**μ-Opioid Receptors on Distinct Neuronal Populations Mediate Different Aspects of Opioid Reward-Related Behaviors**

Amie L. Severino,1,* Nitish Mittal,2,3,* Joshua K. Hakimian,1 Nathaniel Velarde,1 Ani Minasyan,1 Ralph Albert,1 Carlos Torres,1 Nicole Romaneschi,1 Camille Johnston,1 Suchi Tiwari,1 Alex S. Lee,1 Anna M. Taylor,1 Claire Gavériaux-Ruff,6,7,8,9,10 Brigitte L. Kieffer,11 Christopher J. Evans,1,5 Catherine M. Cahill,1,5 and Wendy M. Walwyn1,5

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1Department of Psychiatry and Biobehavioral Sciences, UCLA Semel Institute for Neuroscience and Human Behavior, David Geffen School of Medicine, University of California Los Angeles, Los Angeles, CA 90095, 2Division of Pharmacology and Toxicology, College of Pharmacy, University of Texas at Austin, Austin, TX 7871, 3ZS Associates, San Mateo, CA 94402, 4Department of Pharmacology, University of Alberta, Edmonton, Alberta, T6G 2R3 Canada, 5UCLA Brain Research Institute, University of California Los Angeles, Los Angeles, CA 90095, 6Department of Translational Medicine and Neurogenetics, Institut de Génétique et de Biologie Moléculaire et Cellulaire, Illkirch, France, UM7104, 7Université de Strasbourg, Illkirch, France, 67081, 8Centre National de la Recherche Scientifique, Unité Mixte de Recherche 7104, Illkirch, France, 9Institut National de la Santé et de la Recherche Médicale, Unité 964, Illkirch, France, 10Ecole Supérieure de Biotechnologie de Strasbourg, Illkirch, France, CS10413, and 11Department of Psychiatry, Douglas Mental Health Institute, McGill University, Montreal, Quebec, QC H4H 1R3 Canada

**Abstract**

μ-Opioid receptors (MORs) are densely expressed in different brain regions known to mediate reward. One such region is the striatum where MORs are densely expressed, yet the role of these MOR populations in modulating reward is relatively unknown. We have begun to address this question by using a series of genetically engineered mice based on the Cre recombinase/loxP system to selectively delete MORs from specific neurons enriched in the striatum: dopamine 1 (D1) receptors, D2 receptors, adenosine 2a (A2a) receptors, and choline acetyltransferase (ChAT). We first determined the effects of each deletion on opioid-induced locomotion, a striatal and dopamine-dependent behavior. We show that MOR deletion from D1 neurons reduced opioid (morphine and oxycodone)-induced hyperlocomotion, whereas deleting MORs from A2a neurons resulted in enhanced opioid-induced locomotion, and deleting MORs from D2 or ChAT neurons had no effect. We also present the effect of each deletion on opioid intravenous self-administration. We first assessed the acquisition of this behavior using remifentanil as the reinforcing opioid and found no effect of genotype. Mice were then transitioned to oxycodone as the reinforcer and maintained here for 9 d. Again, no genotype effect was found. However, when mice underwent 3 d of extinction training, during which the drug was not delivered,

**Significance Statement**

μ-Opioid receptors (MORs) mediate the effects of the commonly misused and prescribed opioids. These receptors are expressed in different neurons and pathways mediating reward. Although it is well known that μ receptors in the midbrain regulate dopamine release and are important in mediating reward, little is known of the role of other populations that are expressed in the different neurons of the striatum, a hub of many reward pathways. In this study, we deleted selective populations of these receptors that are enriched in the striatum and studied the effect of each deletion on reward-related behaviors. We found that each population plays a specific role in reward demonstrating a more complex role than previously thought of how these receptors mediate reward.
but all cues remained as during the maintenance phase, drug-seeking behavior was enhanced when MORs were deleted from A2a or ChAT neurons. These findings show that these selective MOR populations play specific roles in reward-associated behaviors.

**Key words:** floxed MOR; hyperlocomotion; intravenous self-administration; morphine; μ-opioid receptor; oxycodone

**Introduction**

μ-Opioid receptors (MORs), the principal target of addictive analgesics are widely expressed in diverse brain regions associated with reward (for review, see Le Merrer et al., 2009). MORs are expressed on the GABAergic neurons that innervate the dopaminergic neurons of the ventral tegmental area (VTA) so are poised to enable dopamine release (Ben Hamida et al., 2017; Charbogne et al., 2017) an important mediator of rewarding behavior. MORs are also expressed in the striatum which controls movement and the formation of behavioral habits associated with reward. These two behaviors, reward and locomotion, are mediated by different dopaminergic signaling profiles in distinct neurons (Howe and Dombeck, 2016) and are often used to generate a profile of reward behavior in mice (Mitchell et al., 2005; Zhang and Kong, 2017).

MORs are widely expressed in the different neuronal populations and subregions of the striatum (Wang et al., 1996, 1997; Wang and Pickel, 1998; Miura et al., 2008; Cui et al., 2014). They are expressed on dopamine 1 (D1) receptor, D2, and adenosine 2a (A2a) subpopulations of medium spiny neurons (Cui et al., 2014; Oude Ophuis et al., 2014). They are also expressed on cholinergic interneurons (Pontiero et al., 2013) and on cortical or thalamic glutamatergic neurons innervating medium spiny neurons (for review, see Miura et al., 2008). Within these different neuronal populations, MORs are differentially expressed in striatal sub-regions. For example, they are expressed in the patches or striosomes where they colocalize with dynorphin-expressing D1 medium spiny neurons (Brimblecombe and Cragg, 2017) but also in the matrix where their expression is less and on D1 or D2 medium spiny neurons (Cui et al., 2014).

Although we do not fully understand the functional role of each of the striatal neuronal populations, we do have insight as to their function from their cellular expression patterns and electrophysiology studies. MORs are expressed presynaptically on glutamatergic afferents projecting to the striatum and postsynaptically on striatal dendrites and dendritic spines (Wang et al., 1996). Activation of these receptors inhibits both glutamatergic afferent activity and that of GABAergic collaterals from medium spiny neurons (Blomeley and Bracci, 2011; Ma et al., 2012; James et al., 2013). In addition, MORs inhibit cholinergic interneurons so regulating local spontaneous dopamine release (Sandor et al., 1992; Pontiero et al., 2013; Pontiero et al., 2018). Presynaptic MORs are also found on low threshold spike interneurons and so modulate their spontaneous activity (Elghaba and Bracci, 2017). At the behavioral level, several studies point toward a role of striatal MOR populations in reward behaviors. Earlier studies showed that ablating MOR-enriched striosomes of the dorsal striatum produces deficits in motor-skill learning (Lawhorn et al., 2009). Forebrain MORs are known to play a role in alcohol, food and heroin reward behaviors (Ben Hamida et al., 2017; Charbogne et al., 2017) and in the hedonic reward value of food reward (Boulos et al., 2019). In addition, MOR re-expression on dynorphin expressing medium spiny neurons in an otherwise null background are sufficient to reinstate some, but not all, opioid reward behaviors (Cui et al., 2014).

