMP signaling. Following a 1 hr interpulse interval (Figure S3D), we found that pretreatment with brefeldin A did not prevent opto-MOR-induced inhibition of cAMP (Figures S3C and S3E). Furthermore, Dyngo-4a pretreated cells also show cAMP inhibition, suggesting that opto-MOR continues to inhibit cAMP from the plasma membrane with no loss of signal when inhibiting dynamin-clathrin pathways (Figures S3C and S3F). These data together suggest that opto-MOR may indeed recycle and use clathrin-dynamin endocytosis, which has been documented in opioid receptors (Gupta et al., 2014) and other GPCR systems (Irannejad et al., 2013; Tsvetanova and von Zastrow, 2014), and that opto-MOR may have some utility in dissecting these diverse signaling dynamics.

Opto-MOR Expresses and Is a Functional Receptor in DRG Neurons

To test whether opto-MOR is expressed and functions outside the context of an artificial heterologous system, we generated a Cre-dependent adeno-associated viral construct (Figure 3A), whereby we inserted opto-MOR into a double inverted open reading frame flanked by LoxP and Lox2227 sites (Zhang et al., 2010). Utilizing this virus (AAV5-EF1 α -DIO-opto-MOR-YFP), we infected cultured dorsal root ganglia (DRG) neurons from Advillin-Cre⁺ mice (da Silva et al., 2011; Zhou et al., 2010). We did not detect any signal in uninfected control neurons (Figure 3B). In Advillin-IRES-Cre⁺opto-MOR DRG neurons, we observed opto-MOR expression in as little as 2 days post-transduction (green, Figure 3C), suggesting rapid Cre-mediated excision in DRG. Strong opto-MOR expression was seen after 5 days (Figure 3D), which partially co-localized with synapsin (Figure 3D), demonstrating that opto-MOR is trafficked efficiently along neuronal processes and to presynaptic terminals. We also show that following photostimulation of opto-MOR-expressing DRGs, there is an increase in pERK compared to untreated controls (Figures 3E and 3H). Following photostimulation (1 min, 1 mW), we observed an increase in pERK levels 5 min following an initial light stimulation that returns to basal levels within 30 min (Figures 3F–3H). We also report evidence for photostimulated receptor internalization in these DRG cultures

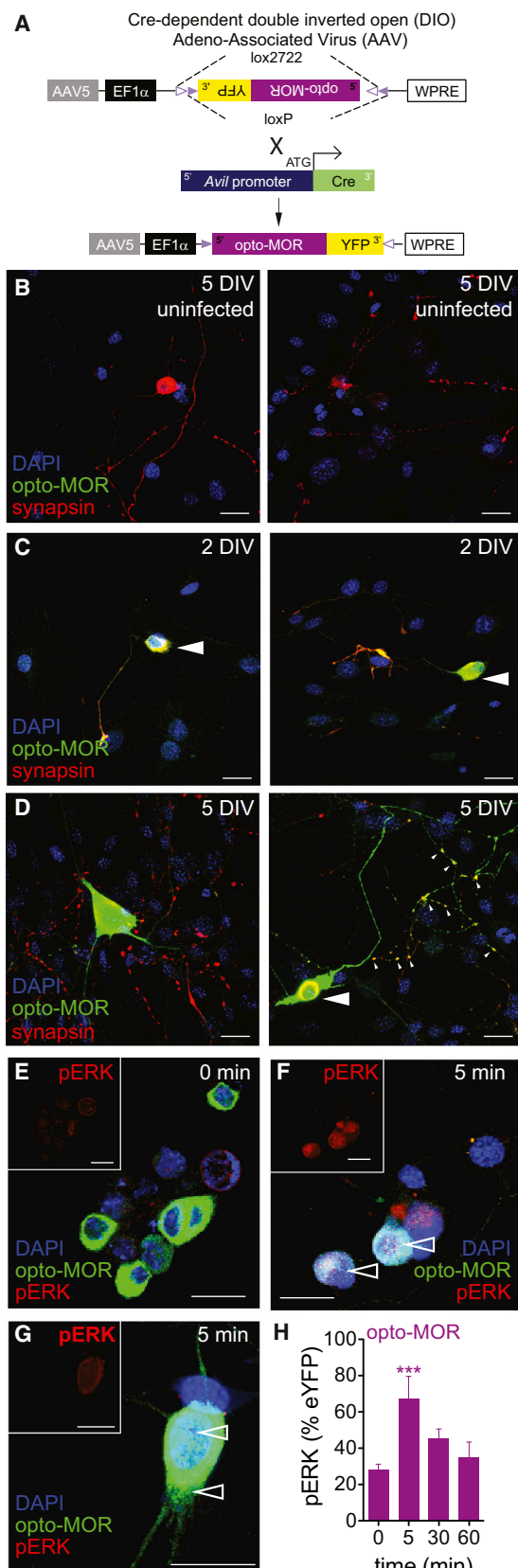


Figure 3. Opto-MOR Expression and Function in DRG Neurons

(A) Opto-MOR packaged into an AAV5-DIO-EF1 α -YFP viral construct. Cre expression is driven from the sensory neuron-specific Advillin promoter following insertion after the initial start codon, resulting in Cre-mediated recombination and inversion of the opto-MOR construct into the appropriate 5'-3' orientation.

(B) Uninfected DRGs colabeled with the nuclear stain DAPI (blue) and synapsin (red).

(C) Opto-MOR expression (green) in DRGs 2 days in vitro (DIV). Large arrowheads point to the soma of infected neurons.

(D) Opto-MOR expression (green) in DRGs 5 DIV. Small arrowheads highlight opto-MOR expression in synaptic terminals colabeled with synapsin (red). Scale bars for (B)–(D), 30 μ m.

(E) Unstimulated opto-MOR expressed in DRGs after 5 DIV labeled with pERK (red and inset).

(F) Opto-MOR expressing DRGs (5 DIV) 5 min following photostimulation and labeled for pERK (red and inset). Open arrows denote internalized punctate receptors.

(G) Higher magnification of opto-MOR expressing DRGs (5 DIV) 5 min following photostimulation and labeled for pERK (red and inset). Open arrows denote internalized punctate receptors.

