Mu and delta opioid receptors oppositely regulate motor impulsivity in the signaled nose poke task.
Mary Olmstead, Abdel-Mouttalib Ouagazzal, Brigitte Kieffer

To cite this version:
Mary Olmstead, Abdel-Mouttalib Ouagazzal, Brigitte Kieffer. Mu and delta opioid receptors oppositely regulate motor impulsivity in the signaled nose poke task.. PLoS ONE, Public Library of Science, 2009, 4 (2), pp.e4410. 10.1371/journal.pone.0004410. inserm-00370156

HAL Id: inserm-00370156
https://www.hal.inserm.fr/inserm-00370156
Submitted on 24 Mar 2009

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Mu and Delta Opioid Receptors Oppositely Regulate Motor Impulsivity in the Signaled Nose Poke Task

Mary C. Olmstead1*, Abdel-Mouttalib Ouagazzal2, Brigitte L. Kieffer2

1 Department of Psychology, Queen’s University, Kingston, Ontario, Canada, 2 Institut de Génétique et de Biologie Moléculaire et Cellulaire, Département Neurobiologie, Illkirch, France

Abstract

Impulsivity is a primary feature of many psychiatric disorders, most notably attention deficit hyperactivity disorder and drug addiction. Impulsivity includes a number of processes such as the inability to delay gratification, the inability to withhold a motor response, or acting before all of the relevant information is available. These processes are mediated by neural systems that include dopamine, serotonin, norepinephrine, glutamate and cannabinoids. We examine, for the first time, the role of opioid systems in impulsivity by testing whether inactivation of the mu- (Oprm1) or delta- (Oprd1) opioid receptor gene alters motor impulsivity in mice. Wild-type and knockout mice were examined on either a pure C57BL6/J (BL6) or a hybrid 50% C57BL6/6J-50% 129Sv/pas (HYB) background. Mice were trained to respond for sucrose in a signaled nose poke task that provides independent measures of associative learning (responses to the reward-paired cue) and motor impulsivity (premature responses). Oprm1 knockout mice displayed a remarkable decrease in motor impulsivity. This was observed on the two genetic backgrounds and did not result from impaired associative learning, as responses to the cue signaling reward did not differ across genotypes. Furthermore, mutant mice were insensitive to the effects of ethanol, which increased disinhibition and decreased conditioned responding in wild-type mice. In sharp contrast, mice lacking the Oprd1 gene were more impulsive than controls. Again, mutant animals showed no deficit in associative learning. Ethanol completely disrupted performance in these animals. Together, our results suggest that mu-opioid receptors enhance, whereas delta-opioid receptors inhibit, motor impulsivity. This reveals an unanticipated contribution of endogenous opioid receptor activity to disinhibition. In a broader context, these data suggest that alterations in mu- or delta-opioid receptor function may contribute to impulse control disorders.

Citation: Olmstead MC, Ouagazzal A-M, Kieffer BL (2009) Mu and Delta Opioid Receptors Oppositely Regulate Motor Impulsivity in the Signaled Nose Poke Task. PLoS ONE 4(2): e4410. doi:10.1371/journal.pone.0004410

Editor: Kenji Hashimoto, Chiba University Center for Forensic Mental Health, Japan

Received September 3, 2008; Accepted December 24, 2008; Published February 9, 2009

Copyright: © 2009 Olmstead et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was funded by the Natural Sciences and Engineering Research Council of Canada (NSERC), the Centre National de la Recherche Scientifique (CNRS), the Institut National de la Santé et de la Recherche Médicale (INSERM), the Université Louis Pasteur (ULP), the National Institute of Drug Abuse (NIH-NIDA #DA 05010) and the European Consortium GENADDICT # LSHM-CT-2004-005166. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: olmstead@queensu.ca

Introduction

Impulsivity is a behavioral trait that varies across the general population. Extreme manifestations of impulsivity are revealed in a variety of pathological conditions including antisocial and borderline personality disorders, attention deficit hyperactivity disorder (ADHD), pathological gambling, eating disorders, obsessive-compulsive disorder and substance abuse [1]. Patients with neurological impairments such as Parkinson’s disease, Schizophrenia, Tourette’s syndrome and frontal lobe dementia also present with clinical features of impulsivity. The pervasiveness of this trait across patient populations, and the fact that it is a significant predictor of therapeutic efficacy for some disorders [2–4], provide compelling arguments for understanding the neuropharmacology of impulsivity.

Altered serotonergic transmission has long been associated with impulse control disorders [5], and an important role for dopamine (DA) has been inferred from the therapeutic efficacy of psychostimulants in the treatment of ADHD [6]. Moreover, in humans, polymorphisms in serotonin (5-HT) and DA transporters or receptors is associated with ADHD and addiction [7]. Animal research has confirmed an important role for both DA and 5-HT in impulsivity, and also implicated norepinephrine, glutamate, and cannabinoid systems [8]. To date, no one has investigated the role of opioid systems in impulsivity using animal models.

