Evaluation of pharmacological mechanisms of antinociceptive effect of *Teucrium polium* on visceral pain in mice

Zendehdel, M.^{1*}; Taati, M.²; Jadidoleslami, M.³ and Bashiri, A.⁴

¹Department of Physiology, Faculty of Veterinary Medicine, University of Tehran, Tehran, Iran; ²Department of Pathobiology, School of Veterinary Medicine, Lorestan University, Khorramabad, Iran; ³Graduated from Faculty of Veterinary Medicine, University of Tehran, Tehran, Iran; ⁴DVM Student, Faculty of Veterinary Medicine, University of Tehran, Iran

***Correspondence:** M. Zendehdel, Department of Physiology, Faculty of Veterinary Medicine, University of Tehran, Tehran, Iran. E-mail: zendedel@ut.ac.ir

(Received 7 Nov 2010; revised version 23 May 2011; accepted 31 May 2011)

Summary

Teucrium polium is used for treatment of visceral pain in Iranian folkloric medicine. In this study antinociceptive mechanisms of *T. polium* hydroethanolic extract were examined by acetic acid-induced writhing test as a model of visceral pain in male NMRI mice. To reveal the antinociceptive mechanisms of the extract, we examined the effects of opioidergic, serotonergic, adrenergic and histaminergic antagonists on extract-induced antinociception. The results of this study showed that pretreatment with naloxone, chlorpheniramine and cimetidine significantly attenuate the antinociceptive effect of the extract. However, cyproheptadine and phentolamine had no effect. Our results clearly show antinociceptive effects of *T. polium* may be mediated by opioidergic and histaminegic H_1 and H_2 receptors.

Key words: Teucrium polium, Opioid receptor, Histamine receptors, Writhing test, Mouse

Introduction

Pain is a sensorial modality and primarily protective in nature, but often causes discomfort. It is the most important symptom that brings a patient to a physician. Analgesics relieve pain as a symptom without affecting its cause (Hasan et al., 2009). Currently available analgesic drugs such as opiates and nonsteroidal antiinflammatory drugs (NSAIDs) are not useful in all cases due to their side effects. The search for new analgesic compounds has been a priority of pharmacologists and pharmaceutical industries (Mattison et al., 1998). Medicinal plants are believed to be an important source of new chemical substances with potential therapeutic effects (Blumenthal, 2000). Thus, the study of plant species that traditionally have been used as pain killers should still be seen as a logical research strategy in the search for new analgesic drugs (Rang et al., 1998).

The T. polium is a wild-growing flowering plant belonging to the family labiatae and is found abundantly in south western Asia, Europe and North Africa (Abdollahi et al., 2003). This plant is used as a visceral pain killer in Iranian folkloric medicine. Previous studies have demonstrated some of the pharmacological effects of T. polium such as its antiinflammatory (Capasso et al., 1983), antinociceptive (Abdollahi et al., 2003), antidiabetes (Gharaibeh et al., 1988) and antihypertensive effects (Suleiman et al., 1988). In this study, to reveal the antinociceptive mechanisms of T. polium we examined the opioidergic, effects of serotonergic, noradrenergic and histaminergic receptor Т. *polium*-induced antagonists on antinociception.

Materials and Methods

Preparation of crude extract

Teucrium polium aerial parts were

collected during the flowering season from Zabol (Sistan and Baloochestan province, Southeastern Iran). Samples of the plant were identified by a botanist from the division of Pharmacognosy, Faculty of Pharmacy, Tehran University of Medical Sciences, Iran. The plant materials were cleaned, shade dried and coarsely ground. The powdered material was soaked in hydroethanolic (70%) solvent for three days with occasional shaking. First, it was filtered through a muslin cloth and then through a filter paper. This process was repeated twice more and the combined filtrated material was evaporated on a rotary evaporator under reduced pressure to a thick semi-solid mass of dark brown color (Subhan et al., 2007). The percentage yields based on the dried starting material was 15.6% for dried hydroalcoholic extract (W/W).