(H) pERK intensity normalized to opto-MOR-YFP intensity at varying times following photostimulation ($n = 3$ –32 DRG neurons per time point; *** $p < 0.001$ via one way ANOVA followed by Dunnett's multiple comparison test to 0 min). Scale bars for panels (E)–(G), 20 μ m. Data are represented as mean \pm SEM.

(Figures 3F–3H), suggesting that the receptor can be dynamically regulated in physiologically relevant systems. These data closely mimic the kinetics of ERK activation seen in HEK293 cells expressing opto-MOR (Figures 1J and 1K) and further support the utility of using opto-MOR to dissect signaling in neuronal preparations.

Opto-MOR Couples to G $\beta\gamma$ -Mediated GIRK Currents in PAG GABAergic Neurons

In order to assess whether opto-MOR couples to similar intracellular signaling pathways following endogenous MOPR activation in neurons, we focused on the periaqueductal gray (PAG), which contains a high proportion of MOPR⁺ neurons (Vaughan et al., 2003). We injected AAV5-EF1 α -DIO-opto-MOR-YFP into vGAT-IRES-Cre⁺ mice (Vong et al., 2011) within the PAG (AP: -5.0 mm, ML: ± 0.5 mm, DV: -2.8 mm) and targeted YFP⁺ GABAergic neurons for whole-cell recordings from acute brain slices 2–3 weeks after injection. Photostimulation with 470 nm LED light (10 mW/mm²) evoked rapid outward currents together with a simultaneous decrease in input resistance that was reversed by bath application of barium (1 mM), a blocker of GIRK currents (Figures 4A and 4B), suggesting that opto-MOR activation leads to downstream activation of GIRK channels, which is consistent with the activation of endogenous MOPR receptors in these neurons (Vaughan et al., 2003). A parallel set of experiments was performed on MOPR positive cells in the PAG. Bath application of DAMGO (1 μ M) showed similar outward currents and decreased input resistance that were also barium sensitive (Figures 4C and 4D). Voltage ramps performed pre- and post-stimulation demonstrated current-voltage curves consistent with the properties of GIRK channels, which were inhibited by barium (Figures 4G and 4H). Peak changes in holding current and input resistance for opto-MOR (LED) and MOPR (DAMGO) were not statistically different following stimulation, or in the presence of barium (Figures 4E and 4F), suggesting

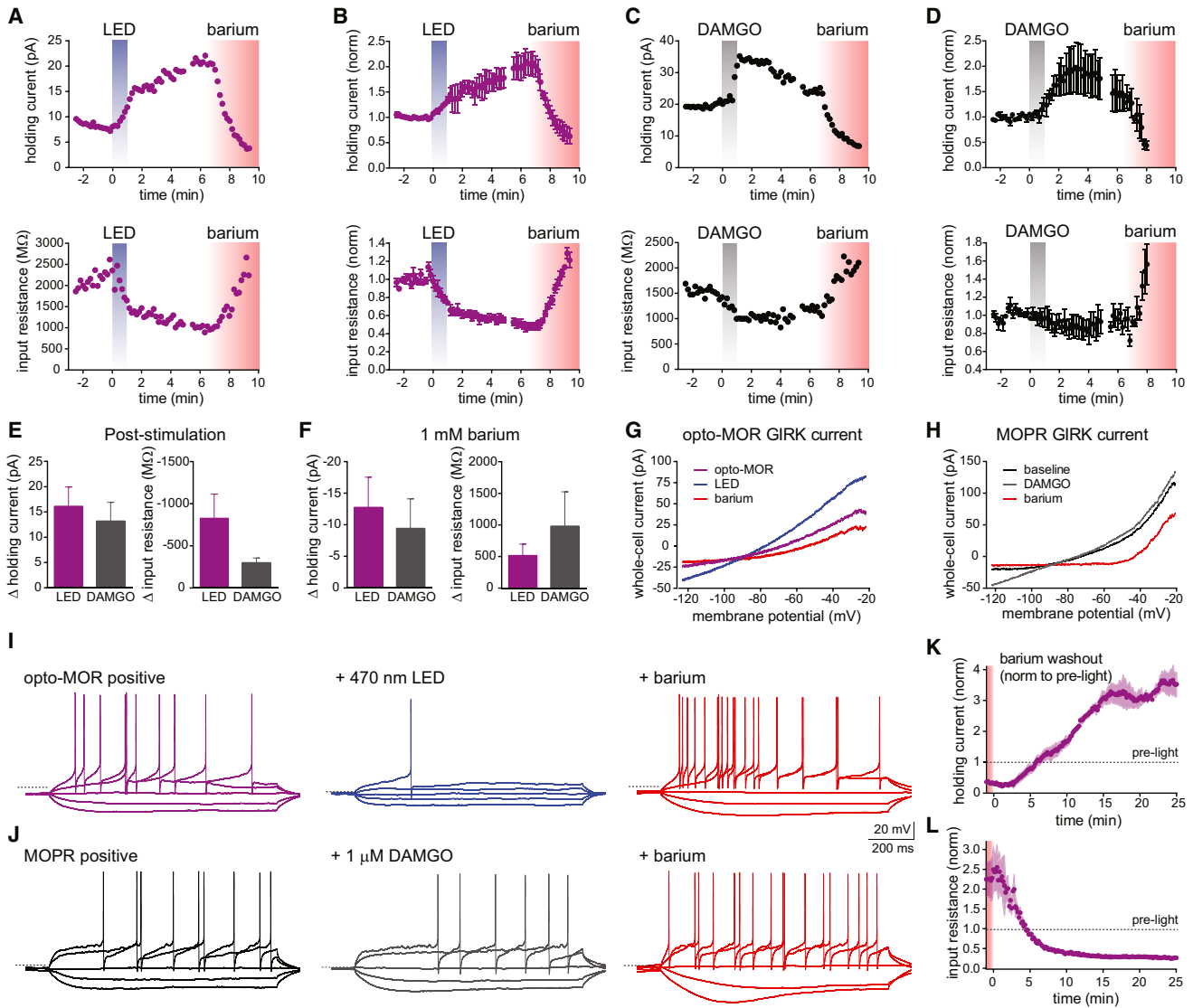


Figure 4. Photostimulation of Opto-MOR and Activation of Endogenous MOPRs Have Similar Effects on Neuronal Physiology and Excitability in GABAergic PAG Neurons

(A) Representative plots from an opto-MOR⁺ neuron in acute PAG slices illuminated with 470 nm LED light (10 mW/mm²) for 60 s. The top trace shows a rapid outward current in response to illumination, while the bottom trace depicts the simultaneous drop in input resistance. Bath application of 1 mM barium (red shading) both blocked the outward current and reversed the change in input resistance.

