Cannabinoid 1 receptor knockout mice display cold allodynia, but enhanced recovery from spared-nerve injury-induced mechanical hypersensitivity

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Abstract

Background: The function of the Cannabinoid 1 receptor (CB1R) in the development of neuropathic pain is not clear. Mounting evidence suggest that CB1R expression and activation may contribute to pain. Cannabinoid 1 receptor knockout mice (CB1R⁻⁻⁻⁻) generated on a C57Bl/6 background exhibit hypoalgesia in the hotplate assay and formalin test. These findings suggest that Cannabinoid 1 receptor expression mediates the responses to at least some types of painful stimuli. By using this mouse line, we sought to determine if the lack of Cannabinoid 1 receptor unveils a general hypoalgesic phenotype, including protection against the development of neuropathic pain. The acetone test was used to measure cold sensitivity, the electronic von Frey was used to measure mechanical thresholds before and after spared-nerve injury, and analysis of footprint patterns was conducted to determine if motor function is differentially affected after nerve-injury in mice with varying levels of Cannabinoid 1 receptor.

Results: At baseline, CB1R⁻⁻⁻⁻ mice were hypersensitive in the acetone test, and this phenotype was maintained after spared-nerve injury. Using calcium imaging of lumbar dorsal root ganlion (DRG) cultures, a higher percentage of neurons isolated from CB1R⁻⁻⁻⁻ mice were menthol sensitive relative to DRG isolated from wild-type (CB1R⁺⁺⁺⁺) mice. Baseline mechanical thresholds did not differ among genotypes, and mechanical hypersensitivity developed similarly in the first two weeks following spared-nerve injury (SNI). At two weeks post-SNI, CB1R⁻⁻⁻⁻ mice recovered significantly from mechanical hypersensitivity, while the CB1R⁺⁺⁺⁺ mice did not. Heterozygous knockouts (CB1R⁺⁺⁻⁻) transiently developed cold allodynia only after injury, but recovered mechanical thresholds to a similar extent as the CB1R⁻⁻⁻⁻ mice. Sciatic functional indices, which reflect overall nerve health, and alternation coefficients, which indicate uniformity of strides, were not significantly different among genotypes.

Conclusion: Cold allodynia and significant recovery from spared-nerve injury-induced mechanical hypersensitivity are two novel phenotypes which characterize the global CB1R⁻⁻⁻⁻ mice. An increase in transient receptor potential channel of melastatin 8 channel function in DRG neurons may underlie the cold phenotype. Recovery of mechanical thresholds in the CB1R knockouts was independent of motor function. These results indicate that CB1R expression contributes to the development of persistent mechanical hypersensitivity, protects against the development of robust cold allodynia but is not involved in motor impairment following spared-nerve injury in mice.

Keywords

Cannabinoid 1 receptor, CB1R knockout, dorsal root ganglion, allodynia, spared-nerve injury, neuropathic pain

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Background

Despite the plethora of studies examining the potential of cannabinoids for the treatment of pain of various etiologies, relatively little information is known about the spatial and temporal changes of the targets of these compounds within the nervous system. One of these targets is the cannabinoid 1 receptor (CB1R), a G-protein coupled receptor localized on neurons in the peripheral and central nervous systems that modulates neurotransmitter release. Because CB1R is one of the most abundant G-protein coupled receptors in the body and is located on key circuits involved in nociception, perturbations of CB1R expression may affect the experience of pain. After injury and during painful states, there are dynamic changes in CB1R expression; the magnitude and direction of these changes appear to be dependent upon the severity of the insult, the type of tissues affected, and the time points examined relative to the initial injury. However, it is not clear how CB1R expression is involved in the development of neuropathic pain. For cannabinoids to be used safely and effectively in the treatment of neuropathic pain, it is necessary to understand how expression of CB1R affects the course of sensory and motor impairment after nerve injury.

