Analgesic Effects of Vilazodone, Indatraline, and Talsupram in a Rat Model of Neuropathic Pain

Levent HACIŞÜLEYMAN*, Bülent SARAÇ, Ziad JOHA
Sivas Cumhuriyet University, Faculty of Medicine, Department of Pharmacology, Sivas, Türkiye

ABSTRACT

Objectives: Drugs that inhibit the reuptake of serotonin, norepinephrine, and/or dopamine are widely used for treating depressive disorders and have emerged as effective drugs for neuropathic pain. They have no substantial anti-nociceptive effects but are considered, with gabapentin/pregabalin, first-line drugs for neuropathic pain.

Materials and Methods: In this study, three different antidepressant agents were used in different doses to investigate their anti-hyperalgesic effects in rat models of neuropathic pain using hot plate and tail flick methods. They have different mechanisms of action; vilazodone hydrochloride is a selective serotonin inhibitor and a 5-HT1A partial agonist; talsupram hydrochloride is a selective noradrenaline inhibitor, and it has a high affinity for noradrenaline transporter (NET), whereas indatraline hydrochloride is a triple reuptake inhibitor that inhibits transporters for 5-HT (SERT), dopamine (DAT), and NET.

Results: All the drugs used in the experiment were found to have an anti-hyperalgesic effect in both tests compared to the sham group. When anti-hyperalgesic effects of the three agents were compared to each other, it was found that talsupram hydrochloride was significantly more effective than the two other drugs in hot plate test. However, there was no statistically significant difference in the tail flick test. Indatraline hydrochloride was more effective than vilazodone hydrochloride at the same doses in the tail flick test.

Conclusion: Our data suggest that three drugs are effective analgesics in rat models of neuropathic pain and inhibition of noradrenaline reuptake represents the cornerstone of analgesic mechanisms of effective antidepressants.

Key words: Neuropathic pain, antidepressant, vilazodone, talsupram, indatraline, hot plate, tail flick, anti-hyperalgesic, sciatic nerve ligation

INTRODUCTION

The International Association for the Study of Pain defined neuropathic pain as "pain caused by a lesion or disease of the somatosensory nervous system".¹ According to a report published in 2011, one-third of Americans experience chronic pain. This exceeds total amount of cardiovascular diseases, diabetes, and cancer cases.² The prevalence rate of chronic pain in Europe is about 25-30%.³ Almost one-fifth of people affected by chronic pain have neuropathic pain.⁴ This high prevalence rates of chronic pain, especially neuropathic pain, are due to the lack of effective drugs. While nociceptive pain can be managed with analgesic drugs such as opioids and non-steroidal anti-inflammatory drugs, the medications used to manage neuropathic pain have a mild effect and in a small percentage of patients. This is mainly because they are unable to target the exact underlying mechanisms; this is why syndromes like fibromyalgia, whose pathophysiological mechanisms are not clear, have lower treatment success rates.⁵ The existing medications for neuropathic pain are non-specific and often inadequately effective.⁶ Other medications such as opioids, on the other hand, have serious side effects. Therefore, there is a persistent need for improved and more specific therapeutic strategies. Before clinicians can prescribe precise medications for neuropathic pain patients, main targets in the pathway must be understood.

The pharmacological treatment of neuropathic pain is complicated and there is no effective treatment for many patients. While a general consensus indicates as to which drugs should be used as first-line medications, controversy over second- and third-line drugs continues, particularly regarding weak and strong opioids. Although opioids are effective in the management of neuropathic pain, they are
Antidepressants have been proven to have analgesic effects in chronic pain even though they were not initially designed to be used as analgesic drugs. Antidepressants have practically no antinociceptive effects, but are considered with pregabalin and gabapentin first choice drugs for neuropathic pain\(^7,8\) and fibromyalgia.\(^9\)

There is no full understanding of how antidepressants are effective in pain management. An early concept of analgesic mechanisms of antidepressants for neuropathic pain was that these drugs could potentiate effectiveness of the descending noradrenergic and serotoninergic inhibitory pathways that extend from the brain stem to the dorsal horn of the spinal cord. This is done by inhibiting the reuptake of serotonin and noradrenaline released into the spinal synapses between the first-order neurons (nociceptors) and the second-order neurons (spinothalamic neurons). The synaptic transmission between these neurons can be inhibited by the neurotransmitters released from the inhibitory descending fibers, such as noradrenaline, which binds \(\alpha\)-2 adrenergic receptors. They can also induce spinal interneurons to release inhibitory materials like GABA and endogenous opioids, such as serotonin at its metabotropic receptors or noradrenaline at \(\alpha\)-1 adrenergic receptors.\(^1\)

Our study aimed to explore the anti-hyperalgesic effect of three different antidepressant drugs at different doses in rat models of neuropathic pain using hot plate and tail flick methods, and to compare analgesic activity of these drugs. Anti-hyperalgesic effects of these agents have not been studied extensively before.