(B) Normalized summary plots showing the response to LED stimulation from additional opto-MOR⁺ neurons as described in (A) (n = 4).

(C) Example traces recorded from a MOPR⁺ neuron stimulated with 1 μM DAMGO (gray bars) showing similar outward currents (top) and decreased input resistance (bottom) compared to opto-MOR⁺ neurons shown in (A) and (B).

(D) Normalized summary plots from additional MOPR⁺ neurons depicting the response to DAMGO as described in (C) (n = 4).

(E) Quantification of the peak changes in holding current and input resistance following LED (purple) or DAMGO (gray) stimulation (n = 4).

(F) Quantification of the peak changes in holding current and input resistance after application of 1 mM barium (n = 4).

(G and H) Representative current-voltage traces from a 250 ms voltage ramp from -20 to -120 mV, in an opto-MOR⁺ neuron (G) before (purple trace) and after 60 s LED stimulation (blue), or a MOPR⁺ neuron (H) before (black) or after stimulation with 1 μM DAMGO (gray). Both currents were reduced by the GIRK channel blocker barium (1 mM, red).

(I and J) Left traces depict voltage traces from current-clamp recordings of an opto-MOR⁺ (purple traces) (I) or MOPR⁺ neuron (black) (J) in response to hyperpolarizing current injections of -20 and -10 pA, and depolarizing current injections of 1- and 2-times rheobase. Middle traces show decreased input resistance and excitability following LED (blue) and DAMGO (gray) in response to the same current injections before stimulation. The right traces demonstrate the increased input resistance and neuronal excitability observed following GIRK channel block with 1 mM barium (red). Dashed lines indicate -60 mV membrane potential.

(K and L) Normalized summary plots of the persistent outward current (K) and decreased input resistance (L) in opto-MOR⁺ neurons following light stimulation and subsequent barium washout (n = 3). These effects were observed >45 min after 60 s LED stimulation. The dashed lines indicate the average response before illumination. Data are represented as mean ± SEM. See also Figure S4.

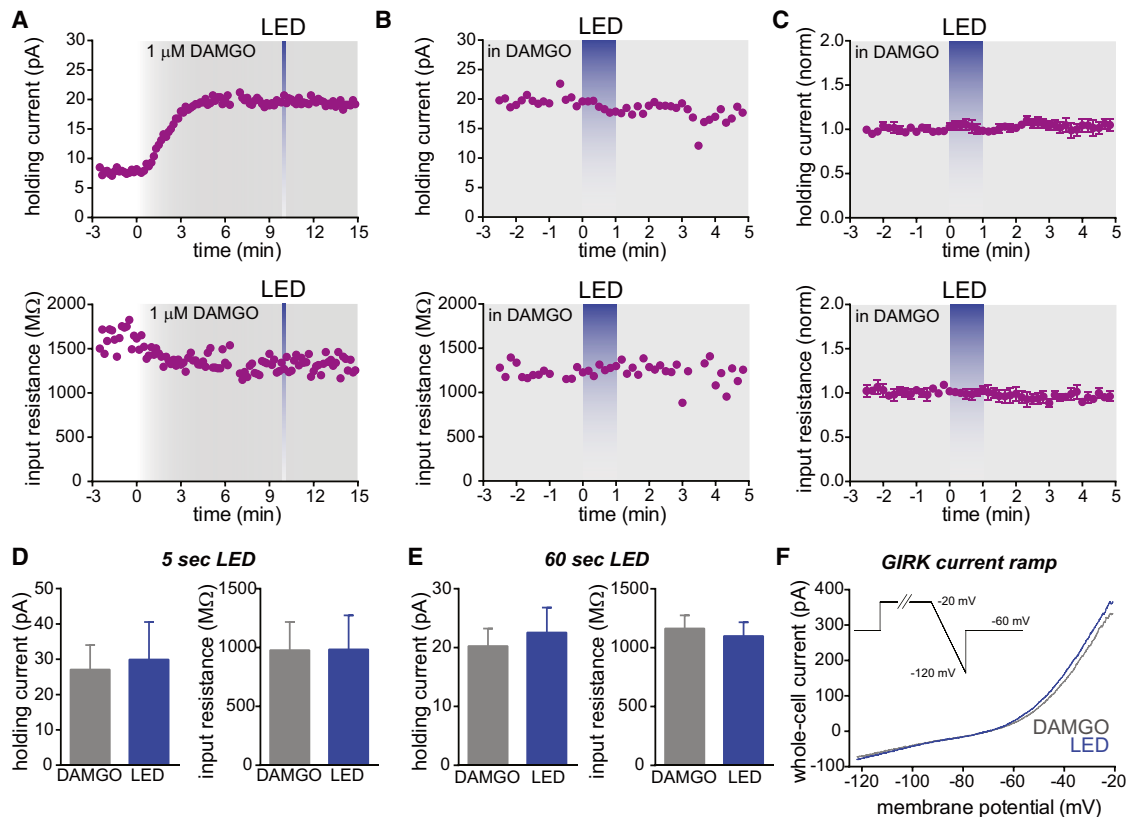


Figure 5. Activation of Endogenous Mu-Opioid Receptors Occludes GIRK Channel Activation by Opto-MORs

(A) Example plots from an opto-MOR and MOPR⁺ GABAergic neuron in the PAG. An outward current and simultaneous decrease in input resistance were observed following prolonged application of 1 μM DAMGO, which plateaued after ~5 min. Brief (5 s) LED illumination did not result in further changes to either of these parameters.