Cannabinoids have been suggested as potential adjuvants for neuropathic pain treatment, a particularly difficult condition to treat in humans since it is often refractory to any treatments. Cannabinoid usage is controversial, partly because of its small therapeutic window before the manifestation of a myriad of side effects associated with stimulation of CB1R in the central nervous system. Moreover, there is evidence to suggest that CB1R stimulation may not be effective in relieving neuropathic pain and may actually contribute to hyperalgesia in experimental pain models in animals and humans. Specifically in preclinical neuropathic pain models, the effectiveness of cannabinoids is inconsistent. For example, in the chronic constriction injury (CCI) model, intrathecal injection of the non-specific CBR agonist WIN 55,212-2 is effective in mitigating mechanical allodynia, and it was postulated that CB1R up-regulation in the spinal cord after injury mediates this pain relieving effect. In contrast, others have demonstrated that an antagonist of CB1R administered systemically significantly reduces mechanical and thermal hyperalgesia induced by CCI in rats and in mice. Several additional studies show that compounds that inhibit CB1R promote pain relief and recovery after peripheral nerve and burn injuries, suggesting that CB1R expression and activation can be maladaptive.

To date, no one has examined the contribution of CB1R expression in the development of spared-nerve injury (SNI)-induced neuropathic pain, since standard analgesics such as morphine and gabapentin are much less effective in rats with SNI. The mixed CBR receptor (CB1R/CB2R) agonist WIN 55,212-2 when administered to mice with SNI is similarly ineffective. Interestingly, low doses of WIN 55,212-2 are antihyperalgesic in several other pain models, suggesting that cannabinoid receptors are differentially modulated depending on the injury. We sought to determine if the lack of CB1R unveils a general hypoalgesic phenotype. SNI was performed on CB1R wildtype (+/+), homozygous knockout (−/−), and heterozygous knockout mice (+/−) generated by Zimmer et al. The electronic von Frey and acetone tests were performed before and after injury to determine whether a global absence of functional CB1R would unveil hypoalgesia to innocuous cold and mechanical stimuli and protect against the development of persistent neuropathic pain.

Methods

Animals

All experiments were approved by and conducted in accordance with guidelines set forth by the New York University Animal Care and Usage Committee. Cannabinoid 1 receptor knockout (CB1R−/−) mice were generated from CB1 heterozygous (+/−) breeding pairs obtained from Dr Andreas Zimmer, University of Bonn, Germany, through the European Mutant Mouse Archive. Genotypes were confirmed with PCR as previously described. Adult male mice (8–12 weeks old) were used for experiments. Food and water were available ad libitum.

Behavioral measurements: Mechanical and thermal thresholds

All behavior tests were performed during the day portion of the circadian cycle (06:00 h to 18:00 h), and the experimenter was blinded to the genotype during the tests.

To measure mechanical thresholds, the electronic Von Frey method was used. Mice were first weighed on a scale before being placed in separate plexiglass boxes on top of a raised wire mesh. They were left to acclimate for approximately 30 minutes prior to testing. Stimulation of the lateral area of the hind paws was done using the 90 g arm and the size 8 filament. The maximum threshold readout that accompanied a paw flick was recorded. Both paws of each animal were stimulated three times, with a 2-minute rest in-between stimulations.

To measure for cold hypersensitivity, the acetone test of evaporative cooling was used. After the von Frey test, mice were allowed to rest for approximately 20 minutes before commencing with acetone applications.
A small drop of acetone was placed on the plantar surface of the hind paws with a pipette, making sure that only the acetone drop touched the paw, and not the pipette tip. The duration of paw withdrawal was recorded for 30 seconds afterwards. As a control, we placed a drop of water at 37°C on the paws. This induced a quick paw flick (<1 s), but no further paw withdrawal (data not shown).

**Spared-nerve injury**

Using aseptic surgical technique, under 2% isoflurane anesthesia, the skin on the lateral surface of the right thigh was incised, and the biceps femoris muscle separated to expose the sciatic nerve and its three terminal branches: the sural, common peroneal, and tibial nerves. The common peroneal and tibial branches were ligated together with a 9-0 silk suture, transected distal to the ligature, and 2 mm of each distal nerve stump was removed. The overlying muscle was closed with an absorbable suture 6.0 PDS Plus Antibacterial (Polydioxanone) by Ethicon, and the skin closed with non-absorbable monofilament, nylon size 5 sutures. After the surgeries, mice were housed separately and monitored for signs of autotomy or distress.

