Dimerization with Cannabinoid Receptors Allosterically Modulates Delta Opioid Receptor Activity during Neuropathic Pain

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Abstract

The diversity of receptor signaling is increased by receptor heteromerization leading to dynamic regulation of receptor function. While a number of studies have demonstrated that family A G-protein-coupled receptors are capable of forming heteromers in vitro, the role of these heteromers in normal physiology and disease has been poorly explored. In this study, direct interactions between CB1 cannabinoid and delta opioid receptors in the brain were examined. Additionally, regulation of heteromer levels and signaling in a rodent model of neuropathic pain was explored. First we examined changes in the expression, function and interaction of these receptors in the cerebral cortex of rats with a peripheral nerve lesion that resulted in neuropathic pain. We found that, following the peripheral nerve lesion, the expression of both cannabinoid type 1 receptor (CB1R) and the delta opioid receptor (DOR) are increased in select brain regions. Concomitantly, an increase in CB1R activity and decrease in DOR activity was observed. We hypothesize that this decrease in DOR activity could be due to heteromeric interactions between these two receptors. Using a CB1R-DOR heteromer-specific antibody, we found increased levels of CB1R-DOR heteromer protein in the cortex of neuropathic animals. We subsequently examined the functionality of these heteromers by testing whether low, non-signaling doses of CB1R ligands influenced DOR signaling in the cortex. We found that, in cortical membranes from animals that experienced neuropathic pain, non-signaling doses of CB1R ligands significantly enhanced DOR activity. Moreover, this activity is selectively blocked by a heteromer-specific antibody. Together, these results demonstrate an important role for CB1R-DOR heteromers in altered cortical function of DOR during neuropathic pain. Moreover, they suggest the possibility that a novel heteromer-directed therapeutic strategy for enhancing DOR activity, could potentially be employed to reduce anxiety associated with chronic pain.

Introduction

Among the many current targets being explored for the treatment of neuropathic pain are opioid and cannabinoid receptors [1,2], which share similar signaling properties and are widely co-distributed in regions of the peripheral and central nervous system associated with ascending pain sensation, descending inhibition of pain, as well as emotional processing [3–10]. Activation of both receptor types can produce analgesia in humans and animals experiencing neuropathic pain [5,11,12] by a variety of mechanisms, including but not limited to, agonist stimulated induction of gene expression, receptor crossactivation, cannabinoid induced release of opioid peptides and opioid induced release of endocannabinoids [13–15]. Opioid and cannabinoid receptor activation can also reduce anxiety and depressive-like phenomena, particularly through selective activation of subtypes of these classes of receptors, including the delta opioid receptor (DOR) subtype [16,17] or cannabinoid type 1 receptor (CB1R) subtype [5,18–20]. Previous studies have shown indirect inhibitory interactions between these two receptors. For example, DOR activity is increased in the brains of CB1R −/− mice [21], while CB1R activity is increased in the brains of DOR −/− mice [22,23], indicating that DOR can modulate CB1R activity and vice versa. Additionally, anxiolytic-like responses induced by Δ9-THC, a CB1R agonist, could be blocked by naltrindole, a DOR antagonist [24], suggesting that supraspinal interactions between DOR and CB1R can modify behavior.

G protein-coupled receptor heteromerization has been previously shown to enhance the repertoire of receptor signaling, thereby dynamically modulating receptor function [25,26]. A recent set of studies examined direct interactions between CB1R and DOR in heterologous systems [22]. Co-expression of tagged DOR and CB1R leads to an increase in BRET signal, whereas co-expression of CCR5 and CB1R does not [27], indicating that DOR and CB1R are in close proximity to each other in cells expressing both receptors. Moreover, DOR and CB1R can be co-immunoprecipitated in cells expressing both receptors [22], demonstrating that CB1R forms receptor heteromers with DOR.

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CB₁R activity is increased in cortical membranes from mice with a genetic deletion of DOR, and CB₁R activity is reduced when cells endogenously expressing CB₁R are transfected with DOR [22], suggesting that these receptors normally inhibit each other’s activity. Moreover, CB₁R signaling is reduced in the presence of low concentrations of DOR ligands in cells co-expressing both receptors [22]. Together, these data suggest that CB₁R directly interacts with DOR, and that occupancy of one receptor may allosterically alter the activity of the other receptor within this receptor heteromer. As such, we examined whether CB₁R and DOR interact in endogenous tissue. Furthermore, given that several studies have shown that opioid and cannabionoid receptor expression is dynamically regulated during neuropathic pain, we also examined the degree to which this interaction regulates receptor activity during a pathologic state, neuropathic pain.

Previous studies, under a variety of experimental conditions in rodents, have focused mostly on pain-associated changes in the expression of these receptors within primary afferents/dorsal root ganglia (DRG) or within spinal cord. Within one or two days after a peripheral nerve lesion, opioid receptor mRNA and protein, particularly for the mu opioid receptor (MOR) and DOR, are upregulated within DRG and within the injured nerve proximal to the lesion site, indicating increased transport of opioid receptors to the periphery [13,28–33]. However, a decrease in receptor levels has been reported at later time points after the lesion [28,34–36]. Within the dorsal horn of the spinal cord, MOR and DOR levels have been reported to increase transiently [37,38] immediately after peripheral nerve lesion, but are unchanged or slightly decreased relative to control levels by 7 days and at time points up to 4 weeks following a lesion [34,36–45]. Decreases in MOR and DOR levels in the spinal cord could be a consequence of receptor activation and internalization in response to enhanced release of endogenous opioid peptides. Unlike opioid receptors, CB₁R levels in DRG and spinal cord are reportedly unchanged immediately after peripheral nerve lesion, but are consistently increased during chronic neuropathic pain [33,46–49]. Interestingly, animals with increased CB₁R expression show enhanced analgesic responses to a CB₁R agonist, suggesting that the increase in receptor levels could have a protective function [46,48,50].