Animals

Male albino NMRI mice weighing 25-30 g from the Pasteur Institute of Tehran were used for all the experiments. Animals were housed in a temperature $(22 \pm 2^{\circ}C)$ and light-controlled room under a 12-h light/12h dark cycle (light on at 7:00 a.m.). Food and water were available ad libitum. The animals were allowed to adapt to the laboratory for at least 2 h before testing and were used only once. To reduce the experimental variation, all experiments were performed during the light phase of the cycle (10:00-17:00). All experimental procedures followed the Guidelines on Ethical Standards for Investigations of Experimental Pain in Animals (Zimmermann, 1983) and were carried out according to a protocol approved by the local Animal Ethics Committee.

Writhing test

Nociception was induced by an intraperitoneal (i.p.) injection of 0.6% acetic acid in a volume of 10 ml/kg in mice. The nociceptive behavior induced is characterized by abdominal contractions known as writhing, described as an exaggerated extension of the abdomen combined with the outstretching of the hind limbs (Koster et al., 1959). The total number of writhing was recorded in the following periods: 0-10, 10-20 and 20-30 min immediately after acetic acid administration.

Evaluation of antinociceptive activity of *T. polium* and pretreatment with antagonists

The *T. polium* hydroethanolic extract was dissolved in saline and administered intraperitoneally at doses of 50, 100 and 200 mg/kg. Indomethacin (5 mg/kg) was dissolved in saline and administered i.p. as the reference drug for comparison (Kozak *et al.*, 1998). Control group received normal saline. Antinociceptive activity was expressed as the percentage of inhibition of abdominal constrictions using the ratio:

 $\frac{(\text{control mean} - \text{treated mean}) \times 100}{\text{control mean}} = \% \quad \text{writhing} \quad \text{pain}$

Animals were pretreated i.p. with either saline, opioidergic receptor antagonist (naloxone, 2 mg/kg), serotonergic receptor antagonist (cyproheptadine, 4 mg/kg), aadrenergic receptor antagonist (phentolamine, 20 mg/kg), histamine H_1 -receptor antagonist (chlorpheniramine, 10 mg/kg) and histamine H₂-receptor antagonist (cimetidine, 10 mg/kg) 15 min before i.p. administration of vehicle or the most effective dose of *T. polium* (200 mg/kg). The writhing test response was tested 30 min after the treatment with either vehicle or extract. Additionally, onset of the first abdominal writhing was recorded as latency time. The time and dose of antagonists used were chosen on the basis of preliminary studies and previous publications (Schmitt et al., 1974; Leza et al., 1990; Zendehdel and Babapour, 2010). All drugs were dissolved in 5% dimethyl sulfoxide (DMSO). Control group received vehicle.

Acute toxicity

Mice were divided into control and test groups (n = 8). The first group served as normal control. *Teucrium polium* extract was administered i.p. to different groups at increasing doses of 400, 800, 1600, 3200 and 6400 mg/kg. After injections of extracts, mice were allowed food and water *ad libitum* and all animals were observed for possible mortality cases and behavioral

changes for 72 h (Lorke, 1983).

Statistical analysis

The data were presented as the mean \pm SEM. Statistical analysis was carried out by one-way analysis of variance (ANOVA) with Tukey's post-hoc test. P-values less than 0.05 were considered to indicate statistical significance.

Results

Evaluation of antinociceptive effects of *T. polium* in writhing test

The results of this study showed that hydroethanolic extract of T. polium at doses of 50, 100 and 200 mg/kg induced significant reduction in pain response in a dose dependent manner when compared to control group (P<0.001). Further, indomethacin significantly decreased the number of writhing as a reference drug (P<0.001). The percentage of the inhibition of writhing response induced by extract at doses of 50, 100 and 200 mg/kg were 43.33, 64.42 and 70.5%, respectively, despite the fact that writhing inhibited indomethacin the response by 80%. Furthermore, indomethacin significantly delayed the onset of first abdominal writhing (latency time) when compared to control group (P<0.001). But other groups had no effect on the onset of first abdominal writhing in comparison with control group (Table 1). The ED_{50} for antinociceptive effects of hydroethanolic extract was 62.7 mg/kg.

