Hemopressin an Inverse Agonist of Cb1 Cannabinoid Receptors Reverses Mechanical Sensitivity on Diabetes-Induced Neuropathy in Mice

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Abstract

Peripheral neuropathy is one of the most common complications of diabetes affecting about 50% of patients with the disease. The most prominent symptoms involve the extremities and occur as both an exaggerated response to noxious stimuli (hyperalgesia), mild or non-painful stimuli (allodynia). Hemopressin (Hp) is a non aepetidine first found in rat brain extracts, which selectively binds CB1 cannabinoid receptors (CB1R) and exerts antinociceptive actions in experimental inflammatory and neuropathic pain models. However, there is no data about its efficacy in neuropathic metabolic-related disease, like diabetes mellitus. The aim of this study was to investigate the role of Hp on mechanical and thermal sensitivity of mice submitted to an experimental model of type 1 diabetes mellitus-induced neuropathy. Mechanical allodynia and thermal sensitivity were assessed by von Frey filaments or plantar test, respectively, 7, 14, 21 and 28 days after streptozocin injection (STZ; 200 mg/kg). Body weight and blood glucose were monitored once a week. Hp was administered orally, once a day (2.5 mg/kg) for 28 days. Hp reversed mechanical allodynia in diabetic mice without changing blood glucose levels or body weight. No effects were observed for thermal sensitivity. These results make hemopressin an attractive approach for the development of cannabinoid-based therapies for the treatment diabetic neuropathic pain.

Keywords: Hemopressin; Diabetic neuropathy; Cannabinoid receptor; Mechanical allodynia; Thermal sensitivity; Mice

Abbreviations: Hp: Hemopressin; STZ: Streptozotocin; Sal: Saline; DPN: Diabetic Neuropathy

Introduction

Diabetic peripheral neuropathy (DPN) caused by diabetes mellitus is one of the most common complications of diabetes affecting about 50% of patients with the disease [1,2]. Among the many symptoms of this neuropathy, the development of chronic pain is one of the major complications on ceits emergence depends on multifactorial components, which are still poorly understood [3]. This neuropathic pain typically involves the extremities and is characterized by spontaneous and evoked pain stimuli with changes in pain perception, increased sensitivity to noxious stimuli (hyperalgesia) and sensitivity to light stimuli or stimuli that previously were not painful (allodynia). These factors strongly affect the quality of life of patients with this syndrome [4-7].

A precise cellular mechanism for the hyperalgesia and allodynia in the DPN is not yet known thus, the existing treatments, including anticonvulsant drugs or tricyclic antidepressant drugs [8-11], are still ineffective and unsatisfactory and only a few patients with DPN benefit from some pain relief [1,4,12,13]. An alternative therapy that has gained clinical acceptance is the use of compounds that modulate cannabinoid receptors, once these receptors are expressed in both neurons and microglia of rats with diabetic neuropathy, at both spinal and supraspinal levels [11,14-16].

The use of derivatives from Cannabis sativa (Δ9-THC) for the treatment of various neurological disorders, including chronic pain, is supported by experimental and clinical data [17-20]. Although they are seen as promising target for the development of medications, clinical and preclinical studies have shown that Δ9-THC and other CB1 ligands generally produce undesirable effect in the Central Nervous System. CB1 agonists are generally at risk for psychoactive effects and dependence, limiting the optimization of doses in clinical trials and preclinical studies [21-23]. Thus, the development of drugs capable of binding to the cannabinoid receptors without psychoactive effects provides therapeutic potential without the risk of adverse effects [22].