**MATERIALS AND METHODS**

**Animals**

Experiments were performed on adult male Wistar albino rats weighing 200–225 g. The animals were kept at 22 ± 1°C, four in each cage, and maintained with a light-dark cycle of 12:12 h and free access to water and food. Sivas Cumhuriyet University Animal Ethics Committee approved all experiment protocols (approval no: 65202830-050.04.04-284).

**Drugs**

5-[4-[4-(5-Cyano-1H-indol-3-yl) butyl]-1-piperazinyl]-2-benzofurancarboxamide hydrochloride (vilazodone hydrochloride) (Tocris Bioscience), 1,3-dihydro-N,3,3-trimethyl-1-phenylbenzo[c]thiophene-1-propanamine hydrochloride (talsupram hydrochloride) (Tocris Bioscience) and \((1R,3S)-rel\)-3-(3,4-dichlorophenyl)-2,3-dihydro-N-methyl-1H-inden-1-amine hydrochloride (indatraline hydrochloride) were diluted in dimethyl sulfoxide. Solutions were freshly prepared on the days of experimentation. Intraperitoneal (I.P.) vilazodone hydrochloride (5-HT1A partial agonist and SSRI 2.5, 5, and 10 mg/kg), talsupram hydrochloride (selective inhibitor of noradrenaline transporters 2.5, 5, and 10 mg/kg), and indatraline hydrochloride (5-HT noradrenalin and dopamine reuptake inhibitor 2.5, 5, and 10 mg/kg) were applied before the analgesia tests.

**The experimental protocol and the analgesia tests**

All experiments were carried out blindly between 10.00 and 16.00 h in normal light and temperature (22 ± 1°C) in a quiet room. The rats were allowed to adapt to the laboratory for at least 2 h before the test and their tails were marked to differentiate the treatment groups. The rats were randomized into 10 groups (3 groups for each drug (1 group for each dose) and 1 group as a sham). Each experimental group had six rats. The same person performed all neuropathic operations and analgesia tests to minimize experimental variability.

**Surgical intervention**

The neuropathic pain model was produced by partial sciatic nerve ligation. Surgical interventions were performed at Sivas Cumhuriyet University Medical Faculty Experimental Animals Laboratory. Anesthesia was performed using intramuscular ketamine (90 mg/kg) and xylazine (3 mg/kg). Under aseptic conditions, a 1 cm incision was applied to biceps femoris and the sciatic nerve was reached in the middle thigh level of the right leg. Then, the sciatic nerve was freed of adherent tissues with careful blunt dissection and the dorsal one-third to half of the nerve was tightly ligated with 4.0 chromic catgut. The incision was closed with 4.0 silk. In sham group of rats, the same intervention was applied but without nerve ligation. After surgery, the rats were returned to their cages and kept for 21 days under the abovementioned same conditions.\(^14,15\)

**Analgesia tests**

To evaluate thermal pain standard tail flick test (May TF 0703 Tail beat unit, Commat) and hot plate test (May AHP 0603 Analgesic HP, Commat) devices were used. In the tail flick test, an intensive light beam was aimed at the animal’s tail and a timer begins. When the animal flocks its tail, the timer is stopped and recorded time (latency) represents the pain threshold. Tail-flick latencies were measured before the administration of the vehicle or investigational drugs to obtain a baseline and 15, 30, 60, 90 and 120 min after the I.P. administration. The maximum response time was set to 15 seconds (cut-off latency) to avoid tissue damage. Rats that did not show a response within 15 seconds were excluded. The hyperalgesic responses in this test reflect the mechanisms of pain in the central nervous system.\(^16,18\)

A hot plate device was used to evaluate thermal pain. In this test, the rats were placed on a hot plate with the temperature set at 53 ± 0.5°C for a maximum time of 30 s to prevent injury. Response time was recorded (when the animals licked their fore and hind paws or jumped) before and 15, 30, 60, 90, and 120 min after I.P. administration of the vehicle or test drugs. The hyperalgesic reactions in this test reflect the mechanisms of pain in both the central and peripheral nervous systems.\(^16,18\)

