

RESEARCH ARTICLE

2 Anticonvulsant Activity of the Aqueous Leaf Extract of Croton zambesicus (Euphorbiaceae) in Mice and 4Rats

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- 7 Received February 23, 2008; Revised July 6, 2008; Accepted August 12, 2008

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

9 ABSTRACT

10 To determine the anticonvulsant activity of the leaf extract of Croton zambesicus in mice and rats, and in 11 order to verify the traditional use of the plant in the treatment of epilepsy, the pentyleneterazole (PTZ) and 12the maximal electroshock seizure (MES) models were used for assessing the anticonvulsant effects of 13the aqueous leaf extract in mice and rats. In the PTZ test, the leaf extract (1000-2000 mg/kg p.o.) pro-14duced a significant (p < 0.05) increase in the onset of seizures in rats and mice compared with the control 15 group. The aqueous extract (1500 and 2000 mg/kg p.o.) produced some protection (42.9%) in rats, while 161000 mg/kg p.o. of that produced significant protection (71.4%) against PTZ-induced convulsion in mice. 17 In the MES test, the aqueous extract (500-1500 mg/kg p.o.) produced a significant (p < 0.05) increase in 18the onset of seizures compared with the control group. At 1500 mg/kg p.o., the extract also produced sig-19 nificant protection (71.4%) against MES-induced convulsions in mice. The results obtained from this study 20 indicate that the aqueous leaf extract of Croton zambesicus may be beneficial in both absence and tonic 21 clonic seizures.

22 **Keywords:** Croton zambesicus, Mice, Rats, Anticonvulsant, Pentylenetetrazole, MES test

Croton zambesicus muell Arg. (Euphorbiaceae) syn 46 study aims to evaluate the anticonvulsant activity of 24C. amabilis muell. Arg., syn. C. gratissumus Burch, is 47 aqueous leaf extract of Croton zambesicus using penty-25an ornamental tree grown in villages and towns of Nige- 48lenetetrazole (PTZ) and the maximal electroshock sei-26ria. It is a Guinea-Congolese species widely spread in 49zure (MES) tests. 27tropical Africa [1]. The leaf decoction is used in Benin 28 Republic as antihypertensive and antimicrobial (urinary 29 infections) [2]. The Ibibios in urunan area of Akwa 30 Ibom state of Nigeria use leaf traditionally as a remedy 31 for malaria [1]. Antidiabetic activity of the ethanolic 32 leave extract has also been reported [3].

34 isolated from dichloro-methane extract of the leaves has 54 University of Jos, Nigeria and was duly approved by the 35 cytotoxic activity on Hela cells [4]. The alkaloidal frac- 55 Faculty Postgraduate Board. 36tions of the leaf have been reported to possess weak 56 Plant Material 37 antimicrobial activity [5]. While the essential oil found 38in the leaves contain p-cymene are linalool and beta- 57 39caryophyllene [6]. Mekkawi [7] also reported that the 58from Bauchi Road, in Jos, Plateau State, Nigeria in July 40 constituents of the essential oil found in the flowering 592006. The plant was identified by Mr. I.A. Kareem, at 41 tops include; pinene, limonene linalool, menthol, car- 60 the Federal College of Forestry, Jos and confirmed at 42 vone, thymol, alpha-humulene and ceisnerolidol.

44 leaves is used in the prevention and treatment of epilep- 63 and then powdered using mortar and pestle. Fifty grams 45tic seizures (Dr. Azija, personal Communication). This 64(50 g) portion of the powdered leaves was extracted by

MATERIALS AND METHODS

51 Institutional Approval

The work was conducted in the Department of The ent-trachyloban-3β-ol, a trachylobane diterpene, 53 Pharmacology, Faculty of Pharmaceutcal Sciences,

The leaves of Croton zambesicus were collected 61 Forestry Research Institute of Nigeria (FRIN), Ibadan. In Jos, Plateau State of Nigeria, the decoction of the 62 The fresh leaves of the plant were shade dried for 8 days 80 | IJPT | January 2008 | vol. 7 | no. 1

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Table 1. Effect of aqueous extract of Croton zambesicus on pentylenetetrazole-induced seizures in rats

Treatment	Onset of convulsion	Number convulsed	Mortality	Protection
	(seconds)	/number used	(%)	(%)
Distilled water	44.3 ± 1.69	7/7	100	0
Phenobarbitone				
30 mg/kg)	-	-	0	100
C. zambesicus extract				
00 mg/kg	45.3 ± 0.47	7/7	100	0
000 mg/kg	45.6 ± 3.22	5/7	71.4	28.6
500 mg/kg	$55.0 \pm 1.22*$	4/7	57.1	42.9
000 mg/kg	$129.5 \pm 6.90*$	4/7	57.1	42.9

Results are expressed as mean \pm S.E.M. and as % mortality and protection (n=7). *p < 0.05 compared with control. One-way ANOVA followed by Duncan post test and Chi square test.

