# **Original Research Article**

The effect of ethanol extract of *Elephantopus scaber* Linn in
decreasing blood uric acid levels of hyperuricemic male rats

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## 6 Abstract

Background: Gout causes attacks of pain and swelling in one or more joints and
control of serum uric acid level has been used as one of the therapeutic methods for
gout. Inhibition method of xanthine oxidase (XO) activity which can oxidize
hypoxanthine to uric acid has been commonly used to decrease serum uric acid level.
On the other hand, *Elephantopus scaber* Linn leaf have been used for treatment of
rheumatoid arthritis as folk medicine by some people in Indonesia

Aims: This study was to determine the hypouricemic effect of ethanol extract of *Elephantopus scaber* leaf by in vivo study in caffeine (PO)-induced hyperuricemic male
rats.

Methodology: The *E. scaber* leaf was obtained from Research Institute for Spices and Medicinal Plants, Bagor, Indonesia. Preparation of *E. scaber* leaf extract was done by cold maceration extraction technique using ethanol 70%. Male rats (Sprague-Dawley) were induced by using caffeine with dose 27 mg/200 g b.w until the levels in the blood of male rats become hyperuricemic. Rats were divided into 6 groups, as a positive

control was used allopurinol 36 mg/200 g b.w, test preparation were used 3 doses, i.e,
175 mg/200 g b.w, 350 mg/200 g b.w and 700 mg/200 b.w which suspended in NaCMC 0.5%

**Results:** The result of a decrease blood uric acid levels in rats, on the 15th days showed that a dose of 350 mg / 200 g b.w gave decrease the highest percentage ie 43%. Statistical analysis on the 15th day showed that all of the test preparation groups of *E. scaber* had effects decrease uric acid in blood serum of rats and significantly different from negative controls ( $p \le 0.05$ ). The dose 350 mg/200 g b.w of *E. scaber*. The dose of 350 mg/200 g b.w of *E. scaber* had the same effect with a dose of 36 mg/200 g b.w allopurinol in reducing uric acid in experimental rats.

31 Conclusions: *E. scaber* is a plant that is quite potential to be used in the treatment of
32 gout

33 Keywords: *Elephantopus scaber*, gout, join pain, rat, uric acid.

# 34 1. INTRODUCTION

Uric acid disease or gout is a type of joint disease that occurs due to too high levels of uric acid in the blood. Under normal conditions, uric acid dissolves in the blood and exits through urine. But under certain conditions, the body can produce excessive amounts of uric acid or experience disruption in removing excess uric acid, so that uric acid builds up in the body. Uric acid is the end product of the metabolism of destruction of purine compounds, a nucleotide that has many roles in the functioning of cells. 41 Normal uric acid levels in humans around 4 mg / dl. Uric acid levels in the blood can 42 increase beyond normal levels and is called hyperuricemia, due to increased production 43 or decreased excretion of uric acid. Increased blood uric acid levels can cause build up 44 of uric acid crystals that form like needles, especially in joints. As a result will cause 45 pain in the joints (1, 2).

46 Gout can be treated with one or more of the following drug, namely:

47	a.Nonsteroidal anti-inflammatory drugs, such as ibuprofen, naproxen and
48	celecoxib etc.
49	b. Colchicine
45	0. Colemenie
50	c. Probenecid
51	d. Allopurinol
52	e. Salicylic acid etc.

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53 Side effect of these gout drugs, generally are drowsiness, headache, diarrhea, vomiting, 54 stomach discomfort, nausea, cramping (3, 4). E scaber is highly potential for treating 55 gout, because *E scaber* contains chemical compounds that have structure activity relationship (SAR) with allopurinol, salicylate and synthetic drugs for other gout drugs, 56 namely the group of sesquiterpenes of lactone, phenolic acids and flavonoids as 57 compounds chemistry of E. scaber (5, 6,), as shown in Fig. 1. Based on this reason, we 58 59 conducted the research by giving ethanol 70% extract of E.scaber orally to rats and 60 determined the effect of the efficacy of *E scaber* leaf extract in rat blood.

61 scarlet coloured seeds are described.