**Statistical analysis**

In all groups for each rat, antinociceptive effects of the drugs were measured as tail flick and hot plate latencies and
transformed into a percentage maximum possible effect (% MPE). MPE was obtained using the formula: \[\text{MPE} = \frac{(\text{post-drug latency} - \text{pre-drug latency})}{(\text{cut-off latency} - \text{pre-drug latency})} \times 100\]. Pre-drug and post-drug \(N\) values were the same in each group. The data were analyzed using one and two-way analysis of variance (ANOVA) and repeated measures ANOVA followed by a Tukey post-hoc test (SPSS 14.0 for Windows) for multiple comparisons between groups. All data are presented as a mean ± standard error of the mean. The significance level was determined as \(p<0.05\).

RESULTS

Determination of neuropathic pain formation by sciatic nerve ligation

The occurrence of neuropathic pain in the rats was detected using the paired student’s \(t\)-test. The post-surgery basal latencies of the rats were considerably lower than the pre-surgery basal latencies in both hot plate and tail flick tests \((p<0.05)\) (Figures 1 and 2).

Effects of vilazodone hydrochloride on neuropathic pain

Vilazodone hydrochloride was applied at three doses: 2.5, 5, and 10 mg/kg. In both tail flick and hot plate tests, the responses were measured before the drug was administered i.P. and after the administration at 15, 30, 60, 90, and 120 min. Maximum % MPE was observed in 90 mins after administration of these three doses. One-way ANOVA test was applied to compare the different doses with the sham group and with each other. In both tail flick and hot plate tests, doses at 5 and 10 mg/kg were found to be effective against neuropathic pain compared with the sham group. Dose at 10 mg/kg was effective from 30 to 120 mins in both tests with a statistically significant difference compared with 2.5 mg/kg dose at 90 min in the hot plate test and in 90, 120 min in the tail flick test. Dose at 5 mg/kg was effective from 30 to 90 min in both tests. Dose at 2.5 mg/kg was not statistically different from the sham group in the hot plate test at all minute points, while it was noticed to be effective only in the tail flick test at 60 and 90 min (Figure 3).

Effects of talsupram hydrochloride on neuropathic pain

Talsupram hydrochloride was applied intraperitoneally at three doses: 2.5, 5, and 10 mg/kg. The maximum percentage MPE was observed 60 min after the drug was administered for all three doses. One-way ANOVA test was used. The first dose of 2.5 mg/kg was effective from 30 to 120 mins in the hot plate test and from 30 to 90 min in the tail flick test. The other two doses at 5 and 10 mg/kg were effective at all

![Figure 1. Tail flick basal latencies of rats before and after surgery (*\(p<0.05\) paired student’s \(t\)-test)](image1)

![Figure 2. Hot plate basal latencies of rats before and after surgery (*\(p<0.05\) paired student’s \(t\)-test)](image2)

![Figure 3. The effect of vilazodone hydrochloride intraperitoneal administration on the neuropathic pain model in the hot plate test (A) and tail flick test (B). This was expressed as a percentage of the maximal possible effect (MPE). Each point represents the mean ± SEM of % MPE for 6 rats *\(p<0.05\) when the groups were compared to the sham group, \(\Psi\): \(p<0.05\) when the groups were compared to the sham and 2.5 mg/kg dose groups, SEM: Standard error of the mean)](image3)
minute points in the hot plate test, while their effectiveness in the tail flick test was noticed from 15 to 90 min compared to dose of 2.5 mg/kg, a statistically significant difference was noticed only with dose at 10 mg/kg at 60 min in the hot plate test (Figure 4).

**Effects of indatraline hydrochloride on neuropathic pain**

Indatraline hydrochloride was administered intraperitoneally at 3 doses; 2.5, 5, and 10 mg/kg. Maximum MPE% was observed at 60 min after the drug was administered for all three doses. One-way ANOVA test was applied. The drug was effective at all three doses in both tests. After the first dose 2.5 mg/kg was administered, the anti-hyperalgesic effect was statistically significant in 60 min in the hot plate test and from 30 to 90 min in the tail flick test. The second dose at 5 mg/kg was effective at 30 and 60 mins in the hot plate test and from 15 to 90 min in the tail flick test. The anti-hyperalgesic effect for the third dose of 10 mg/kg from 30 to 90 min was statistically significant compared to the sham group, and at 60 min the first 2.5 mg/kg dose in the hot plate test. While in the tail flick test, anti-hyperalgesic effect of 10 mg/kg dose was clear from 15 to 90 min compared to the sham group (Figure 5).