Opioid receptors have been studied extensively in relationship to drug abuse. The opioid system consists of endogenous neuropeptides that produce effects by acting at mu, delta and kappa opioid receptors. Deletion of the mu-opioid receptor gene (Oprm1) in mice reduces or eliminates the rewarding properties of opioids as well as non-opioid drugs such as ethanol, cocaine, nicotine and cannabinoids. The consistency of behavioral effects across different classes of abused drugs led to the hypothesis that mu-opioid receptors represent a gateway to drug addiction [9]. Unlike mu-opioid receptor knockout mutants, mice lacking the delta opioid receptor gene (Oprd1) show intact cannabinoid-induced reward [10] and increased self-administration of ethanol [11]. These mice exhibit increased depressive-like behaviors and higher basal anxiety levels [12], the latter of which is reversed following ethanol intake [11]. Delta opioid receptors, therefore, influence emotional responses and this, in turn, impacts on drug taking behaviors.
Drug addiction is a complex state which results from gradual adaptations of the brain to repeated drug exposure. The rewarding properties of drugs promote initial drug use when drugs are consumed voluntarily but, as the disorder develops, addicts lose control of their behavior and drug intake becomes independent of drug reward [13,14]. Impulsivity is a critical feature of this state in that addicts are unable to inhibit their drug-taking responses even if the subsequent rewarding effects of the drug are minimal. The question then arises, whether mu- and delta-opioid receptors, in addition to their roles in reward and emotional processing, also regulate impulsivity. Mu and delta knockout animals exhibit marked opposing phenotypes in a number of tasks (locomotor activity, dark-light box, elevated plus maze, forced swim); in contrast, kappa receptor knockout mice are comparable to wild-type controls in these tests [12]. Thus, as a first step in studying the role of opioid receptors to motor impulsivity, we elected to examine mu and delta mutant lines. To this aim, we tested whether deletion of the Oprm1 or the Opend gene alters motor impulsivity in mice. We used a signaled nose poke task [15] that provides independent measures of impulsivity (the inability to withhold a prepotent response) and conditioned reward (approach responses to a light previously paired with a sucrose reward). Mice are required to withhold responding for a liquid sucrose reward until a visual cue is presented (Fig. 1). Impulsivity in this task is comparable to premature responding in the 5-choice serial reaction time task or disinhibition in the go/no-go task, two standard measures of impulsivity in rodents. Compared to these other tests, however, the signaled nose poke task is acquired rapidly and optimal performance does not rely as heavily on attentional (5-choice) or discriminative (go/no-go) abilities. Importantly, performance in the signaled nose poke task seldom reaches asymptotic levels, allowing both increases and decreases in impulsivity to be easily detected. Given the role of endogenous opioids in the neuropharmacological effects of ethanol [16], and the common assumption that ethanol intoxication is associated with impulsive behavior in humans [17], we performed one additional test in which we evaluated whether ethanol alters performance of wild-type or knockout mice in the signaled nose poke task.

Materials and Methods

Subjects

Mu- and delta-opioid receptor knockout mice and their wild-type controls were bred in-house and genotyped a few days after birth. Mice were bred on either a hybrid 50% 129SVPas 50% C57Bl6j background (HYB), (the original background of opioid receptor knockout mice) or backcrossed onto the C57Bl6j background (classically used in mouse behavioral studies) for more than 12 generations (BL6). A detailed description of the construction of these lines and their genotyping has been described previously [12,18].

At 112–126 days of age, mice were transferred from the breeding facility to the phenotyping centre where they were group housed on a 12-h light/dark cycle with lights on at 700 am. Food was available ad libitum throughout the experiment; with the exceptions noted below; access to water was restricted to 2 h per day beginning 2–4 h after testing. Behavioral testing commenced at 17–20 weeks of age and finished at 26–33 weeks of age. All animals were weighed twice per week; there were no significant group differences in weight across the experiment. Animal care was conducted in accordance with the European Communities Council Directive of 24 November 1986 (86/609/EEC) and the experiments were approved by the Comité régional d’éthique en matière d’experimentation animale de Strasbourg, CREMEAS, 2003-10-08-[1]-58.

Apparatus

Impulsivity testing was conducted in operant boxes (26.5×22.0×20.0 cm) housed in a sound-attenuating chamber (Coulbourn Instruments, Allentown PA). Each box was fitted with two nose poke response holes, a cue light, a houselight and a liquid dipper that was accessed through a recessed magazine. Sucrose delivery was signaled by an infrared light located 4 cm above the magazine. Nose pokes and entries to the magazine were detected by infrared beams crossing each opening vertically. The control of stimuli and recording of responses were managed by an IBM-type computer using Graphic State Notation 2 software.

Sucrose and water consumption were measured in 24 automated chambers (11×22×19 cm) made of wire mesh floor and Plexiglas sidewalls (Imetronic, Pessac, France). Each cage was fitted with infra-red captor located 2 and 8.5 cm from the floor, allowing measurement of ambulatory locomotor activity and rears, respectively. Water and sucrose consumption were measured using an automated lickometer.

Behavioral procedures

Signaled nose poke task. Mice were trained in the signaled nose poke task as described previously [15] with minor modifications. These included the cue signaling reward availability (5 s visual stimulus rather than a 3 s auditory stimulus) and the reward (5 s presentation of liquid sucrose rather than the delivery of a single sucrose pellet). The sessions were increased from 30 to 40 min to accommodate these changes.

Figure 1. The progression of each trial during phase 4 of the signaled nose poke task. Each trial started with a 20 s ITI; responses during this period had no consequence. Nose pokes during the subsequent pre-CS period (VT 1–8 s) reinitialized the ITI. If mice refrained from responding during the pre-CS period the trial progressed to the 5 s cue presentation. Nose pokes in the pre-assigned reinforcement hole during the cue presentation turned off the cue and immediately elevated the sucrose dipper. Nose pokes in the non-reinforced hole had no consequence.

CS = conditioned stimulus; ITI = inter-trial interval; VT = variable time.

doi:10.1371/journal.pone.0004410.g001
One day prior to magazine training, and following 21 h of water deprivation, a sucrose solution (25% w/v) was presented in the home cage for one hour. Beginning the next day, all mice underwent magazine training (2 sessions) consisting of 30 presentations of liquid sucrose, presented at random intervals approximately every 30 seconds. During sucrose presentation, the houselight was turned off and the magazine light was turned on. Magazine entries were recorded and used as an indicator that mice had learned the location of the sucrose reward. The liquid dipper remained elevated for 5 s after a magazine entry to a maximum of 20 s per sucrose presentation.