# 62 2. MATERIAL AND METHODS

*E. scaber* leaf was obtained from Research Institute for Spices and
Medicinal Plants (BALITRO) Bogor, Indonesia and to determine plant
authentication was carried out in Biology research center, Indonesian
Institute of Sciences, Bogor, Indonesia.

A total of 500 mg leaf powder of *E. scaber* was extracted by repeated maceration method using 70% ethanol solvent and performed occasional shaking. The process was carried out for 3 weeks, where once in two days the solvent was replaced and filtered to obtain liquid extract, then the liquid extract was evaporated with a vacuum rotary evaporator to obtain a viscous extract and dried using a freeze dryer.

## 73 2.1. Compound and phytochemical screening of *E.scaber* extract

Compound and phytochemical screening of *E.scaber* extract was done based on
Harbone and Farnswoth methods, namely for the groups of alkaloid, flavonoid,
saponin, steroid, triterpenoid, tannin, quinone and essential oil (7, 8), . as follows:

#### 77 2.1.1. Determination of the Alkaloid Group

A total of 0.5 grams of *E. Scaber* extract was dissolved in 1% hydrochloric acid and filtered. The filtrate was divided into two parts, one part was dropped with Mayer's reagent and the other was dropped with Dragendorf reagent. Positive results were

81 indicated by the formation of white precipitation with Mayer reagent and red82 precipitation with Dragendorf reagent

#### 83 2.1.2. Determination of the Flavonoid Group

A total of 0.5 grams of *E. Scaber* extract was dissolved with 2 mL of 70% ethanol and 3
drops of NaOH solution were added. The change in the intensity of the yellow color
becomes colorless on the addition of sulfuric acid indicating the presence of flavonoids

#### 87 2.1.3. Determination of the Saponin Group

As much as 0.5 gram of extract *E. Scaber* was put into the test tube, 10 ml of hot water was added, cooled and then shaken vertically for 10 seconds. The formation of foam as high as 1–10 cm which was stable for not less than 10 minutes showed the presence of saponins. At the addition of 1 drop of 2N HCl, foam was not lost

## 92 2.1.4. Determination of Terpenoid Groups and Steroids

The contents of the secondary metabolites of the terpenoid group and the steroid extract of *E. scaber* were determined by using Liebermann-Burchard reagent. A total of 0.5 gram of extract was added 5 mL of chloroform, then acetic anhydride was added and a few drops of concentrated sulfuric acid. The test results were positive for terpenoids when dark green was formed. Positive test results for steroids if pink or red were formed As much as 0.5 gram of E scaber extract was dissolved in 5 mL aquadest then drops of
10% iron (III) chloride solution, if it was formed in blue or blackish green color showed
the presence of tannins

103 **2.2. Treatment of test preparations in experimental animals** 

The male white rats, strain of Sprague-Dawley with 3-4 months old (weight 190-250 g) 104 were acclimatized for two weeks and maintained on 12hours light, 12hours dark cycle 105 on temperature 25°C. Procedure maintenance of rats and conducting experiments on 106 107 animals were done based on norms of Committee for the Purpose of Control and 108 Supervision on Experiments on Animals /CPCSEA, 2003(9). The rats qualified for the 109 experiment were divided into 6 groups. The number of rats per group was calculated 110 based on Federer's formula (10), where for 6 group were greater than 15, therefore the 111 number of rats per group was obtained:

112	(n-1).(t-1) = (6-1).(4-1) > 15. Or each group consists of 4 rats.
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114		
115	No	Groups
116		$\mathbf{\nabla}$
117	1	Normal control was given only solution Na-CMC 0.5%
118	2	Negative control was given caffeine 27 mg/200 g BW in solution of Na-CMC
119		0.5%

