

ORIGINAL ARTICLE

²The Involvement of Non Opioidergic Mechanism in the Antinociceptive and Antilocomotive Activity of Bacopa monnieri

5MUZAFFAR ABBAS, FAZAL SUBHAN, KHALID RAUF, IKRAM-UL-HAQ, and 6SYED NADEEM-UL-HASSAN MOHANI

7 For author affiliations, see end of text.

Received February 12, 2011; Revised May 9, 2011; Accepted June 27, 2011

This paper is available online at http://ijpt.iums.ac.ir

10 ABSTRACT

11A hydroethanolic extract (HE-ext) of Bacopa monnieri (BM) was studied for antinociceptive effect in the 12 animal models of acetic-acid-induced writhing test and antilocomotive effect in mice. Standard centrally-13 acting analgesic, morphine (MP), and peripherally-acting one, diclofenac (Diclo), were also tested along 14 with the extract for comparison. The extract exhibited significant antinociceptive effect (p < 0.001) in this 15 test, not antagonized by the opioid receptor antagonist, naloxone (NLX) in a fashion similar to diclofenac. 16 This excluded the involvement of opioids in the mediation of antinociceptive response of Bacopa 17 monnieri. Moreover, the BM HE-ext exhibited highly significant antilocomotive (p < 0.0001) that was also 18 unaffected by naloxone. These results indicate that Bacopa monnieri possesses antinociceptive and 19 antilocomotive effect that may be mediated through non-opioidergic mechanism.

20 Keywords: Bacopa monnieri, Hydroethanolic extract, Antinociceptive activity, Acetic-acid-induced 21 writhing test, Antilocomotive effect

Bacopa monnieri (family: Scrophulariaceae) [1] also 41 23known as Bacopa monniera, water hyssop, Herpestis 24monnieri is a perenial creeping, succulent herb found in 42Bacopa monnieri 25 marshy areas of Indo-Pak subcontinent [2]. In India, It 26 is commonly known as "Brahmi" as an ancient and 27 renowned medicinal plant with legendary reputation as 28a memory vitalizer [3]. Bacopa monnieri is held in high 29 repute to be the brain booster and is highly valued in 30 conditions affecting CNS. In ancient traditional system 31 of medicine, it is often prescribed for epilesy, insomnia, 32 and psychiatric disorders such as mental breakdown in 33 Alzheimer's disease [4], neuralgia, and memory loss 34[15]. It is known to possess cardiotonic, sedative, 35 analgesic, anticonvulsant, anti-inflammotry [6], 36 antioxidant [7], anticancer, antipyretic, laxative, 52 37 diuretic, antistress [8], and anxiolytic [9] properties. In 53 shade and coarsely grinded. The coarsely-ground 38 this study, we have examined Bacopa monnieri for 54 material was extracted with 70% ethanol and was 39 antinociceptive and antilocomotive activity in animal 55 concentrated on rotary evaporator at 60 °C, and then to 40 models.

MATERIALS AND METHODS

Bacopa monnieri was collected from Ramli stream 44 near Quaid-e-Azam University Islamabad, Pakistan and 45 authenticated by Dr. Muhammad Ibrar, Professor of 46Botany University of Peshawar. A reference specimen 47 was submitted to the herbarium of the Botany 48Department, University of Peshawar and a voucher 49 specimen (029006/Bot. University of Peshawar) was 50 obtained.

51 Preparation of Bacopa monnieri extract

Aerial parts were separated from roots, dried under 56 semisolid form (% yield: 37.25).

16 | IJPT | January 2012 | vol. 11 | no. 1

57 Chemicals and Drugs

Ethanol was obtained from Khazana Sugar Mills 59 Mardan through proper channel. Diclofenac sodium was 60 gratefully donated by Zinta Pharmaceutical Ptv, 61Peshawar, Pakistan. Morphine was secured through 62proper channel (PDH Lahore, Pakistan). Opioid 100 constrictions of treated drug / Mean no. of abdominal 63 antagonist, naloxone was purchased from Sigma, USA. 64For experiments, all drugs and extracts were dissolved 65 in water for injection.