**Dissociated dorsal root ganglia primary cultures**

Primary cultures of dorsal root ganglion (DRG) from lumbar areas of the spinal column were extracted from CB1R+/-+and CB1R-/- mice at 3–4 months of age. The tissues were enzymatically digested with trypsin and collagenase at 37°C, subjected to a gradient to remove myelin and plated on poly-D-lysine coated coverslips (12.5 μg/ml or 25 μg/ml). Cells were grown for two days in Neurobasal A media supplemented with 2% B27, 1% Pen/Strep, 0.5 mM Glutamax, and NGF (100 ng/ml) and GDNF (10 ng/ml), as previously described.

**Calcium imaging**

To measure changes in intracellular calcium levels in DRG cultures derived from CB1+/- and CB1-/- mice, the ratiometric calcium indicator dye FURA-2 AM and the PTI (Photon Technology International, Inc. Lawrenceville, NJ, USA) imaging system were used. All measurements were performed at 48 hours in vitro within a temperature range of 30°C to 32°C. FURA-2 was excited at 340 nm and 380 nm by using a high-speed multi-wavelength illuminator alternated every 0.5 seconds. The emitted fluorescence of single cells was filtered with a fluorescence barrier filter at 515 nm, collected with a Cool-Snap HQ2 camera, and analyzed with ImageMaster 5 software. Cells were continually perfused with a HEPES buffer (pH 7.4) containing (in mM): 140 NaCl, 5 KCl, 5 NaHCO₃, 10 HEPES, 1 MgCl₂, 1.5 CaCl₂, and 10 glucose. Cells were “pulsed” using a pressurized superfusion system (AutoMate Scientific, Inc) with the following compounds (dissolved in HEPES buffer): KCl (50 mM) for 5 seconds to confirm the presence of viable neurons and menthol (250 μM) for 7 seconds to stimulate transient receptor potential channel of melastatin 8 (TRPM8) channels. Cells were allowed to recuperate for 10 minutes between stimulations.

**Data analysis**

For behavioral tests, a total of three stimulations per paw were administered to each animal on the testing day. Averages per animal were calculated from these data and pooled according to genotype. Data are presented as the mean ± SEM. Data were pooled, and t tests performed using Graph Pad Prism 6 software. A p value less than 0.05 was considered statistically significant. Percentage decreases in mechanical thresholds post-SNI were calculated from baseline. The two-way ANOVA test was used for time course of mechanical and cold-hypersensitivity following SNI, with genotype and time as main factors; Tukey’s post hoc test with multiple comparisons was used.

For the imaging experiments, calcium traces from the 340 nm and 380 nm signals, and the derived ratio were exported from the Image Master 5 software into Microsoft Excel. Traces were analyzed in pClamp 10.5.2.6 and Graph Pad Prism 6. Neurons were identified on the basis of their increased cytoplasmic response to 50 mM KCl. Neurons were considered menthol sensitive (MS) if cytoplasmic calcium levels increased at least 5% in the period after drug administration relative to the pre-stimulation baseline. Neurons that were not responsive were designated as menthol insensitive (MI). Differences in menthol sensitivity between genotypes were assessed using the two-tailed, Fisher’s exact test.