Table 1. Groups of test animals

120	3	Positive control was given caffeine 27 mg/200 g BW and alopurinol 36 mg/200
121		g BW in solution of Na-CMC 0.5%
122	4	Low dose was given caffeine 27 mg/200 g BW in solution of Na-CMC 0.5%
123		and 175 mg/200 g BW extract E. Scaber in solution of Na-CMC 0.5%
124	5	Middle dose was given caffeine 27 mg/200 g BW in solution of Na-CMC 0.5%
125		and 350 mg/200 g BW extract <i>E. Scaber</i> in solution of Na-CMC 0.5%
126	6	High dose was given caffeine 27 mg/200 g BW in solution of Na-CMC 0.5%
127		and 700 mg/200 g BW extract E. Scaber in solution of Na-CMC 0.5%
128		
129	In this	experiment were calculated, the dose of allopurinol as a positive control used for
130	human	s was 200 mg / day. The conversion factor from human to rat was 0.018 and the
131	pharm	acokinetics factor used is 10 Therefore dose for rat was 200 mg x 0,018 x $10 =$

132 36 mg / 200 g b.w. The dose of caffeine that used for humans is 150 mg / day. The 133 conversion factor from human to rat was 0.018 and the pharmacokinetics factor used is 134 10. (b). Therefore dose for rat is 150 mg x 0,018 x 10 = 27 mg/ 200 g b.w. As shown in 135 table 1.

In this case before the experiment was done, the rats were fasted for 12 hours. To
increase uric acid levels in rats blood, the rats were induced with caffeine 27 mg/200 g
b.w. Based on experiment that was done by Azizahwati et al (2005) uric acid levels in
rats blood were already high on the sixth day (11).

On the seventh day, the treatment was given based on each group every day. Caffeine
was also administered to all groups except the normal group. Measurement of blood
uric acid levels continued on the day, 9th, 12th and 15th.

The measurement level of uric acid in the blood was done by taking the blood on therats tail and measured with equipment (Easy Touch) by using uric acid strip.

The test results data were analyzed using data processing software SPSS-19 and presented in the mean and standard deviation of each group. The data were processed using statistical analysis with normality test, homogeneity test, One Way ANOVA and Kruskal-Wallis Test.

### 149 **3. RESULTS AND DISCUSSION**

The results of determination of plant taxonomy was done by Herbarium Bogoriense,
Biological Research Center, Indonesian Institute of Sciences, indicating that the plant
used was *E.scaber*.

The result extraction of 500 mg simplicia powder (*E. scaber*) was obtained 92.6 g extract or the yield of simplicia was 92.6 g divided by 500 g was 18.52%.

The results of phytochemical screening of *E.scaber* was done based on the Harbone and Farnswoth methods, *E. scaber* contains groups of chemical compounds, as shown in table 2.

- 158 The result of Phytochemical Analysis for the identification of the chemical constituents
- 159 present in the 70% ethanol extract based on Harborne and Farnswoth methods were

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160	obtained chemical compound groups as shown in table 2. In this study we used 70%
161	ethanol solvent, because 70% ethanol solvent was a more powerful solvent in isolating

162 chemical compounds in natural products compared to other organic solvents (7).

163 Table 2. The results of phytochemical screening of *E scaber* 70% ethanol extract. Group of chemical Screening results compounds

	compounds	
	a. Alkaloid	+
	b. Flavonoid	+
	c. Saponin	+
	d. Steroid/triterpenoid	+
	e. Tannin	+
	f. Quinone	-
	g. Essential oil	+
	h. Qoumarin	<u> </u>
164	Note:	
165	(+) contain chemical co	<b>1</b>
166	(-) does not contain che	mical compounds
167		
168	The result of chemical compo	unds in this research was obtained the group of alkaloid,
169	flavonoid, saponin, steroid/tr	terpenoid, tannin, essential oil, as shown in Table 2.
170	As shown in Table 3 and Figu	re 1, the results of rats were induced with caffeine cause
171	increase uric acid level in the	blood on day sixth on all groups of rats and differed
172	significantly from normal rats,	namely rats group that were not induced with caffeine (p
173	≤ 0.05).	