(B) Example plots from a different neuron positive for both opto-MOR and MOPR in the presence of 1 μM DAMGO. The plateaued response to DAMGO is not shown in this example. Prolonged LED illumination (blue, 60 s) did not alter the holding current or input resistance.

(C) Normalized summary graph showing the occluded responses to LED illumination following stimulation of endogenous MOPRs with DAMGO (n = 3).

(D) Quantification of the holding current and input resistance in the presence of DAMGO and following 5 s of LED stimulation (n = 3).

(E) Same quantification as in (D), but with 60 s illumination (n = 3).

(F) Representative GIRK channel ramp in the presence of DAMGO and following LED stimulation. Data are represented as mean ± SEM.

that activation of both receptors is coupled to similar downstream effectors. We also found that activation of both opto-MOR and MOPR substantially decreased neuronal excitability in response to step current injections, which was also reversed by application of barium (Figures 4I and 4J). We did, however, notice a persistent outward current and significantly reduced input resistance following washout of barium (Figures 4K and 4L), indicating that brief activation of opto-MOR can couple to downstream effectors to suppress neuronal activity for an extended period of time.

We next asked whether opto-MOR and MOPR activation engages converging intracellular signaling cascades leading to GIRK channel activation, or whether these two receptor populations signal through parallel subcellular pathways. To test this, we recorded from MOPR⁺/YFP⁺ GABAergic neurons in the PAG and applied saturating concentrations of DAMGO (1 μM). We observed sustained outward currents coupled with a decrease in input resistance (Figure 5A). After this response plateaued, we stimulated neurons with either a brief LED light pulse

(5 s, Figure 5A) or prolonged illumination (1 min, Figures 5B and 5C). We found that opto-MOR coupling to GIRK channel currents and changes in input resistance were occluded by sustained activation of endogenous mu-opioid receptors (Figures 5D–5F), suggesting that activation of opto-MORs converge onto the same pool of intracellular signaling cascades as endogenous mu-opioid receptors. In addition, we injected AAV5-EF1α-DIO-opto-MOR-YFP into the rostromedial tegmental nucleus (RMTg) (AP: -3.9 mm, ML: -0.1 mm, DV: -4.5 mm) of a new cohort of vGAT-IRES-Cre mice and targeted YFP⁺ interneurons for whole-cell recordings. Photostimulation with 470 nm LED light (10 mW/mm²) evoked outward currents together with a decrease in the input resistance (Figures S4A and S4B), suggesting opto-MOR activation is coupled to GIRK channels, which is consistent with the endogenous MOPR function in these neurons (Matsui and Williams, 2011; Matsui et al., 2014). Voltage ramps performed pre- and post-stimulation demonstrated current-voltage curves consistent with the properties of GIRK channels (Lüscher et al., 1997) (Figure S4C).

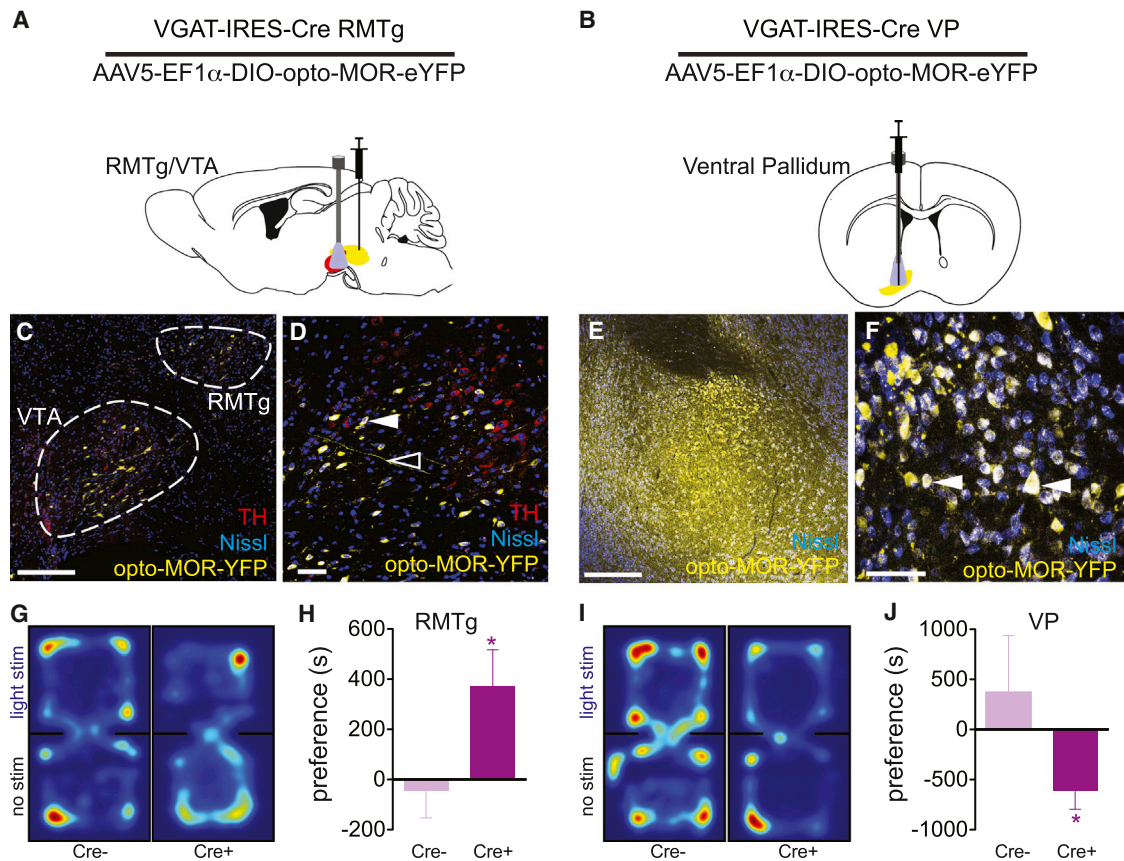


Figure 6. Photostimulation of Opto-MOR Causes MOR-like Behavioral Profiles In Vivo

(A and B) Mice with AAV5-DIO-opto-MOR-YFP injected into the RMTg (yellow) (A) with a fiber optic implanted into the VTA (red) and the ventral pallidum (yellow) (B).