**Footprint analysis**

For footprint analysis, mice performed several pre-training trials during which they were habituated to a runway. A total of N = 4 mice per genotype were used. The start of the runway was brightly lit and the end contained a darkened enclosed chamber. Immediately after the pre-training sessions, footprints were recorded on a white paper on the runway floor after applying non-toxic tempera paint to the paws. Front and rear paws were distinguished by their differing paint color. Between 1 and 3 sets of footprints were collected for each animal in order to obtain footsteps that provided complete print of the foot as well as contiguous steps. From these trials,
the highest quality footprints were chosen and used for analysis for each. The footprint patterns were analyzed using the program Image J (NIH). Toe spread was defined as the distance between the most extreme digits. Paw length was measured beginning from the tip of the longest digit to the end of the heel. Sciatic functional indices (SFI) were calculated from the footprint patterns as previously described, by using the following formula: $SFI = 118.9 \left( \frac{ETS-NTS}{NTS} \right) - 51.2 \left( \frac{EPL-NPL}{NPL} \right) - 7.5$, where ETS is the experimental toe spread, NTS is the normal toe spread, EPL is experimental toe length, and NPL normal paw length. An SFI near “0” is a normally functioning nerve, whereas a value of −10 indicates complete impairment.

The distance from the R → L (horizontal distance of the step) foot was measured in addition to the R → R (forward distance of the step). An N of three measurements for each horizontal and forward was obtained when possible. The alternation coefficient was then derived by calculating the absolute value of 0.5 minus the ratio of the average horizontal to average forward distance for each mouse, as previously described. The purpose of the alternation coefficient is to show the level of variability or uniformity between each step. A gait which is completely in tandem will have a horizontal to forward ratio of 0.5 resulting in an alternation coefficient of 0. The coefficient was calculated pre-operatively in the week prior to injury, as well as on the following post-operative days (POD): 1, 3, 7, 14, 21, and 28. The coefficients were pooled by genotype and averaged for each day of the study. The average coefficients by genotype and day were then plotted on an $x$–$y$ line graph in Graph Pad Prism 6. Standard errors of the mean were calculated for each genotype by day.

**Results**

**CB1R$^{-/-}$ mice are hypersensitive to evaporative cooling, but exhibit no differences in mechanical thresholds**

Before performing the peripheral nerve injury, baseline cold responses, mechanical thresholds, body weights, and footprint patterns were measured in adult male CB1R$^+/+$, CB1$^+/-$, and CB1R$^{-/-}$ mice to determine the extent to which CB1R expression affected these parameters. The innocuous cooling agent acetone was used to measure cold sensitivity. The homozygous (CB1R$^{-/-}$) knockouts, but not the heterozygous knockouts (CB1R$^{+/+}$), displayed significant cold allodynia in both paws relative to the CB1R$^{+/+}$ mice (Figure 1(a)). The average duration of paw withdrawals for the CB1R$^{-/-}$ versus the CB1R$^{+/+}$ mouse was 3.75 ± 1.01 versus 1.26 ± 0.41 s in the right paw. While there was a significant difference in cold responses, mechanical thresholds, measured with the electronic von Frey, were the same among the genotypes (Figure 1(b)).

Thresholds in the left and right paws of the mice were 4.39 ± 0.16 and 4.26 ± 0.19 in CB1R$^+/+$ mice; 4.20 ± 0.25 and 4.05 ± 0.24 in CB1R$^{+//-}$ mice; and 4.14 ± 0.33 and 3.81 ± 0.25 in the CB1R$^{-/-}$ mice (Figure 1(b)). The heterozygous and homozygous knockouts weighed significantly less than the CB1R$^{+/+}$ mice (Figure 1(c)), but this finding did not affect the mechanical thresholds (Figure 1(b)). As a measure of potential gait abnormalities, footprint patterns were measured. No significant differences were observed in toe spreads, paw lengths, or step widths (Figure 1(d)).