174 Statistical test of one-way ANOVA and Least Significance Different (LSD) on day 9th, 175 the levels of blood uric acid on all groups were still significantly different ( $p \le 0.05$ ) 176 with the normal control group. This was because the work of caffeine to increase uric 177 acid was stronger than the work of test preparation to decrease uric acid levels in all 178 groups of rats.

experiment (mg / dL)						
Days	Normal Control	Negative Control	Positive Control	Low Dose	Medium Dose	High Dose
0	1.65	1.48	1.30	1.60	1.25	1.53
6	1.50	2.90	2.78	2.80	3.00	2.80
9	1.43	3.33	2.45	2.48	2.33	2.58
12	1.50	3.55	1.78	2.30	2.23	1.95
15	1.43	3.85	1.15	1.75	1.70	1.65

179 Table 3. The mean measurements of uric acid blood levels of the test animals during the 180

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	.0	ч

182 Statistical test of one-way ANOVA and LSD on day 12th, the levels of blood uric acid on high dose and positive control were not significantly different ( $p \ge 0.05$ ) with the 183 normal control group. This was because the work of caffeine to increase uric acid was 184 almost same with the work of test preparation to decrease uric acid levels in these 185 186 groups of rats

Statistical test of one-way ANOVA and LSD on day 15th, the levels of blood uric acid 187 on positive control, low dose, middle dose and high dose were not significantly 188 189 different ( $p \ge 0.05$ ) with the normal control group. This was because the work of caffeine to increase uric acid was almost same with the work of positive control, low 190 dose, middle dose and high dose to decrease uric acid level in the blood of these groups. 191

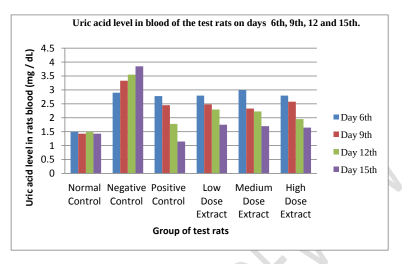




Figure 1. Uric acid level in blood of the test rats on days 6th, 9th, 12th and 15th after
administration test preparation.

196 While uric acid level in the blood of negative control was still high, because on this 197 group caffeine still increase uric acid level and there was no drug or chemical compound that work to increase uric acid level in the blood. Whereas on positive 198 199 control, low dose, middle dose and high dose, the work of test preparation can increase 200 of uric acid reduce uric acid which was enhanced by caffeine or in other words, the gout 201 on rats and human can be treated by E. scaber extract. As shown in Table 3 and Figure 202 1. Percentage (%) of increase and decrease of uric acid levels after was induced with 203 caffeine and administration of test preparations on the 9th, 12th and 15th days, 204 compared to 6th day, ie before the administration of the test preparation, can be shown in Table 4 & Figure 2. 205

Table 4. Percentage (%) of increase and decrease of uric acid levels after was induced
 with caffeine and administration of test preparations on the 9th, 12th and 15th days,

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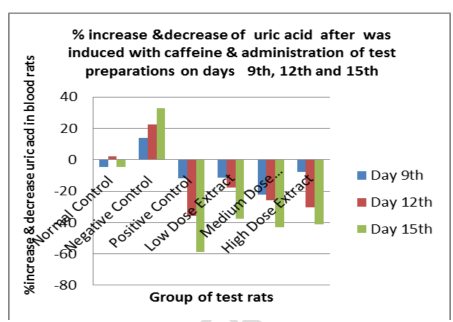
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Days	Normal	Negative	Positive	Low	Medium	High Dose
	Control	Control	Control	Dose	Dose	
9	(-)4.66	13.79	-11.87	-11.43	-22.33	-7.86
12	0.02	22.41	-35.97	-17.86	-25.67	-30.36
15	(-)4.66	32.75	-58.63	-37.50	-43.00	-41.00

209

210 In this case on normal controle group because there was no administration caffeine and 211 test preparation, uric acid level just experience a slight fluctuation value, decrease on 212 day 9th about 4.66% and increase + 0,02 % on day 12th and increase again on day 213 15th. On negative controle group because there was administration caffeine 27 mg/200 214 g b.w every day from first day until day 15th of experiment has caused uric aci level in 215 the blood increase and increase from day to day far beyond uric acid levels in the normal control group, which reaches levels 3.85 mg/dL on the day 15th. While uric acid 216 level in the blood of normal control just about 1.50 mg/dL. Percentage (%) decrease 217 happened on day 15th with value for Positive Control, Low dose, Medium Dose, High 218 Dose (-)58.63%, (-)37.50%, (-)43.00% and (-)41.00% respectively. 219

As shown in Table 4 and Figure 2, Negative control. Group always increase from day
6th until day 15th, caused was induced with caffeine, while Positive Control, Low dose,
Medium Dose, High Dose always decrease from day 9th until day 15th, caused the
word of test preparation.