Subsequent training was conducted in 4 phases with one training session per day. In Phase 1, mice were reinforced for nose poking on a fixed-ratio (FR1) schedule of reinforcement on either the right or left side (assignment of reinforced hole counterbalanced across groups). The discriminative stimulus (i.e., the visual cue) was turned on throughout these sessions, except during presentation of the reinforcer (5–20 s elevation of the sucrose dipper). Training continued until 25 reinforcers were presented in one session. In Phase 2, response requirements were increased to an FR3 schedule and the elevation of the sucrose dipper was limited to 5 s. Training continued until 25 reinforcers were presented in one session. In Phase 3, the discriminative stimulus was reduced to 5 s and presented 50 times with a fixed inter-trial-interval (ITI) of 30 s; only those responses occurring on the reinforced side during the cue were reinforced (FR1) by a 5 s elevation of the sucrose dipper. Training progressed to Phase 4 when animals made 10 reinforced nose pokes (i.e., during the cue) in one session. Animals that did not reach the training criterion in Phases 1, 2, or 3 after 20 sessions were removed from the experiment, and data from these sessions was not included in the statistical analyses. In Phase 4, the ITI was set at 20 s, followed by a pre-cue interval of 1–8 s. Phase 4 testing continued for 10 sessions. Nose pokes in the presence of the discriminative stimulus immediately turned off the visual cue and elevated the sucrose dipper (5 s). In one final session, the duration of the cue presentation was increased to a maximum of 30 s per trial. Thus, mice had up to 30 s to detect and respond to the visual cue before the next ITI was initiated.

Ethanol administration. Following the completion of Phase 4 testing, mice were retested in the signaled nose poke task following an injection of ethanol (0, 0.75, 1.25, 1.75 g/kg i.p.; 20% w/v in isotonic saline). Doses were administered in ascending or descending order (counterbalanced within groups) and all animals were tested twice at each dose. A minimum of 48 h separated each test.

Sucrose Preference. At the end of all behavioral tests, sucrose and water intake were assessed for 24 h, beginning at 11:00 during the light cycle (lights off at 19:00). Water and sucrose (25%) were provided in two bottles at the front of the cage and the number of licks was automatically scored for each bottle and measured as a function of time. Regular chow was freely available throughout the session. Sucrose preference was calculated for each phase as a percentage of total fluid intake (sucrose and water combined). Locomotor activity was measured as horizontal beam breaks at the front and back of the cage.

Statistical analyses

The same wild-type mice on a BL6 background were used as controls for both mu and delta opioid receptor knockout mice; separate groups of HYB wild-type animals were used as controls of Oprm1<sup>-/-</sup> and Oprd1<sup>-/-</sup> groups on a HYB background. Data were then analyzed separately (Oprm1<sup>-/-</sup> and Oprd1<sup>-/-</sup>) using an analysis of variance (ANOVA) with genotype (wild-type versus knockout) and background (BL6 versus HYB) as a between-subjects factors and session as a within-subjects factor (Magazine training in Phase 4, and Ethanol testing sessions). Post-hoc analyses were conducted using Tukey’s test. Whenever there were violations of sphericity, P-values from the Greenhouse-Geisser correction to the within-group degrees of freedom were reported for all ANOVA tests. During Magazine training, the total number of magazine entries, magazine entries during sucrose presentation and the % of total magazine entries that occurred during sucrose presentations were analyzed. The dependent variable in Phases 1–3 was days to criterion. In Phase 4 and during Ethanol testing, the conditioned responses (CR; reinforcers/cue presentations) and efficiency ratio (reinforcers/nose pokes) were dependent variables. Appetitive learning is reflected in the CR measure whereas impulsivity is reflected in the efficiency ratio. Data from the ethanol sessions were analyzed with genotype and background as between-subjects factors and dose as a repeated factor. There was no effect of repeated testing on efficiency ratios or CR so data were collapsed across this factor. In all sessions, the number of responses on the non-reinforced side, the number of magazine entries during reward and non-reward periods, the latency to enter the magazine following elevation of the sucrose dipper, were recorded. Sucrose preference data were analyzed using a two-way ANOVA (genotype x background).

Results

Training

Mice from the six groups progressed through the training phases of the signaled nose poke task at the same rate (Table 1). All animals learned the significance of the reward presentation, evidenced by an increase in the proportion of magazine entries that occurred during sucrose availability; there were no genotype or background differences on this measure. Nor were there any genotype, background or interaction effects on the days to criterion measure in Phases 1–3 (all F<sub>s</sub><1.2; NS), indicating that deletion of neither mu-opioid nor delta-opioid receptors affected learning on this task.

Testing

Deletion of the mu opioid receptor gene decreases motor impulsivity. Fig. 2 shows the effect of mu-opioid receptor deletion on testing in the signaled nose poke task. In contrast to the training data, Oprm1<sup>-/-</sup> mice on both genetic backgrounds performed significantly better than wild-type controls during Phase 4 testing. Efficiency ratios (rewards/nose pokes) increased across 10 sessions [F(9,351) = 59.82, p<0.01] verifying that all groups were capable of learning the task (Fig. 2A and 2E). The significant 3-way interaction [F(9,351) = 2.05, p<0.05] indicated that the difference in efficiency ratios between wild-type and knockout mice depended on the session and the background strain. Pairwise comparisons revealed that HYB, but not BL6, knockout animals differed from their wild-type controls on session 6. In sessions 7–10, BL6 knockout mice performed better than controls, whereas HYB knockout mice showed higher efficiency ratios only on session 8. The significant genotype [F(1,39) = 8.79, p<0.01] and background [F(1,39) = 6.73, p = 0.05] differences were due to higher efficiency ratios in both Oprm1<sup>-/-</sup> groups compared to their own wild-type control, as well as higher efficiency ratios in the HYB background.

As seen in Fig. 2B and 2F, responses on the rewarded side during the pre-CS period (variable time 1–8 s) showed a general decline across sessions, [F(9,351) = 19.49, p<0.01] and an interaction of this measure with genotype [F(9,351) = 5.73, p<0.01]. There were significant genotype [F(1,39) = 15.5, p<0.01] and background
Responses on the rewarded side during the 20-s inter-trial interval (ITI) showed a similar pattern [F(9,351) = 17.11, p<.01], but the rate of decrease differed across groups [F(9,351) = 2.77, p<.05] (Fig. 2C and 2G). Responding was similar in all four groups during the initial sessions, but declined to a lower level in the Opi1−/− groups by the end of the sessions. The significant main effects of genotype [F(1,39) = 8.2, p<.01] and background [F(1,39) = 4.81, p<.05] reflected lower overall responses in the Opi1−/− groups and in the HYB strain.