**An increased percentage of CB1R$^{-/-}$ DRG neurons are sensitive to menthol relative to CB1R$^{+/+}$ neurons**

Before SNI, the only significant behavioral difference found to be affected by a global absence of CB1R was cold sensitivity. This was suggestive of a change in the function of a temperature sensitive channel within primary sensory neurons of the DRG. A drop of acetone on the hindpaws is an innocuous cold stimulus, and the TRPM8 is activated upon innocuous and noxious cooling; hence, we tested whether TRPM8 sensitivity in DRG neurons is correlated with the cold allodynia in the CB1R$^{-/-}$ mice. Primary dissociated cultures generated from the lumbar DRG of CB1R$^{+/+}$ and CB1R$^{-/-}$ mice were challenged with the TRPM8 agonist menthol. A significantly higher percentage of neurons from the CB1R$^{-/-}$ cultures were sensitive to menthol compared to CB1R$^{+/+}$ (39% (47/119) versus 16% (25/158) as measured by Ca$^{2+}$ imaging ($p<0.0001$, Figure 2(a)). Interestingly, the mean amplitude of the menthol responses in the CB1R$^{-/-}$ neurons was significantly lower compared to CB1R$^{+/+}$ neurons (Figure 2(b)), but more neurons with a diameter of 20 μm and above responded to menthol in CB1R$^{-/-}$ cultures (Figures 2(e) to (f)). Neurons that were MI had a similar diameter frequency distribution between the genotypes (Figure 2(f)). Overall, lumbar DRG cultures from both genotypes contained neurons with similar diameter distributions, with a mean of 20.4 ± 0.5 μm in CB1R$^{+/+}$ cultures and 21.4 ± 0.7 μm in CB1R$^{-/-}$ cultures (Figure 2(g)). Similar frequencies across a range of diameters, but different responses to menthol stimulation indicate that CB1R genotype affected TRPM8 sensitivity rather than a preferential survival of a subset of neurons in culture. An increase in the percentage of menthol-sensitive DRG neurons (Figure 2) is related to the observed cold allodynia phenotype displayed by CB1R$^{-/-}$ mice (Figure 1(a)).
CB1R expression protects against the development of cold allodynia, but contributes to persistent mechanical hypersensitivity following SNI in mice

After collecting pre-surgery data, mice were subjected to SNI as described in methods. Following SNI, the high number of cold responses was maintained in the ipsilateral paws of the CB1R+/−/− mice for the four-week observation period (Figure 3(a)), but not in the contralateral paws (Figure 3(b)). At POD 7, 14, and 21, the contralateral paws lost the cold allodynia phenotype, but regained it by POD 28 (Figure 3(b)). Interestingly, though CB1R+/− protected against cold hypersensitivity before surgery (Figure 1(a)), this genotype was sufficient to produce transient, significant cold hypersensitivity at POD 7 comparable to CB1R−/− mice. The cold allodynia phenotype never developed in the CB1R+/+ mice (Figure 3(a)). However, in all the mice, the contralateral paws displayed a decrease in their cold sensitivity between POD 7 and 21 (Figure 3(b)), which resulted in significant differences between the ipsilateral and contralateral paws in CB1R+/− (POD7, p < 0.0001) and CB1R−/− mice (POD7, p < 0.05).

With respect to mechanical thresholds, all genotypes developed a similar magnitude of hypersensitivity in the ipsilateral paws, indicated by a significant decrease in thresholds immediately after the injury (Figure 3(c)). However, beginning at POD 14, the threshold of the CB1R+/− and CB1R+/−/− mice began to significantly increase relative to CB1R+/+ mice, reaching approximately 50% recovery (Figure 3(c)). The contralateral paws of mice from each genotype never developed mechanical hypersensitivity (Figure 3(d)). Although CB1R+/+ mice weighed, on average, more than the knockouts at every time point, animals in all groups maintained their weight following SNI, indicating that the surgery did not limit their access to and interest in food (data not shown). Moreover, for a given animal per
genotype, changes in mechanical thresholds over time are not due to differences in body weight.

**CB1R expression is not involved in SNI-induced motor impairment**

Following SNI, toe spread significantly decreased for each genotype relative to their baselines, throughout the four week observation period (Figure 4(a)). There was a tendency for the injured paw length to increase over time, but was only significantly higher in CB1R\(^{+/+}\) at POD14 relative to pre-surgery length (Figure 4(b), \(p < 0.05\)). Step width also significantly increased immediately after SNI, but resolved to pre-surgical distances by POD28 (Figure 4(c)). A two-way ANOVA was conducted that examined the effect of genotype and time on toe-spread, paw length, or step width. There was no statistically significant interaction between genotype and time, on these parameters. As an overall measure of motor nerve health, SFIs significantly decreased to a similar extent after SNI in all genotypes. This decrease was maintained over the entire observation period (Figure 4(d)). Moreover, uniformity of strides, measured by alternation coefficients was similar between genotypes (Figure 4(e)).