Given the broad but diverse distribution of MORs on different neuronal subtypes throughout the striatum, we set out to determine the contribution of these MORs to opioid reward behaviors. In order to do this, we bred fMOR mice, in which exons two and three of the MOR gene (oprm1) are flanked by LoxP, with four different Cre-recombinase mice (D1cre, D2cre, A2acre, ChATcre). We first verified these deletions using RNAseq in *situ* mRNA hybridization and quantitative PCR. We then assessed opioid-induced hyperlocomotor, sensitization of this effect, and intravenous opioid self-administration (IVSA). From these studies, we conclude that each of these MOR-expressing populations are required for distinct aspects of opioid reward-related behaviors.

**Materials and Methods**

**Experimental design**

**Subjects**

All procedures were authorized by the Institutional Animal Care and Use Committee (IACUC) and are in compliance with the Policies on the Use of Animals in Research as outlined by this journal. All transgenic mice...
used in this study were bred by the Animal Breeding Colony. D1flMORs, D2flMORS, A2afliMORs and choline acetyltransferase (ChAT) flMORs were generated by breeding flMOR mice (loxP sites flanking exons 2–3 of the oprm1 gene on a 50:50 C57BL/6J:129Sv background, stock #030074, The Jackson Laboratory) with four Cre driver lines to obtain Cre recombinase on one (D1cre; stock #030989-UCD, D2cre; 032108-UCD, A2acre; 036158-UCD, MMRRC, NIH, DHHS, 100% C57BL/6J) or two (ChAT-IRE5-Cre; stock #028861, The Jackson Laboratory, 100% C57BL/6J) alleles and flMOR on both alleles. Control flMOR mice of the same background were generated as littermates from the breeding strategies used. Mice lacking all MORs (stock #007559, 100% C57BL/6J, The Jackson Laboratory) were bred as heterozygous pairs to generate knock-out (KO) and wild-type (WT) littermates. Male and female transgenic mice were used between age 8–32 weeks and 20–36 g of body weight. Animals were maintained on a 12/12 h light/dark cycle with ad libitum access to food and water, and experiments were conducted at ZT4–ZT8 (Zeitgeber Time). All mice were group housed for the duration of the experiment except for the IVSA experiments during which mice were singly housed in an enriched environment after surgery.

**Compounds**

All Schedule II drugs, remifentanil, oxycodone, cocaine, and morphine, were obtained from the NIDA Drug Supply Program (RTI).

**RNA in situ hybridization and light sheet fluorescent microscopy**

Mice were euthanized, their brains removed and flash frozen. All equipment and surfaces were cleaned with RNase inhibitor solution and ISH (Advanced Cell Diagnostics) performed as previously described (Severino et al., 2018). To characterize MOR knock-down in the D1-, D2-, A2a-, and -flMOR mice, the following riboprobes were used: oprm1 (catalog #315841, Atto 550), drd2 (catalog #406501-C2, Alexa Fluor 488), and drd1a (catalog #406491-C3, Atto 647). To characterize MOR knock-down in ChAT-flMOR mice, the same oprm1 and drd1a riboprobes were used as well as a ChAT riboprobe (catalog #408731-C2, Alexa Fluor 488). RNA in situ hybridization was imaged using a 63× oil immersion objective on a Leica SP8 stimulated emission depletin microscope (STED, Leica Microsystems) at the Advanced Light Microscopy Core. The images were compiled in Adobe Illustrator 2019 and brightness and contrast and the tonal adjustments feature uniformly applied across the entire composite image. To determine the extent to which MOR was deleted from specific neuronal types within each of the mouse lines generated, we counted the number of drd1a, drd2, or ChAT-positive cells and then determined the number of these cells that were MOR positive (having a minimum of three grains). The data are expressed as the percentage of MOR-positive cells within each of the subgroups (drd1a, drd2, or ChAT).

**Quantitative reverse transcription-PCR (qPCR)**

qPCR was performed in flMOR, D1-, D2-, A2a-, and ChAT-flMOR mice to define the relative expression levels of oprm1, drd1a, and drd2 using the primers shown in Table 1 and methodology as previously described (Hakimian et al., 2017). Relative ratios comparing conditional KOs to flMOR expression for each gene of interest were calculated by using β-actin as reference gene and the 2^(-ΔΔCt) method to evaluate differential expression levels.

**Open-field locomotion**

Fiberglass open field boxes (28 × 28 × 18 cm) were placed on a horizontal glass pane 71 cm above an infrared camera (caA1300-60gm Basler ace camera) at 250 lux. After 2 d of habituation, mice were placed in the chamber for 15 min followed by a subcutaneous injection of saline or drug and placed back in the chamber for 60 min and their locomotion activity recorded (Ethovision XT10, Noldus). This was repeated at the same time of day for three consecutive days.

**IVSA**

An intravenous catheter (0.2 mm i.d., 0.4 mm o.d., Norfolk Access) was inserted into the right jugular vein of mice under sterile conditions as previously described (James et al., 2013; Storey et al., 2016; Mittal et al., 2017). After 3 d of recovery, the mice began daily self-administration in operant chambers (Med-Associates) for 2 h or 50 reinforcers, whichever came sooner. A two-lever design was used in which the active cue and drug-paired lever, or the inactive lever, was randomly assigned. An active lever press resulted in an intravenous drug infusion (0.67 μl/g body weight) and the presentation of a 10-s tone and visual light cue. Each reinforcer was followed by a 10-s “timeout” period during which no reinforcers could be delivered but presses could be made on either lever. On the first 2 d of this protocol, mouse exploration of the levers was facilitated by placing a drop of 20% sweetened condensed milk on both the active and inactive levers (3 x per session). The mice initially underwent 5–5 d of acquisition training using remifentanil (0.05 mg/kg/infusion) at a fixed ratio of one (one lever press resulted in one infusion, FR1). Oxycodone (0.25 mg/kg/infusion) was then used as the reinforcer for nine consecutive days, the maintenance phase, on the same FR1 schedule. This was followed by extinction training over 3 d during which the mice underwent the same FR1 schedule to a maximum of 50 reinforcers or 2 h, but saline was delivered through the catheter. Catheter patency was tested using an infusion of propofol (20 μl of 1% propofol w/v in saline) every 5 d.