While most studies have focused on neuropathic pain-associated changes in receptor expression within peripheral nerves, DRGs or spinal cord, few have examined alterations within supraspinal regions, despite clear evidence that neural activity within thalamus, cerebral cortex, and amygdala, is altered during neuropathic pain [51,52]. Among these brain regions, the cerebral cortex and its subregions play a prominent role in pain perception and response. Altered opioid and cannabinoid receptor activity in cerebral cortex may contribute to changes in neuronal signaling and mood states during neuropathic pain. Previous studies have shown that activation of these receptors can affect affective states. For example, administration of DOR agonists or enkephalinase inhibitors (the latter leading to increases in endogenous opioids) reduces depressive-like and anxiety-like behavior [16,17,53,54], while systemic or cingulate cortex injection of DOR antagonists heightens anxiety [16,55,56]. In addition, genetic deletion of DOR increases levels of anxiety and depressive-like behavior [18,19]. Similarly, systemic or prefrontal cortical (PFC) injection of CB₁R agonists produces antidepressant effects and extinction of conditioned fear responses [18,19], while low doses of CB₁R agonists [50] or inhibitors of endocannabinoid degrading enzymes [59,60] can induce anxiolytic effects. Based on this work, it is clear that cortical activation of DOR and CB₁R can positively modulate affective states. Therefore, these receptors are suitable targets for alleviation of depression, anxiety and fear during neuropathic pain.

A few studies have examined changes in supraspinal opioid and cannabinoid receptor expression and activity during neuropathic pain. In the case of DOR, one study reported no change in DOR activity in lower midbrain and limbic forebrain [61] while another study found reduced DOR activity in the frontal cortex of nerve lesioned mice [55]. Stimulation of DOR is anxiolytic, so a reduction in DOR activity may underlie a heightened anxiety state. In the case of CB₁R, no change in receptor binding was observed in anterior singulate cortex following peripheral nerve lesion, but CB₁R activity was reduced 10 days after injury [62]. It has recently been demonstrated that, in the anterior singulate cortex, CB₁R protein levels initially decrease 7 days after spinal cord injury, then increase by 42 days [63]. Therefore, a clear picture of the time course and degree to which cortical opioid and cannabinoid receptor expression and activity are changed during neuropathic pain has not yet emerged. To address this question, we examined changes in cortical DOR and CB₁R expression, activity and interaction at several time points subsequent to a peripheral nerve lesion. We observed that, while cortical CB₁R and DOR protein levels increase after peripheral nerve lesion, CB₁R activity increases but DOR activity decreases. Because reduced cortical DOR activity during neuropathic pain may underlie heightened anxiety states, we questioned whether heteromerization between CB₁R and DOR would diminish DOR activity, and whether this could be restored pharmacologically by CB₁R ligands. We found that low, non-signaling doses of CB₁R ligands (agonist or antagonist) could allosterically enhance DOR binding and activity, and, given that this effect is blocked by a heteromer specific antibody, that this enhancement is mediated by the CB₁R-DOR heteromer. These results are consistent with allosteric modulation of DOR activity by CB₁R within the CB₁R-DOR heteromer, and suggest that cortical CB₁R-DOR heteromer is a suitable target for blockade of neuropathic pain-associated negative mood states.

Results

Neuropathic Pain Model

We used a common peripheral nerve lesion model, L5 spinal nerve ligation/transaction (L5SNT) [64,65], to produce neuropathic pain in adult male Sprague-Dawley rats. In this model, the right L5 spinal nerve is exposed, ligated and transected 1 mm distal to the ligation. For “sham” animals, the L5 spinal nerve is exposed, but is neither ligated nor transected. Using the Von Frey test [66,67], we measured the presence of mechanical allodynia before and up to 14 days after surgery and found that lesioned animals show a lower threshold of response within two days after surgery (Fig. 1). This decrease in response threshold is maintained in lesioned animals throughout the 14 day testing period; sham animals do not demonstrate altered mechanical pain thresholds (Fig. 1). Therefore, L5SNT surgery produces neuropathic pain that is long-lasting.

Altered receptor expression and activity in cortex of lesioned animals

Next, changes in levels of CB₁R and DOR in various brain regions of animals with peripheral nerve lesion were investigated. We found that, when compared to sham animals, CB₁R levels are increased in cortical membranes prepared from brains of lesioned animals 14 days after surgery (Fig. 2A–C) as measured by Western blot, ELISA and RT-PCR. The increase in CB₁R levels is only evident 14 days after surgery, and not at 3 or 7 days
days. This increase is only evident 14 days after surgery, not at 3 or 7.

We further found significant increases in CB1R levels in some, but not all, brain regions. For example, there was an increase in hypothalamus and midbrain, but not in striatum, hippocampus and cerebellum (Table 2). Taken together, these results indicate that CB1R is selectively upregulated in specific brain regions in response to nerve lesion. Next, we focused on the cortex and examined whether the relative distribution of CB1R was altered in response to nerve lesion. This was evaluated 14 days after nerve lesion by immunohistochemistry. In layer II/III of PFC, a region of interest as it normally expresses high levels of CB1R, a synaptic- and neuritic-like staining pattern is evident in sections from both sham and lesioned animals (Fig. 3A–B). Furthermore, the density of CB1R expression was found to be higher in the cortex of lesioned animals (Fig. 3B–C). Interestingly, lesioned animals show increased CB1R expression in cell bodies, which suggests that the receptor is redistributed and/or new receptors are synthesized during neuropathic pain.

Next, we examined if the L5SNT nerve lesion led to changes in DOR levels. As in the case of CB1R, we find increases in DOR levels in cortical membranes prepared from the brains of lesioned animals, but not sham animals, 14 days after surgery (Fig. 4A–B). This increase is only evident 14 days after surgery, not at 3 or 7 days (Table 1), similar to what was found for CB1R. DOR levels are also differentially regulated in other brain regions 14 days after surgery; we observed significant increases in hypothalamus and in striatum, but not in hippocampus, cerebellum and midbrain (Table 2). Taken together, these results indicate that DOR levels are also selectively increased in specific brain regions in response to nerve lesion.

