

1735-2657/11/101-1-5 **IRANIAN JOURNAL OF PHARMACOLOGY & THERAPEUTICS** Copyright @ 2011 by Razi Institute for Drug Research (RIDR) IJPT 10: 1-5, 2011

**RESEARCH ARTICLE** 



# <sup>2</sup>Antibacterial Activity and Cytotoxicity Screening of Sumatran Kaduk (Piper sarmentosum Roxb.)

4E. ATIAX, F. AHMAD, H. M. SIRAT and D. ARBAIN

5 For author affiliations, see end of text.

6 Received December 15, 2009; Revised July 3, 2010; Accepted September 28, 2010

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

# 8 ABSTRACT

Phytochemical investigations of *Piper sarmentosum* Roxb., yielded four compounds; three amides, ioidentified as 3-(3',4',5'-trimethoxyphenylpropanoyl) pyrrolidine, 3-(4'-methoxyphenylpropanoyl) pyrrole, *N*-11 (3-phenylpropanoyl) pyrrole and a sterol namely  $\beta$ -sitosterol. 3-(4'-Methoxyphenylpropanoyl) pyrrole was 12 found for the first time in this Piper species. All chemical constituents were tested for their antibacterial 13 activity using disk diffusion method and cytotoxicity screening using sul-forhodamine B (SRB) assay. All 14of the compounds were found only active towards gram-positive bacteria except 3-(4'-15 methoxyphenylpropanoyl) pyrrole with no activity against both gram-positive and gram-negative bacteria. 16 Meanwhile, the cytotoxicity screening using SRB assay indicated that none of these compounds was 17 active as an anticancer agent.

18 Keywords: Piperaceae, P. sarmentosum, Amides, Antibacterial, Cytotoxicity

The study of medicinal plants opened the door to the 47them for a long time in a variety of medicinal capacities 20 development of purified and defined chemical 48[5]. 21 compounds as dose-controlled medicines. Natural 49 22 compounds can become central players in the treatment 50 which is known as kaduk, sirih duduk or mengkadak in 23 of disease and in the understanding of disease 51 Indonesia. Traditionally, it was used as a remedy for 24 mechanisms. Compounds that emerged from the study 52 tooth-ache and for fungoid dermatitis on the feet, for 25 of ethnobotanic extracts became important as medicines 53 treatment of coughs, influenza and rheumatism [6]. A 26 and were enabling as pharmacologic tools in the 54 decoction of the leaves is drunk to treat malarial fever 27 elucidation of disease mechanisms [1]. Piperaceae 55[7] and the crushed leaves are mixed with water and 28 family has provided many past and present civilizations 56 used for bathing to treat kidney stones and difficulty in 29 with a source of diverse medicines and food grade spice 57 urination [8]. Previous chemical constituents on this 30[2]. This plant is distributed pantropically. The earliest 58 plant have resulted in the isolation of a number of 31 classification of the Piperaceae family recognized 59 compounds [9-13]. We now describe the isolation of an 32 between 7 to 15 genera and five of them such as Piper, 60 additional amide from the aerial part of this plant which Peperomia, Lepianthes, Macropiper, and Trianopiper 61 was collected from Sumatra, Indonesia and also its 34 are only accepted as the principle genera of Piperaceae. 62 antibacterial and cytotoxicity activities. 35 This genus contains over 1000 species in the world [3]. 36 This plant can be recognized by three main features: 37 articulate stem, asymmetrical or cordate leaves, and 63 38 axillary spikes of little round berry-like fruits [4]. 64 General Experimental Procedures 39 According to Jaramilo [3], Asian tropic has 340 species 40 of Piper, including Sumatra tropical rainforests. This 65 41 species takes the form of shrubs, herbs, lianas, and 66 Gallen III apparatus. IR spectra were recorded on a 42 mostly woody climbers. They are common in the warm, 67 Perkin-Elmer 1650 FTIR spectrophotometer. NMR <sup>43</sup>humid region and in the lowland of wet forests. The <sup>68</sup>spectra were recorded on a Bruker Avance 300 <sup>44</sup>leaves are typically aromatic or have pungent smell. <sup>69</sup>Spectrometer, <sup>1</sup>H NMR spectra were measured at 300 45 This genus consists of a large family of plants 70 MHz and <sup>13</sup>C NMR spectra were measured at 75 MHz. 46 indigenous to the tropic and native people have used 71 Deuterated solvent of chloroform (CDCl<sub>3</sub>) was used as