**Table 1: The primer sequences used in the qPCR validation of MOR knock-down**

| Gene primer sequences (5′–3′) | OPRM1 FWD | OPRM1 RVS | DRD1 FWD | DRD1 RVS | DRD2 FWD | DRD2 RVS | ACTB FWD | ACTB RVS |
|-------------------------------|------------|------------|----------|----------|----------|----------|----------|----------|
| OPRM1: μ-opioid receptor, DRD1: dopamine 1 receptor, DRD2: dopamine 2 receptor, ACTB: β-actin control, FWD: forward, RVS: reverse. | TCAAGGCCCTGAGTTTCCGTACCC | CGGGGACACAAATGGCGAAAGAGA | CTTCATGTCGCCGTCGTCGACGGG | GGCATGACAAAGACAACGCCAACAGG | TTGTTCTGGTGTTGTTACTC | TATAGATGATGTTGGTACGC | TGTGCACCTTTATTTTGCTC | GATGATGACGGCTTTTGTGC |

Note: The primer sequences were as follows: OPRM1 (5′-TCAAGGCCCTGAGTTTCCGTACCC-3′), DRD1 (5′-CGGGGACACAAATGGCGAAAGAGA-3′), DRD2 (5′-CTTCATGTCGCCGTCGTCGACGGG-3′), ACTB (5′-GGCATGACAAAGACAACGCCAACAGG-3′), and ACTB (5′-TTGTTCTGGTGTTGTTACTC-3′). The primer sequences were designed to amplify specific regions of the target genes.
Statistical analysis

Power analyses of prior data indicate that power is 0.8 or greater with cell means of $n = 8$ ($n = 12$ used for experiments where animal drop-out rates are expected because of jugular cannula failure, etc.). For experiments where we lacked sufficient prior data for an a priori power analysis, we used prior experience with similar methods to guide us. Although we did use male and female mice, we did not analyze sex as a biological factor as we did not have sufficient power to do so. All experiments included both genotypes with males and females representing 46% and 53%, respectively, of the total number of mice used.

Several analytical methods were used

ANOVA

One-way or two-way ANOVA was used to analyze data obtained from the RNA ISH, qPCR, total locomotion and the intrasession IVSA datasets using Prizm v8 (GraphPad) with further details provided in the results and statistical tables.

Linear mixed models (LMM)

LMM were used to analyze the intrasession locomotion data so as to examine the slope and so rate of change over time of this dataset. We also used LMM with coefficients accounting for random slope or intercept within subjects to define and interpret the intersession IVSA datasets. We used the ImeRTest (Kuznetsova et al., 2017) package in R to run LMM. The linear models were used to assess the effect of time, treatment group, or an interaction of these factors on each variable. The resulting model is a regression equation where the intercept or the slope is allowed to vary for each subject:

$$Y_{\text{Characteristic}} = \beta_0 + \beta_{\text{Group}} X_{\text{Group}} + \beta_{\text{Day}} X_{\text{Day}} + U_{\text{Subject}},$$

where $Y_{\text{Characteristic}}$ is the characteristic being modeled (e.g., distance traveled, lever presses, etc.), each predictor variable is represented by its subscripted $X$, $U_{\text{Subject}}$ represents the random intercept or slope associated with each individual subject. The coefficients ($\beta$) are estimated and assessed for significance. Whenever a significant effect was observed, an ANOVA against a reduced null model was used to assess the impact of the respective factor.

Results

Validation of the selectivity and extent of MOR knockdown in striatal subpopulations

We first defined the selectivity of the loxP/Cre recombinase system by RNA in situ hybridization to examine cell-specific knock-down of the MOR encoding gene ($oprml$) in the dorsolateral striatum. We found that, for cells labeled with the $drd1$ probe, oprml and $drd1$ colocalization was reduced in D1flMORs (representative image, Fig. 1A; quantified expression, Fig. 1B; $p < 0.001$, Table 2, item a) and enhanced in D2flMORs (Fig. 1B; $p < 0.05$, Table 2, item a). For cells labeled by the $drd2$ probe, oprml and $drd2$ colocalization was reduced in D2flMORs (representative image, Fig. 1A; quantified expression, Fig. 1B; $p < 0.01$, Table 2, item b). A2afIMORs showed oprml expression in $drd1^+\text{ cells}$ and a deletion from some, but not all $drd2^+\text{ cells}$ (representative image, Fig. 1A; quantified expression, Fig. 1B; $p < 0.0001$, Table 2, item c).

qPCR was performed to determine overall striatal expression levels of oprml, $drd1$, and $drd2$ in fIMOR in the conditional knock-down strains. We found a loss of oprml $cDNA$ in D1flMORs ($p = 0.0001$) and D2flMORs ($p = 0.015$; Fig. 1C; Table 2, item d) but no other line. There was no compensatory effect of these MOR deletions on $drd1$ (Fig. 1C; Table 2, item e) or $drd2$ (Fig. 1C; Table 2, item f) expression in the different lines.

Selective MOR deletions define specific roles of D1 and A2a MOR populations in opioid-induced hyperlocomotion

Oxycodone

As the analgesic effects of oxycodone may be non-specific (Yang et al., 2016), we first examined the locomotor effect of oxycodone (10 mg/kg, s.c.) in mice lacking MORs in all cells, a global MOR KO, and their WT littermates, (Fig. 2A) over three consecutive days. On day 1, we found no effect of oxycodone in MOR KOs compared with WTs ($p < 0.01$), a lack of effect that did not differ from WTs injected with saline ($p = 0.92$, Table 3, item a). By the third day, the oxycodone locomotor response had sensitized in WTs ($p < 0.001$) but no change was observed in KOs ($p = 0.97$, Table 3, item b). The 5-min timebins of the intrasession data further show oxycodone-induced hyperlocomotion in WT but not KOs and sensitization of this response in only WTs over time (Fig. 2B, $p < 0.01$; Table 3, item c).

We then examined the dose–response relationship of oxycodone using 0 (saline), 1, 3, and 10 mg/kg subcutaneously in each of the genotypes (Fig. 3A). We found no effect of genotype following saline suggesting no effect of these deletions on basal locomotion (Table 4, item a). However, a significant dose by genotype interaction was found following oxycodone ($p < 0.001$, Table 4, item b). (1) Dose. When compared with the saline group of the same genotype, the 1 mg/kg dose of oxycodone had no effect, but 3 and 10 mg/kg of oxycodone induced hyperlocomotion in control fIMORs ($p = 0.007$ and $p = 0.0002$, respectively), A2afIMORs ($p < 0.0001$ for both doses), and following 10 mg/kg in D2fIMORs ($p < 0.0001$) and ChATfIMORs ($p = 0.0013$). However, there was no effect of oxycodone in the D1fIMORs (Table 4, item b). (2) Genotype. D1fIMORs showed a decreased response compared with fIMORs at 10 mg/kg ($p = 0.04$) whereas A2afIMORs showed a greater locomotor response than fIMORs ($p < 0.0001$ for both doses). Neither ChATfIMORs nor D2fIMORs differed from fIMORs (Table 4, item c).