Finally, conditioned responding (rewards/signals) increased across sessions [F(9,351) = 119.51, p<.01], but there were no genotype or background differences on this measure and none of the interactions were statistically significant (Fig. 2D and 2H).

**Deletion of the delta opioid receptor gene increases motor impulsivity.** Fig. 3 shows the effect of delta opioid receptor deletion on testing in the signaled nose poke task. Similar to the mu-opioid receptor knockout animals, efficiency ratios increased across the 10 testing sessions of Phase 4 [F(9,351) = 26.76, p<.01], indicating that all groups were capable of learning the task (Fig. 3A and 3E). The genotype x session interaction [F(9,351) = 3.78, p<.01] and the main effect of genotype [F(1,35) = 11.53, p<.01] were the result of both Opi1−/− groups achieving lower efficiency ratios (i.e., increased impulsivity) compared to their respective control groups (post-hoc tests p<0.05). In addition, as in the Opi1−/− experiment, there was a significant difference in the efficiency ratios of the two wild-type strains, with the HYB animals exhibiting significantly higher efficiency ratios in sessions 3–10. The session x background and session x genotype interactions were not statistically significant.

Both the genotype [F(1,35) = 4.32, p<.05] and background [F(1,35) = 87.34, p<.01] differences in responding were apparent during the 1–8 s pre-CS period when animals must learn to inhibit their responses (Fig. 3B and 3F). Pre-CS responses declined across sessions [F(9,351) = 14.4, p<.01] and the effect was more pronounced in BL6 mice of both genotypes [F(9,351) = 6.3, p<.01]. Responses also decreased during the ITI period [F(9,351) = 20.24, p<.01] but this factor did not interact with group or genotype (Fig. 3C and 3G). Unlike pre-CS responses, there were no group differences in responding during the ITI period [F(1,35) = 3.15, p>.05], although the HYB animals continued to respond at higher rates during this period [F(1,35) = 61.45, p<.01]. Thus, increased responding in the signaled nose poke task by Opi1−/− mice was confined to the pre-CS period indicating that these mice are incapable of refraining from making an anticipated response.

The proportion of CRs also increased across sessions [F(9,315) = 123.29, p<.01] (Fig. 3D and 3H), but there were no significant genotype or background differences on this measure and none of the interactions were statistically significant. Thus, in contrast to the impulsivity measure, the ability to learn a conditioned responses was not disrupted by deletion of the Opi1 gene.

In sum, mice lacking mu-opioid receptors were significantly better at inhibiting a motor response, whereas mice lacking the delta-opioid receptor were significantly worse.

It is unlikely that these changes in efficiency ratios simply reflect a general reduction in responding because there were no genotypic differences in: 1) responses on the non-reinforced side throughout training and testing (Table 2, left columns) or 2) responses on the reinforced side during the final sessions of Phases 1–3 or the first session of Phase 4 (Table 2, right columns). In addition, conditioned responses to the reward-paired cue were similar across genotype and strain suggesting that differences in associative learning did not impact on task performance. To evaluate possible differences in attentional processes, we increased the duration of the visual cue from 5 to 30 s during a final test session. This manipulation did not affect efficiency ratios or conditioned responses in any group (data not shown). The mean latency to respond to the visual cue in this final session was 2.28 s and less than 2% of the cue presentations reached the maximum 30 s duration; there were no genotype or background differences on either measure. Thus, it is unlikely that the decreased impulsivity in Opi1−/− mice or the increased impulsivity in Opi1+/− mice reflects alterations in attentional processes. Finally, sucrose preference was similar in wild-type and knockout mice, and there were no strain differences in this measure, ruling out the possibility that differential responses to the reward influenced our findings (mean sucrose preference across groups = 94.7%).

**Ethanol administration alters performance in delta but not mu opioid receptor knockout mice.** Ethanol administration decreased efficiency ratios and conditioned responses in wild-type mice but had no effect on mice lacking mu-opioid receptors (Fig. 4).

As seen in Fig. 4A, the effect of ethanol on efficiency ratios was dependent on dose [F(3,246) = 5.95, p<.01] and genotype

| Group         | Magazine Training | Phase 1: FR1 | Phase 2: FR3 | Phase 3: ITI |
|---------------|-------------------|--------------|--------------|--------------|
|               | Proportion Reward | Days to Criterion | Days to Criterion | Days to Criterion |
| wild-type HYB | 0.38±.06          | 3.5±.35      | 3.20±.31     | 5.70±.41     |
| Opm1−/− HYB   | 0.44±.05          | 4.0±.49      | 2.70±.47     | 5.20±.67     |
| Opr1−/− HYB   | 0.47±.10          | 2.88±.64     | 3.87±.64     | 6.12±.58     |
| wild-type BL6 | 0.51±.06          | 4.0±.57      | 3.17±.70     | 6.25±.52     |
| Opm1−/− BL6   | 0.52±.03          | 4.4±.60      | 3.80±.39     | 5.50±.62     |
| Opr1−/− BL6   | 0.53±.03          | 3.33±.41     | 4.00±.67     | 6.00±.73     |

The first two columns display the proportion (±SEM) of magazine entries that occurred during elevation of the sucrose dipper across two sessions of magazine training. The last three columns display the days to criterion measure for Phases 1–3 during task training. With fixed ratio (FR) responding in Phases 1 and 2, the criterion to progress to the next phase was 25 reinforceers earned per 40-min session. With the introduction of the inter-trial interval (ITI) in Phase 3, this was reduced to 10 reinforceers per session. There were no significance differences between knockout and wild-type animals on any measure.