Effects of cyproheptadine and phentolamine on the antinociceptive action of *T. polium*

The results showed that the *T. polium* (200 mg/kg) induced significant reduction in pain response when compared to control group (P<0.001). Pretreatment with cyproheptadine and phentolamine had no effect on the antinociceptive properties induced by the extract. Moreover, none of the drugs had any effect on the onset of first abdominal writhing in comparison with control group (Tables 2 and 3).

Effects of naloxone, chlorpheniramine and cimetidine on the antinociceptive action of *T. polium*

Intraperitoneal injection of T. polium

Treatment	Dose (mg/kg, i.p.)	Latency time (sec)	Writhing count (Mean±SEM)	Inhibition (%)	P-value			
Control	10 (ml/kg, ip)	189 ± 25	74.50 ± 2.68					
T. polium	50	178 ± 23	43.17 ± 5.19	43.33	<0.001 vs. contol			
T. polium	100	206 ± 32	26.50 ± 1.22	64.42	<0.001 vs. contol			
T. polium	200	202 ± 30	21.00 ± 1.84	70.50	<0.001 vs. contol			
Indomethacin	5	$783\pm67^{*}$	14.50 ± 2.15	80	<0.001 vs. contol			
*D 0 001 up another barrier Carteria and r. Sfor each group								

*P<0.001 vs. control group, Saline: Control, and n = 8 for each group

Table 2: Effect of cyprohepta	line on T. polium-induced antinociception writhing with acetic acid	in
mice		

Treatment	Dose (mg/kg, i.p.)	Latency time (sec)	Writhing count (Mean±SEM)	Inhibition (%)	P-value
Control	10 (ml/kg)	179 ± 18	75.17 ± 3.17		_
Vehicle + T. polium	200	200 ± 27	23.13 ± 2.97	69.2	<0.001 vs. control
Cyproheptadine	4	188 ± 31	81.5 ± 2.66		_
Cyproheptadine + T. polium	4 + 200	191 ± 29	28.83 ± 2.23	61.64	<0.001 vs. control

Vehicle is 5% DMSO, Vehicle + Saline: Control, and n = 8 for each group

Treatment	Dose (mg/kg, i.p.)	Latency time (sec)	Writhing count (Mean±SEM)	Inhibition (%)	P-value
Control	10 (ml/kg)	181 ± 27	78.66 ± 1.81		
Vehicle + T. polium	200	190 ± 32	24.03 ± 1.63	69.08	<0.001 vs. control
Phentolamine	20	199 ± 36	85.33 ± 1.78		_
Phentolamine + T. polium	20 + 200	177 ± 19	32.16 ± 2.53	59.11	<0.001 vs. control

Vehicle is 5% DMSO, Vehicle + Saline: Control, and n = 8 for each group

Treatment	Dose	Latency time	Writhing count	Inhibition	P-value
	(mg/kg, i.p.)	(sec)	(Mean±SEM)	(%)	
Control	10 (ml/kg)	185 ± 24	76.17 ± 2.33	_	_
Vehicle + T. polium	200	210 ± 35	27.83 ± 1.22	70.02	<0.001 vs. control
Naloxone	2	193 ± 20	84.83 ± 2.54		_
Naloxone + T. polium	2 + 200	201 ± 26	45.33 ± 4.50	40.5	<0.001 vs. control
					<0.01 vs. T. polium

Table 4: Effect of naloxone on T. polium-induced antinociception writhing with acetic acid in mice

Vehicle is 5% DMSO, Vehicle + Saline: Control, and n = 8 for each group

 Table 5: Effect of chlorpheniramine on T. polium-induced antinociception writhing with acetic acid in mice

Treatment	Dose (mg/kg, i.p.)	Latency time (sec)	Writhing count (Mean±SEM)	Inhibition (%)	P-value
Control	10 (ml/kg)	186 ± 20	76.50 ± 1.87		
Vehicle + T. polium	200	201 ± 27	24.83 ± 1.07	67.5	<0.001 vs. control
Chlorpheniramine	10	197 ± 25	75.67 ± 1.94	—	_
Chlorpheniramine + T. polium	10 + 200	215 ± 36	38.33 ± 2.24	49.89	<0.001 vs. control
					<0.05 vs. <i>T. polium</i>