Morphine

Our first experiments examined the dose-dependent locomotor effects of morphine using 0 (saline), 3, 10, and...
**Figure 1.** Validation of the selectivity and extent of MOR knock-down in striatal subpopulations. **Ai,** Representative RNA *in situ* hybridization images for MOR (oprm1 in white), dopamine receptor 1 (D1 or drd1a in red) and dopamine receptor 2 (D2 or drd2 in green), are shown from the dorsolateral striatum of control, flMOR, and D1flMOR, D2flMOR, and A2aflMOR mouse lines. White arrows without a tail demonstrate D1-expressing cells and yellow arrows with a tail show D2-expressing cells. The cells marked by pink arrows in the A2aflMOR images show cells that are oprm1 and drd2 positive. **Aii,** Representative RNA *in situ* hybridization images of the dorsolateral striatum showing oprm1 (white), ChAT (green), and D1 (red) labeling in flMOR and ChATflMOR lines. Arrows highlight ChAT+ cells. Scale bar = 20 μm **(Ai, Aii).** **Bi,** Oprm1 and drd1a expression were quantified and presented as the % colocalization for each genotype of MOR with D1+ cells in **Bi,** MOR with D2+ cells in **Bii,** and MOR with ChAT+ or D1+ cells in **Biii;** *p* < 0.05, **p** < 0.01, and ***p*** < 0.001 versus flMORs of the same experiment. **C,** qPCR for oprm1, drd1, and drd2 cDNA levels in striatal tissue shows reduced oprm1 expression in D1flMORs (*****p*** < 0.001) and D2flMORs (*p* < 0.05) compared with flMORs. There was no effect of these MOR deletions on drd1 (**Cii**) or drd2 (**Ciii**) expression. Refer to Table 2 for statistical analyses. All data are shown as mean ± SEM, and the individual datapoints are shown in Extended Data Figure 1-1, for which this legend also applies.
15 mg/kg (subcutaneously; Fig. 3B). We observed a dose × genotype interaction (p < 0.001, Table 4, item d) as follows. (1) Dose. When compared with the group receiving saline of the same genotype, we found that 15 mg/kg morphine, but not any lower doses, induced hyperlocomotion in flMORs (p = 0.003) and D2flMORs (p < 0.0001). D1flMORs and ChATflMORs showed no response at any dose (Table 4, item d) whereas A2aflMORs showed hyperlocomotion after both 10 and 15 mg/kg (p < 0.0001 for both doses), but not 3 mg/kg. (2) Genotype. Between genotype analysis (Table 4, item e) showed a similar effect of genotype following morphine as oxycodone treatment in that, when compared with flMORs, A2aflMORs showed an enhanced response at the higher doses used, 10 (p = 0.0001) and 15 (p = 0.004) mg/kg, whereas D1flMORs showed a reduced response at 15 mg/kg (p = 0.004), but not 10 mg/kg. Both D2- and ChAT-flMORs were not different from flMORs.

Cocaine

To assess whether the changes in opioid-induced locomotor responses were generalizable to other drug classes, we determined the effect of genotype on cocaine-induced locomotion (15 mg/kg, s.c.; Fig. 3C). We found cocaine-induced locomotion in all genotypes (Fig. 3C, p ≤ 0.001; Table 4, item f) but this effect was enhanced in ChATflMORs (p < 0.0005, Table 4, item g).

Locomotor sensitization

Repeated opioid exposure is well known to induce a sensitization of the initial hyperlocomotor response (Tao et al., 2017). This occurs concurrently with an increase in the incentive motivational properties of a drug and has been considered as a window into this property of drug-seeking behavior (Robinson and Berridge, 1993). To assess the role of each of these MOR populations in this phenomenon, we examined sensitization to oxycodone (10 mg/kg, s.c.), morphine (15 mg/kg, s.c.), and saline, over three consecutive days of drug exposure in all genotypes. The data were analyzed by two-way ANOVA to assess the effect of day and drug on the first and last days of the test. The flMORs showed a genotype × day interaction as both oxycodone (p = 0.002) and morphine (p = 0.02), but not saline, induced sensitization (Fig. 3D; Table 4, item h). The D1flMORs showed no sensitization effect following oxycodone or morphine and this response was not different from saline (Fig. 3E; Table 4, item i). The D2flMORs were similar to flMORs as they sensitized to both oxycodone (p < 0.0001) and morphine (p < 0.0001) but not saline (Fig. 3F; Table 4, item j). The A2aflMORs sensitized to oxycodone (p < 0.0001) but not to morphine or saline (Fig. 3G; Table 4, item k). The ChATflMORs similarly sensitized to oxycodone (p < 0.0001), but not morphine or saline (Fig. 3H; Table 4, item l).

Intrasession locomotor activity

We then defined the locomotion profile induced by each drug with each session using linear mixed model analysis to assess the effect of time and genotype. This was done using 5-min timebins on day 1 and day 3 of 10 mg/kg oxycodone or 15 mg/kg morphine. (1) There was a genotype × time interaction on day 1 of oxycodone (Fig. 3I, p < 0.0001; Table 4, item m). The D2flMORs (Table 4, item n, p = 0.02) and A2aflMORs (p < 0.0001), but not flMORs, D1- or ChAT-flMORs showed a change in locomotor activity within the session. (2) We did not find a timebin × genotype interaction (Table 4, item o) on day 3 of oxycodone. However, the D1flMORs showed decreased activity over time (Fig. 3J, p < 0.001; Table 4, item p), but no other change in activity over time was observed in other lines. (3) There was a genotype × time interaction on day 1 of morphine (Fig. 3K, p < 0.0001; Table 4, item q), with D1flMORs showing a different locomotor profile than flMORs (p = 0.002). Further post hoc analyses showed that flMORs (Table 4, item r, p < 0.0001), D2flMORs (p < 0.0001), A2aflMORs (p < 0.0001), and ChATflMORs (p < 0.002), but not D1flMORs, increased their locomotor activity during the session. (4) We also observed a significant genotype × time interaction on day 3 (Fig. 3L, p < 0.0001; Table 4, item s), with both D1flMOR (p < 0.0001) and ChATflMOR (p < 0.001) showing less activity during the session than

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**Table 2: Statistical analyses of MOR knock-down in D1-, D2-, A2a-, and ChAT-flMOR lines by RNA ISH and qPCR (Fig. 1)**

| Item | Figure | Experiment | Statistical test | Effect or interaction | Main effect | flMOR | D1flMOR | D2flMOR | A2aflMOR | ChATflMOR |
|------|--------|------------|-----------------|----------------------|-------------|-------|---------|---------|----------|-----------|
| a    | 1Bi    | RNA in situ hybridization | One-way ANOVA | Genotype, oprm1 and drd1 probes | F(3,18) = 29.14, p < 0.0001 | Reference genotype | n = 5 | p = 0.0001 | n = 5 | p = 0.02 | N.S. | N.A. |
| b    | 1Bi    | RNA in situ hybridization | One-way ANOVA | Genotype, oprm1 and drd2 probes | F(3,18) = 8.76, p = 0.0013 | Reference genotype | n = 5 | p = 0.004 | n = 5 | p = 0.004 | N.S. | N.A. |
| c    | 1Bi    | RNA in situ hybridization | One-way ANOVA | Genotype, oprm1, drd1 and ChAT probes | F(3,12) = 15.95, p = 0.0001 | Reference genotype | n = 4 | N.A. | N.A. | ChAT+ vs -; p < 0.0001, n = 4 | D1+ vs -; NS |
| d    | 1Ci    | qPCR | One-way ANOVA | Genotype, oprm1 probe | F(3,20) = 8.59, p < 0.001 | Reference genotype | n = 13 | p = 0.0002 | n = 6 | p = 0.015 | N.S. | N.S. |
| e    | 1Ci    | qPCR | One-way ANOVA | Genotype, drd1 probe | F(3,20) = 0.339, p = 0.797, N.S. | Reference genotype | n = 6 | N.S. | n = 5 | N.S. | N.S. | N.S. |
| f    | 1Ci    | qPCR | One-way ANOVA | Genotype, drd2 probe | F(3,20) = 0.925, N.S. | Reference genotype | n = 10 | N.S. | n = 7 | N.S. | N.S. | N.S. |