66 Animals

Balb-C mice bred in the animal house of the 68Department of Pharmacy, University of Peshawar, were 69used in this study. Animals were housed in groups of 70 eight in cages with sawdust bedding. Experiments were 71 carried out during the light phase between 9.00 am and 723.00 pm strictly in accordance with procedures laid 73 down under the Animal Scientific Procedure Act 74(1986). Both anti-nociceptive and locomotive studies 75 were carried out on mice of either sex weighing 18-22 76g. Control animals received equal volume of normal 77saline (0.9% NaCl). Animals were marked for their 114 between 14 50 minutes are so minutes after drug 78 proper identification.

79 Procedures

80 Acetic-acid-induced writhing test

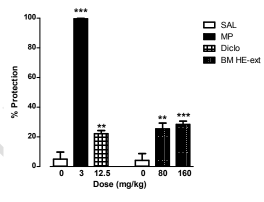
Balb-C mice of either sex (n=8) weighing 18-22 g 82were used. Animals were withdrawn from food and 119 83 water 2 hours before the start of experiment. Writhing 120 variance (ANOVA) with post hoc tests for multiple 84 behavior was tested, in which 1% acetic acid (AA) was 21 comparisons and Student's t test. Effects were stadministered intraperitoneally (i.p.) and number of 12 considered significant at p < 0.05. 86 abdominal constrictions occurring over the period of 20

87 minutes were counted just after 1% AA (10 mL/kg)

89diclofenac (12.5 mg/Kg) or normal saline (SAL) were 25 hydroethanolic extract of Bacopa monnieri in

90 administered i.p. 30 minutes before 1% AA₁₂₆ acetic-acid-induced writhing test 91 administration. However, hydroethanolic extract (80,127 As shown in the Fig 1, hydroethanolic extract of 92160 mg/kg) were administered orally (PO) 1 hour before 128 Bacopa monnieri (80, 160 mg/Kg Body weight),

93 administering 1% AA. For antagonism, naloxone (0.5129 morphine (3 mg/Kg body weight) and diclofenac (12.5



94mg/kg body weight) was administered subcutaneously 95(s.c.) 5 minutes before AA administration. All drugs 96 were administered in the volume of 0.1 mL/20.g i.p and 97s.c. and 0.2 mL/10 g PO. Percent analgesia was 98 calculated with the help of following formula:

% Protection = (1 - Mean no. of abdominal)101 constrictions of control) 100

102 Locomotor activity

Balb-C mice of either sex (n=8) weighing 22 ± 2 g 104were used. Animals were acclimatized under red light (40 Watt red bulb) one hour before the start of experiment in laboratory with food and water available ad libitum. The locomotor activity arena measured 50 x 840 cm and the floor was divided by lines into 4 equalsized rectangular zones. Doses of BM HE-ext (80 mg/kg), or morphine (10 mg/kg), or saline were administered intraperitoneally and animals were placed 2 in the recording apparatus 30 minutes later. Group mean 3 line crossing counts were subsequently recorded 14between 1- 30 mins. For antagonism, naloxone (0.25 116 administration. All drugs were administered in the 117volume of 0.1 mL/10 g i.p. and 0.1 mL/20 g s.c.

18 Statistical analysis

Results were analyzed by one-way analysis of

RESULTS

88administration [10,11]. Morphine (3 mg/kg) or 124 Antinociceptive effect of morphine, diclofenac and

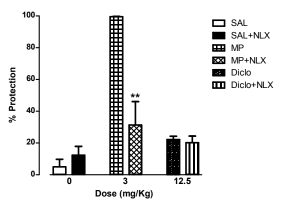


Fig 1. Antinociceptive effect of diclofenac, morphine and hydroethanolic extract of Bacopa monnieri calculated as percent protection in acetic acid induced writhing test in mice. Each column represents mean \pm S.E.M. (n=8). **p < 0.01, ***p < 0.001, Difference between treatment groups and saline control was analyzed by one way analysis of variance with Dunnett's post-hoc test.