doi:10.1371/journal.pone.0004410.t001

---

**Table 1.** Training on the signaled nose poke task.

*Opioids and Impulsivity*
Opioids and Impulsivity

Figure 2. Oprm1−/− mice perform better in the signaled nose poke task. Performance on Phase 4 of the signaled nose poke task for mu-opioid receptor knockout mice and their wild-type controls on a HYB (A–D) and BL6 (E–H) backgrounds. A and E: Efficiency ratios (rewards/nose pokes) increased across sessions with Oprm1−/− mice on both genetic backgrounds performing significantly better than wild-type controls. Mice lacking the Oprm1 gene exhibited lower levels of responding throughout Phase 4 testing; this phenotypic difference was most apparent during the pre-CS period (Fig. 2B and 2F), when animals must learn to inhibit responding in order to maximize the number of rewards earned. C and G: Responses on the rewarded side during the 20-s inter-trial interval (ITI) decreased across sessions with lower overall responses in the Oprm1−/− and HYB strain. D and H: Conditioned responding (rewards/signals) increased across sessions with no genotype or background differences. Oprm1−/− HYB, n = 11; Oprm1−/− BL6, n = 12; Oprm1+/+ HYB, n = 10; Oprm1−/− BL6, n = 11.

doi:10.1371/journal.pone.0004410.g002

Discussion

This study demonstrates that mice lacking mu-opioid receptors exhibit decreased motor impulsivity whereas those lacking delta-opioid receptors show increased motor impulsivity. Thus, our results reveal an unforeseen role of endogenous opioid receptor activity in disinhibition and suggest that mu opioid receptors promote, whereas delta opioid receptors inhibit, impulsive behaviors. Future studies, using a similar approach, will determine whether kappa receptor activity also influences motor impulsivity.

The increased ability to withhold a motor response in Oprm1−/− mice occurred when reward-related learning remained intact. The latter observation seems to contrast other reports of reward reduction in these animals [9]. These studies, however, examined the effect of mu-opioid receptor gene deletion on the hedonic properties of abused drugs whereas our study tested nosepoking responses to a food reward which is not altered in mu-opioid receptor knockout mice [19]. Importantly, the fact that mu-receptors modulate motor impulsivity independently from conditioned reward in the signaled nose poke tasks is consistent with the notion that reward and impulsivity are mediated by different neural systems [15]. Changes in drug reward and impulsivity, therefore, are separate processes that interact to influence the development of addiction: combined with previous evidence, our findings highlight the fact that mu-opioid receptors play a key role in both processes. These receptors, therefore, not only mediate initial drug reward but may also be implicated in further behavioral changes, such as loss of control, that occur in chronic drug abusers.

Our findings that mice lacking mu-opioid receptors are less impulsive has direct relevance to heroin addiction. At least a small population of heroin addicts exhibit greater binding potential of mu-opioid receptors [20], suggesting that these individuals should display the opposite behavior to that of our knockout mice (i.e., increased impulsivity). The evidence for this hypothesis is mixed: although heroin addicts show clear deficits in cognitive or choice impulsivity [21–23], their performance on tests that measure motor impulsivity is less clear. Some of these discrepancies may reflect drug history as pure heroin abusers [24,25], but not heroin addicts who also abuse other substances [23], exhibit these deficits.

Motor deficits in opiate abusers may also depend on current drug use because heroin addicts maintained on methadone show motor impulsivity deficits that are not apparent in abstinent abusers [26]. Although the relationship between the behavior of Oprm1−/− mice and heroin addicts is not straightforward, our results fit well with evidence that opioid antagonists enhance control of motor responses in healthy participants [27] and may be effective treatment in impulse-control disorders [28].

The decreased ability of Opod−/− mice to withhold a motor response is intriguing for two reasons. First, these data add to increasing evidence that mu and delta opioid receptors have opposing roles in many behavioral responses. These include anxiety and depressive-like behaviors [12], ethanol self-administration [29] and a conditioned place preference to cannabinoids [10]. The two receptors also differentially regulate mesolimbic DA tone [30]. Second, clinical research and practice have probed the function of mu opioid receptors using morphine-derived compounds for decades. In contrast, delta opioid receptor pharmacology is less well developed and many aspects of delta receptor function remain unexplored. Our findings suggest that, in addition to their anxiolytic and antidepressant effects, delta receptor agonists are of potential interest for impulse control disorders.

With regards to drug abuse, delta opioid receptors may minimize changes in emotional state and impulsivity, both of which develop under chronic drug abuse and contribute to relapse.

Consistent with previous evidence [31,32], strain differences in impulsivity emerged in that BL6 mice were more impulsive than the HYB groups. The fact that basal motor impulsivity was distinct in the two wildtype strains emphasizes the importance of examining knockout mice on both background. Strain conferred a performance advantage in all groups, with the exception of Oprm1−/− mice, probably because these animals were already responding at optimal levels. Indeed the performance of Oprm1−/− mice was so efficient that they made, on average, fewer than 3 pre-CS responses during the 10th training session. Compared to other studies using the same task [15,33], our animals performed exceptionally well with efficiency ratios of wild-type animals...
Figure 3. *Oprd1*−/− mice are impaired in the signaled nose poke task. Performance on Phase 4 of the signaled nose poke task for delta-opioid receptor knockout mice and their wild-type controls on a HYB (A–D) or BL6 (E–H) background. A: Efficiency ratios (rewards/nosepokes) increased across 10 sessions indicating that all groups were capable of learning the task. Both HYB and BL6 *Oprd1*−/− mice exhibited lower efficiency ratios (i.e., more impulsivity) than their wild-type controls. The significant background effect reflected the fact that both knockout and wild-type BL6 mice were more impulsive than mice on a HYB background. B and F: Responses during the pre-CS period also declined across sessions although the effect was not as pronounced in *Oprd1*−/− mice. Again, responses of HYB mice were increased compared to mice on a BL6 background. C and G: Responses during the 20-s inter-trial interval (ITI) decreased across sessions and the rate of decline was consistent across groups. Nonetheless, HYB mice respond at higher rates than BL6 mice of both genotypes throughout the sessions. D and H: Conditioned responding (rewards/signals) increased across sessions, but there were no genotype or background differences on this measure and no statistically significant interactions. *Oprd1*−/− HYB, n = 8; *Oprd1*−/− BL6, n = 10.