Next, we examined if increased receptor levels were accompanied by increases in receptor activity using the [35S]GTPγS assay, which measures coupling between the receptor and G-protein. We find that CB1R can be activated by a specific ligand, Hu-210, in cortical membranes from sham and lesioned animals, and that maximal activity is higher in cortical membranes from lesioned animals (~178% for sham and ~224% for lesioned) (Fig. 5A). Thus, 14 days after surgery, lesioned animals demonstrate increased CB1R mRNA, protein levels and receptor activity in cortex.

Changes in DOR activity using [D-Pen2,D-Pen5]Enkephalin, [D-Pen2]Enkephalin (DPDPE) as a receptor specific ligand were also examined. Unexpectedly, we found a decrease in maximal DOR activity in cortical membranes from lesioned animals when compared with sham animals (~171% for sham and ~127% for lesioned) (Fig. 5B). This result was somewhat surprising, as DOR activity was expected to increase in the cortex of lesioned animals commensurate with the documented increases in receptor protein. One possible explanation for the decrease in DOR activity despite an increase in DOR protein is that DOR activity is suppressed by altered interactions with other proteins. This is consistent with previous reports that demonstrated direct, antagonistic interactions between CB1R and DOR in vitro [22,27].

Neuropathic pain-associated alterations in CB1R-DOR levels

Our data suggest that the abundance of CB1R-DOR heteromers may increase after nerve ligation surgery, given that the levels of both receptors individually increase in the cortex of lesioned animals. In order to directly probe the presence of and changes in the levels of CB1R-DOR heteromers, a specific mouse monoclonal antibody directed against the CB1R-DOR heteromer was generated using a previously employed subtractive immunization strategy [25,26]. Among twelve clones that exhibited specificity towards CB1R-DOR we selected one for further characterization. Hereafter this antibody will be referred to simply as CB1R-DOR mAb (Fig. 6). The CB1R-DOR mAb only detects an epitope in cells expressing both CB1R and DOR, but not in cells expressing the individual receptors (Fig. 6A). More importantly, the antibody does not recognize an epitope when CB1R is co-expressed with receptors other than DOR and vice versa (Fig. 6A). Finally, the CB1R-DOR mAb only detects an epitope in cortical membranes from wild type, but not CB1R−/− or DOR−/− mice (Fig. 6B). Next we examined CB1R-DOR levels in various brain regions 14 days after surgery and found that they are upregulated in some, but not all, brain regions: significant increases were observed in cortex (Fig. 6C), hypothalamus and midbrain, and no changes in striatum, hippocampus and cerebellum (not shown).

Allosteric modulation of DOR by interaction with CB1R during neuropathic pain

Allosteric modulation of CB1R on DOR activity was specifically explored by examining if a CB1R agonist could restore the suppressed DOR activity in cortical membranes from lesioned animals. For this, low, non-signaling doses of a CB1R agonist were used to examine whether occupancy, but not activation, of CB1R is sufficient to alter DOR activity. In the presence of Hu-210, a CB1R agonist, DOR activity in cortical membranes from lesioned animals increased (Fig. 7A & B). However Hu-210 had no effect on DOR activity in cortical membranes from sham animals (Fig. 7B). Whether occupancy of CB1R by an antagonist is sufficient to restore DOR activity was also examined. We found that the CB1R antagonist PF-541273 also enhanced DPDPE-simulated DOR activity in the cortex of lesioned animals (Fig. 7C). Neither MOR, nor kappa opioid receptor (KOR) specific ligands altered cortical DOR activity (Fig. 7D). Furthermore, treatment with a low, non-signaling dose of Hu-210 did not enhance DOR activity in membranes from hippocampus (Fig. 8; see panel c), a brain region in which CB1R and DOR levels do not change during neuropathic pain (Table 2). These results are consistent with the idea that the decrease in DOR activity in the cortex of neuropathic animals is due to an interaction with CB1R, and that this antagonistic interaction can be reversed by occupancy of CB1R. Together, the data suggests that occupancy of CB1R by an antagonist or a low, non-signaling doses of selective agonist) in

![Figure 1](image)
The cortex of lesioned animals is sufficient to allosterically modulate DOR activity.

CB1R-DOR heteromer-specific antibody blocks enhancement of DOR activity by CB1R ligands

In order to directly test whether the allosteric modulation of DOR activity by CB1R ligands is specific to the receptors within the CB1R-DOR heteromer, the CB1R-DOR heteromer selective antibody was used. The CB1R-DOR mAb selectively blocked Hu-210-mediated increases in DOR activity (Fig. 8A). In contrast, antibodies directed against other heteromer pairs (including MOR-DOR mAb [25] or CB1R-AT1R mAb [26]) or antibodies to CB1R or DOR alone did not block Hu-210 induced increases in DOR activity (Fig. 8A). The CB1R-DOR mAb did not alter basal [35S]GTPγS binding, nor did it alter DPDPE-stimulated DOR activity (Fig. 8B). Finally, CB1R-DOR mAb had no effect on DOR activity in the absence or presence of Hu-210 in membranes from hippocampus, a region in which neither CB1R, DOR, nor CB1R-DOR expression changed 14 days after lesion (Fig. 8C; Table 2). Together, these results support the idea that the potentiation of DOR activity by a low dose of Hu-210 is specific to regions in which CB1R-DOR heteromer expression is enhanced during neuropathic pain; the CB1R-DOR mAb only blocked heteromer-mediated signaling in those regions that showed an increase in CB1R-DOR expression.

DOR binding is enhanced by interaction with CB1R within CB1R-DOR heteromer

Next, the allosteric modulation of DOR activity by occupancy of CB1R using ligand binding assays was examined. In the cortical membranes from sham animals, a low dose of Hu-210 did not alter [3H]DPDPE binding to DOR (Fig. 9). However, in cortical membranes from lesioned animals, the same dose of Hu-210 significantly enhanced [3H]DPDPE binding to DOR (Fig. 9). These results show that the occupancy of CB1R leads to an allosteric modulation of DOR conformation that allows increased DOR binding by its selective ligand.