(C) Viral expression in VTA and RMTg.

(D) Viral expression in fibers (open arrows) projecting into the VTA (identified by tyrosine hydroxylase staining in red) as well as GABAergic interneurons of the VTA (closed arrows).

(E) Viral expression in VP.

(F) Viral expression in VP GABA neurons (closed arrows).

(G and H) Mice expressing opto-MOR in the RMTg-VTA ($n = 8$) display significantly increased real-time preference behavior compared to controls (Cre-littermates, $n = 7$) as shown by representative position heatmaps (G) and the mean preference for the stimulation-paired chamber (H).

(I and J) Mice injected into the VP ($n = 16$) display the converse aversion behavior in the place preference assay compared to controls (Cre- littermate controls, $n = 5$) spending less time in the light-stimulated chamber also shown by position heatmaps (I) and mean place preference results (J). * $p < 0.05$ via unpaired t test. Data are represented as mean \pm SEM. See also Figure S5.

Taken together, these results from two different brain regions suggest that opto-MOR couples to canonical MOR G protein signaling pathways, yet retains rapid photo-engagement of opioid signal transduction pathways, making it an ideal tool for spatiotemporal approaches to study MOR signaling in real time in vivo.

In Vivo Photostimulation of Opto-MOR in Selected GABAergic Neurons Mediates Preference or Aversion

To demonstrate the utility of opto-MOR for studying opioid neural circuits in behavior, we locally infused AAV5-EF1 α -DIO-opto-MOR-YFP into the brains of two naive groups of vGAT-IRES-Cre mice, one group in the RMTg (Figure 6A) and one group in the ventral pallidum (VP) (Figure 6B), and waited at least 3 weeks for optimal viral expression at the soma (Figures 6C–6F). Fiber

optic implants (Sparta et al., 2012) were chronically implanted above the VTA and the VP, respectively. Mice were placed into black opaque boxes and allowed to explore both chambers freely to evaluate preference or aversion behavior using the real-time place preference assay. During testing, mice received constant 10 mW 473 nm blue laser light upon entry into the “light stimulation” side (Figures 6G and 6I). Preference and aversion were calculated as the difference in time spent in the light-stimulated box compared to the box with no stimulation, with preference defined as more time spent in the photostimulated box and aversion defined as less time in the photostimulated chamber. vGAT-IRES-Cre+^{opto-MOR:RMTg} mice displayed a significantly greater preference for the stimulation than their Cre(-)-injected opto-MOR littermate controls (Figures 6G and 6H, $t_{13} = 2.231$, $p < 0.05$; Figures S5A and S5B). In addition, based on our data

demonstrating a power response relationship with opto-MOR in activating cAMP and pERK (Figures 1H and S2I), we posited that different light powers might produce different behavioral phenotypes. As such, we focused on real-time place preference in vGAT-IRES-Cre⁺opto-MOR:RMTg mice. We found that animals receiving no light stimulation in the VTA show no preference for either chamber (Figure S5I). However, when we increase the power to 1 mW, animals begin to show a place preference and spend more time in the conditioned chamber than those receiving no light stimulation, but less time than those receiving 10 mW photostimulation (Figure S5I), thus demonstrating that opto-MOR signaling can be more finely manipulated in vivo.

In contrast, mice injected with opto-MOR into the VP (vGAT-IRES-Cre⁺opto-MOR:VP) displayed significant place aversion to photostimulation, as compared to Cre(-) littermate controls (Figures 6I and 6J; $t_{19} = 2.174$, $p < 0.05$; Figures S5E and S5F). Importantly, there were no differences in locomotor activity throughout the session for either group (Figures S5C and S5G), and light stimulation did not induce acute changes in locomotion in the stimulation chamber (Figures S5D and S5H). Finally, as a positive control, and to compare the efficacy of our opto-MOR experiments for a more traditional optogenetic strategy, we tested a new cohort of mice in which we inhibited GABAergic neurons in the VTA using halorhodopsin (eNpH3.0) in the real-time preference assay (Figures S5J–S5L) as previously reported (Jennings et al., 2013). These data were strikingly similar to inhibiting RMTg GABAergic neuronal population with opto-MOR (Figure S5A), suggesting that inhibition of these populations has comparable behavioral phenotypes with alternative optogenetic tools. Taken together, these behavioral results indicate that opto-MOR functions to engage signaling and neuronal inhibition with the ability to elicit robust real-time behavioral responses.

DISCUSSION

Here we describe the development of a new opto-XR receptor based on the mu-opioid receptor. We show that this opto-MOR behaves functionally like its wild-type MOPR counterpart in engaging canonical opioid G protein signaling pathways and receptor internalization. Furthermore, we found that selective expression of opto-MOR in GABAergic neurons within two distinct brain regions promotes diverse behavioral responses consistent with those previously observed with local MOPR agonist infusion. This opto-MOR receptor could potentially also be used alongside chemogenetic approaches, in peripheral studies to assess nociceptive circuits, or as a robust way to inhibit neuronal activity without the limits of the currently used Cl⁻ and H⁺ pump approaches that include halorhodopsin or archaerhodopsin (Berndt et al., 2014; Raimondo et al., 2012; Wietek et al., 2014).