**Discussion**

In this study, we found that cold allodynia and significant recovery from SNI-induced mechanical hypersensitivity are two novel phenotypes which characterize the global CB1R\(^{-/-}\) mice. In the acetone test of evaporative cooling, the CB1R\(^{-/-}\) mice lift their paws significantly longer in response to this innocuous stimulus. The cold allodynia is related to an increase in functional TRPM8 channels in DRG neurons, as assessed by calcium imaging of menthol-stimulated cultures. After SNI, the ipsilateral paw maintained cold hypersensitivity for at least four weeks, but the contralateral paws lost this phenotype at weeks 1, 2, and 3. Thus, central CB1R expression may also be contributing to the sensory phenotypes following SNI. Moreover, it was also found that CB1R expression and constitutive activity do not affect baseline mechanical thresholds or the level of injury-induced mechanical hypersensitivity during the first two weeks following SNI. However, the presence of CB1R appears to contribute to persistent mechanical allodynia in the SNI model, since significant recovery was evident only in mice lacking CB1R. Collectively, these results indicate that CB1R may protect against the development of some types of pain such as cold alldodynia, but it may...
contribute to the development of other types of pain such as nerve injury-induced mechanical hypersensitivity. While global deletion of CB1R may promote significant recovery from SNI-induced mechanical hypersensitivity in mice, it does not promote significant motor recovery.

With respect to the thermal phenotype of CB1R−/− mice, because the acetone test measures responses to innocuous cooling, the TRPM8 channel is a likely candidate involved in the behavioral responses that were observed. TRPM8 within the DRG contributes to the perception of cold sensation under normal conditions, and increases in this channel’s activity has been associated with the development of cold allodynia. In order to further probe this behavioral finding, menthol stimulation of DRG primary cultures was used to determine whether TRPM8 channel sensitivity differs between the genotypes. It was found that CB1R−/− DRG cultures have a significantly greater percentage of neurons sensitive to menthol, including larger diameter neurons; however, the peak menthol-induced calcium responses are lower in the knockouts. Menthol responses in the larger diameter neurons from CB1R−/− are suggestive of a phenotypic switch, since in vivo TRPM8 expression is found in approximately 10% to 15% of the total adult DRGs and mostly in small diameter, non-peptidergic, and un-myelinated neurons of C and Aδ fibers. Our data are the first to demonstrate that levels of CB1R in sensory neurons significantly affect menthol responses and thus TRPM8 channel activity. An increase in the proportion of sensory neurons with functional TRPM8 channels correlates with the presence of cold allodynia in CB1R−/− animals. In prior studies using TRPM8-transfected HEK cells, a relationship between CB1R activity and TRPM8 channel function has been suggested. While cannabinoids can directly inhibit TRPM8 channel activity, this inhibition is augmented with increased CB1R expression. These findings are consistent with our observations in that a complete down-regulation of CB1R expression (CB1R−/−) results in an increase in TRPM8 activity.

Figure 3. CB1R expression affects the development of cold and mechanical hypersensitivity following SNI in mice. (a) Cold responses in ipsilateral paws of CB1R−/− mice are significantly higher than CB1R+/+ responses (*p < 0.05; **p < 0.01; ***p < 0.001) and CB1R−/− mice develop this phenotype at day 7 (^p < 0.05 vs. CB1R+/+). Two-way ANOVA with Tukey’s post hoc test with multiple comparisons. (b) Cold responses in the contralateral paws: *p < 0.05, **p < 0.01 CB1R−/− vs. CB1R+/+; #p < 0.05 CB1R−/− vs. CB1R+/+. Significances between ipsilateral and contralateral paws are indicated in results. (c) Percentage change in mechanical thresholds calculated from baseline before surgery (POD -3). All genotypes develop mechanical hypersensitivity, but beginning at POD14 up to POD28, thresholds are significantly higher in the knockouts (CB1R−/− and CB1R+/−) compared to CB1R+/+. *p < 0.05; @ p < 0.01; $ p < 0.001; #p < 0.0001 vs. baseline; *p < 0.05; ***p < 0.001 vs. baseline at particular time point. (d) Percentage change of contralateral paw mechanical thresholds.
Presently, we cannot rule out compensatory changes of other ion channels in CB1R−/− that may contribute to the cold responses in the mice and increased menthol sensitivity in the DRG sensory neurons.15,16 However, CB1R expression levels indeed appear to be important for cold hypersensitivity and mechanical thresholds, since we also show that partial CB1R knockdown (CB1R+/−) is sufficient to induce transient cold allodynia after SNI and promote recovery from mechanical allodynia.