Vehicle is 5% DMSO, Vehicle + Saline: Control, and n = 8 for each group

	7	ption writhing with acetic acid in mice
Table 6. Effect of cimetidine on T	noluum_induced enfineeicei	ntian writhing with seatic seid in mics
I able v. Effect vi chilenume vii I.	<i>vouum</i> -muuteu anunotite	

Treatment	Dose (mg/kg, i.p.)	Latency time (sec)	Writhing count (Mean±SEM)	Inhibition (%)	P-value
Control	10 (ml/kg)	191 ± 23	78.50 ± 2.76	_	—
Vehicle + T. polium	200	183 ± 18	30.33 ± 1.23	66.4	<0.001 vs. control
Cimetidine	10	189 ± 27	71.33 ± 3.20		
Cimetidine + T. polium	10 + 200	211 ± 33	50.50 ± 3.06	35.66	<0.001 vs. control
					<0.001 vs. T. polium

Vehicle is 5% DMSO, Vehicle + Saline: Control, and n = 8 for each group

extract (200 mg/kg) induced significant reduction in pain response when compared to control group (P<0.001). Pretreatment naloxone, chlorpheniramine with and significantly attenuated cimetidine the antinociceptive effects of extract (from 70.02% to 40.5%, 67.5% to 49.89% and 66.4% to 35.66%, respectively) (P<0.01, and P<0.001. P<0.05 respectively). Furthermore, none of the drugs had any effect on the onset of first abdominal writhing in comparison with control group (Tables 4, 5 and 6).

Acute toxicity

Teucrium polium extract, at doses of 400-6400 mg/kg i.p. given to mice, had no effect on their behavioral responses and no mortality during the observation period of 72 h after administration. Therefore, it can be indicated that *T. polium* extract has no toxicity profile.

Discussion

In the present study, no mortality case

up to the dose of 6.4 g/kg of *T. polium* extract (i.p.) was observed. Therefore, we may suggest that the extract has no lethal toxicity in mice.

In this study, i.p. injection of T. polium showed а dose-dependent extract antinociceptive effect in writhing test. Previous studies reported that this effect of the plant is due to flavonoids, and the main flavonoids of this plant are quercetine and apigenine (Yazdanparast and Ardestani, 2009). Furthermore, to reveal the antinociceptive mechanisms of T. polium, the involvement of opioidergic, possible serotonergic, noradrenergic and histamine receptor antagonists on T. polium-induced antinociception were examined. Our findings showed that pretreatment with naloxone significantly attenuates the antinociceptive effects of the extract (from 70.02 to 40.5%), thus it reverses analgesic activity of T. polium to some extent. In this regard, Anjaneyulu and Chopra (2003) reported that quercetin probably acts through modulation of opioidergic mechanisms. Furthermore, Opioid agents exert their

analgesic effects via supraspinal (m_1, k_3, d_1, s_2) and spinal (m_2, k_1, d_2) receptors (Reisine and Pasternack, 1996). As the antinociceptive activity of the extract was inhibited by naloxone, the extract likely acts on spinal opioid receptors such as m_2 , k_1 and d_2 receptors, although other mechanisms of action such as inhibition of cyclooxygenases are also possible.

In the present study, pretreatment with chlorpheniramine, H₁ receptor antagonist, and cimetidine, H_2 receptor antagonist, attenuated the extract inducedantinociception (from 67.5% to 49.89% and 66.4% to 35.66%, respectively) but cyproheptadine and phentolamine had no effect. Thus, our results suggest that the part of T. polium-induced antinociceptive effect is mediated by H_1 and H_2 receptors, whereas serotonergic and adrenergic receptors had no effect. However, previous studies reported that serotonin and norepinephrine play an important role in the modulation of pain response (Taherianfard and Khazaee, 2006; Zendehdel and Babapour, 2010).