Opto-MOR Mimics Mu-Opioid Receptor Signaling In Vitro

A few other opto-XR receptors have been reported (Airan et al., 2009; Barish et al., 2013; Kim et al., 2005), including an opto- β_2 adrenergic receptor as well as the recently described 5HT1A-like opsin-chimera using the C-terminal components of the wild-type

receptor (Masseck et al., 2014). However, few reports have directly examined the intracellular signaling dynamics of opto-XR signaling as they relate to their wild-type receptor counterpart, nor have these receptors been utilized in native circuits where their endogenous complementary receptor types are expressed. This report shows side-by-side that GPCR Class A rhodopsin-like chimeras can be generated to mimic neuropeptide receptor function in vitro. We found that opto-MOR indeed engages inhibition of cAMP, couples to GIRK outward currents in neurons, and activates ERK rapidly following light pulse exposure (within 2 min), and finally that opto-MOR internalizes rapidly in response to photostimulation. Furthermore, we show the opto-MOR receptors functionally desensitize (Figures 2A and 2B), in a manner similar to that of their wild-type MOPR counterparts, suggesting that these receptors couple to native down-regulation signaling pathways including G protein-coupled receptor kinases (GRKs) and arrestins. In addition, our data suggest that similar to other previously reported GPCRs, these receptors may potentially signal at both membrane and intracellular domains (Irannejad et al., 2013). The possibility therefore exists that opto-MOR may be capable of optical activation while localized intracellularly, not only highlighting an important caveat, but also illustrating the potential utility of this chimera to elucidate components of intracellular signaling cascades. However, future studies to examine opto-MOR regulation of signaling dynamics in native neurons are required, and mutation of the opto-MOR receptor at C-terminal sites previously associated with receptor regulation could potentially provide advantageous optogenetic tools in dissecting MOPR-related signaling and tolerance in vivo.

In Vivo Utility of Opto-MOR for Behavioral Optogenetic Studies

We primarily generated this unique tool to facilitate mimicry of opioid signaling in vivo for use in optogenetic studies. The ability to time-lock activation of mu-opioid-like signaling in an awake-behaving animal provides an alternate approach for mapping the role of opioid receptor signaling in discrete neural circuits and cell populations in real time. In order to demonstrate the utility of this approach in vivo, we chose two brain regions widely known to express high levels of endogenous MOPR that have previously been linked to mu-opioid-dependent reward and aversion. In a first series of experiments, we examined the effects of opto-MOR expression in VTA/RMTg GABAergic neurons using real-time place testing. Recent work had demonstrated that mu-opioid receptors are expressed on both local VTA interneurons as well as a large proportion of nearby RMTg GABAergic cells. It is hypothesized that these receptors likely act to disinhibit dopaminergic activity resulting in MOPR-mediated reward behavior. We demonstrate here that photostimulation of opto-MOR^{vGATCre+} receptors in RMTg neurons produces real-time place preference and reward seeking behavior. This result is consistent with reports using mu-opioid agonists into this region (Jalabert et al., 2011; Zhou et al., 2012; Matsui et al., 2014). In addition, prior reports have demonstrated that optogenetic activation or inhibition of VTA and RMTg GABAergic neurons drives real-time place preference or aversion behavior (Tan et al., 2012; van Zessen et al., 2012), further

suggesting that these VTA neuronal populations are poised to regulate dopaminergic output and behavior.

The ventral pallidum (VP) region is well known to express mu-opioid receptors, and regional activation has been reported to lead to both reward and aversive behavioral responses (Hjelmstad et al., 2013). We therefore expressed opto-MOR in VP GABAergic neurons and selectively targeted coordinates corresponding to the previously reported opioid “hedonic cold spot” (Smith and Berridge, 2005; Smith et al., 2009) in order to determine if activation of opto-MOR signaling in this region could elicit real-time aversion. Consistent with prior reports showing that MOPR agonism in this anterior central region decreases reward seeking (Smith et al., 2009), we observed that opto-MOR activation within these neurons produced avoidance of the photostimulation chamber. These two behavioral assays demonstrate that opto-MOR can elicit robust real-time opioid-like behavioral responses in vivo and highlight their utility for future studies examining circuit and cell-type functions of this pathway.

In addition, these in vitro and in vivo findings coupled with our evidence of coupling to GIRK signaling highlight the utility of opto-MOR for more generalizable optogenetic inhibition studies, such as those that are routinely conducted using halorhodopsin or archaerhodopsin. Opto-MOR couples to native signaling pathways to regulate neuronal activity via GIRKs and might be preferred since it does not come with the potential caveats that the photosensitive chloride pumps (eNpHR3.0) bring with them, including alterations in the reversal potential of the GABA_A receptor, that lead to changes in synaptically evoked spiking activity in the period after photoactivation (Raimondo et al., 2012). Future work utilizing opto-MOR in other neuronal cell types and circuits is warranted, however, for determining its utility as a broadly generalizable optogenetic inhibition approach.

Limitations and Future Directions

Although we have developed a photosensitive mu-opioid-like receptor for the interrogation of opioid neural circuits in vitro and in vivo, there are some potential limitations of this approach that are worth noting. We chose to use vertebrate rhodopsin as a backbone for generation of opto-MOR because it has previously been shown to couple to Gi/o-related signaling (Li et al., 2005) and thus made for an ideal chimeric approach. However, this choice of rhodopsin has limitations since the receptor is very photosensitive, as indicated in our in vitro studies and in vivo behavior. This is an important consideration when using the tool in vitro for slice physiology and circuit analysis, as well as in vivo, when wanting to elicit discrete activation in a particular anatomic subregion. To work around these caveats, we were careful to use the tool under red light conditions in vitro, and while in vivo we kept the fiber implants capped. To resolve this, future variants might be warranted with mutations that render opto-MOR less photosensitive and/or use other short-wavelength opsins as part of the chimeric design.

It is also important to note that the receptor is only partially a mu-opioid receptor and thus might display a different protein-interactome than the wild-type MOPR. Indeed, we show that the receptor utilizes a pool of intracellular signaling similar to that of MOPR (Figures 1 and 2) and traffics to synapses (Figure 3); however, recent work has suggested a complex regulatory pro-

teome of MOPR interactions (Al-Hasani and Bruchas, 2011), and thus we are careful to interpret our data in light of known transmembrane and extracellular loop residues that are important for MOPR function in neurons.

Future studies using the opto-MOR in side-by-side experiments with other exciting recent developments (Banghart and Sabatini, 2012) in optical control of neuropeptide function should provide researchers with a means to better dissect the role of opioid neuropeptide signaling in complex neural circuits and in freely moving mice (Kim et al., 2013). Understanding the relative dynamics of neuropeptide release and receptor signaling is a key question in neuroscience, and opto-XRs that mimic endogenous signaling pathways hold promise in delineating some of these mechanisms. Furthermore, there is a growing need to better understand the molecular-cellular mechanisms of opioid tolerance in pain, and thus the optodynamic properties of this approach may have utility in this arena. The use of this tool will provide an additional layer of spatiotemporal specificity when used with traditional pharmacology, physiology, and behavioral approaches.