doi:10.1371/journal.pone.0004410.g003

Table 2. Nosepoke responses on the signaled nose task.

| Group            | Phase 1 | Phase 2 | Phase 3 | Phase 4 |
|------------------|---------|---------|---------|---------|
|                  | non-RF  | Total   | non-RF  | Total   |
| Oprm1−/− HYB     | 19.09 (3.08) | 67.27 (7.73) | 11.55 (2.48) | 144.45 (15.32) |
| Oprm1−/− BL6     | 21.75 (3.77) | 76.40 (9.49) | 13.16 (3.03) | 136.10 (16.89) |
| Oprm1−/− + BL6   | 25.50 (5.80) | 57.52 (8.04) | 16.93 (3.65) | 115.17 (12.5) |
| Oprm1−/− + BL6   | 22.10 (4.29) | 51.50 (8.49) | 17.61 (4.74) | 116.70 (15.42) |
| Oprm1−/− + BL6   | 20.75 (3.98) | 69.00 (8.75) | 15.00 (2.0) | 87.13 (9.17) |
| Oprm1−/− + BL6   | 25.50 (5.80) | 57.52 (8.04) | 16.93 (3.65) | 115.17 (12.5) |
| Oprm1−/− + BL6   | 18.20 (3.78) | 66.90 (9.69) | 19.70 (3.33) | 135.7 (15.36) |

The two columns under each phase display mean nose poke (SEM) in the non-reinforced (non-RF) hole (left column) and total nose pokes (right column) during the final session of Phases 1–3 (i.e., the day the animals met criterion) and the first session of Phase 4. Note that the number of sessions in Phases 1–3 varied for each animal, depending on the days to reach criterion. There were no significance differences between knockout and wild-type animals on any measure.

Oprm1−/− HYB, n = 11; Oprm1−/− BL6, n = 12;
Oprm1−/− + BL6, n = 11;
Oprd1−/− HYB, n = 9; Oprd1−/− BL6, n = 8;
Oprd1−/− + BL6, n = 12;
Oprd1−/− BL6, n = 10.

doi:10.1371/journal.pone.0004410.t002

ranging from .31 to .5 at the end of training. In addition to strain differences, we used a visual rather than an auditory cue and liquid rather than pellet sucrose reward. Either or all of these factors may have increased the performance of wild-type mice and make it even more surprising that we observed such a dramatic enhancement in mice lacking the *Opml* gene.

We also provide compelling evidence that *Opml*−/− mice are insensitive to the acute effects of ethanol in a signaled nose poke task. This fits with evidence that mice lacking the *Opml* gene show reduced responses to ethanol in other behavioural tests [34] which may be mediated by the reduction in ethanol-stimulated DA release in the ventral striatum in *Opml*−/− mice [35]. We also found that strain differences, which were apparent during impulsivity testing, disappeared under the influence of ethanol. This probably reflects increased experience with the task rather than a drug-induced dampening of strain differences because BL6 and HYB wild-type mice showed the same level of performance following vehicle injections. Thus, although strain may influence impulsivity measures, the effect is not nearly as robust as the *Opml*−/− and *Oprd1*+/+ difference that was apparent in both drug-free and drug-tested mice.

Higher activity under the influence of ethanol probably accounts for the increased responses during both pre-CS and ITI periods, producing lower efficiency ratios in both wild-type and *Oprd1*−/− mice. The reduction in conditioned responses could reflect a cognitive deficit that is manifested as an inability to detect and/or respond to the visual stimulus. Even if this is true, mice were not completely disoriented because responses on the non-reinforced side did not change with ethanol administration (mean responses per session <10 for all groups). In light of these dramatic effects of ethanol in wild-type and *Oprd*−/− mice, the most striking finding is that the drug was completely ineffective in altering the behavior of mice lacking the *Opml*−/− gene.

Mechanisms by which mu- and delta-opioid receptors regulate motor impulsivity remain to be elucidated: there are several potential neural substrates for opioid-controlled disinhibition. The decreased impulsivity we observed in *Opml*−/− mice could be mediated through an interaction with the subthalamic nucleus (STN), dopamine (DA) D2 receptors in the ventral striatum (VS), and/or excitatory projections from the prefrontal cortex (PFC) to the striatum. Lesions of either the PFC [36] or STN [37] increase impulsivity suggesting that deletion of the *Opml* gene may increase activity in either structure. With regards to the STN, this hypothesis is plausible as activation of mu-opioid receptors inhibits excitatory inputs to the STN [38]; the absence of mu-opioid receptors would be associated with increased excitation of STN neurons. The possibility that mice lacking mu-opioid receptors exhibit increased activity in PFC-striatal circuits is less convincing because activation of mu-opioid receptors in the PFC inhibits GABA interneurons that synapse on PFC projections [39]. Removal of opioid-induced inhibition of GABA neurons would decrease activity in PFC projections. On the other hand, a small population of mu-opioid receptors is located on PFC...
projection neurons [40] so deletion of the Oprm1 gene could increase PFC activity by directly removing this inhibitory input. Finally, decreased availability of DA D2-like receptors in the VS is associated with high levels of trait impulsivity [41] suggesting that knockout mice may exhibit increased signaling in DA D2 pathways. This idea fits with evidence that mice lacking the Oprm1 gene show increased DA D2 receptor binding in the striatum and increased activity levels when the receptor is stimulated [42]. Much less is known about delta receptor function in these brain areas, or about D2/delta receptor interactions. In addition, the effect of selective opioid receptor ligands in the STN, PFC or VS on impulsive responding has not been tested directly. The deletion of the Oprm1 and Oprd1 genes may, together, alter inhibitory mechanisms through one or more of these neural systems.