While pre-surgical mechanical thresholds are similar among genotypes, significant recovery from SNI-induced mechanical hypersensitivity occurs only in the CB1R−/− and CB1R+/− mice. To our knowledge, this is the first study to demonstrate these phenotypes. The data suggest that constitutive activity of CB1Rs or their stimulation by endocannabinoids following SNI contributes to persistent mechanical hypersensitivity. In contrast, studies using global CB1R−/− mice and other models of peripheral nerve injuries fail to demonstrate any differences in mechanical threshold changes relative to wildtype.42–44 These studies include partial sciatic nerve ligation (PSL) performed on the same mouse line used in our study43; and CCI of the sciatic nerve performed on global CB1R−/− mouse generated on a C57Bl/6 background9 and on a CD-1 background.44 PSL and CCI involve ligating different proportions of sciatic nerve fibers at the mid-thigh level.45 The SNI model used here involves the ligation, transection, and partial removal of two of the three terminal branches of the sciatic nerve distal to the ligation.19 In PSL and CCI, there is intermingling of the injured and intact axons both proximal and distal to the injury, but in SNI, this interaction only occurs proximal to the injury. During the behavioral testing, the SNI-evoked responses are from the uninjured sciatic nerve fibers and possibly from the sprouting sympathetic and saphenous nerves, whereas responses in the PSL and CCI models include both the intact and regenerating sciatic nerve axons.22 We show in the SNI model that CB1R−/− and CB1R+/− promotes significant recovery of mechanical thresholds. Collectively, irrespective of mouse strain, in peripheral-nerve injuries that do not involve transection of sciatic nerve fibers (i.e., PSL and CCI), global CB1R expression perturbation does not significantly contribute to sensory changes. In contrast, hypersensitivities resulting from sciatic nerve fiber transection injuries (i.e., SNI) appear to be significantly affected by CB1R expression. It is, therefore, possible that there are different mechanisms by which CB1R is involved in the development of nerve injury-evoked functional changes. In fact, it has been suggested that analgesic sensitivity is dependent on the nerve injury model,22 and this may partly explain why CBR agonists have been reported as ineffective in SNI-induced neuropathic pain in mice.8
Because the data presented in the current study are derived from global CB1R−/− mice, the distinct contributions of peripheral and central nervous system regions are not clear. Several mouse lines have been generated in which CB1R is conditionally deleted from a particular type of neuron; these include deletions in principal forebrain neurons, GABAergic neurons in the brain, GABAergic neurons in the dorsal horn, and in peripheral nociceptors. Of those, SNI has only been performed on the mice with conditional CB1R deletion in peripheral nociceptors. This mouse line targeted DRG that also co-expressed IB4, TRPV1, Substance P, and Na+,1.8 channels; however, CB1R expression was not affected in large diameter A neurons expressing NF200. In these mice, pre-surgical mechanical thresholds were lower while acetone responses were the same relative to control wild-type mice. After SNI, no differences in mechanical thresholds were evident in the three week observation period and only the conditional knockouts had significant cold allodynia two weeks after injury. However, within the DRG, CB1R expression is not limited to small diameter, nociceptors; CB1R is expressed across of range of DRG neurons in vivo, including those expressing calcitonin gene related peptide (CGRP). Though mice with a conditional deletion of CB1R on CGRP expressing sensory neurons have not been generated, selective ablation of the CGRP neuron population caused increased responses in the acetone test, but had no effect on mechanical thresholds pre- and post-SNI. Taken together, CB1R expressing cells play a complex role in somatic sensation, since global or conditional deletion produces varying phenotypes, when assessed before and after SNI. Hence, the distinct pain phenotypes of CB1R knockout mice observed in this study may involve both peripheral and central mechanisms.