Piper sarmentosum Roxb. is one of the Piper genus

# **MATERIALS AND METHODS**

Mps. (uncorr.) were determined using the Leica

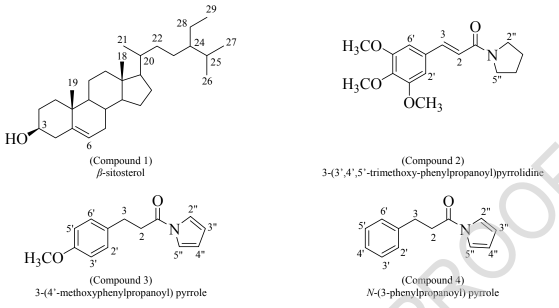


Fig 1. Chemical structure of the isolated compounds of P. sarmentosum

72 solvents. Mass spectra data were obtained from Kent108 concentrated and further purified by recrystallization 73Mass Spectrometry Services, United Kingdom. CC:109 from hexane to yield 3-(3',4',5'-trimethoxy-74 silica gel (Merck 70-230 mesh and 230-400 mesh). 110 phenylpropanovl) pyrrolidine (Fig 1- Compound 2) 75 Spots on TLC were visualized by UV (254 and 365 nm)111 (135.6 mg,  $8.5 \times 10^{-3}$ %) as colourless crystalline solids 76 and vanillin-sulphuric acid reagent. Streptomycin 112 with mp 158-159°C (lit. [15] 156-157°C).

77 sulphate standard was purchased from Oxoid 11 78(Hampshire, UK).

# 79 Plant Material

14 VLC using hexane, mixture of hexane-CH<sub>2</sub>Cl<sub>2</sub> and 115CH<sub>2</sub>Cl<sub>2</sub> by step gradient polarity technique to yield 116 seven fractions (PSEA-PSE G). Fraction PSEB (1.033

The aerial parts of P. sarmentosum were collected 17 g) was purified by CC over SiO<sub>2</sub> (70-230 mesh, 75 g) 81 from Desa Sariak, Sungai Pua, about 11 km from 118 with hexane-CH2Cl2 (50: 50) as eluent to give four 82Bukittinggi, West Sumatra, Indonesia in 2005. The119 fractions. Fractions 25-124 were combined and 83 sample (EM-01/1205) was identified by Mr. Rusdi120 concentrated to yield 3-(4'-methoxyphenylpropanoyl) 84 Tamin and Ms. Nurainas and specimen was deposited at 121 pyrrole (Fig 1- Compound 3) (18.9 mg, 0.0012%) as 85the Andalas Herbarium (ANDA), University of 122 yellow crystalline solids with mp 83-84°C (lit. [16] 86-86 Andalas, Padang, Indonesia. 12387°C).

124

## 87 Extraction and Isolation

89*sarmentosum* (1.6 kg) was 90 successively with 3.5 L of each hexane and ethyl acetate 128 rechromatographed by CC over SiO2 with CH<sub>2</sub>Cl<sub>2</sub> as 91 for 18 hours. The solvent of each extract was evaporated 129 eluent to yield N-(3-phenylpropanoyl) pyrrole (Fig 1-92in vacuo to afford the crude hexane, PSH (28.9 g, 1.81130 Compound 4) (34.5 mg, 0.0022%) as colourless liquid. 93%) and ethyl acetate, PSE (24.9 g, 1.55%). The crude 131 Antibacterial Assay (disk diffusion method) 94PSH extract (10 g) was fractionated by VLC over silica 95gel (230-400 mesh, 250 g) and eluted with gradient132 96 solvent system of hexane, hexane- CH<sub>2</sub>Cl<sub>2</sub> and CH<sub>2</sub>Cl<sub>2</sub>133 were tested against gram-negative; Escherichia coli, 97 to afford 10 fractions (PSHA-PSHJ).