65 macerating with distilled water for 24 hours and then 87 1000, 1500, and 2000 mg/kg p.o.) was administered. 66 boiled for 15 minutes, allowed to cool and filtered. The 88 After a pretreatment time of 60 minutes, PTZ (85 69C prior to use.

70 Animals

Albino mice (20-25 g) and albino rats (60-80 g) of 93 PTZ- induced convulsion in mice 72 either sex were obtained from the animal house of the 73 Department of Pharmacology, University of Jos, Nige-74ria.

75 Drugs

The drugs used were supplied from the stock of De-77 partment of Pharmacology laboratory, University of Jos, 78Nigeria and include; pentylenetetrazole (Sigma) and 79 phenobarbitone (Merck).

80 PTZ-induced convulsion in rats

Six groups, each containing seven rats were used to 82 test for the effect of aqueous extract on PTZ-induced 83 seizures. They were treated as follow;

- Group I (control): distilled water (0.5 ml p.o.).
- Group II: phenobarbitone (30 mg/kg i.p.).
- Groups III-VI: graded doses of aqueous extract (500,108 of seven animals each and treated.

67 extract was evaporated to dryness at a temperature of 89 mg/kg i.p.) was administered to the six groups of ani-6840-45°C. A yield of 3.90 g was obtained and kept at 4 ° 90 mals. The onset of convulsion, number of animals that 91 convulsed and number of animals that were protected 92 were recorded [8].

A total of forty-two mice were divided into six 95 groups of seven animals each. They were treated as fol-96low;

Group I (control): distilled water (0.5 ml p.o.).

Group II: phenobarbitone (30 mg/kg i.p.).

Groups III-VI: graded doses of aqueous extract (500, 001000, 1500, and 2000 mg/kg p.o.) was administered.

After a pretreatment time of 60 minutes, PTZ (85 102mg/kg i.p.) was administered to the six groups of ani-103 mals. The onset of convulsion, number of animals that 104convulsed and number of animal that were protected 05 were recorded [8].

106 Electrically-induced seizure in mice

Thirty-five male mice were allotted into five groups

Table 2. Effect of aqueous extract of Croton zambesicus on pentylenetetrazole-induced seizures in mice

Treatment	Onset of convulsion	Number convulsed/	Mortality	Protection
	(seconds)	number used	(%)	(%)
Distilled water	42.0 ± 0.85	7/7	100	0
Phenobarbibitone				
(30 mg/kg)	-	0/7	0	100
C. zambesicus extract				
500 mg/kg	$47.9 \pm 0.89*$	7/7	100	0
1000 mg/kg	120.5 ± 0.50	2/7	28.6	71.4*
1500 mg/kg	$62.5 \pm 2.50*$	4/7	57.1	42.9
2000 mg/kg	62.5 ± 2.50 *	4/7	57.1	42.9

Results are expressed as mean \pm S.E.M. and as % mortality and protection (n=7). *p < 0.05 compared with control. One-way ANOVA followed by Duncan post test and Chi square test

Table 3. Effect of aqueous extract on maximal electroshock-induced seizures in mice

Treatment	Onset of convulsion	Number convulsed/ num-	Mortality	Protection
	(seconds)	ber used	(%)	(%)
Distilled water	9.86±0.67	7/7	100	0
Phenobarbitone				
(30 mg/kg)	-	0/7	-	100
C. zambesicus				
500 mg/kg	25.0±1.15*	3/7	42.9	57.1*
1000 mg/kg	39.0±4.36*	3/7	42.9	57.1*
1500 mg/kg	27.0±1.00*	2/7	28.6	71.4*

Results are expressed as mean ± S.E.M. and as % mortality and protection (n=7). *p < 0.05 compared with control. One-way ANOVA followed by Duncan post test and Chi square test.

Group I (control): distilled water (0.5 ml p.o.).

Group II: phenobarbitone (30 mg/kg i.p.).

Groups III-V: extract (500, 1000 and 1500 mg/kg149mice (Table 3).

112**p.o.).**

After a pretreatment time of 60 minutes, a CFP 114stimulator (model 8048) was used to deliver a stimulus 150 115 of 50 Hertz at 20 volts via ear electrodes to the different 116 groups. The animals were observed for 2 minutes. The 151 117 onset of tonic hind limb extension and number of ani-152 the threshold of PTZ-induced convulsion in rats and 118 mals protected was recorded [9].