Though we found significant genotypic differences in sensory tests, gross motor function was not differentially affected following SNI. The motor findings are not completely unexpected. In nerve transection injuries proper synapse formation is hindered and motor function continues to be significantly impaired at two months. In contrast, if the sciatic nerve is crushed, sensory and motor functions are completely restored within one month. The SNI model in the mouse further ensures prolonged denervation of sciatic fibers because the tibial and peroneal branches are also ligated before being transected, and pieces of the nerves are also removed. Importantly, SNI does not preclude the capacity for motor recovery since effective pharmacological interventions have been found. However, by computing sciatic functional indices and assessing alternation coefficients, we showed that expression of CB1R does not significantly improve motor function, and is therefore not involved in SNI-induced motor impairment.

Given that cannabinoids acting at CB1R are attractive therapeutic compounds for the relief of pain, yet neuropathic pain conditions can manifest as a complex mixture of mechanical and thermal allodynia and hyperalgesia, further studies are warranted to understand how acute and chronic treatments with CB1R agonists affect the distinct symptoms experienced by pain patients. Our data add to the findings of several neuropathic pain animal models, but emphasize that differences in the regulation of endocannabinoid system exist across models and likely across humans. Within the spectrum of neuropathic pain disorders in patients, there appears to be mechanistic heterogeneity which may invariably contribute to the efficacy of cannabinoids.

At this point, it is difficult to extrapolate for which conditions and symptoms CB1R activation will be most efficacious. With respect to thermal sensitivity, our present findings in the knockout mice, other pre-clinical work and experimental pain models in humans suggest that CB1R activation may oppositely modulate the thermo-TRP channels TRPM8 and TRPV1. Additional clinical studies are needed to measure the effect of CB1R activation specifically on thermal and mechanical thresholds. Our data also suggest that CB1R antagonism may prevent the development or ameliorate established nerve-injury induced mechanical hypersensitivity. While animal studies show this intervention to be promising, further drug development of CB1R neutral antagonists or negative allosteric modulators may be key, since severe psychiatric side effects can precipitate in vulnerable patients taking CB1R inverse agonists.

When discussing the place of individual cannabinoids, whole plant, or extracts of marijuana in pain management, it is of great importance to emphasize that medical marijuana throughout the USA and in other countries is not universally regulated with respect to cannabinoid content and ratio, or the form and route of administration. Marijuana contains many cannabinoids and other chemicals that interact to produce their effects in the body. Marijuana is as heterogeneous and complex as the pain patient, urging both caution and continued efforts to elucidate pain mechanisms that will ultimately inform optimal, personalized cannabinoid therapy.

Conclusion

Compounds acting at CB1R may have therapeutic potential for the relief of pain, including neuropathic origin. It is important to decipher how CB1R expression correlates with behavioral endpoints. To our knowledge, the data presented here are the first to demonstrate that global CB1R−/− mice are hypersensitive to evaporative cooling and recover significantly from SNI-induced mechanical hypersensitivity. Our data
suggest that CB1R expression contributes to distinct aspects of persistent neuropathic pain in the SNI model in mice.

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Authors’ contributions

AS performed the surgeries, the behavioral tests, calcium imaging, data analysis, and wrote the manuscript; BP assisted in primary culture preparation, calcium imaging, and footprint analysis; LR assisted in initial surgeries, behavioral testing, and footprint analysis; MN taught the surgeries and assisted in behavioral testing; TJJB and ERP were involved in conceptualizing the study and providing support. All authors read and contributed to the final version of the manuscript.

Declaration of Conflicting Interests

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