Conclusions

In summary, here we provide and characterize a novel tool for neuroscience that allows for spatiotemporal control of opioid receptor signaling in vitro and in vivo. Opto-MORs functionally couple to canonical intracellular signaling pathways and have utility in dissecting opioid contributions to behavioral effect. The combination of opto-XR approaches with chemogenetic and traditional optogenetic channel approaches expands our toolbox for understanding neuropeptide signaling within neural circuits, within real-time, freely moving animals. Furthermore, advances in receptor chimeras such as opto-MOR will facilitate the generation of additional related approaches for other GPCR classes.

EXPERIMENTAL PROCEDURES

Construct Design and Cloning

Using the constraint-based multiple alignment tool (COBALT, NCBI; http://www.st.va.ncbi.nlm.nih.gov/tools/cobalt/re_cobalt.cgi), rat rhodopsin 4 (RO4) sequences were aligned against rat MOR. RO4 extracellular domains, including the retinal binding site, and transmembrane domains were identified and conserved. HindIII/EcoRI sites were added to the N terminus while XhoI/BamHI sites were added to the C terminus of the chimeric sequence. The stop codons were replaced with leucine residues, and silent mutations were added to remove an endogenous XhoI restriction digest site 990 bp into the gene. The construct was synthesized by IDT in a non-expressing pSMART vector (Lucigen) then cloned without stop codons into pcDNA3-YFP (Invitrogen) containing ampicillin resistance cassettes and CMV promoter regions for expression in mammalian cells. Restriction enzyme digest sites EcoRI and XhoI were used to excise the opto-receptor construct from the pSMART vector and purified by gel extraction. pcDNA3-YFP was also digested with EcoRI, and XhoI and T4 Ligase were used to ligate opto-MOR into the vector. *E. coli* was transformed with pcDNA3-opto-MOR-YFP plasmids, plated on ampicillin-containing plates. GeneWiz verified chimeric sequence integrity. Rat MOR-GFP was kindly donated from the lab of Dr. Charles Chavkin at University of Washington, Department of Pharmacology (Celver et al., 2004).

Real-Time Measurement of cAMP Dynamics

Stable HEK cell lines containing opto-MOR and MOR were transfected with the pGloSensor-22F cAMP plasmid (Promega E2301) using JetPrime

(Polyplus) transfection reagent per manufacturer's instructions. Stable co-transfected cells were maintained under both G418 (400 $\mu\text{g/ml}$) and hygromycin (200 $\mu\text{g/ml}$) selective pressure. The day before an experiment, cells were plated on 96-well tissue culture-treated plates (Costar), 60K–100K cells/well, and allowed to recover overnight at 37°C, 5% CO₂. The next day, media was replaced with 2% GloSensor reagent (Promega) suspended in CO₂-independent growth medium (GIBCO) and incubated for 2 hr at 37°C. For real-time cAMP, a baseline was first obtained with no treatment by recording relative luminescent units (RLUs) every 6 s for 1 min using a SynergyMx microplate reader (BioTek). 1 μM forskolin (dashed line in each figure) was applied to each well, and RLUs were recorded for 3 min. Cells were then treated with either a 20 s, 1 mW, blue LED pulse or 1 μM DAMGO (Tocris) and RLUs recorded for an additional 10 min. For data expressed as cAMP (fold baseline), RLUs for 1 min of baseline were averaged and all subsequent RLUs were then divided by this average. For data expressed as cAMP (percent max), raw RLUs were entered into GraphPad Prism (v5.0d, GraphPad Software), and the normalization function was used to assign the lowest RLU a value of 0% and the highest RLU a value of 100%. Time constants were calculated in GraphPad Prism using one-phase association ($Y = Y_0 + (\text{Plateau} - Y_0) \cdot (1 - \exp(-K \cdot x))$) nonlinear regression analyses yielding a time constant value (τ_{on}). For cAMP (percent control) data, area under the curve for each treatment group was normalized to its corresponding experimental control. Each data point is a minimum of three wells per experimental replicate.

Immunoblotting

Immunoblotting for MAPK activation was performed as previously described (Bruchas et al., 2011). Opto-MOR and MOPR expressing cells were grown to 100% confluency in 24-well plates. Cells were serum starved in plain DMEM for at least 4 hr before drug or LED treatment using Plexon LED Driver LD-1 (465 nm). Cells were harvested in 500 μl of lysis buffer containing (in mM) 50 Tris-HCl, 300 NaCl, 1 EDTA, 1 Na₃VO₄, 1 NaF, 10% glycerol, 1% Nonidet P-40, and 1:100 of phosphatase and protease inhibitor mixture set (Calbiochem, EMD Millipore). Lysates were sonicated for 10 s and spun down for 20 min (14,000 $\times g$, 4°C). 25 μg protein was run in each well, and blots were transferred to nitrocellulose (Whatman), blocked with 5% FBS albumen for 1 hr, then incubated with 1:1,000 goat anti-rabbit pERK1/2 (phospho Thr-202/Tyr-204; Cell Signaling) and 1:20,000 goat anti-mouse β -actin (Abcam) antibodies at 4°C overnight. Membranes were washed 4 times for 10 min each in Tris-buffered saline and 0.1% Tween-20 (TBST) and incubated with 1:10,000 IRDyeTM 800- and 700-conjugated affinity-purified anti-rabbit or anti-mouse IgG in a 1:1 mixture of 5% milk/TBS and Li-Cor blocking buffer (Li-Cor Biosciences) for 1 hr at room temperature. Membranes were washed 4 times for 10 min each with TBST, then 2 times with TBS to remove Tween. Blots were visualized, background subtracted, and quantified with Odyssey Infrared Imager (Li-Cor Biosciences). pERK bands were normalized to β -actin from each sample and calculated as fold increase from baseline or as percent of maximal ERK phosphorylation as appropriate.