119 Statistical analysis

121 were statistically analyzed using one-way analysis of 157 blocked by drugs that reduce T-type calcium currents 122 variance (ANOVA), followed by Duncan's multiple 158 (ethosuximide) and drugs that enhance inhibitory neuro-123 range post test and Chi square test. Values of $p < 0.05_{159}$ transmission by GABAA receptors (benzodiazepine, 124 were considered significant.

RESULTS

126 The effect of aqueous extract on PTZ-induced 127 convulsion in rats

129 clonic convulsions with 100% mortality in the control₁₆₇ increased the threshold of seizures and offered protec-130 group. The aqueous extract (1000 and 1500 mg/kg p.o.)₁₆₈tion in the MES test. It has been found empirically that 131 significantly (p < 0.05) increased the onset of convul-169 drugs which inhibit PTZ-induced convulsions and raise 132 sion in rats compared with the control group. Extract 170 the threshold for production of electrically-induced sei-133(1500 and 2000 mg/kg p.o.) offered 42.9% protection₁₇₁ zures are generally effective against absence seizures, 134 against PTZ- induced convulsion in rats (Table 1).

135 The effect of aqueous extract on PTZ-induced 136 convulsion in mice

1380.05) increased the threshold of PTZ-induced convul-177 of Croton zambesicus contains p-cymene, linalool and 139 sion in mice compared with the control group. At 1000178 beta-caryophyllene [6]. Psychopharmacological evalua-140 mg/kg p.o., the extract produced significant protection 179 tion of linalool in mice revealed that this compound has 141 (71.4%) against PTZ-induced convulsion in mice (Table 180 dose-dependent marked sedative effects at the CNS,

143 The effect of aqueous extract on MES-induced 144 convulsion in mice

DISCUSSION

147 sion in mice compared with the control group. At 1500

148 mg/kg p.o., the extract produced (71.4%) protection in

The aqueous extract of Croton zambesicus increased 153 offered protection against PTZ-induced convulsion. The 154 protection offered against PTZ-induced convulsion in 155 mice (71.4%) was significant compared to that produced The data are expressed as mean \pm S.E.M. The data 156 in rats (42.9%). Clonic seizures induced by PTZ are 160 phenobarbital and valproate) [10]. Convulsants whose 161 actions previously were unexplained (including penicil-162lin and PTZ) may act as relatively selective antagonist 163 of the action of GABA [11,12]. The fact that the extract 164 protected animal against PTZ-induced seizures may 165 suggest that the plant extract contains compound(s) that Intraperitoneal administration of PTZ induced tonic-166 facilitate GABAergic transmission. The extract also 172 whereas those that reduce the duration and spread of 173 electrically-induced convulsions are effective in tonic-174clonic seizures [13].

The anticonvulsant activity of the extract, are similar The extract (500-2000 mg/kg p.o.) significantly (p < 176 to those of linalool. The essential oil found in the leaves 181 including protection against PTZ, picrotoxin ,quinolic 182 acid and electroshock induced convulsions [14].

The results of this study show that the aqueous leaf 184 extract of Croton zambesicus possess anticonvulsant The extract (500-1500 mg/kg p.o.) significantly (p < 185 properties which are possibly mediated partly via facili-1460.05) increased the threshold of MES-induced convul-186 tation of GABA transmission. These results suggest that

187 the leaves of *Croton zambesicus* will be beneficial in the 215 188 management of absence and tonic-clonic seizures. 217**7. ACKNOWLEDGEMENT** 2198 We would like to thank the Department of Pharma-221 191 cology, Faculty of Pharmaceutical Sciences, University 2229. 192 of Jos, Nigeria for supplying the drugs used in this 223 193 study. 22510. 194 REFERENCES 22711. Okokon JE, Ofodum KC, Ajibesin KK, Danladi B, Gamaniel 228 KS. Pharmacolgical screening and evaluation of antiplasmodial activity of Croton zambesicus against Plasmodium berghei berghei infection in mice. Ind J Pharmacol 2005; 37:243-246. 23112. Adjanohaun EJ, Ajakidje V, de Souza S. Contribution to Ethno-232 1992. bothanical and Floristic Studies in Benin Republic ,1989, Vol. 1. Agency for Cultural and Technical Cooperation. 2023. Okokon JE, Bassey AL, Obot J. Antidiabetic activity of ethanolic leaf extract of Croton zambesicus muell.(Thunder plant) in $^{235\,14}$. Alloxan diabetic rats. African J Trad Complement Alt Med²³⁶

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