The RNA probes used were: oprm1 (MOR), drd1 (D1 receptor), drd2 (D2 receptor), and ChAT (cholineacetyltransferase) in flMOR, D1-, D2-, A2a-, and ChAT-flMOR lines. The qPCR probes used were: oprm1 (MOR), drd1 (D1 receptor), drd2 (D2 receptor) in flMOR, D1-, D2-, A2a-, and ChAT-flMOR lines. RNA ISH: RNA in situ hybridization, N.S.: not significant, N.A: not applicable.
the fIMORs. Further post hoc analyses showed that fIMORs (Table 4, item t, \( p < 0.0001 \)), D2fIMORs (\( p < 0.0001 \)), A2afIMORs (\( p < 0.0001 \)), and ChATfIMORs (\( p = 0.03 \)), but not D1fIMORs, increased their locomotor activity during this session.

Selective MOR deletions define specific roles of A2a and ChAT MOR populations in opioid IVSA

Although opioid-induced locomotion and sensitization of this response have been used as an index of reward behaviors (Robinson and Berridge, 1993; Stewart and Badiani, 1993), IVSA is considered as a more direct measure of reward seeking and addiction (Everitt et al., 2018). We therefore examined whether deleting MORs from these neurons altered opioid IVSA through an indwelling jugular catheter under a short-access FR1 schedule. Each of the phases of the IVSA protocol (remifentanil acquisition, oxycodone maintenance and extinction) were analyzed separately and results presented for each of the following four parameters; active and inactive lever presses, reinforcers earned and lever choice as shown by the percent of active lever/total lever presses made.

Remifentanil acquisition

Remifentanil, a fast-acting opioid, was used to establish the association of an active lever press with an opioid infusion and associated cues. During this short acquisition phase, we did not find a genotype \( \times \) day interaction or any main effect of genotype on any of the four parameters measured; (Fig. 4A–D, respectively). However, we found a main effect of day on active lever presses made (\( p < 0.0001 \), \( \chi^2 = 21.017 \); Table 5, item a), reinforcers earned (\( p < 0.0001 \), \( \chi^2 = 19.132 \); Table 5, item b), and percentage active lever presses (\( p < 0.01 \), \( \chi^2 = 9.730 \); Table 5, item c), but not inactive lever presses, showing that all lines acquired this self-administration behavior but there was no effect of genotype.

Oxycodone maintenance

The mice were then transitioned to oxycodone self-administration for 9 d. Compared with those on saline, mice receiving oxycodone made more active lever presses (Fig. 4E, \( p < 0.001 \); Table 5, item d), earned more reinforcers (Fig. 3G, \( p < 0.0001 \); Table 5, item e), and had a higher percentage active lever presses (Fig. 4H, \( p < 0.0001 \); Table 5, item f). There was no difference in the inactive lever presses made between the saline and

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Table 3: Statistical analyses of the hyperlocomotor effects of oxycodone in MOR KO mice (Fig. 2)

| Item | Figure | Experiment | Statistical test | Effect or Interaction | Main effect | WT | \( \mu \) KO |
|------|--------|------------|------------------|----------------------|-------------|-----|------------|
| a    | 2A     | Oxycodone; total locomotion on day 1 | Two-way ANOVA | Genotype, \( \mu \) KO vs WT | \( F_{(2,26)} = 10.78, p < 0.0004 \) | \( n = 7 \), day 1 | \( p < 0.01 \) vs WT oxycodone, \( n = 9 \) |
|      |        |            |                  |                      |             |     |            |
| b    | 2A     | Oxycodone sensitization; total locomotion | Two-way ANOVA | Genotype, \( \mu \) KO vs WT | \( F_{(2,26)} = 5.6, p = 0.0095 \) | \( p < 0.01 \) day 1 vs \( p = 0.97 \) day 1 vs 3, \( n = 9 \) |
|      |        |            |                  |                      |             |     |            |
| c    | 2B     | Oxycodone sensitization; intrasession analysis | Two-way ANOVA | Genotype, \( \mu \) KO vs WT | \( F_{(2,20)} = 1.67, p = 0.015 \) | \( p < 0.01 \) day 1 vs \( p = 0.001 \) day 1 and 3, \( n = 7 \) |

N.S.: not significant, N.A.: not applicable.
oxycodone groups (Fig. 4F). There was no effect of genotype on any parameter.

**Extinction**

Extinction has been shown to increase drug-seeking behavior following oxycodone self-administration (Hakimian et al., 2019). We similarly found that, when compared with saline, all genotypes showed a treatment × day interaction in the number of active lever presses made (p < 0.01; Fig. 4E; Table 5, item g) and reinforcers earned (p < 0.01; Fig. 4G; Table 5, item h), but not inactive lever presses (Fig. 4F) or percentage active lever presses (Fig. 4D) between the last day of oxycodone maintenance and the first day of extinction. Post hoc analyses showed an effect of oxycodone in that mice receiving oxycodone made more active lever presses (p < 0.0001; Table 5, item i), inactive lever presses (p < 0.0001, Table 5, item j) and earned more reinforcers (p < 0.0001; Table 5, item k) on the first day of extinction.
### Table 4: Statistical analyses of the hyperlocomotor effects of oxycodone, morphine, and cocaine in all lines (Fig. 3)