Fig 2. The effect of naloxone on morphine and diclofenac induced antinociception calculated as percent protection in acetic acid induced writhing test in mice. Each column represents the mean ± S.E.M. (n=8), **p < 0.01, values showed significant antagonism by naloxone as compared to morphine treated groups when analyzed by Student's t test

Antinociceptive/Antilocomotive B monnieri

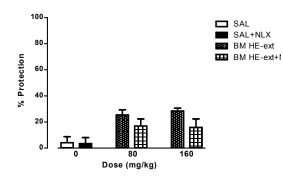


Fig 3. Effect of naloxone on BM HE-extract induced antinociception calculated as percent protection in acetic acid induced writhing test in mice. Each column represents mean ± S.E.M. (n =8). Student's t-test revealed no significant difference between two comparison groups (p > 0.05).

130 mg/Kg body weight) exhibited antinociceptive effect by 163 131 decreasing the number of acetic-acid-induced 132 abdominal constrictions in mice that was statistically 104 116 hockeept to top use the liberation of histamine, 165 writhing test results from the liberation of histamine, 133 significant.

134 Antagonism of Bacopa monnieri morphine- and 135 diclofenac-induced antinociception with naloxone

As depicted in Fig 2, pretreatment with naloxone 137(0.5 mg/kg, s.c.) reversed the antinociceptive response 138 of morphine (3 mg/Kg body weight) significantly (**p17 139< 0.01). However, the antinociceptive effect of 140 diclofenac (12.5 mg/Kg, i.p.) was unaffected with 141 naloxone (0.5 mg/Kg, s.c) pretreatment.

142 Antagonism of Bacopa monnieri hydroethanolic 143 extract induced antinociception with naloxone.

As shown in the Fig 3, naloxone did not antagonize 145 the antinociceptive effect of hydroethanolic of Bacopa 146 monnieri administered PO at the dose level of 80, 160 182 acetic acid [15,16]. Acetic-acid-induced writhing assay 147 mg/Kg body weight.

148 Effect of acute administration of morphine and 149 hydroethanolic extract of Bacopa monnieri on 150 locomotor activity in mice

As depicted in the Fig 4, acute administration of 152morphine (10 mg/Kg, i.p.) or hydroethanolic extract (80 153mg/Kg, i.p.) significantly reduced locomotor activity when compared to control (***p < 0.0001).

155 Effect of naloxone pretreatment on morphine and bydroethanolic extract of Bacopa monnieri induced 157 locomotor activity in mice

As shown in the Fig 5, in contrast to morphine (10 159 mg/Kg B.w.), the antilocomotive effect of 160 hydroethanolic extract of *Bacopa monnieri* (80 mg/Kg) 161 was not antagonized with naloxone (0.25 mg/Kg, s.c.) 162 pretreatment.

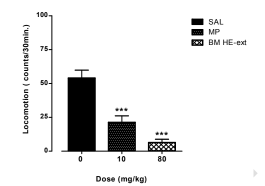


Fig 4. Effect of morphine and hydroethanolic extract of Bacopa monnieri after acute administration on locomotor activity in mice. Each column denotes mean line crossings \pm S.E.M. (n=8). ***p < 0.0001, values were significantly different as compared to control (ANOVA with Dunnett's post hoc test).

DISCUSSION

The nociceptive response in the acetic-acid-induced 166kinins, Prostaglangins, serotonin and substance P. The 167 nociceptive activity of acetic acid may be due to 168 cytokine release, such as TNF- α , inteleukin-1 β and interleukin-8, by resident peritoneal macrophages and mast cells [12]. It has been reported that intraperitoneal administration of acetic acid causes an increase in the concentration of glutamate and aspartate in the 73 cerebrospinal fluid [13].