The involvement of histamine in the inflammatory pain of chemicals (e.g. formalin-induced) is well-documented. Peripheral histamine specifically activates and sensitizes itch-specific nociceptive C fibers (Schmelz et al., 1997), while it has emerged that central histamine plays an important role in antinociception (Robertson et al., 1988). Central injection of histamine shows an analgesic effect in several experiments including the tail-flick and hotplate tests (Thoburn et al., 1994). Previously, several lines of evidence have demonstrated that systemic or central injection of histamine or histamine agonist produces antinociception, which suggest an important role in the regulation of antinociception (Chung et al., 1984). Furthermore, it has been reported that blockade of the H_1 and H_2 receptors attenuate the antinociception induced by nefopam, decursinol and restraint (Girard et al., 2004). Both H_1 and H_2 receptor when applied intracerebroantagonists ventricularly or into the periaqueductal gray have been shown to block histamine-induced antinociception (Thoburn et al., 1994). Broad functional overlap, but also a striking molecular specificity anatomical and

characterizes these distinct sensations (Ikoma et al., 2006). Most convincing seems to be the evidence implicating histamine H_2 receptors in the periaqueductal gray in the histamine mediated antinoception (Thoburn et al., 1994), although H_1 receptors may be important in other areas such as the spinal cord (Suh et al., 1996). Taken together, our results suggest that T. polium may be effective on the inflammatory pain even at the central nervous system level. Since opioid, H₁ and H₂ receptors antagonist could not completely reverse T. polium analgesic activity, it is possible that other mechanisms influence its antinociceptive effects. According to these findings, there is evidence to suggest that the central effect and the peripheral one may result in different extract compounds, polar compound(s) which act centrally through the opioid and histaminergic system, and the other class of compounds which act peripherally (Bittar et al., 2000).

In conclusion, the present study proposes that *T. polium* possesses a strong antinociceptive property, acting on, at least, the central nervous system level and opioidergic, histamine H_1 and H_2 receptors appear to be involved in the production of *T. polium*-induced antinociception.

Acknowledgement

This research was supported by a grant from the Research Council of the Faculty of Veterinary Medicine, University of Tehran.

References

- Abdollahi, M; Karimpour, H and Monsef-Esfehani, HR (2003). Antinociceptive effects of *Teucrium polium* L. total extract and essential oil in mouse writhing test. Pharmacol. Res., 48: 31-35.
- Anjaneyulu, M and Chopra, K (2003). Quercetin, a bioflavonoid, attenuates thermal hyperalgesia in a mouse model of diabetic neuropathic pain. Prog. Neuro-Psychopharmacol. Biol. Psychiatry. 27: 1001-1005.
- Bittar, M; de Souza, MM; Yunes, R; Lento, RA; Delle-Monache, F and Cechinel-Filho, V (2000). Antinociceptive activity of I3, II8 Binaringenin, a biflavonoid present in plants of the Guttiferae. Planta. Med., 66: 84-86.