| Item | Figure | Experiment | Statistical test | Effect or interaction | Main effect | LMM genotype | Genotype | Reference genotype | N.S. n = 8 | N.S. n = 8 | N.S. n = 8 | N.S. n = 9 |
|------|--------|------------|-----------------|----------------------|-------------|---------------|----------|------------------|------------|------------|------------|------------|
| a    | 3A     | Oxycodone dose response: dose response (0 mg/kg or saline) | One-way ANOVA | Genotype | $F_{1,149} = 2.54, p = 0.056$ | Reference genotype | n = 8 | N.S. n = 8 | N.S. n = 8 | N.S. n = 8 | N.S. n = 9 |
| b    | 3A     | Oxycodone dose response: dose response | Two-way ANOVA | Genotype $\times$ dose | $F_{1,149} = 3.76, p < 0.001$ | N.S. n = 8–11 | 3 mg; p = 0.56 vs 0, 10 mg; p = 0.0002 vs 0, N = 8–12 | N.S. n = 8 | N.S. p < 0.0001 vs 0, 10 mg; p = 0.0013 | N.S. n = 6–9 |
| c    | 3A     | Oxycodone dose response: genotypetype | Two-way ANOVA | Genotype $\times$ dose | $F_{1,149} = 3.76, p < 0.001$ | Reference genotype | 10 mg; p = 0.04 | N.S. | 3 mg; p = 0.0001, 10 | N.S. | mg; p < 0.001 |
| d    | 3B     | Morphine dose response: dose response | Two-way ANOVA | Genotype $\times$ dose | $F_{1,149} = 5.7, p < 0.001$ | N.S. n = 8–11 | 10 mg; N.S. 15 mg; p = 0.003 vs 0 n = 8–11 | N.S. n = 8–11 | 10 mg; p = 0.0001, 15 mg; p < 0.001 | N.S. n = 6–9 |
| e    | 3B     | Morphine dose response: genotype | Two-way ANOVA | Genotype $\times$ dose | $F_{1,149} = 5.7, p < 0.001$ | Reference genotype | 15 mg; p = 0.004 | N.S. | 10 mg; p = 0.0001, 15 mg; p = 0.004 |
| f    | 3C     | Cocaine; dose response | Two-way ANOVA | Treatment $\times$ genotype | $F_{1,33} = 3.77, p < 0.0073$ | N.S. n = 9; saline n = 9 | p < 0.001 | Cocaine, n = 9; saline n = 9 | N.S. n = 9 | p < 0.0005 cocaine, n = 9; saline n = 9 |
| g    | 3C     | Cocaine; genotype | Two-way ANOVA | Treatment $\times$ genotype | $F_{1,33} = 3.77, p < 0.0073$ | Reference genotype | N.S. for saline and cocaine | N.S. for saline and cocaine | N.S. | Saline; N.S. Cocaine; |
| h    | 3D     | Locomotor sensitization: fIMORs | Two-way ANOVA | Genotype $\times$ day | $F_{2,218} = 3.9, p = 0.049$ | Oxycodone; p = 0.002 morphine; p = 0.018 n = 11 for both saline n = 8 | N.S. for saline and cocaine | N.S. for saline and cocaine | Saline; N.S. Cocaine; | p < 0.0005 |
| i    | 3E     | Locomotor sensitization; D1IMORs | Two-way ANOVA | Genotype $\times$ day | $F_{2,218} = 0.6, p = 0.56$ | N.S. oxycodone n = 8, morphine n = 11 | Oxycodone; p < 0.0001, n = 7, morphine; p < 0.0001 n = 11, saline n = 8 |
| j    | 3F     | Locomotor sensitization; D2IMORs | Two-way ANOVA | Genotype $\times$ day | $F_{2,218} = 12.24, p = 0.0002$ | Oxycodone; p < 0.0001, n = 7, morphine; p < 0.0001 n = 11, saline n = 8 |
| k    | 3G     | Locomotor sensitization; A2aIMORs | Two-way ANOVA | Genotype $\times$ day | $F_{2,218} = 8.23, p = 0.0018$ | Oxycodone; p < 0.0001, n = 11, morphine; p = 0.015, n = 9, saline n = 8 |
| l    | 3H     | Locomotor sensitization; CHATIMORs | Two-way ANOVA | Genotype $\times$ day | $F_{2,218} = 11.53, p = 0.0003$ | Oxycodone; p < 0.0001, n = 8, morphine; N.S., n = 9, saline; n = 9 |
| m    | 3I     | Oxycodone intrasession analysis: Day 1 | LMM | Genotype $\times$ timebin | $\chi^2 = 11.882, p = 0.018$ | N.A. | N.S. n = 9 | N.S. n = 8 | N.S. n = 11 | N.S. n = 9 |
| n    | 3J     | Oxycodone intrasession: Day 1 | LMM | Timebin | $\chi^2 = 31.215, p < 0.0001$ | N.S. n = 11 | p = 0.076, n = 9 | p = 0.006, n = 8 | p < 0.0001, n = 11 | N.S. n = 9 |
| o    | 3K     | Oxycodone intrasession analysis: day 3 | LMM | Genotype | $\chi^2 = 31.215, p < 0.0001$ | N.S. n = 11 | p = 0.076, n = 9 | p = 0.006, n = 8 | p < 0.0001, n = 11 | N.S. n = 9 |
| p    | 3K     | Oxycodone intrasession analysis: day 3 | LMM | Timebin | $\chi^2 = 12.66, p = 0.027$ | N.S. n = 9 | p < 0.001, n = 9 | N.S. n = 8 | N.S. n = 11 | N.S. n = 9 |
| q    | 3L     | Morphine intrasession analysis: day 1 | LMM | Genotype $\times$ timebin | $\chi^2 = 21.239, p < 0.001$ | N.A. | p = 0.002 | N.S. | N.S. |
| r    | 3L     | Morphine intrasession analysis: day 1 | LMM | Timebin | $\chi^2 = 54.796, p < 0.0001$ | N.S. | p < 0.0001 | p < 0.0001 | p = 0.002 |
| s    | 3M     | Morphine intrasession analysis: day 3 | LMM | Genotype $\times$ timebin | $\chi^2 = 32.962, p < 0.0001$ | N.A. | p < 0.0001 | N.S. | p < 0.0001 |
| t    | 3M     | Morphine intrasession analysis: day 3 | LMM | Timebin | $\chi^2 = 64.194, p < 0.0001$ | N.S. | p < 0.0001 | p < 0.0001 | p = 0.03 |

N.S.: not significant, N.A.: not applicable.
versus the last day of maintenance. No such transition effect was observed across any parameter in the saline group.