The production of prostaglandins [14,15] results 175through the action of the constitutive enzyme 176 cyclooxygenase-1 (COX-1) and its isoform COX-2 177 which produce pain [15,16]. Induction of this 178 mechanism through COX enzymes and stimulation of 179 these sensory pathways in the mouse peritoneum incites 80 a viscero-somatic reflex and the abdominal constrictions 81 observed in response to an algogenic agent such as 183 is sensitive procedure to evaluate peripherally and

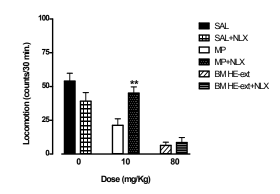


Fig 5. Effect of naloxone pre-treatment on morphine and BM HEext induced locomotor activity in mice. Each column denotes mean line crossings ± S.E.M. (n=8). Student's t-test revealed significant difference between two comparison groups (**p < 0.01).

ijpt.iums.ac.ir 17 **ARTICLE IN PRESS**

18 | IJPT | January 2012 | vol. 11 | no. 1

184 centrally acting analgesics [17-22] and sensory afferents 240 REFERENCES

185 in the peritoneum carry $\alpha_{1/2}$ -adrenoceptors, β -2411. 186 adrenoceptors and opioid receptors on their terminals²⁴² 187 [23]. When activated by appropriate agonists, these 2432. 188 receptors depress the generation of pain impulses, in²⁴⁴ 189 some instances there being an interaction between α -2453. 190 adrenoceptors and opioid receptors in the mouse²⁴⁶ 191 peritoneum [10,11,23,24].

2484. In this study, morphine, diclofenac and 193 hydroethanolic extract of Bacopa monnieri produced 250 194 significant antinociceptive effect in acetic-acid-induced²⁵¹ 195 writhing method. In order to investigate further the 2525. 196 mechanism of antinociceptive effect, the extract of 253 197 Bacopa monnieri, and standards diclofenac and 254 198 morphine were examined in the presence of non_{2556}^{2556} . 199 selective opioid receptor antagonist, naloxone. In 250 200 contrast to morphine, the antinociceptive effects of HE-2587 201 ext and diclofenac were not antagonized with naloxone. 259 202 The fact that hydroethanolic extract of Bacopa monnieri260 203 inhibits chemical-induced nociception and that 2618. 204 nociception is not antagonized with naloxone suggests²⁶² 205that the extract does not possess opioid-mediated 264 206 antinociceptive activity. This finding is in contrast to as 265 207 reported by vohora et al. 1997 [25].

Opioids have been known to possess sedative effect267 209[26,27] and that is believed to due their action at opioid²⁶⁸ 210 receptors within the central nervous system [28].26910. 211Naloxone has been known to antagonize the sedative 212 effect of opioid by acting on opioid receptors [29]. Our 213 study has also revealed that hydroethanolic extract of 11 214 Bacopa monnieri was able to promote a motor 215 depressant effect in mice. Thus, administered acutely at 2 216the dose level of 80 mg/Kg body weight, the extract2 7612 217 exerted significant decrease in locomotor activity. 218 indicating sedative properties of the extract. 219Furthermore, the anti-locomotor effect of the extract₂₈₀ 220 was not antagonized with naloxone, excluding the 28113 221 involvement of opioid receptors in the mediation of 282 222antilocomotor activity of the extract. However, 283 223 naloxone pretreatment antagonized the antilocomotive 28414. 224 activity of morphine at the dose of 10 mg/kg.