Hemopressin (Hp), a nonapeptide (PVNFKFLSH) derived from the hemoglobin α1 chain was previously shown to target CB1 receptor, and to modulate its signaling [24-26]. Hp exhibits antinociceptive effect in inflammatory pain models [26,27]. Hp inhibits carrageen-an-induced hyperalgesia only at the injured paw; without presenting antinociceptive effect in the contralateral, uninflamed paw, indicating that the effect of Hp is limited to tissue injury induced pain [26]. Also, intrathecal administration of Hp induces significant antinociception in the first and second phases of the formalin test [27]. The effects of Hp on carrageenan-induced hyperalgesia are independent of route of administration (oral, local, or intrathecal) [26]. More interesting is the fact that neurological side effects that are typically associated with antinociceptive doses of CB1 receptor ligands, including hypothermia, catalepsy and hypoactivity, were not reported with antinociceptive doses of Hp [26]. This, taken with the fact that the effects of Hp on carrageenan-induced hyperalgesia were found to be independent of route of administration, raises the possibility that Hp could be developed as a novel class of drug that modulates CB1 receptor for the treatment of pain. This study aimed to examine the effect of Hp...
treatment on mechanical and thermal sensitivity in a mouse model of diabetic neuropathy induced by streptozotocin (STZ).

Materials and Methods

Animals

Male C57BL6 mice weighing 20-25 g, age-matched, were used throughout this study. Animals were maintained under controlled light cycle (12/12 h) and temperature (21 ± 2°C) with free access to food and water. Throughout the experiments, animals were managed using the principles and guidelines for the care of laboratory animals in studies involving pain and were approved by the Ethics Committee on the Use of Animals at University of São Paulo (CEUA, protocol number 157/2011). Animals were divided in four groups: control saline group (Sal+Sal), streptozotocin group (STZ+Sal), Hp group (Sal+Hp) and STZ/Hp group (STZ+Hp)

Diabetic Neuropathy (DPN)

Mice received a single intraperitoneal injection of streptozotocin (STZ; 200 mg/Kg, body weight; Sigma, St. Louis, MO) dissolved in sterile 0.9% saline after 4 hours of food restriction. Control animals received only sterile 0.9% saline injection. The fasting blood glucose levels were measured, after a fasting period of 4 h, by glucometer (ACCU-CHEK, Roche Diagnostics) before STZ or Sal injection and again on days 7, 14, 21 and 28 after STZ. Mice whose blood glucose levels exceeded 300 mg/dl were considered diabetic [28].

Behavioral analysis

Mechanical allodynia - von Frey test: Testing for mechanical allodynia (Von Frey filaments - Touch-Test’ Sensory Evaluators - North Coast Medical) was performed according to the method of Chaplan [29]. Mice were placed individually in plastic cages with a wire bottom, which allowed access to their paws. To reduce stress, mice were habituated to the experimental environment one day before the first measurement. At the day of the test, the animals were placed in the cages 30 min before the beginning of each measurement. The area tested was the mid-plantar left paw.

Thermal sensitivity - Planar test: Assessment of sensitivity to the thermal stimulus was based on the method of Hargreaves et al. [30] with instrumentation provided by IITC Life Science. Animals were placed for 30 minutes in compartments of 15 cm² surface with a transparent glass for the habituation process. To reduce stress, mice were habituated to the experimental environment one day before the first measurement. The glass allows the passage of the light beam, which can lead to painful thermal stimuli, adjusted to 30% of the maximum intensity of heating. Test was conducted after a pre-habitation thermal sensitivity baseline. The beam was directed to the plantar surface of the left hind paw (the thermal stimulus was applied for a maximum period of 20 sec to prevent tissue damage in the animal’s paw). Latency in seconds to withdraw the paw was considered as nociceptive response. Three measurements were made for each hindpaw, with an interval of 5 minutes between them. The mean of the measurements was used as the thermal nociceptive response.

Statistics

Results are presented as the mean ± standard error of the mean (SEM). Statistical analyses of data were generated using GraphPad Prism, version 4.02 (GraphPad Software Inc., San Diego, CA, USA). Statistical comparison of more than two groups was performed using analysis of variance (ANOVA), followed by Bonferroni's test. Statistical comparison for treatment over time was performed using two way ANOVA followed by Bonferroni’s test. In all cases, p ≤ 0.05 was considered statistically significant.