**Comparison of the anti-hyperalgesic effects of vilazodone hydrochloride, talsupram hydrochloride, and indatraline hydrochloride on neuropathic pain**

We used two-way variance analysis followed by Tukey HSD test in this comparison. In hot plate test, MPE% values obtained from doses of 2.5, 5, 10 mg/kg of talsupram hydrochloride dose were significantly higher than values obtained from the same doses of vilazodone hydrochloride and indatraline hydrochloride. Even at lower doses, talsupram was more effective than the other two drugs. MPE% of talsupram at 2.5 mg/kg was > vilazodone 5 mg/kg and indatraline 10 mg/kg, while at 5 mg/kg, MPE% was > vilazodone 10 mg/kg (p<0.05). Whereas in the tail flick test, MPE% values obtained from the different doses of talsupram hydrochloride were not statistically different from those of the same doses of vilazodone hydrochloride and indatraline hydrochloride (p>0.05), except for MPE% of 2.5 mg/kg dose of indatraline hydrochloride, which was higher than that of the same dose of talsupram hydrochloride (p<0.05). There was a statistically significant difference between the MPE% values obtained from the same doses of vilazodone hydrochloride and indatraline hydrochloride in favor of the latter in the tail flick test (p<0.05), whereas no statistically significant difference

![Figure 4](image1.png)

*Figure 4. The effect of talsupram hydrochloride intraperitoneal administration on the neuropathic pain model in the hot plate test (A) and tail flick test (B). This was expressed as a percentage of the maximal possible effect (MPE). Each point represents the mean ± SEM of % MPE for 6 rats. *p<0.05 when the groups were compared to the sham group, Ψ: p<0.05 when the groups were compared to the sham and 2.5 mg/kg dose groups, SEM: Standard error of the mean.

![Figure 5](image2.png)

*Figure 5. The effect of indatraline hydrochloride intraperitoneal administration on the neuropathic pain model in the hot plate test (A) and tail flick test (B). This was expressed as a percentage of the maximal possible effect (MPE). Each point represents the mean ± SEM of % MPE for 6 rats. *p<0.05 when the groups were compared to the sham group, Ψ: p<0.05 when the groups were compared to the sham and 2.5 mg/kg dose groups, SEM: Standard error of the mean.*
was found between them in the hot plate test (p>0.05) (Figure 6).