Whether DOR binding was allosterically modulated by CB1R was also examined using a cell culture model that allowed for manipulation of CB1R expression. In N2A-DOR cells, which endogenously express CB1R and are stably transfected with DOR, the binding of DOR ligand was enhanced in the presence of Hu-210 in a dose dependent fashion (Fig. 10A). This enhancement was blocked by the CB1R-DOR mAb (Fig. 10A) or by knocking down CB1R expression (Fig. 10B). This indicates that enhancement of DOR binding by Hu-210 requires the presence of CB1R.

Figure 2. CB1R levels increase in cortex of lesioned animals. A, Representative Western Blot and quantification for CB1R and CNX (Calnexin) in cortical membranes from 3 individual sham or lesioned animals, 14 days after surgery. Data represent Mean ± SEM (n = 6 animals/group). B, ELISA data for CB1R detection using a CB1R specific antibody in cortical membranes from sham and lesioned animals 14 days after surgery. Data represent Mean ± SEM (n = 4 animals/group). Statistically significant differences between sham and lesion groups are indicated*, p<0.05; **, p<0.01 (t-test). C, RT-PCR for CB1R (measured relative to GAPDH) in cortical preparations from sham and lesioned animals 14 days after surgery. Data represent Mean ± SEM (n = 5 animals/group). Statistically significant differences between sham and lesion groups are indicated *, p<0.05 (t-test). doi:10.1371/journal.pone.0049789.g002
and is mediated through the CB1R-DOR heteromer. We also examined if CB1R occupancy is sufficient to alter DOR binding, and found that treatment with the CB1R antagonist PF-514273 also enhanced [3H]DPDPE binding in a dose dependent fashion, and was blocked by addition of 1 μg of CB1R-DOR mAb (Fig. 10C). Together, these results indicate that enhancement of DOR activity requires the presence of CB1R, occupancy of CB1R, and is mediated through the CB1R-DOR heteromer.

Discussion

The present study reveals that neuropathic pain-induced suppression of DOR activity can be reversed through allosteric modulation of the CB1R-DOR heteromer. Specifically, we showed that 1) DOR and CB1R expression increased in select brain regions 14 days after peripheral nerve lesion; 2) DOR activity decreased, while CB1R activity increased in the cortex of lesioned animals, suggesting that CB1R activity may suppress DOR activity; 3) Treatment with a CB1R antagonist or a low, non-signaling dose of a CB1R agonist could restore suppressed cortical DOR activity back to normal levels, an indication that, under certain conditions, CB1R can allosterically modulate DOR activity and binding; and 4) Modulation of DOR activity by CB1R occurs within the CB1R-DOR heteromer, as a heteromer-specific antibody was able to block allosteric enhancement of DOR activity. These results demonstrate an important role for CB1R-DOR heteromer formation in cortex during neuropathic pain.

| Table 1. Time course of changes in CB1R and DOR levels in membranes from sham vs. lesioned animals. |
| Days post-surgery | CB1R | DOR |
|-------------------|------|-----|
| 3 days            |      |     |
| Sham              | 100.0 ± 21.32 | 100.0 ± 46.67 |
| Lesion            | 102.4 ± 17.84 | 50.37 ± 18.50 |
| Statistics        | p = 0.9369 | p = 0.3789 |
| 7 days            |      |     |
| Sham              | 100.0 ± 12.73 | 100.0 ± 13.02 |
| Lesion            | 102.1 ± 20.68 | 100.1 ± 40.77 |
| Statistics        | p = 0.9348 | p = 0.9982 |
| 14 days           |      |     |
| Sham              | 100.0 ± 8.766 | 100.0 ± 21.03 |
| Lesion            | 132.5 ± 4.801 | 280.9 ± 24.50 |
| Statistics        | p = 0.0314 (*) | p = 0.0050 (**) |

Levels of CB1R or DOR were measured by Western blot analysis as described in "Methods". p-values from t-tests comparing mean CB1R or DOR levels (percent relative to CNX and normalized to sham) in cortical membranes of sham vs. lesioned animals (n = 6 animals/group) at different time points after surgery. Statistically significant differences between sham and lesion groups are indicated.
* p < 0.05,
** p < 0.01 (t-test).

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Table 2. Changes in CB1R, DOR and CB1R-DOR levels in membranes from sham vs. lesioned animals.

| Region   | CB1R  | DOR  | CB1R-DOR |
|----------|-------|------|----------|
| Sham     | 100.0±4.74 | 100.0±12.27 | 100.0±11.02 |
| Lesion   | 147.0±11.02 | 127.4±17.78 | 127.4±17.78 |
| Statistics | p = 0.0572 | p = 0.0173 | p = 0.2737 |
| Hippocampus |      |      |          |
| Sham     | 100.0±5.91 | 100.0±23.41 | 100.0±5.91 |
| Lesion   | 85.5±8.05 | 84.76±5.84 | 88.26±6.08 |
| Statistics | p = 0.1367 | p = 0.7240 | p = 0.3636 |
| Cerebellum |       |      |          |
| Sham     | 100.0±26.41 | 100.0±4.96 | 100.0±5.76 |
| Lesion   | 116.2±21.58 | 114.3±31.39 | 155.4±37.20 |
| Statistics | p = 0.0124 | p = 0.0471 | p = 0.0020 |
| Hypothalamus |       |      |          |
| Sham     | 100.0±4.34 | 100.0±4.96 | 100.0±5.76 |
| Lesion   | 127.3±4.59 | 128.4±8.72 | 155.5±5.14 |
| Statistics | p = 0.0124 | p = 0.0471 | p = 0.0020 |
| Midbrain |       |      |          |
| Sham     | 100.0±11.65 | 100.0±16.65 | 100.0±8.20 |
| Lesion   | 169.4±14.39 | 131.1±1.70 | 156.0±4.09 |
| Statistics | p = 0.0200 | p = 0.1368 | p = 0.0036 |

Changes in CB1R, DOR or CB1R-DOR levels were detected in different brain regions by ELISA as described in “Methods”. p-values from t-tests comparing mean CB1R, DOR or CB1R-DOR levels (normalized to sham, percent of sham listed) in membranes from various brain regions of sham vs. lesioned animals (n = 6 animals/group) 14 days after surgery are shown. Statistically significant differences between sham and lesion groups are indicated.

* p < 0.05,
** p < 0.01,
*** p < 0.001 (t-test).