We then assessed the change in drug-seeking behavior over the 3 d of extinction in mice that had received oxycodone using LMM analysis. We found no genotype × day interaction, however there was a main effect of genotype on reinforcers earned (Fig. 4G, p < 0.05; Table 5, item n) with ChATIMORs (p < 0.01) and A2afIMORs (p < 0.05) earning more reinforcers than flMOR mice over these 3 d. There was a trend toward a main effect of genotype for
Table 5: Statistical analyses of the IVSA profile in all lines (Fig. 4)

| Item | Figure | Experiment | Statistical test | Effect or interaction | Main effect | fMOR | D1MOR | D2MOR | A2aMOR | ChATfMOR |
|------|--------|------------|------------------|----------------------|-------------|------|-------|-------|--------|----------|
| a    | 3A     | Remifentanil acquisition | LMM | AL day effect | \( p < 0.0001, \chi^2 = 21.017 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 12 \) | \( n = 12 \) |
| b    | 3C     | Remifentanil acquisition | LMM | RNFS earned day effect | \( p < 0.0001, \chi^2 = 19.132 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 12 \) | \( n = 12 \) |
| c    | 3D     | Remifentanil acquisition | LMM | Percent AL presses day effect | \( p < 0.01, \chi^2 = 9.730 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 12 \) | \( n = 12 \) |
| d    | 3E     | Oxycodone maintenance | LMM | AL presses × Treatment | \( p < 0.001, \chi^2 = 10.926 \) | \( n = 16 \) | \( n = 14 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) |
| e    | 3G     | Oxycodone maintenance | LMM | RNFS × treatment | \( p < 0.0001, \chi^2 = 9.480 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| f    | 3G     | Oxycodone maintenance | LMM | Percent AL presses treatment × day | \( p < 0.0001, \chi^2 = 29.255 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| g    | 3F     | Extinction; transition | LMM | AL presses treatment × day | \( p < 0.01, \chi^2 = 8.296 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| h    | 3G     | Extinction; transition | LMM | RNFS treatment × day | \( p < 0.01, \chi^2 = 7.547 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| i    | 3E     | Extinction; transition | LMM | AL presses day effect | \( p < 0.0001, \chi^2 = 29.255 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| j    | 3F     | Extinction; transition | LMM | IAL presses day effect | \( p < 0.01, \chi^2 = 9.396 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| k    | 3G     | Extinction; transition | LMM | RNFS day effect within oxycodone treated | \( p < 0.0001, \chi^2 = 25.725 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| l    | 3G     | Extinction | LMM | AL presses × genotype within oxycodone treated | \( p = 0.05, \chi^2 = 9.480 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| m    | 3G     | Extinction | LMM | IAL presses × genotype within oxycodone treated | \( p = 0.05, \chi^2 = 9.148 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| n    | 3G     | Extinction | LMM | RNFS earned × genotype within oxycodone treated | \( p = 0.01, \chi^2 = 11.934 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| o    | 3G     | Extinction | LMM | RNFS earned day effect | \( p < 0.0001, \chi^2 = 12.979 \) | \( n = 15 \) | \( n = 12 \) | \( n = 10 \) | \( n = 8 \) | \( n = 7 \) |
| p    | 3/     | Last day AL | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 0.45, \chi^2 = 3.135 \) | Reference N.S. | N.S. | N.S. | \( p = 0.031 \) | \( p = 0.007 \) |
| q    | 3L     | Last day oxycodone | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 0.2, \chi^2 = 2.26 \) | Reference N.S. | N.S. | N.S. | \( p = 0.05 \) | N.S. |
| r    | 3J     | First day extinction AL | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 1.44, \chi^2 = 3.946 \) | Reference N.S. | N.S. | N.S. | \( p = 0.05 \) | N.S. |
| s    | 3M     | First day extinction RNFS | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 1.66, \chi^2 = 2.02 \) | Reference N.S. | N.S. | N.S. | \( p = 0.05 \) | N.S. |
| t    | 3J     | Third day extinction AL | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 0.42, \chi^2 = 2.02 \) | Reference N.S. | N.S. | N.S. | \( p = 0.05 \) | N.S. |
| u    | 3M     | Third day extinction RNFS | Two-way | Genotype × time | ANOVA | \( F_{(55)} = 0.2, \chi^2 = 2.26 \) | Reference N.S. | N.S. | N.S. | \( p = 0.05 \) | N.S. |

N.S.: not significant, N.A.: not applicable.

active lever presses (Fig. 3E, \( p = 0.0507 \); Table 5, item l) and percentage active lever presses (Fig. 4H, \( p = 0.0571 \); Table 5, item m), with A2aMORs showing increased active lever presses (\( p < 0.05 \)) and percentage active lever presses (\( p < 0.05 \)) made over these 3 d than the flMORs. We also observed a main effect of day on reinforcers earned (Fig. 4G, \( p < 0.0001 \); Table 5, item o) with all mice showing a decrease in reinforcers earned over the 3 d of extinction with no effect of genotype. No other effects were found for active lever presses, inactive lever presses and percentage active lever presses across these 3 d.

**Intrasession analysis**

We also analyzed the cumulative frequency of active lever presses and reinforcers earned during the 2-h test on three specific days of the IVSA protocol (Fig. 4f–n). The first of these days, day 9 of the maintenance phase and the last day of oxycodone self-administration, showed a lack of genotype effect on either the cumulative active lever presses (Fig. 4f; Table 5, item p) or reinforcers earned (Fig. 4j; Table 5, item q). However, on the next day assessed, extinction day 1, A2aMORs showed an increase in cumulative active lever presses (Fig. 4k, \( p < 0.05 \); Table 5, item r) and reinforcers earned (Fig. 4l, \( p < 0.05 \); Table 5, item s). ChATfMORs also earned more reinforcers than flMORs on this day (Fig. 4l, \( p < 0.05 \); Table 5, item s). By the third day of extinction, there was no effect of genotype on cumulative active lever presses (Fig. 4m; Table 5, item t), but there was an effect of genotype on cumulative reinforcers earned with ChATfMORs earning more reinforcers than flMORs during the last 40 min of the test (Fig. 4n; Table 5, item u).
Discussion

These findings outline distinct roles for MORs on neuronal populations in behaviors associated with opioid-induced locomotion and reward behaviors. These are that selective ablation of MORs from D1 receptor-expressing neurons prevents opioid-induced locomotor hyperactivity as well as locomotor sensitization but has no effect on opioid IVSA. Second, removal of MORs from A2a neurons enhances opioid-induced hyperlocomotion, locomotor sensitization and drug-seeking behaviors during extinction following opioid IVSA. Third, ablation of MORs from ChAT neurons results in an agonist-dependent hyperlocomotor effect whereby morphine fails to elicit dose-dependent locomotor hyperactivity or sensitization yet oxycodone-induced effects are similar to control flMOR mice. These mice also show an increase in drug-seeking behavior during extinction. Fourth, despite the common theory that A2a receptor expression is equivalent to D2 receptor expression in medium spiny neurons, our data suggests that the A2a cre deletes MORs from only a subset of D2 medium spiny neurons and, that, in stark contrast to MOR deletion from A2a neurons, MOR deletion from D2 neurons results in no discernible change in these reward-based behaviors (Fig. 5A).

Our study shows that MORs on D1 neurons are required for the initial locomotor and sensitization response to morphine and oxycodone. The effect of morphine is in line with a previous study in which the expression of MORs in only D1 neurons in striatal patches in an otherwise null background reinstated morphine-induced locomotion (Cui et al., 2014). Together these 2 findings demonstrate both the requirement and necessity of this MOR population for this striatal-mediated output. This may be a result of MORs on D1 recurrent collaterals inhibiting D2 neurons to reduce striatal output and attenuate the motor effect of opioids, as modeled in Figure 5B. Another possibility is that these receptors are required for the release of dopamine in the VTA (Cui et al., 2014), which is required for this response (Steidl et al., 2017). In regards our IVSA findings, the lack of effect of the D1 MOR deletion in the acquisition of oxycodone IVSA is in contrast with previous work (Cui et al., 2014), perhaps as other MOR populations such as those within the matrix, are also involved in the acquisition phase of this behavior. It is also possible that this is an example of an opioid-specific effect in which the faster-acting opioid, remifentanil, used in (Cui et al., 2014), results in greaterleverpressing behavior than oxycodone.