Results

Hp inhibits mechanical sensitivity on diabetic mice

STZ induced mechanical alldynia in mice from the 7th day after STZ injection up to 28 days (n=6) (Figure 1). Once confirmed the presence of neuropathic pain, at the 7th day groups of animals (n=6) received oral treatment with Hp for 28 days and presented a full reversion of mechanical sensitivity in all evaluated days, which was not observed in the group of mice treated with saline (n=6) (Figure 1). Furthermore, a control saline group that was also treated with Hp showed no changes in pain sensitivity when compared to the initial measurement or with the Saline+Saline group (n=6) (Figure 2).

Treatement with Hp does not interfere with thermal sensitivity of diabetic mice

Mice treated with STZ (n=5) showed a loss on pain sensitivity (hypoalgesia) to the thermal stimulation in all evaluated times when compared with Sal+Sal group (n=5) (Figure 2). Also, treatment with Hp was not able to reverse STZ-induced thermal hypoalgesia (n=5) (Figure 2).

Thermal sensitivity of mice was measured before the injection of STZ or Saline (time 0), 7 days after STZ or Sal and 28 days after treatment with Hp or Sal.
by itself didn’t cause any changes in control animals (n=11) (Figure 3).

Furthermore, Hp treatment did not interfere with the increase on blood glucose levels of STZ mice (n=11). Hp treatment did not interfere with the increase on blood glucose levels of animals when compared to non-diabetic (saline) mice, in all evaluated times. This result is consistent with previous reports from literature demonstrating that diabetic mice show a low pain threshold when compared to non-diabetic (saline) mice.

Glucose levels were measured prior to any treatment (time 0), 7 days after STZ or Sal injection and after 7, 14, 21 and 28 days after HP or Sal treatment. STZ-injected mice (n=11) presented a decrease on body weight that was not modified by Hp-treatment (n=11) (Figure 4). Also, Hp did not compromise the weight gain in the control group (n=11) (Figure 4).

**Discussion**

In this study, it was evaluated the antinociceptive effect of Hp, a CB receptor ligand, Hp in a mice model of diabetic-induced neuropathy [28]. Results presented herein demonstrate that STZ-induced diabetic mice showed a decrease on mechanical pain threshold when compared to non-diabetic (saline) mice, in all evaluated times. This result is consistent with previous reports from literature demonstrating that diabetic mice show a low pain threshold, thus characterizing the diabetic neuropathy as a model of neuropathic pain [21]. We found that 28 days of oral treatment with Hp could completely block signs of mechanical pain from the 7th day after Hp administration, supporting the idea that Hp induces true antinociception in this neuropathic pain model. These data corroborate with data obtained by our group 

During the experiments, animals were weighed before any treatment (time 0), after 7 days of streptozotocin (STZ) or saline (Sal) injection, and after treatment with hemopressin (Hp) or Sal. Hp was administered once a day, orally at a dose of 2.5 mg/kg for 28 days. Data are mean ± S.E.M. 11 animals per group. ** p <0.01 and *** p <0.001 compared to group Sal+Sal and Sal+Hp. (Two-way ANOVA followed by Bonferroni post-test).

**Effect of Hp treatment on blood glucose levels of diabetic mice**

Serum glucose levels were checked before any treatment (time 0), 7 days after STZ or Sal injection to verify the presence of hyperglycemia or normal glucose levels, and after 7, 14, 21 and 28 days of Hp treatments.