An important remaining question is whether deletion of the Oprm1 or Oprd1 genes affect other measures of impulsivity, such as delay-discounting or reflection impulsivity. The latter is particu-

Figure 4. Oprm1−/− mice are not affected by ethanol in the signaled nose poke task. A: Efficiency ratios (rewards/nosepokes) decreased with increasing doses of ethanol, but only in wild-type mice. B: The same effect was observed during the pre-CS period. There were no significant strain differences on any behavioral measure during ethanol testing, nor were there any strain-dose interactions. C: Ethanol increased nose pokes that occurred during the inter-trial interval (ITI) in wild-type but not knockout animals. D: Ethanol also altered conditioned responses in Oprm1+/*, but not Oprm1−/− mice. The highest dose of ethanol (1.75 g/kg) produced the greatest effect on conditioned responses in wild-type mice. Oprm1+/* HYB, n = 11; Oprm1−/− BL6, n = 12; Oprm1−/− HYB, n = 10; Oprm1−/− BL6, n = 11.

doi:10.1371/journal.pone.0004410.g004
Figure 5. Oprd1<sup>−/−</sup> and Oprd1<sup>+/+</sup> mice are impaired by ethanol in the signaled nose poke task. A: Ethanol produced a dose-dependent decrease in efficiency ratios (rewards/nosepokes) in all groups of animals, with knockout mice displaying lower efficiency ratios throughout testing. B: Ethanol increased nose pokes that occurred during the 1–8 s pre-CS period and during the inter-trial interval (ITI) (C). Both wild-type and knockout mice on a BL6 background exhibited higher rates of responding than HYB mice during the ITI but this effect did not interact with dose or with group. D: Ethanol altered conditioned responses in both Oprd1<sup>−/−</sup> and Oprd1<sup>+/+</sup> mice, although only the highest dose (1.75 g/kg) produced an effect. 

Oprd1<sup>+/+</sup> HYB, n = 9; Oprd1<sup>−/−</sup> BL6, n = 12; Oprd1<sup>+/−</sup> HYB, n = 8; Oprd1<sup>−/−</sup> BL6, n = 10.

doi:10.1371/journal.pone.0004410.g005

Acknowledgments

We thank Celia Goeldner for assistance with behavioral testing and data analysis.

Author Contributions

Conceived and designed the experiments: MCO AMO. Performed the experiments: MCO. Analyzed the data: MCO AMO. Wrote the paper: MCO BLK.

larily interesting because it represents a cognitive marker for substance dependence that does not recover with prolonged abstinence and is associated with multiple drugs of abuse [43]. The importance of impulsivity in addiction is emphasized further by a relationship between level of abuse and treatment retention [44]. Finally, beyond addiction, the role of the Oprm1 and Oprd1 genes in impulsivity has implications for understanding and treating attention deficit hyperactivity disorder, eating disorders, gambling and other disorders of impulse control.
References

1. American Psychiatric Association (1994) Diagnostic and Statistical Manual of Mental Disorders. Amherst, NY: Prometheus Books.

2. Nederkoorn C, Braet C, Van Eeys S, Tanghe A, Jansen A (2006) Why obese children cannot resist food: the role of impetuosity. Eating Behavior 7: 315–322.

3. Garavan H, Hester R (2007) The role of cognitive control in cocaine dependence. Neuropsychology Review 17: 337–345.

4. Moeller FG, Dougherty DM, Barratt ES, Oderinde V, Mathias CW, et al. (2002) Increased impulsivity in cocaine dependent subjects independent of antisocial personality disorder and aggression. Drug and Alcohol Dependence 68: 105–111.

5. Limneos M, Virkkunen M, George T, Higley D (1993) Impulse control disorders. International Clinical Psychopharmacology 8: 33–56.

6. Pliakas AM (2005) The neuropsychopharmacology of attention-deficit/hyperactivity disorder. Biological Psychiatry 57: 1385–1390.

7. Kreek MJ, Nielsen DA, Butelman E, LaForge S (2005) Genetic influences on impetuosity, risk taking, stress responsivity and vulnerability to drug abuse and addiction. Nature Neuroscience 8: 1450–1457.

8. Pattij T, Vanderschuren LJ (2008) The neuropharmacology of impulsive behavior. Trends in Pharmacological Sciences in press.

9. Contet C, Kieffer BL, Befort K (2004) Mu opioid receptors: a gateway to drug addiction. Current Opinion in Neurobiology 14: 370–378.

10. Ghozland S, Mathews HWD, Simonin F, Filliol D, Kieffer BL, et al. (2002) Motivational effects of cannabinoids are mediated by mu-opioid and kappa-opioid receptors. Journal of Neuroscience 22: 1146–1154.

11. Roberts AJ, Gold LH, Polis I, McDonald JS, Filliol D, et al. (2001) Increased ethanol self-administration in delta-opioid receptor knockout mice. Alcoholism: Clinical and Experimental Research 25: 1249–1256.

12. Filliol D, Ghozland S, Chiba J, Martin M, Mathews HWD, et al. (2000) Mice deficient for delta- and mu-opioid receptors exhibit opposing alterations of emotional responses. Nature Genetics 22: 195–200.

13. Everitt BJ, Robbins TW (2003) Neural systems of reinforcement for drug addiction: from actions to habits to compulsion. Nature Neuroscience 8: 1481–1489.

14. Robinson TE, Berndt KC (1995) The neural basis of drug craving: an incentive-sensitization theory of addiction. Brain Research Reviews 18: 247–291.

15. Bowers BJ, Weinher JM (2001) Ethanol consumption and behavioral impulsivity are increased in protein kinase Cε null mutant mice. Journal of Neuroscience 21: RC180.

16. Sanchez-Segura C, Spanagel R (2006) Behavioural assessment of drug reinforcement and addictive features in rodents: an overview. Addiction Biology 11: 38.

17. Critchlow B (1986) The powers of John Barleycorn: Beliefs about the effects of alcohol on social behavior. American Psychologist 41: 751–764.