225 In conclusion, this study has demonstrated that $^{200}_{287}$ 226 hydroethanolic extract of *Bacopa monnieri* possesses $^{28815}_{287}$ 227 antinociceptive effect and inhibited the locomotor $^{289}_{289}$ 228 activity involving a non opioidergic mechanism as the $^{290}_{299}$ 229 both activities were not affected by opioid receptor $^{291}_{290}$ 230 antagonist, naloxone. $^{29216}_{29216}$

ACKNOWLEDGMENTS

The authors are gratefully thankful to Dr.29818. Muhammad Ibrar, Department of Botany, University of Pashawar, Pakistan for the identification of the plant³⁰⁰¹⁹. In the support of the Ministry of Health and Ministry of narcotic control, Pakistan for granting Pashawar, Pakistan to acquire morphine for the study. We are Ministry of narcotic control, Lahore for Base thankful to Punjab Drug House (PDH), Lahore for Sign to morphine.

Stewart RR. Flora of West Pakistan. Fakhri printing press, Karachi. 1972, p: 646.

Nadkarni KM. Indian Materia Medica. Popular Prakashan Private, Bombay. 1976:624–625.

Anonymous. The wealth of India: Raw materials, Council of scientific and industrial research, New Delhi, vol.2, 1988, p: 2-3.

Salil KB, Ashok K, Shibnath G. Effect of Bacopa monnieri on animals models of Alzheimer's disease and perturbed central cholinergic markers. *Molecular Aspects of Asian Med* 2001; 1:21-32.

Roodenrys S, Booth D, Bulzomi S, Phipps A, Micallef C, Smoker J. Chronic effects of Brahmi (Bacopa monnieri) on human memory. *Neuropsychopharmacology* 2002; 27:279-81.

Channa S, Dar A, Anjum S, Yaqoob M. Atta-ur Rahman. Antiinflammatory activity of Bacopa monniera in rodents. *J Ethnopharmacol* 2006; 104:286–89.

Tripathi YB, Chaurasia S, Tripathi E, Upadhyay A, Dubey GP. Bacopa monneira as an antioxidant: mechanism of action. *Indian J Exper Biol* 1996; 34:523-6.

Chowdhuri DK, Parmar D, Kakkar P, Shukla R, Seth PK, Srimal RC. Antistress effects of bacosides of Bacopa monniera: modulation Hsp70 expression, Superoxide dismutase and cytochrome P450 activity in rat brain. *Phytother Res* 2002; 16:639-45.

Salil KB, Ghosal S. Anxiolytic activity of a standardized extract of Bacopa monniera: an experimental study. *Phytomedicine* 1998; 5:77-82.

Gray AM, Spencer PSJ, Sewell RDE. The involvement of the opioidergic system in the antinociceptive mechanism of action of antidepressant compounds. *Br J Pharmacol* 1998; 124:669-74.

Gray AM, Nevinson MJ, Sewell RDE. The involvement of opioidergic and noradrenergic mechanisms in nefopam antinociception. *Eur J Pharmacol* 1999; 365:149-57.

Ribeiro RA, Vale ML, Thomazzi SM, Paschoalato ABP, Poole S, Ferreira SH, Cunha FQ. Involvement of resident macrophages and mast cells in the writhing nociceptive response induced by zymosan and acetic acid in mice. *Eur J Pharmacol* 2000; 387:111-8.

Feng Y, Cui M, Willis W. Gabapentin markedly reduces acetic acid induced visceral nociception. *Anesthesiology* 2003; 98:729-33.

Berkenkopf JW, Weichmann BM. Production of prostaglandin in mice following intraperitoneal injection of acetic acid, phenyl benzoquinone and zymosan: Its role in the writhing response. *Prostaglandins* 1988; 36:693-709.

Matsumoto H, Naraba H, Ueno A, Fujiyoshi T, Murakami M, Kudo I, Oh-ishi S. Induction of cycloxygenase-2 causes an enhancement of writhing response in mice. *Eur J Pharmacol* 1998; 352:47-52.