Receptor Internalization and Confocal Microscopy

Opto-MOR-YFP and MOPR-GFP-expressing cells were plated to approximately 50% confluence on poly-D-lysine-coated coverslips (Becton Dickinson). Following LED (1 W/cm², 1 min) or DAMGO (1 μM) treatment, cells were washed 3 times with PBS and then fixed with 4% paraformaldehyde. Coverslips were then mounted with Vectashield Hard Set mounting medium with DAPI (Vector Laboratories) and imaged on a Leica TCS SPE confocal microscope (Leica), and Application Suite Advanced Fluorescence (LAS AF) software calculated receptor internalization (inside fluorescence/total fluorescence $\times 100$).

Animal Subjects

Group-housed adult (25–35 g) male vGAT-IRES-Cre mice (Vong et al., 2011) and Cre(–) littermate controls were used for all in vivo experiments. For eNpH3.0 VTA GABA experiments (see Supplemental Materials), GAD2-Cre mice were used. All mice were maintained on a 12 hr light:12 hr dark cycle and given ad libitum access to food and water. The Washington

University in St. Louis Animal Studies IACUC approved all experimental methods.

Viral Injections and Surgical Procedures for Fiber Optic Placement

All surgeries were performed under isoflurane anesthesia (Piramal Healthcare, Maharashtra, India). For slice physiology experiments, adult mice were injected bilaterally with AAV5-EF1 α -DIO-opto-MOR-eYFP. For behavior experiments, adult male mice were injected unilaterally with 500 nl of AAV5-EF1 α -DIO-opto-MOR-eYFP virus (WUSTL Hope Center Viral Core, St. Louis, MO) into the RMTg (AP: –3.9 mm, ML: –0.1 mm, DV: –4.5 mm) or the VP (AP: 0.7 mm, ML: 1 mm, DV: –5.25 mm). Fiber optic ferrules were chronically implanted above the VTA (AP: –3.4 mm, ML: –0.5 mm, DV: –4.0 mm) or the VP (AP: 0.7 mm, ML: 1 mm, DV: –4.8 mm), respectively, and dental cement (Lang Dental, Wheeling, IL) was applied to hold the ferrules in place (Sparta et al., 2012). Mice were allowed to recover for at least 3 weeks following infusion of virus and fiber placement prior to further behavioral testing.

Slice Physiology

Acute brain slices were prepared using a protective cutting and recovery method (Ting et al., 2014). Anesthetized mice were transcardially perfused with NMDG-substituted aCSF containing (in mM) 93 NMDG, 2.5 KCl, 1.25 NaH₂PO₄, 30 NaHCO₃, 20 HEPES, 25 glucose, 5 ascorbic acid, 2 thiourea, 3 Na-pyruvate, 12 N-acetyl-L-cysteine, 10 MgSO₄, 0.5 CaCl₂ (pH = 7.3). 200 μm thick coronal sections of the PAG were cut using a Vibratome VT1000s (Leica) and transferred to an oxygenated recovery chamber containing NMDG aCSF for 5–10 min at 32°C–34°C before being transferred to a holding chamber filled with modified aCSF containing (in mM) 92 NaCl, 2.5 KCl, 1.2 NaH₂PO₄, 30 NaHCO₃, 20 HEPES, 25 glucose, 2 CaCl₂, 2 MgCl₂ (pH adjusted to 7.3 with NaOH), and Osm 300-315 and incubated in the dark at room temperature for >1 hr before recording. 225 μm thick sagittal sections were prepared for experiments in RMTg. Additional details can be found in the Supplemental Experimental Procedures.

Behavior

Real-Time Place Preference and Aversion, rtPP/A

Mice were placed into a custom-made black acrylic two-chambered box (52.5 \times 25.5 \times 25.5 cm) and allowed to explore each of two chambers for 60 min. Using Noldus Ethovision hardware controller connected to a master 9 functional generator, light stimulation (473 nm, 10 mW) was delivered through fiber optic implants during the duration of their time spent in the “conditioned” side of the chamber, and mice received no stimulation on the “unconditioned” side. The experimental animals were counterbalanced for both group and conditioning side. Preference and aversion scores in each experiment were determined by comparing the amount of time spent in the conditioned versus unconditioned sides during the real-time testing phase. Preference (RMTg) and aversion (VP) were calculated by comparing time spent in the light stimulation box with time spent in the no stimulation box during the final 30 min of the test session. All behavioral data (locomotion, velocity, entries) were analyzed using Noldus Ethovision (v9.5) with a ZR900 Canon camera.

Immunohistochemistry

As previously described (Kim et al., 2013), after the conclusion of behavioral testing, mice were anesthetized with sodium pentobarbital and transcardially perfused with ice cold PBS, followed by 4% phosphate-buffered paraformaldehyde. Brains were removed, post-fixed overnight in paraformaldehyde, and saturated in 30% phosphate-buffered sucrose. 50 μm sections were cut, washed in 0.3% Triton X-100/5% normal goat serum in 0.1 M PBS, stained with fluorescent Nissl stain (1:400 Neurotrace, Invitrogen) for 1 hr, and mounted onto glass slides with Vectashield (Vector Laboratories). VTA sections were stained with rabbit anti-tyrosine hydroxylase (1:1,000, Millipore) overnight at 4°C and AlexaFluor 633 goat anti-rabbit for 2 hr at room temperature (1:1,000, Molecular Probes) prior to the Nissl step. Opto-MOR expression was verified using fluorescence (Olympus) and confocal microscopy (Leica Microsystems). Images were produced with Leica Application Suite Advanced Fluorescence software. Animals that did not show targeted expression were excluded.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and five figures and can be found with this article online at <http://dx.doi.org/10.1016/j.neuron.2015.03.066>.

AUTHOR CONTRIBUTIONS

E.R.S., B.A.C., and M.J.S. designed and performed experiments, collected and analyzed data, and wrote the manuscript. M.A.B., R.A., S.C.F., and J.G.M. designed and performed experiments and collected and analyzed data. W.J.P. provided technical support and facilitated design, construction, and cloning of the receptor chimera and virus. R.W.G. helped design and oversee experiments. M.R.B. helped design, analyze, and oversee experiments and wrote the manuscript.

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