134 Pseudomonas aeruginosa and gram positive bacteria; Fraction PSHD and PSHF were combined (421.5135 Bacillus subtilis, and Staphylococcus aureus. Agar 99mg) and subjected to CC over SiO<sub>2</sub> (70-230 mesh, 40 g)136 cultures of the test microorganisms were prepared 100 to yield nine fractions. The seventh fraction was 137 according to Mackeen et al. [17]. Samples were 101 concentrated and recrystallized from CH<sub>2</sub>Cl<sub>2</sub>-hexane to 138 dissolved in MeOH (1 mL). The test samples (10 µL) 102 give  $\beta$ -sitosterol (Fig 1- Compound 1) (95.1 mg, 139 were loaded onto each Whatman filter paper disks (0.6 1030.0059%) as white crystalline needles with melting140mm) and evenly placed on the agar surface previously 141 inoculated with the suspensions of microorganism to be 104point (mp) 133-134°C (lit. [14] 138-139°C). Fraction PSH I (248.5 mg) was chromatographed142tested. Standard disk of streptomycin sulphate (10 106 over SiO<sub>2</sub> (30 g) CC with CH<sub>2</sub>Cl<sub>2</sub> (100%) as eluent to 143µg/disk) was used as the positive control and DMSO

107 give 156 fractions. The combined fractions 17-39 was 144 was used as the negative control. The plates were

The crude EtOAc extract (10 g) was fractionated by

125 over SiO<sub>2</sub> (70-230 mesh, 40 g) with hexane-CH<sub>2</sub>Cl<sub>2</sub> The powdered of aerial plant parts of P.126(10:90) as eluent to afford 102 fractions (PSED1-PSE

Fraction PSE D (481.7 mg) was subjected to CC

The chemical constituents from P. sarmentosum

soxhlet-extracted 127D9). Fraction PSE D4 (83.6 mg) was further

www.SID.ir

Atiax et al.

### Antibacterial activity of Sumatran Kaduk

Table 1. Antibacterial activity of the isolated compounds of P. sarmentosum

Compounds		Zone of Inhibition (mm)					
	Gram-Positi	ve Bacteria	Gram-Negative Bacteria				
	B. s	<i>S. a</i>	Р. а	Е. с			
(1)	9.7 <u>+</u> 0.52	10.3 <u>+</u> 0.82	-	-			
(2)	$12.3 \pm 0.52$	-	-	-			
(3)	-	-	-	-			
(4)	-	$10.2 \pm 0.41$	-	-			
SS	19.3 <u>+</u> 1.21	21.3 <u>+</u> 0.81	19.1 <u>+</u> 0.54	18.8 <u>+</u> 0.74			

Data represent mean ± standard deviation of three independent experiments performed in duplicate. (-): no activity; B.s: Bacillus subtilis; S.a: Staphylococcus aureus; E.c. Escherichia coli; P.a. Pseudomonas aeruginosa.

		MIC (µg/mL)				MBC (µg/mL)			
Compounds	Gram-Posi	Gram-Positive bacteria		Gram-Negative Bacteria		Gram-Positive Bacteria		Gram-Negative Bacteria	
	<i>B. s</i>	<i>S. a</i>	Р. а	Е. с	<i>B. s</i>	S. a	<i>P. a</i>	<i>E. c</i>	
(1)	-	500	-	-	-	500	-	-	
(2)	500	-	-	-	1000		-	-	
(3)	-	-	-	-	-	-		-	
(4)	-	125	-	-	-	125		-	
ŜŚ	3.91	3.91	3.91	3.91	7.81	7.81	7.81	7.81	

MIC: Minimal inhibitory concentration, MBC: minimal bacterial concentration, B.;, Bacillus subtilis; S.a: Staphylococcus aureus; E.c: Escherichia coli; P.a: Pseudomonas aeruginosa.