DISCUSSION

Neuropathic pain, a pain syndrome caused by a lesion or disease of the somatosensory system, is a main public health issue and becoming a global burden. An epidemiological study indicated that the prevalence rate of neuropathic pain is in the range of 6.9% to 10% and increases year after year. Patients with neuropathic pain report significantly lower levels of health-related quality of life. The high rate of comorbidity between pain and depression has led to the wide use of antidepressants in chronic pain treatment. Tricyclic antidepressants (TCA), particularly desipramine, amitriptyline, nortriptyline, and imipramine, are the most effective antidepressants in neuropathic pain management. TCAs have effects on various targets. This lack of selectivity is related to their efficacy. For instance, amitriptyline has a local anesthetic effect by blocking voltage-gated sodium channels. TCAs are effective in many neuropathic conditions. However, these multiple actions of TCAs also contribute to many adverse effects that limit their use, in particular their anticholinergic effects that increase the risk of cardiotoxicity, orthostatic hypotension, mouth dryness, constipation, and urinary retention. To avoid these issues, serotonin-norepinephrine reuptake inhibitors (SNRI), particularly duloxetine, have been suggested in the management of neuropathic pain. SNRIs, e.g. duloxetine, have shown consistent efficacy in several neuropathic syndromes, including painful polyneuropathy, post-herpetic neuralgia, low back pain, and painful diabetic neuropathy. Opioids are recommended to be used as second- and third-line treatments because of their adverse effects. Tramadol and FDA-approved tapentadol are used in second-line treatment, while the strong opioids, oxycodone, and morphine are used in the third-line treatment. Therefore, there is still a need for more effective drugs with less serious adverse effects for neuropathic pain. In this study, we investigated anti-hyperalgesic effects of three different antidepressant drugs at different doses in rat models of neuropathic pain using the hot plate method. These drugs have different mechanisms of action; vilazodone hydrochloride is a selective serotonin inhibitor, while talsupram hydrochloride is a selective noradrenaline inhibitor, and it has a high affinity for the noradrenaline transporter, whereas indatraline hydrochloride inhibits transporters for 5-HT, dopamine, and noradrenaline. All the drugs used in the experiment were found to have an anti-hyperalgesic effect compared to the sham group. These results support the evidence for role of noradrenaline, serotonin, and probably dopamine in the analgesic effects of antidepressants on neuropathic pain and corroborate a previous study highlighted that indatraline has analgesic profile in neuropathic mice. Some preclinical studies on animals have indicated the important roles of noradrenaline and serotonin in the processing of pain. Experimental studies have demonstrated that intrathecal administration of serotonin and norepinephrine receptor agonists inhibits pain behavior. Other data indicate that serotonin agonists such as fenfluramine trigger the neuronal release of substance P and thus pain behavior. Furthermore, intrathecal administration of serotonin receptor antagonists such as ondansetron inhibits the experimental pain response in rats. 5-HT1A, 5-HT2A/2C, 5-HT3, and 5-HT7 receptors that highly contribute to the transmission of nociceptive messages are expressed in the dorsal horn of spinal cord. It seems that serotonin both inhibits and enhances pain sensation by various physiological mechanisms, contrary to norepinephrine, which is essentially inhibitor. A review of studies on SSRIs showed inconsistent efficacy for migraine, diabetic neuropathy, and fibromyalgia; however, some studies of SSRI treatment for mixed-chronic pain are positive. When anti-hyperalgesic effects of three agents were compared to each other, it was found that talsupram hydrochloride was significantly more effective than vilazodone hydrochloride and indatraline hydrochloride in the hot plate test. This could be related to the high affinity of talsupram hydrochloride for norepinephrine transporters and the more important role of noradrenaline in the anti-hyperalgesic activity compared to serotonin and dopamine. However, there was no statistically significant difference in the tail flick test. While the response in the tail flick test is a spinal reflex rather than an indication of pain behavior involving higher brain

Figure 6. The effect of vilazodone hydrochloride, talsupram hydrochloride and indatraline hydrochloride on the neuropathic pain model (in hot plate and tail flick tests) was expressed as a percentage of the maximal possible effect (MPE). All drugs were administered intraperitoneally. Each point represents the mean of % MPE for 6 rats.
centers, the response in hot plate test is considered to integrate supraspinal pathways. Therefore, comparative results suggest that analgesic effect of talsupram hydrochloride is more effective than that of vilazodone hydrochloride and indatraline hydrochloride at the supraspinal level. MPE% values obtained from indatraline hydrochloride were more than values obtained from vilazodone hydrochloride at the same doses in the tail flick test. This could be due to a greater anti-hyperalgesic effect of the inhibition of the reuptake of noradrenaline, serotonin, and dopamine compared to the inhibition of reuptake of serotonin alone. Although vilazodone hydrochloride is less effective against neuropathic pain than the other drugs, its relatively benign sexual side effect profile may be worth considering because in addition to serotonin reuptake inhibition, it acts as 5-HT\textsubscript{1A} partial agonist.

**CONCLUSION**

In conclusion, our data suggest that the three drugs used in this study are effective analgesics in rat models of neuropathic pain. Inhibition of noradrenaline reuptake represents the cornerstone of analgesic mechanisms of effective antidepressants. Although SSRIs have a more tolerable side effect profile and the SSRI used in our experiment, i.e. vilazodone hydrochloride, was effective in a rat model of neuropathic pain, the evidence to support the use of SSRIs in the clinical management of chronic pain is still not convincing.

**ACKNOWLEDGMENTS**

This study was funded by Sivas Cumhuriyet University Scientific Research Project (T-866, Doctoral Thesis Project, CUBAP, Türkiye).

**Ethics**

**Ethics Committee Approval:** All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted. Animal experimental procedures were approved by the Animal Ethical Committee at Sivas Cumhuriyet University (Sivas, Türkiye) (approval no: 65202830-050.04.04-284).

**Informed Consent:** Not human subject research.

**Authorship Contributions**


**Conflict of Interest:** No conflict of interest was declared by the authors.

**Financial Disclosure:** The authors declared that this study received no financial support.

**REFERENCES**