 Ballou LR, Botting RM, Goorha S, Zhang J, Vane JR. Nociception in cyclooxygenase isozyme-deficient mice. *Proc Nat Acad Sci* 2000; 97:10272-6.

 29517. Murray CW, Porreca F, Cowan A. Methodological refinement in the mouse paw formalin test: an animal model of tonic pain. J Pharmacol Method 1988; 20:175-86.

Tjosen A, Berg DG, Hunskaar S, Rosland JH, Hole K. The formalin test: an evaluation of the methods. *Pain* 1992; 51:5-17.

Gaertner M, Muller L, Roos JF, Cani G, Santos ARS, Niero R, Calixto NB, Yunes RA, Manache FD, Cechinel-Filho V. Analgesic triterpenes from sebastiania schottiana roots. *Phytomed* 1999; 6:41-4.

Ikeda Y, Ueno A, Naraba H, Oh-ishi S. Involment of vanilloid receptor VR1 and prostanoids in the acid–induced writhes responses of mice. *Life Sci* 2001; 69:2911-9.

Antinociceptive/Antilocomotive B monnieri

- 30721. Chan TF, Tsai HY, Tian-Shang W. Anti-inflammatory and 32828. analgesic activities from the roots of Angelica pubescens. Planta 329
- Medica 1995: 61:2-8. 31022. Bentley GA, Newton SH, Starr J. Evidence for an action of 33129.
- morphine and enkephalins on sensory nerves endings in the332 mouse peritoneum. Br J Pharmacol 1981; 73:325-32.
- Bentley GA, Newton SH, Starr J. Studies on the antinociceptive 333 CURRENT AUTHOR ADDRESSES 31323.
- 314 action of α -agonist drugs and their interaction with opioid mechanisms. Br J Pharmacol 1983; 73:125-34.
- Gray AM, Pache DM, Sewell RDE. Do $\alpha_2\text{-adrenoceptors play}^{335}$ 31624. an integral role in the antinociceptive mechanism of action of 336 antidepressant compounds? Eur J Pharmacol 1999; 378: 161-8. 318
- Vohora SB, Khanna T, Athar M, Ahmad B. Analgesic activity of 338 Dr. Fazal Subhan, Department of Pharmacy, University of Peshawar, 31925. bacosine, a new triterpene isolated from Bacopa monniera.339
- Fitoterapia 1997; 4:361-5. 32226. Eduardo RM, Adolfo RJ, Hemir MQA, Daniela C, Juliany GQ.341
- methadone, butorphanol or tramadol, in combination with 343
- acepromazine, in dogs. Vet Anaesth Analg 2009; 36:25-33
- Stacey YM, Christine M. Measurement of Opioid induced345 32627. Sedation. Pain Manag Nurs 2; 2001:132-49.

- Young-McCaughan S, Miaskowski C. Definition of and mechanism for opioid-induced sedation. Pain Manag Nurs 2001; 2:84-97
- Jay AA. Reversal Agents in Sedation and Anesthesia: A Review. Anesth Prog 1988; 35:43-7
- 334 Muzaffar Abbas, Department of Pharmacy, Sarhad University of Science and Information Technology, Peshawar, Pakistan, Email: mabbas14@yahoo.com, Mob. No. +923435224679, Fax: +92-91-5841460 (Corresponding Author)
 - Peshawar, Pakistan, E-mail: fazal_subhan@upesh.edu.pk
- 340 Khalid Rauf, Department of Pharmacy, University of Peshawar, Peshawar, Pakistan, E-mail: khalidrauf@upesh.edu.pk
- Comparative study on the sedative effects of morphine, 342 Ikram-ul-Haq, Department of Pharmacy, University of Peshawar, Peshawar, Pakistan, E-mail: ikram pharmacist@yahoo.com
 - 344 Syed Nadeem-ul-Hassan Mohani, Department of Pharmacy, Sarhad University of Science and Information Technology, Peshawar, Pakistan, E-mail: nadeem.fls@suit.edu.pk