145 inverted and incubated for 18 hours at 37°C. Clear 182 Cytotoxicity Screening 146 inhibition zones around the discs indicated the presence 147 of antimicrobial activity.

The positive results then followed by the 149 determination of Minimum Inhibitory Concentration 150(MIC) by the micro-titer broth dilution method [18]. 151 This test was performed in a sterile 96-well micro titer 152 plates. The test samples (1 mg) were dissolved in 153 methanol to obtain 1000 µg/mL stock solution. Each 154 methanolic stock samples (10  $\mu$ L) was transferred to 155 micro titer plate well in duplicate at row A. A number of 156 wells were reserved in each plate for positive and 157 negative controls. Sterile broth (100  $\mu$ L) was added to 158 each micro-titer plate well from row B to row H. Then, 159the suspensions of microorganisms (200 µL) were added 160 to the samples at row A. Mixture from row A (100  $\mu$ L) 161 was transferred to each micro titer plate well in order to 162 obtain a twofold serial dilution of stock samples 163(concentration of 500  $\mu$ g/mL to 3.9  $\mu$ g/mL) plates were 164 then incubated for 18 hours at 37°C. Bacterial growth 200 Statistical Analysis 165 was indicated by the presence of turbidity and a pellet at 166 the bottom of the well. The lowest concentrations, 167 which did not show any growth of tested 168 microorganisms after macroscopic evaluation were<sup>203</sup> triplicate samples.

169 determined as MIC values.

170 The MIC values were confirmed by the204 171 determination of Minimal Bactericidal Concentration 172 (MBC) values according to method developed by Arias  $^{203}$  rule hexane extract of *P. sarmentosum* identified 173 *et al.* [19]. All wells in the MIC study, which did not 207 as  $\beta$ -sitosterol (Compound 1) and 3-(3',4',5'-174 show any growth of bacteria after incubation period<sub>208</sub> trimethoxyphenylpropanoyl)-pyrrolidine (Compound 2) 175 were first diluted in fresh nutrient broth (1:4) and then 209 and two amides have also been isolated from the crude 176 sub-cultured onto the surface of freshly prepared<sub>210</sub>EtOAc 177 nutrient agar plates ( $\theta$ ,15 mm). The plates were<sub>211</sub> methoxyphenylpropanoyl)pyrrole (Compound 3) and N-178 incubated for 18 hours at 37°C. The MBC were<sub>212</sub>(3-phenylpropanoyl)pyrrole (Compound 4). 179 recorded as the lowest concentration of the sample that<sub>213</sub> The antibacterial activity using disk diffusion 180 did not permit any visible bacteria colony growth on the214 method, followed by the determination of MIC and 181 appropriate agar plate after the incubation period.

The cytotoxicity screening was carried out according 184to sul-forhodamine B (SRB) method described by Houghton et al. [20]. This method relies on the uptake of the negatively charge pink aminoxanthine dye, <sup>7</sup> sulphorhodamine B by basic amino acids in the cells. The greater the number of cells, the greater the amount of dye is taken up and, after fixing, when the cells are lysed, the released dye will give a more intense colour and greater absorbance.

The screening of cytotoxicity test for isolated scompounds from this Piper species against four 94 cancerous cell lines i.e. human breast carcinoma cell lines (MCF-7 and MDA-MB-231), human intestine epithelial cell line (HT-29) and human ovarian 7 carcinoma cell line (SKOV-3), was carried out by Mr. 8Cheah Yew Hong from the Institute for Medical 9Research (IMR), Malaysia.

Statistical analyses were performed using Sigma plot 2028.0. Data is presented as means standard error of

### RESULTS

Two chemical constituents have been isolated from extract namely 3-(4'as

215 MBC were presented in Table 1 and Table 2. The

Archive of SID

# **ARTICLE IN PRESS**

## 4 | IJPT | January 2011 | vol. 10 | no. 1

Atiax et al.