- Blumenthal, M (2000). *Herbal medicine*. 1st Edn., Austin, Texas, Integrative Medicine Communications, Austin. PP: 419-423.
- Capasso, F; Cerri, R; Morrica, P and Senatore, F (1983). Chemical composition and antiinflammatory activity of an alcoholic extract of *Teucrium polium*. Boll. Soc. Ital. Biol. Sper., 59: 1639-1643.
- Chung, YH; Miyake, H; Kamei, C and Tasaka, K (1984). Analgesic effect of histamine induced by intracerebral injection into mice. Agents Actions. 15: 137-142.
- Gharaibeh, NM; Elayan, HE and Salhab, AS (1988). Hypoglycemic effects of *Teucrium polium*. J. Ethnopharmacol., 24: 93-99.
- Girard, P; Pansart, Y; Coppe, MC; Verniers, D and Gillardin, JM (2004). Role of the histamine system in nefopam-induced antinociception in mice. Eur. J. Pharmacol., 503: 63-69.
- Hasan, SMR; Jamila, M; Majumder, MM; Akter, R; Hossain, MM; Mazumder, MEH; Alam, MA; Jahangir, R; Rana, MS and Rahman, S (2009). Analgesic and antioxidant activity of the hydromethanolic extract of *Mikania scandens* (L.) willd. Leaves. Am. J. Pharmacol. Toxicol., 4: 1-7.
- Ikoma, A; Steinhoff, M; Stander, S; Yosipovitch, G and Schmelz, M (2006). The neurobiology of itch. Nat. Rev. Neurosci., 7: 535-547.
- Koster, R; Anderson, M and De Beer, EJ (1959). Acetic acid analgesic screening. Fed. Proc., 18: 418-420.
- Kozak, W; Archuleta, I; Mayfield, KP; Kozak, A; Rudolph, K and Kluger, MJ (1998).
 Inhibitors of alternative pathways of arachidonate metabolism differentially affect fever in mice. Am. J, Physiol., 275: 1031-1040.
- Leza, JC; Lizasoain, I and Lorenzo, P (1990). H₁- and H₂-histamine receptor blockers and opiate analgesia in mice. Methods Find. Exp. Clin. Pharmacol., 12: 671-678.
- Lorke, DA (1983). A new approach to acute toxicity testing. Arch. Toxicol., 54: 275-287.
- Mattison, N; Trimple, AG and Lasagna, I (1998). New drug development in the United States, 1963 through 1984. Clin. Pharmacol. Ther., 43: 290-301.
- Rang, HP; Dale, MM and Ritter, JM (1998). *Pharmacology*. 4th Edn., New York, Churchill Livingston. PP: 614-616.
- Reisine, T and Pasternack, G (1996). Opioid analgesics and antagonists. In: Hardman, JG and Limbird, LE (Eds.), Goodman and Gilman's, the pharmacological basis of therapeutics. 9th Edn., New York, McGraw-

Hill. PP: 521-526.

- Robertson, JA; Hough, LB and Bodnar, RJ (1988). Potentiation of opioid and nonopioid forms of swim analgesia by cimetidine. Pharmacol. Biochem. Behav., 31: 107-112.
- Schmelz, M; Schmidt, R; Bickel, A; Handwerker, HO and Torebjork, HE (1997). Specific C receptors for itch in human skin. J. Neurosci., 17: 8003-8008.
- Schmitt, H; Le Douarec, JC and Petillot, N (1974). Antagonism of the antinociceptive action of xylazine, an α -sympathomimetic agent, by adrenoceptor and cholinoceptor blocking agents. Neuropharmacology. 13: 295-303.
- Subhan, F; Khan, M; Ibrar, M; Islam, NU; Khan, A and Gilani, AH (2007). Antagonism of antinociceptive effect of hydro-ethanolic extract of hypericum perforatum Linn, By a non selective opioid receptor antagonist, Naloxone. Pak. J. Biol. Sci., 10: 792-796.
- Suh, HW; Song, DK; Choi, YS and Kim, YH (1996). Effects of intrathecally injected histamine receptor antagonists on the antinociception induced by morphine, betaendorphin, and U50, 488H administered intrathecally in the mouse. Neuropeptides. 30; 485-490.
- Suleiman, MS; Abdul-Ghani, AS; Al-Khalil, S and Amir, R (1988). Effect of *Teucrium polium* boiled leaf extract on intestinal motility and blood pressure. J. Ethnopharmacol., 22: 111-116.
- Taherianfard, M and Khazaee, Z (2006). Effect of xylazine and yohimbine on the phasic pain during the estrous cycle in the rat. Iranian J. Vet. Res., 7: 33-39.
- Thoburn, KK; Hough, LB; Nalwalk, JW and Mischler, SA (1994). Histamine-induced modulation of nociceptive responses. Pain. 58: 29-37.
- Yazdanparast, R and Ardestani, A (2009). Suppressive effect of ethyl acetate extract of *Teucrium polium* on cellular oxidative damages and apoptosis induced by 2-deoxy-D-ribose: role of *de novo* synthesis of glutathione. Food Chemistry. 114: 1222-1230.
- Zendehdel, M and Babapour, V (2010). Study of antinociceptive effects of *Ziziphora tenuior* and its interference on opioidergic and serotoninergic systems. J. Vet. Res., 65: 57-60.
- Zimmermann, M (1983). Ethical guidelines for investigations of experimental pain in conscious animals. Pain. 16: 109-110.