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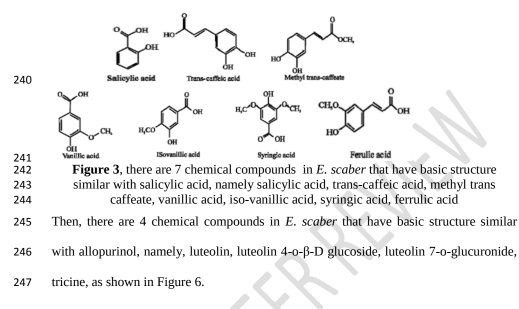


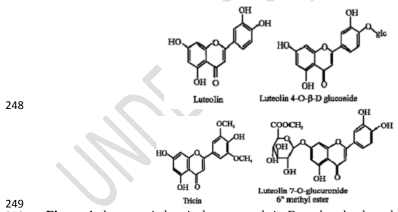
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**Figure 2.** % increase & decrease of uric acid after was induced with caffeine & administration of test preparations on days 9<sup>th</sup>, 12<sup>th</sup> &15<sup>th</sup>. compared day 6th

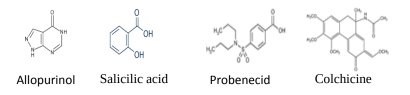
Structure Activity Relationships (SAR) are relations between the molecular structure
and biological or physicochemical activity of chemicals or in pharmacology, chemical
compounds that have the same chemistry and differ in functional groups, will have the
same properties but differ in potential efficacy (12, 13, 14, 15).

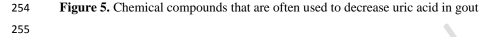
Chemical compound that work in decrease uric acid level in blood of rats, probably derived from Some phenolic acid and flavonoid compound. According to Kabeer and Prathapan (2014), Chang et al (2011) were obtained some chemical compound of phenolic acid and flavonoid compound in *E. scaber* as shown in Figure 3. and Figure 4, (5, 6). These chemical compounds have the same basic structure with chemical compounds that are currently widely used to treat gout, as shown in Figure 5.





- **Figure 4.** there are 4 chemical compounds in *E. scaber* that have basic structure similar with allopurinol, namely, luteolin, luteolin 4-o- $\beta$ -D glucoside, luteolin 7-o-glucuronide, tricine.
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There is also a strong possibility that the chemical compound in *E. scaber* in Figure 3 and Figure 4 above have the work of decrease uric acid better than the drugs for gout that exist today.

The results of this study are similar to the results of research by Jennie et al (2016), where they used *E. scaber* powdered leaves extracted using methanol and distilled water to obtain test extracts. The administration of the test preparation was carried out by using a single subcutaneous injection of 0.1 ml Complete Freund Adjuvant emulsion (CFA). Their experiments also found that *E. scaber* worked significantly in reducing uric acid in gout (16, 17)

## 265 **4. CONCLUSIONS**

*E.scaber* has the effect decrease uric acid on gout. Effect of ethanol extract 70% of *E. scaber* with dose 175 mg / 200 g b.w on rat have the same effect with dose allopurinol
of 36 mg / 200 g b.w rat.

There are about 7 of the salicylic acid derivatives and there are about 4 of the phenolic acids and flavonoid derivatives in *E. scaber*, these chemical compounds are strongly suspected have properties for decrease uric acid in the blood, because their basic

272 structure is very similar with drugs to decrease uric acid. Further research is needed, to

find out what chemicals are better for treating gout in *E. Scaber*.

## 274 CONFLICTS OF INTEREST

275 The authors declare no conflicts of interest.

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#### 277 COMPETING INTERESTS DISCLAIMER: 278

Authors have declared that no competing interests exist. The products used for this research are commonly and predominantly use products in our area of research and country. There is absolutely no conflict of interest between the authors and producers of the products because we do not intend to use these products as an avenue for any litigation but for the advancement of knowledge. Also, the research was not funded by the producing company rather it was funded by personal efforts of the authors.

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