18. Mathies HWD, Maldonado R, Simonin F, Valverde O, Sloew S, et al. (1996) Loss of morphine-induced analgesia, reward effect and withdrawal symptoms in mice lacking the mu-opioid-receptor gene. Nature 383: 819–823.

19. Papalou F, Kieffer BL, Tabarina A, Contarino A (2007) Decreased motivation to reinforcer in mu-opioid-receptor-deficient mice. European Journal of Neuroscience 21: 3598–3605.

20. Sharpe LG, Pilote NS, Shippennberg TS, Goodman CB, London ED (2000) Autoradiographic evidence that prolonged withdrawal from intermittent cocaine reduces mu opioid receptor expression in limbic regions of the rat brain. Synapse 37: 292–297.

21. Kirby KN, Petry NM, Bickel WK (1999) Heroin addicts have higher discount rates for delayed rewards than non-drug using controls. Journal of the Experimental Analysis of Behavior 72: 78–87.

22. Bickel WK, Marsch DM (2001) Toward a behavioral economic understanding of drug dependence: delay discounting processes. Addiction 96: 73–86.

23. Verdejo-Garcia A, Perales JC, Perez-Garcia M (2007) Cognitive impulsivity in cocaine and heroin polysubstance abusers. Addictive Behaviors 32: 950–966.

24. Lee TM, Pau CW (2002) Impulse control differences between abstinent heroin users and matched controls. Brain Injury 16: 341–360.

25. Pau CW, Lee TM, Chan SF (2002) The impact of heroin on frontal executive functions. Archives of Clinical Neuropsychology 17: 663–670.

26. Verdejo A, Toribio I, Orozco C, Puente KL, Perez-Garcia M (2005) Neuropsychological functioning in methadone maintenance patients versus abstinent heroin abusers. Drug and Alcohol Dependence 78: 283–288.

27. Mitchell JM, Tavares VC, Fields HL, DeSposito M, Boettinger CA (2007) Endogenous opioid blockade and impulsive responding in alcoholics and healthy controls. Neuropsychopharmacology 32: 439–449.

28. Kim SW (1990) Opioid antagonists in the treatment of impulse-control disorders. Journal of Clinical Psychiatry 59: 159–164.

29. Roberts AJ, McDonald JS, Hyser CJ, Kieffer BL, Mathews HWD, et al. (2000) Mu-opioid receptor knockout mice do not self-administer alcohol. Journal of Pharmacology and Experimental Therapeutics 295: 1000–1008.

30. Chefer VL, Kieffer BL, Shippenberg TS (2003) Basal and morphine-evoked dopaminergic neurotransmission in the nucleus accumbens of MOR- and DOR-knockout mice. European Journal of Neuroscience 18: 1915–1922.

31. Patel S, Stoeterman IP, Asherson P, Shayter F (2006) Attentional performance of C57BL/6 and DBA/2 mice in the 5-choice serial reaction time task. Behavioural Brain Research 170: 197–203.

32. Isles AR, Humby T, Walters E, Wilkinson LS (2004) Common genetic effects on variation in impulsivity and activity in mice. Journal of Neuroscience 24: 6753–6760.

33. Keller JB, Keller AB, Bowers BJ, Weinher JM (2005) Performance of alpha7 nicotinic receptor null mutants is impaired in appetitive learning measured in a signaled nose poke task. Behavioural Brain Research 162: 143–152.

34. Ghozland S, Chu K, Kieffer BL, Roberts AJ (2005) Lack of stimulant and anxiolytic-like effects of ethanol and accelerated development of ethanol dependence in mu-opioid receptor knockout mice. Neuropharmacology 49: 495–501.

35. Job MO, Tang A, Hall FS, Sora I, Uhl GR, et al. (2007) Mu (mu) opioid receptor regulation of ethanol-induced dopamine response in the ventral striatum. Evidence of genotype specific sexual dimorphic epistasis. Biological Psychiatry (in press).

36. Kovacs KM, Szakall I, O'Brien D, Wang R, Vinod KY, et al. (2005) Decreased oral self-administration of alcohol in kappa-opioid receptor knockout mice. Alcoholism: Clinical and Experimental Research 29: 730–738.

37. Uslaner J, Robinson TE (2006) Subthalamic nucleus lesions increase impulsive action and decrease impulse choice - mediation by enhanced incentive motivation? European Journal of Neuroscience 24: 2345–2334.

38. Shen KZ, Johnson SW (2002) Presynaptic modulation of synaptic transmission by opioid receptor in rat subthalamic nucleus in vitro. Journal of Physiology 541: 219–230.

39. Witkowski G, Szulczyk P (2006) Opioid mu receptor activation inhibits sodium currents in prefrontal cortical neurons via a protein kinase A- and C-dependent mechanism. Brain Research 1049: 92–1006.

40. Schmidt P, Schmelke C, Musshoff F, Prohaska C, Menzen M, et al. (2001) Numerical density of mu opioid receptor expressing neurons in the frontal cortex of drug related fatalities. Forensic Science International 115: 219–229.

41. Dalley JW, Fyer TD, Richard L, Robinson ESJ, Theobald DE, et al. (2007) Nucleus accumbens D2/D3 receptors predict trait impulsivity and cocaine reinforcement. Science 315: 1267–1270.

42. Tien LT, Park Y, Fan LW, Ma T, Loh HH, et al. (2003) Increased dopamine D2 receptor binding and enhanced apomorphine-induced locomotor activity in mu-opioid receptor knockout mice. Brain Research Bulletin 61: 109–113.

43. Clark L, Robbins TW, Ersche KD, Sahakian BJ (2006) Reflection impulsivity in current and former substance abusers. Biological Psychiatry 60: 515–522.

44. Moeller FG, Dougherty DM, Barratt ES, Oderinde V, Mathias CW, et al. (2002) Increased impulsivity in cocaine dependent subjects independent of antisocial personality disorder and aggression. Drug and Alcohol Dependence 68: 103–111.