As D2 receptors are expressed on cholinergic interneurons (Weiner et al., 1991), the A2a cre line has been used to selectively target D2 medium spiny neurons (Fink et al., 1992; Rosin et al., 2003; Wang et al., 2019). Our findings show that this A2a-MOR population is an apparent subset of D2 medium spiny neurons that controls the locomotor sensitivity to oxycodone and morphine and drug-seeking behavior during extinction. These inhibitory receptors may be on some D2-D1 collaterals (Taverna et al., 2008), where their deletion allows an earlier threshold to be reached to increase striatal motor output, as modeled in Figure 5B. As MORs on cholinergic interneurons remains intact and, surprisingly, MORs are also present on some D2 striatal neurons, their deletion displays a remarkably different and striking phenotype from D2flMORs. This could reflect a role of this striatal population or an extrastriatal neuronal population that expresses both A2a and μ opioid but not necessarily D2 receptors. As regards MORs on D2+ neurons, we find that these receptors influence neither opioid-induced locomotion nor opioid IVSA.

While deleting MORs from D1, D2 and A2a neurons was performed to identify their role in GABAergic striatal neurons, deleting MORs from cholinergic interneurons examines the role of these receptors in altering cholinergic neuronal activity. These neurons form 1–3% of the striatal population yet they are remarkably influential in controlling striatal circuits (Gritton et al., 2019) and output, and both MORs and δ-opioid receptors strongly inhibit their activity to affect behavior (Bertran-Gonzalez et al., 2013; Ponterio et al., 2013). Activation of MORs could affect glutamate or acetylcholine release and subsequent dopamine release from nearby terminals (Yorgason et al., 2017) to alter the activity of local circuits (for review, see Clarke and Adermark, 2015; Berke, 2018). Omission of an expected reward induces a dip in dopamine release, a negative reward prediction error (RPE) accompanied by a pause in cholinergic interneuron activity (Hart et al., 2014) to affect local D1 and D2 medium spiny neuron activity (Mamaligas and Ford, 2016). Deleting MORs from these neurons may prevent the encoding of an RPE and facilitate drug-seeking, as shown by an increase in cue-induced reinforcers earned, but not active lever presses, during extinction (Fig. 4M,N).

The rapid increase in hyperlocomotion following oxycodone and the sustained, gradual increase in hyperlocomotion following morphine (Fig. 3F–L) is likely because of the different plasma-kinetic (PK) profiles of these two drugs. Oxycodone has a higher percentage of unbound drug in the blood and a 100-fold greater influx rate than morphine (Boström et al., 2008). This results in a 6-fold higher ratio of unbound oxycodone in the brain: blood and a higher unbound steady state in the brain (Boström et al., 2006, 2008) likely explaining the larger increase in dopamine release following intravenous oxycodone than intravenous morphine (Vander Weele et al., 2014). The ligand-dependent and genotype-dependent effect of morphine but not oxycodone in ChATflMORs further suggests that this receptor population is more sensitive to the PK profile of each ligand. This could be because of a time-dependent effect of these receptors in modulating intrinsic cholinergic interneuron activity and the control of local circuitry.

There are several limitations of this study. One is that we have used the loxP-Cre recombinase system to achieve developmental deletion of MORs from various neuronal populations (Gong et al., 2007). For the most part these populations are striatal where the co-expression of MORs with D1 or D2 receptors can be used to define different medium spiny neuron populations (Gerfen et al., 1990; Weiner et al., 1991). However, dopamine neurons project to various brain regions in addition to the striatum, the hippocampus, amygdala, and prefrontal...
cortex. The behavioral outcomes in this study may therefore be influenced by MOR expression on dopamine circuits outside the striatum. For example, MOR expression on the intercalated neurons of the amygdala (Gregoriou et al., 2019), and in the globus pallidus (Weiner et al., 1991; Delfs et al., 1994) may influence these reward-related behaviors. MOR expression on cholinergic neurons of the medial habenula (Gardon et al., 2014) may also influence...
reward behaviors (Boulos et al., 2020) and MORs and ChAT co-expression in secretomotor neurons of the colon suggests gut function may be altered in ChAT/MORs (Galligan and Akbarali, 2014). Further studies could also assess the role of MORs in different striatal subregions such as in patches or matrix, dorsal ventral striatum and co-expression with both D1 and D2 receptors (Soares-Cunha et al., 2016). An additional limitation is that we did not assess the effect of the cre insertion alone as this would have required further back-crossing of all lines.

Striatal D1 and D2 neurons are traditionally considered to have opposing effects on striatal motor patterns resulting in a coordinated motor activity. In this simple model, activating D1 neurons of the direct pathway increases striatal output to facilitate movement whereas activating D2 neurons of the indirect pathway inhibits competing motor patterns and inhibits movement (Kravitz et al., 2010). This model has been expanded and developed to include several interacting factors that influence the threshold of these outputs by recurrent collaterals between D1 and D2 neurons (Bahuguna et al., 2015), regulation by different interneurons (Taverna et al., 2008), and the regional and compartmental expression patterns of D1 and D2 (Cui et al., 2014; Oude Ophuis et al., 2014).

Nevertheless, the opposing and complimentary effects of medium spiny neuron activation remains a central component of their activity. We show that the effect of deleting MORs from D1 and A2a neurons resembles such complementation, albeit the inverse, as it is the absence of MORs from D1 or A2a neurons that reduces or facilitates motor output, respectively. We propose that this can be explained by the presence of these Gβγ-coupled receptors on recurrent medium spiny neuron collaterals, as shown by the schematic model in Figure 5B. The roles of D1 and D2 medium spiny neurons in mediating reward are also seen as divergent yet complementary in that D1 neurons mediate drug reinforcement and positive reward behaviors, whereas the D2s mediate aversion or ambivalence and are active during withdrawal (Koo et al., 2014; Cole et al., 2018). In addition, D1 and D2 receptors also play complementary but opposing roles in learning value-based and motivated behaviors, an important component of the change in reward value during extinction (Verharen et al., 2019). In regards the role of MORs on these neurons, we show that rather than mediating positive reinforcement during the initial stages of opioid reward, that it is MORs on A2a or ChAT neurons that are important in controlling drug seeking during extinction, a period of increased anxiety and negative affect (Carmack et al., 2019). Additional studies to further define the effect of these deletions on A2a or ChAT neurons under different physiological conditions such as an increase in stress following periods of abstinence, or chronic pain, are needed to enhance our understanding of the complex and interrelated roles of these MOR populations.

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