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Figure 4. DOR levels increase in cortex of lesioned animals. A, Representative Western Blot and quantification for DOR and CNX in cortical membranes from 3 individual sham or lesioned animals. Data represent Mean ± SEM (n = 6 animals/group). Statistically significant differences between sham and lesion groups are indicated **, p < 0.01 (t-test). B, ELISA data for DOR in cortical membranes from sham and lesioned animals. Data represent Mean ± SEM (n = 4 animals/group). Statistically significant differences between sham and lesion groups are indicated ***, p < 0.001 (t-test).

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Figure 5. CB1R activity increases while DOR activity decreases in cortex of lesioned animals. A, [35S]GTPγS binding was carried out with cortical membranes from sham and lesioned animals. Membranes from cortices were prepared as described in “Methods” and treated with 0.1 pM – 10 μM Hu-210 for 1 hour. [35S]GTPγS binding to membranes was detected using a scintillation counter. Basal [35S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 3 individual animals in triplicate). B, Membranes from cortices of sham and lesioned animals were treated with 1 pM – 10 μM DPDPE for 1.5 hours. [35S]GTPγS binding to membranes was detected using a scintillation counter. Basal [35S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 3 individual animals in triplicate).

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Though further studies will be necessary, the present findings suggest that a heteromer-directed therapeutic strategy for enhancing DOR activity could potentially be utilized to reduce neuropathic pain and associated anxiety behaviors.

The LSSTN peripheral nerve lesion model leads to the production of neuropathic pain, and our results show that mechanical allodynia in lesioned animals develops quickly – within 2 days – and is persistent, lasting at least 14 days. Previous studies have shown the development of mechanical allodynia [47,66–68], thermal hyperalgesia [45], central sensitization [69], and spontaneous C-fiber activity [70] using this pain model. Interestingly, a recent study found that spinal nerve ligated rodents do not develop anxiety or depression related behaviors until 15 days after ligation, although sensory hyperalgesia develops much earlier [71]. Our finding, that changes in cortical DOR and CB1R expression are not observed at 3 or 7 days after surgery but only at 14 days, highlight the possibility that changes in cortical receptor expression and activity underlie the development of behavioral phenomena such as anxiety and depression.

We also observed increases in DOR and CB1R levels using cortical membrane preparations. While this is the first study to examine changes in cortical levels of these receptors during neuropathic pain, previous human studies found increases in CB1R levels in prefrontal cortex during depression or after suicide [72,73], suggesting that CB1R upregulation in this region could underlie negative mood states. It is unclear whether increased receptor levels reflect increased receptor synthesis or redistribution of receptors to a particular membrane compartment. Previous studies have shown that pain-induced release of inflammatory mediators causes increased plasma membrane insertion of DOR in DRGs and spinal cord [11,74], though these changes occurred early (after 3 days) and likely underlie the effectiveness of DOR agonists as analgesic, not as anxiolytic or antidepressant agents.

The long term nature of DOR and CB1R upregulation in our studies (both DOR and CB1R levels are increased after 14 days, but not after 3 or 7 days), is suggestive of increased receptor synthesis. Moreover, our finding that lesioned animals showed increases in receptor staining in cortical cell bodies and processes suggests that new receptors are being synthesized and trafficked to neuronal processes. Previous studies have examined CB1R expression throughout the brain and found that the receptor is mostly localized to axons and synaptic terminals (reviewed in [75]). Our finding, that CB1R is localized in cell bodies, in addition to neuritic-like processes and synaptic-like puncta, suggests a new pattern of distribution for the receptor during a disease state. It is also possible that changes in DOR expression might be driving changes in CB1R localization (and vice versa). Studies in heterologous cells demonstrated that a lack of (or a decrease in)
DOR expression leads to intracellular trafficking of CB₁R. Conversely, increasing the level of DOR results in localization of CB₁R to the plasma membrane [22]. A similar mechanism might underlie the distribution and localization of CB₁R in the cortex of animals with peripheral nerve lesion.

In addition to changes in receptor expression and localization in cortex during neuropathic pain, we found that CB₁R, DOR and CB₁R-DOR were upregulated in hypothalamus and midbrain during neuropathic pain. Stimulation of either CB₁R or DOR in the midbrain, particularly in the PAG, can activate fibers that descend into the dorsal horn of the spinal cord and inhibit pain signals. Therefore, upregulation of these receptors in this brain region may be protective and may promote analgesia during neuropathic pain, although no studies have examined pain-associated changes in receptor activity in this region. The exact role of cannabinoid and opioid receptor expression in the hypothalamus is unclear, although activation of receptors in this region may promote feeding behavior and analgesia [75,76].

In this study we provide evidence supporting the idea that CB₁R-DOR heterodimer is formed in cortex during neuropathic pain. For this, we developed a CB₁R-DOR heteromer-specific monoclonal antibody using the subtractive immunization strategy and show that the antibody is both highly selective and is able to quantify heteromer levels in the cortices of animals experiencing neuropathic pain. Hence the heteromer-selective antibody serves as a unique and powerful tool to demonstrate heteromer-specific activity.

Allosteric modulation of receptor activity by dimerization has increasingly been recognized [77]. In this study, we found that the occupancy of CB₁R is sufficient to increase DOR ligand binding to its receptor. This indicates that the ligand occupancy-induced changes in CB₁R conformation alters the interaction between CB₁R and DOR such that DOR is better able to bind its own ligand.