Table 3. Percentage of cells survival on cytotoxicity assay of isolated compounds of P. sarmentosum by SRB assay

	Percentage of cell survival (%) at 20 µg/ml of samples					
Compounds	Cell lines					
-	MCF-7	SKOV3	HT-29	MDA-MB-231		
(1)	100.16	128.59	102.54	110.43		
(2)	97.39	110.26	133.71	113.79		
(3)	93.67	105.23	102.49	100.46		
(4)	69.00	122.31	91.53	94.28		

SRB = sulphorhodamin B, MCF-7 and MDA-MB-231 (human breast cancer cell lines), SKOV3 (human ovarian carcinoma cell lines), HT-29 (human colon/intestinal carcinoma cell lines).

2692.

216 isolated compounds were also screened for their 265 REFERENCES217 cytotoxic assay using SRB assay. Their activities are 2661.218 given in Table 3.267

Rishton JM. Natural Products as a Robust Source of New Drugs and Drug Leads: Past Successes and Present Day Issues. *Am J Cardiol* 2008; 101: 43D-49D.

### DISCUSSION

The isolated compounds were identified based on 2723. 221 the physical, chemical and spectroscopic properties and 221 222 comparison with data of the literatures. This is the first<sub>274</sub> 223 reported of the isolation of 3-(4'-2754 224 methoxyphenylpropanoyl) pyrrole from **P.**276 225 sarmentosum. It was reported previously from Piper 2775. 226 lolot C.DC., from Vietnam [16]. Occurrence of 278 227 chemical constituents of a plant species depends on 279 228 several factors, such as location or environment that will<sup>2806</sup>. 229probably give variation in constituents. Geographical 282 230 distribution, seasons, different plant parts and 2837 231 morphology, climate as well as ecological conditions 232may also influence the biosynthesis of the secondary 2858. 233 metabolites of the plants. This is may be the reason why 286 234 the chemical constituents of this species are different 2879. 235 from the same species which were reported previously. 288 As shown in Tables 1 and 2, all isolated compounds<sup>289</sup> 9010. 237 were found active towards Gram positive bacteria<sup>2</sup> 3-(4'-methoxyphenylpropanoyl)pyrrole 238except that 239 shown no activity against both Gram negative and 29311. 3-(3',4',5'-Trimethoxyphenyl-294 240**positive** bacteria. 241 propanoyl)pyrrolidine showed significant activity 295 242against B. subtilis (MIC 500 µg/ml, MBC 1000 µg/ml)29612. 243 followed by  $\beta$ -sitosterol (MIC and MBC 500 µg/ml) and 297 244N-(3-phenylpropanoyl)pyrrole (MIC and MBC 125<sup>298</sup> 245µg/ml) against S. aureus. But, activity of these 246 compounds is not as good as the activity of positive  $_{30013}^{30013}$ . 247 control streptomycin sulphate (MIC 3.91 µg/ml and 302 248MBC 7.81 µg/ml). All isolated compounds exhibited no<sub>30314</sub>. 249 activity towards Gram negative bacteria.

In the toxicity screening using SRB assay showed 30515. 251 that all isolated compounds have the percentage of the 306 252 cell survival was higher than 50%. Thus, indicated that 307 253 none of these compounds was active as anticancer agent 254 (Table 3). 310

255 As the conclusion, geographical distribution, 256 location or environment, seasons, different plant parts, 257 climate as well as ecological conditions may influence 31317. 258 the biosynthesis of the secondary metabolites of the 259 plants species which portray the variation in chemical 260 constituents. Investigations on the methanolic extracts 31718. 261 of these *Piper* species should be carried out. Different 262 models of biological activities should be performed on 31919. 263 the crude extracts and pure compounds to verify the 264 mode of action of the active candidates.

Scott IA, Jensen HR, Philogene BJR, Arnoson JTA. Review of *Piper* spp. (Piperaceae) Phytochemistry, Insecticidal Activity and Modes of Action. *Phytochem Rev* 2008; 7:65-75.