STZ (n=11) was effective in inducing diabetes on mice observed by the increase on blood glucose levels of animals when compared with Sal-treated mice (n=11). Hp treatment did not interfere with the increase on blood glucose levels of STZ mice (n=11). Furthermore, Hp by itself didn’t cause any changes in control animals (n=11) (Figure 3).
Conflicting data have been reported concerning the thermal nociceptive threshold in STZ-induced diabetes model. While some studies reported thermal hyperalgesia, others observed thermal hypalgesia or normal thermal thresholds after STZ injections. This may be in part due to methodological details such as use of different species, limbs, and methods of heat application. In this sense it is demonstrated that insulin-deficient diabetic rats with the plantar surface of the hind paw exposed to a temperature ramp rising from 30°C at a rate of 1°C/s over 20 s, exhibit a transient thermal hyperalgesia during the first few weeks of diabetes. It progresses to thermal hypalgesia within 2–3 months [34]. This is also common in mice submitted to the STZ model. Some mice injected with streptozotocin (50–200 mg/kg) does not become hyperglycemic but exhibit a transient thermal hyperalgesia when evaluated in the hot plate test, that returned to normal within weeks. This contrasts with hyperglycemic mice that progressed from hyperalgesia to hypalgesia [13,28]. In our present study, mice displayed thermal hypalgesia 1 week after injection of streptozotocin and subsequent induction of diabetes. This phenomena was maintained up to 5 weeks of observation. This result is similar to data from human diabetic patients where there is a progression from painful to degenerative painless neuropathy, although at present it is not clear that thermal hypalgesia in diabetic rats or mice coincides with any loss of epidermal thermal nociceptors, as appears to be the case in diabetic patients [35]. In this work we demonstrate that Hp treatment was not able to interfere with thermal hypalgesia. However, Hp treatment didn’t modify thermal sensitivity on control mice injected with saline, thus reforcing that Hp is specific in treating mechanical hyperalgesia observed on diabetes.

Another interesting finding was the fact that although treatment with Hp has been effective in reversing the painful picture, no changes were observed regarding the blood glucose levels or body weight of animals, which continued to progress with the development of diabetes. Moreover, Hp treatment by itself did not cause any change in the control group of animals, demonstrating that Hp is specific on treating painful neuropathy.

Recent data demonstrated that Hp treatment interfere with the transmission of neuropathic pain message to the central nervous system, reducing nociceptor activation in spinal cord [31]. Also, Hp inhibits calcium mobilization in dorsal root ganglia neurons from both normal and neuropathic rats, reinforcing the idea that Hp modulates primary afferent nociceptive signal by inhibiting sensory neurons [31]. Recent data showed that hemopressin activates distinct neuronal substrates within the brain, focused mainly on the feeding-related circuits of the mediobasal hypothalamus and in nociceptive regions of the periaqueductal grey (PAG) and dorsal raphe (DR). In contrast to AM251, there is a distinct lack of activation of the brain reward centres, such as the ventral tegmental area, nucleus accumbens and orbitofrontal cortex, which normally form a functional activity signature for the central action of synthetic CB1 receptor inverse agonists. Thus, hemopressin modulates the function of key feeding-related brain nuclei of the mediobasal hypothalamus, and descending pain pathways of the PAG and DR, and not higher limbic structures. Thus, hemopressin may offer behaviourally selective effects on nociception and appetite, without engaging reward pathways [36]. These data complement the existing data in the literature showing that neuronal responses that are typically associated with antinociceptive doses of CB1R ligands, including hypothermia, catalepsy and hypoactivity, were not reported with antinociceptive doses of Hp [26]. Another interesting point is that the effects observed for Hp depend on its aminoacid sequence, once it was demonstrated that central administration of an extended form of Hp, VD-PVNFKFLSH (VD-Hp) induces dose-dependent antinociception in mice but also induces undesired effects on the Central Nervous System such as tolerance to antinociception and conditioned place aversion [37]. Taken together, these data raises the possibility that Hp could be developed as a novel class of drug that modulates CB1R for the treatment of pain.

In conclusion, our data demonstrates that Hp exhibits antinociceptive properties on diabetes-induced peripheral neuropathy. This effect is specific for the treatment of chronic neuropathic pain. Although the mechanisms involved in the effects of hemopressin need to be further characterized, the results obtained so far suggest a role for a cannabinoid-like compound in regulating chronic neuropathic pain that could be further explored to develop therapeutic drugs based on the hemopressin sequence.

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