**Figure 7.** DOR activity is enhanced in the presence of CB₁R ligands in cortical membranes from lesioned animals. A, Membranes from cortices of lesioned animals were treated with 10 pM – 10 μM DPDPE in the absence of presence of 1 pM Hu-210, or with 1 pM Hu-210 alone for 1.5 hours. [³⁵S]GTPγS binding to membranes was detected using a scintillation counter. Basal [³⁵S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 3 individual animals in triplicate). Statistically significant differences between 10 μM DPDPE alone and 10 μM DPDPE+1 pM Hu-210 are indicated ***, p<0.001, (t test). B, Membranes from cortices of sham and lesioned animals were treated with 10 μM DPDPE in the absence of presence of 1 pM Hu-210, or with 1 pM Hu-210 alone for 1.5 hours. [³⁵S]GTPγS binding to membranes was detected using a scintillation counter. Basal [³⁵S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 4 individual animals in triplicate). Statistically significant differences between 10 μM DPDPE alone and 10 μM DPDPE+1 pM Hu-210 are indicated **, p<0.01, (t test). C, Membranes from cortices of lesioned animals were treated with 10 μM DPDPE in the absence of presence of 1 μM PF-514273, or with 1 μM PF-514273 alone for 1.5 hours. [³⁵S]GTPγS binding to membranes was detected using a scintillation counter. Basal [³⁵S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 3 individual animals in triplicate). Statistically significant differences between 10 μM DPDPE alone and 10 μM DPDPE+1 μM PF-514273 are indicated *, p<0.05, (t test). D, Membranes from cortices of lesioned animals were treated with 10 μM DPDPE in the absence of presence of 1 μM Hu-210, or 10 nM DAMGO or 10 nM U69593 for 1.5 hours. [³⁵S]GTPγS binding to membranes was detected using a scintillation counter. Basal [³⁵S]GTPγS binding in vehicle treated membranes is taken as 100%. Data represent Mean ± SEM (n = 3 individual animals in triplicate). Statistically significant differences between 10 μM DPDPE and 10 μM DPDPE-ligand are indicated **, p<0.01, (t test).

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ligand. Previous studies have demonstrated that cannabidiol can reduce the rate of dissociation of [3H]naltrindole (a DOR selective ligand) from DOR [78], suggesting that high doses of cannabidiol or low doses of Hu-210 stabilize similar CB1R conformations. Finally, recent studies reported that the DOR ligands allosterically modulate MOR agonist binding and signaling, and this effect could be blocked by the application of a MOR-DOR-selective antibody [25,79]. Together, these studies emphasize the allosteric nature of the interaction between receptor protomers that could be harnessed for the development of combination therapies [80,81]. Based on our results, several pharmacological strategies for targeting CB1R-DOR during neuropathic pain could be proposed:

1) A combination of low doses of CB1R ligands with DOR agonists to increase the potency of selective DOR agonists in reducing anxiety during neuropathic pain; 2) A heteromer specific compound (e.g. bivalent ligand) composed of a weak CB1R ligand and full DOR agonist as a potent anxiolytic for neuropathic pain; and 3) Biologicals that block heteromer activity (such as, CB1R-DOR heteromer specific antibody or a peptide that selectively blocks CB1R-DOR interaction) to reduce neuropathic pain or possibly symptoms of anxiety and depression that associated with neuropathic pain. Strategy 1 (a low dose of DOR antagonists to enhance morphine effects) was employed successfully to enhance morphine analgesia in animal models [82,83,84]. Additionally,
using animal models, strategy 2 (bivalent ligands composed of a DOR antagonist and MOR agonist) was shown to result in more potent analgesics that induced less tolerance than morphine [85]. Strategy 3 (biologics that block heteromer activity) has been used to disrupt MOR-DOR heteromers [86] or D1-D2 heteromers [87] or to block DOR-KOR heteromers [88]. We hypothesize that cortical injection of a peptide disrupting CB1R-DOR interactions or a drug specifically blocking this heteromer would prevent the decrease in DOR activity and concomitantly reduce the development of long-term anxiety in neuropathic animals. However, additional studies will be needed in order to fully test this hypothesis.

In summary, we have identified CB1R-DOR as a novel entity regulating receptor activity during a disease state. Furthermore, we have identified allosteric modulation of one receptor’s activity by the other as a molecular mechanism to enhance the activity of the receptor compromised by the pathology. Such a mechanism can be targeted to develop therapeutics for enhancing the activity of disease-specific heteromers, particularly for those receptors whose activation is physiologically protective, but whose activity is suppressed by its partner receptor.

**Figure 9.** DOR binding is enhanced in the presence of low dose CB1R ligand in cortical membranes. Membranes from cortices of sham and lesioned animals were treated with 0.5 nM [3H]DPDPE in the presence or absence of 1 pM Hu-210 for 1 hour. Ligand binding assay was carried out as described in “Methods” and [3H]DPDPE binding to membranes was detected using a scintillation counter. Data represent Mean ± SEM (n = 7 animals/group). Statistically significant differences between [3H]DPDPE alone and [3H]DPDPE+1 pM Hu-210 are indicated ***, p < 0.001, (t test).

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**Figure 10.** DOR binding is enhanced by CB1R ligands in membranes of cells expressing CB1R and DOR. A, Membranes from N2A-DOR cells were incubated with 100 fM – 10 μM Hu-210 in the presence of 0.5 nM [3H]DPDPE + 1 μg CB1R-DOR monoclonal antibody for 1 hour. Ligand binding assay was carried out as described in “Methods” and [3H]DPDPE binding to membranes was detected using a scintillation counter. Data represent Mean ± SEM (n = 3 experiments in triplicate). B, Membranes from N2A-DOR cells in which CB1R expression was decreased by siRNA transfection were treated with 100 fM – 10 μM Hu-210 in the presence of 0.5 nM [3H]DPDPE. Ligand binding assay was carried out as described in “Methods” and [3H]DPDPE binding to membranes was detected using a scintillation counter. Data represent Mean ± SEM (n = 3 experiments in triplicate). C, Membranes from N2A-DOR cells were treated with 100 fM – 10 μM PF-514273 in the presence of 0.5 nM [3H]DPDPE along with the presence or absence of 1 μg CB1R-DOR monoclonal antibody for 1 hour. Ligand binding assay was carried out as described in “Methods” and [3H]DPDPE binding to membranes was detected using a scintillation counter. Data represent Mean ± SEM (n = 3 experiments in triplicate).

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Methods

Ethics Statement

Animal studies were carried out according to protocols approved by the Mount Sinai School of Medicine Institutional Animal Care and Use Committee (Permit # 02-0805).

Animals

Rats. Adult male Sprague-Dawley rats (200–250 g) were used for all experimental groups (sham and lesion). Animals were maintained on a 12-hour light/dark cycle and were allowed to acclimatize to their environment one week prior to experimentation.

Surgery

Rats subjected to L5 spinal nerve transection (L5SNT) were assessed on the VonFrey test on the day of surgery (prior to surgery) to establish a baseline response profile (see below for VonFrey protocol). Thereafter, animals were tested behaviorally every two days, and sacrificed 3, 7 or 14 days following L5SNT or sham surgery. Ten minutes prior to surgery, animals received an IM injection of ketamine (60 mg/kg)/xylazine (7.5 mg/kg) (Sigma-Aldrich, St. Louis, MO, USA). The surgical area was shaved and then cleaned with ethanol. The right paraspinal muscles (at lumbar levels L4–L6) were separated, and the L6 transverse vertebral process was exposed and partially removed. The L4 and L5 spinal nerves, which run below the L6 transverse process, were separated. The L5 spinal nerve was tightly ligated with a 5-0 silk thread; skin at the incision site was sutured (3-0 nylon). Topical connective tissue were sutured (3-0 nylon) into their anatomical layers; skin at the incision site was sutured (3-0 nylon). Topical antibiotic (bacitracin, neomycin, and polymyxin B) was applied to the incision site. Animals were monitored until awakening from the anesthetic and then returned to their home cages. For sham animals, the L5 spinal nerve was exposed but was not ligated or transected. All other surgical steps were identical between animals in these two groups. Experimental group size was 5–6 animals.

Von Frey Test

For mechanical allodynia, all animals (lesioned and sham) were placed onto a testing platform containing a metal, perforated floor (Stoelting Co., Wood Dale, IL, USA). After 15 min of acclimatization to the testing chamber, mechanical allodynia was assessed on both hindpaws using calibrated filaments (Stoelting Co., Wood Dale, IL, USA). All trials began with the 2.04 g filament, and proceeded using up-down trial design and standard quantitative methods [89,90]. Each filament was applied to the hindpaw six times per filament size. A positive response, defined as hind limb withdrawal or flinching on three of six tests or on two consecutive tests, to one filament was followed by the use of a smaller filament. Conversely, a negative response was followed by the use of larger filament. Each animal was tested with six filaments using the up-down method to determine the threshold of the mechanical response in the hindpaw.

Membrane Preparation

Membranes were prepared from cortices of control, sham and L5SNT lesioned animals. Animals were sacrificed with CO2 gas at 3, 7 or 14 days after surgery and then rapidly decapitated. Brains were removed and the following anatomical subregions were dissected out: hypothalamus, prefrontal cortex, rest of neocortex, striatum, hippocampus, PAG, midbrain, rostral ventral medulla and cerebellum. To prepare membranes, dissected tissue was manually homogenized in cold sucrose buffer containing 50 mM Tris (pH 7.4), 100 mM NaCl, 0.2 mM EDTA, 3 mM MgCl2, 1 mM DTT and 250 mM sucrose. The homogenate was centrifuged at 1000 g for 5 min, the pellet was discarded and the supernatant was then centrifuged twice at 40,000 g for 10 min. The resulting pellet was resuspended in homogenizing buffer containing 50 mM Tris-Cl (pH 7.4), 100 mM NaCl, 3 mM MgCl2, and 0.2 mM EGTA. Membranes were stored at −80°C. Membranes from DOR +/− mice [57] or CB,R −/− mice [92] were used as controls in ELISA experiments for validation of CB,R-DOR antibody.

Western Blotting

40 μg of proteins from membranes (prepared as described above) were added to 6× Laemmli sample buffer. Proteins were resolved by 10% SDS-PAGE and were subjected to Western blot analysis using a rabbit polyclonal antibody against the C-terminus of CB,R (Cayman Chemical, Ann Arbor, MI, USA) (1:300 dilution), a rat polyclonal antibody against DOR [22,93] (1:1000 dilution), or a rabbit polyclonal antibody against Calnexin (Cell Signaling, Boston, MA, USA) (1:10,000 dilution), a resident ER membrane protein. Blotting and imaging with the Odyssey imaging system (LI-COR, Lincoln, NE, USA) was performed as per the manufacturer’s protocols. The following secondary antibodies were utilized for visualization of the primary antibodies: goat-anti-rabbit IRDye 680 or goat-anti-rat IRDye 800 (both at a working dilution of 1:10,000) (Rockland Immunchemicals, Gilbertsville, PA, USA). Band intensity was densitized using the Odyssey imaging system (LI-COR, Lincoln, NE, USA) software.

CB,R-DOR antibody generation

The CB,R-DOR antibody was generated using a subtractive immunization strategy, essentially as described for the generation of our CB,R-AT1R antibody [26]. Mice were made tolerant to immunogenic epitopes in N2A cell membranes endogenously expressing CB,R by intraperitoneal injection. Tail bleeds were monitored for titers of antibody being produced. Upon exhibiting a decrease in antibody titer, as measured by ELISA, mice were then given intraperitoneal injections of membranes from N2A cells expressing DOR, along with booster injections over the next 15 days. Spleen cells from animals secreting antibody were then fused with SP-20 myeloma cells to generate monoclonal antibodies. Clones secreting monoclonal antibodies were screened by ELISA as described [82] against the following cell lines: untransfected N2A cell membranes that endogenously expressed CB,R, N2A cell membranes co-expressing CB,R with either DOR, MOR, KOR, CB,R or AT1R and HEK293 membranes expressing either DOR, KOR and DOR, or MOR and DOR. Cell lines were screened using 1:10 hybridoma supernatant and 1:500 horseradish peroxidase labeled anti-mouse IgG. Only hybridoma supernatants from clones that gave a specific signal with CB,R-DOR were further purified as described [82] and screened for specificity against cortical membranes prepared from wild-type, CB,R −/− and DOR −/− animals.

ELISA

Quantitation of levels of receptors following peripheral lesion was carried out by ELISA. Membranes (from n = 7 animals per group) were diluted in PBS (5 μg/100 μL) and coated onto 96-well plates (5 μg/well). All experiments were carried out in triplicate. Membranes were fixed with 4% paraformaldehyde (PFA) in phosphate buffered saline (PBS) for 20 min and then washed five times in PBS, followed by incubation in blocking buffer (1%
bovine serum albumin (BSA) in PBS) for 90 min. Membranes were incubated in primary antibody overnight at 4°C. Primary antibodies used included rabbit polyclonal antibody against C-term CB1-R (Cayman Chemical, Ann Arbor, MI, USA) (1:300), a rat polyclonal antibody directed against DOR ([68]) (1:1000) and a mouse monoclonal antibody directed against CB1-R-DOR (1:100). After washing in PBS, horse radish peroxidase conjugated secondary antibody was added for 90 min. After final washes in PBS, colorimetric substrate was added and the plate was scanned at 490 nm.

Quantitative Real-Time RT-PCR

The relative levels of receptor mRNA were quantified by quantitative PCR [26,91]. Total RNA was isolated from various brain regions using the TRIzol method [Invitrogen, Carlsbad, CA, USA]. RNA (1.0 µg) was reverse transcribed in 20 µL of buffer containing 50 µM oligo(dT)20, 25 mM MgCl2, 0.1 M dithiothreitol, 40 U/µL RNaseOUT, and 200 U/µL SuperScript III RT for 50 min at 50°C. The reaction was stopped by incubating the samples at 85°C for 5 min, after which 40 µL of nuclease free water was added. Real-time PCR was performed by using the Brilliant SYBR Green QPCR Master Mix (Agilent Technologies, Santa Clara, CA, USA). The PCR template source was either 30 ng of first-strand cDNA or purified DNA standard. Primers used were:

(i) CTTCCGTACCATCACCACAG (forward) and GAAGGCACCTCCGTAAGG (reverse) or (ii) TGTCTCCCATTT-CAAGCAAG and GGTGATGGTACGGAAGGTG (reverse). Amplification was performed with a spectrofluorometric thermal cycler (Stratagene, La Jolla, CA, USA). After an initial denaturation step at 95°C for 10 min, amplification was performed using 40 cycles of denaturation (95°C for 30 s), annealing (56°C for 1 min), and extension (72°C for 1 min). To standardize mRNA levels, GAPDH, a housekeeping gene, was used as an internal control. Gene expression was normalized by calculating the ratio between the number of cDNA copies of CB1-R and that of GAPDH in both sham and lesion conditions.

Immunohistochemistry

Localization of CB1-R within the cortex of sham and lesioned rats, 3, 7 and 14 days after surgery, was evaluated by immunohistochemistry. Rats were deeply anesthetized with 1% normal donkey serum. CB1R was labeled with a rabbit antibody at RT. Images were acquired with a Zeiss LSM510 Meta confocal microscope (Carl Zeiss, Thornwood, NY, USA). Typical sampling for this analysis was ~4 microscopy fields (acquired at 1024×1024 pixel resolution, with a z-step of 0.1 µm) and ~2 tissue sections equally spaced through the cortical layer of interest. For each sample, average intensity values were determined using ImageJ [NIH] software.

[^35]GTPγS Binding

Peripheral nerve lesion-induced changes in receptor activity were measured using [^35]GTPγS binding. Briefly, membranes (n = 6–7 animals per group) from sham or lesioned animals (14 days post-surgery) were incubated with increasing concentrations of Hu-210 (0.1 pM to 10 µM) or DPDPE (1 pM to 10 µM) in the presence of 2 mM GDP and 0.5 nM [^35]GTPγS as described in [94–96]. Basal binding in the presence of GDP and an absence of agonist and cold GTPγS was also determined. Non-specific binding was determined by the addition of 10 µM cold GTPγS to a parallel set of tubes. The radioactivity bound to membranes was separated by filtration and quantified by scintillation counting. Dose dependent activation of [^35]GTPγS binding by DPDPE was also measured in the presence of a non-activating concentration of Hu-210 (1 µM), or FF-514273 (1 µM) in cortical membranes from sham or lesioned animals. [^35]GTPγS binding was analyzed by calculating EC50 and Emax values for each set of experiments. Activation of [^35]GTPγS binding by 10 µM DPDPE ± 1 µM Hu-210 was also measured in the presence of a non-activating concentration of DAMGO (10 nM) or U69593 (10 nM) or in the presence of 1 µg of the following antibodies (CB1-R mAb, CB-R-DOR mAb, CB-R-AT1R mAb [26], MOR-DOR mAb [25], CB-R Ab, DOR Ab or non-specific IgG (Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA) in cortical membranes from lesioned animals.

Radioligand Binding

Membranes were prepared from cortices of sham and lesioned rats, as well as from N2A cells stably expressing DOR [22] or N2A-DOR cells in which CB1-R expression was knocked down by siRNA transfection (pooled siRNAs against CB1-R; from Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA). For all ligand binding experiments, membranes were added to cold assay buffer containing 50 mM Tris, 1 mg/ml fatty acid-free BSA, 10 mM MgCl2 and 0.5 mM DTT. Non-specific binding was assessed using 10 µM DPDPE. Total binding was measured using 0.5 nM [^3H]DPDPE in the absence or presence of indicated concentrations of Hu-210 or PF-514273 in the absence or presence of 1 µg of CB1-R-DOR monoclonal antibody (mAb). Binding assays were carried out for 120 min at 30°C. Membranes were filtered and radioactivity was measured using a liquid scintillation counter.

Statistical Methods and Analysis

For all experiments, changes in group differences were evaluated by using a repeated measures ANOVA followed by a Student’s post-hoc test. A p-value of <0.05 was considered to be statistically significant for all tests. All statistical operations were performed using GraphPad Prism, Version 5.0 [GraphPad Software, La Jolla, CA, USA]. For the analysis of [^35]GTPγS binding results, values from each experimental condition (e.g. DPDPE dose response curve ± 1 µM Hu-210) were compared using a one-way ANOVA. Here again, a p-value <0.05 was considered to be statistically significant. The reported r2 value was calculated from linear correlation analysis.

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Author Contributions
Conceived and designed the experiments: IB LAD. Performed the experiments: IB AG SDS LKM. Analyzed the data: IB AG SDS LKM.

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