Jaramillo MA, Manos PS. Phylogeny and Pattern of Floral Diversity in the Genus *Piper* (Piperaceae). *Am J Bot* 2001; 88:706-16.

Wiart C. Medicinal Plants of Asia and the Pacific. Boca Raton, London, New York: Taylor & Francis. 2006.

Parmar VS, Jain SC, Bisht KS, Jain R, Taneja P, Jha A, Tyagi OD, Prasad AK, Wengel J, Olsen CE, Boll PM. Phytochemistry of the Genus *Piper*. *Phytochemistry* 1997; 46:597-503.

Rahman NNNA, Furuta T, Kojima S, Takane K, Mohd MA. Anti-malarial Activity of Extracts of Malaysian Medicinal Plants. *J Ethnopharmacol* 1999; 64:249-54.

Ong HC, Nordiana M. Malay Ethno-medico Botany in Machang, Kelantan, Malaysia. *Fitoterapia* 1999; 70: 502-13.

Ong HC, Norzalina J. Malay Herbal Medicine in Gemenceh, Negeri Sembilan, Malaysia. *Fitoterapia* 1999; 70:10-4.

Masuda T, Inazumi A, Yamada Y, Padolina WG, Kikuzaki H, Nakatani N. Antimicrobial Phenylpropanoids from *Piper sarmentosum. Phytochemistry* 1991; 30:3227-8.

Likhitwitayawuid K, Ruangrusi N. Structural Elucidation and Synthesis of New Components Isolated from *Piper sarmentosum* (Piperaceae). *Tetrahedron* 1987; 43:3689-94.

Stohr JR, Xiao PG, Bauer R. Isobutylamides and a new methylbutylamide from *Piper sarmentosum*. *Planta Med* 1999; 65:175–7.

Rukachaisirikul T, Siriwattanakit P, Sukcharoenphol K, Wongvein C, Rattanaweang P, Wongwattanavuch P. Suksamrarn A. Chemical Constituents and Bioactivity of *Piper sarmentosum. J Ethnopharmacol* 2004; 93:173-6.

Tuntiwachwuttikul P, Phansa P, Pootaengon Y, and Taylor WC. Chemical Constituents of the Roots of *Piper sarmentosum*. *Chem Pharm Bull* 2006; 54:149-51.

Gunston FD. Fatty Acid and Lipid Chemistry. London, Glasgow, New York: Blacky Academic & Professional 1996.

Achenbach H, Fietz W, Woerth J, Waibel R, Portecop J. Constituents of Tropical Medicinal Plants IXX GC/MSinvestigations of the Constituents of *Piper amalago-30*, New Amides of the Piperine-type. *Planta Med* 1986; 1:12-7.

Luger P, Weber M, Dung NX, Luu VT, Rang DD, Tuong DT, Ngoc PH. The Crystal Structure of 3-(4'-Methoxyphenylpropanoyl)pyrrole of *Piper lolot* C.DC., from Vietnam. *Crystal Res Tehnol* 2002; 37:627-33.

Mackeen MM, Ali AM, El-Sharkawy SH, Manap MY, Salleh KM, Lajis NH, Kawazu K. Antimicrobial and Cytotoxic Properties of Some Malaysian Traditional Vegetables. *Int J Pharmacog* 1997; 35:174-8.

Zavala MA, Perez GS, Perez GRM. Antimicrobial Screening of Some Medicinal Plants. *Phytother Res* 1997; 11:368-71.

Arias ME, Gomez JD, Cudmani NM, Isla MI. Antibacterial Activity of Ethanolic and Aqueous Extracts of *Acacia aroma* Gill. Ex Hook et Arn. *Life Sciences*, 2004; 75:191-202.

### Antibacterial activity of Sumatran Kaduk

- 32220. Houghton P, Fang R, Techatanawat I, Steventon G, Hylands PJ, 331 F. Ahmad, Department of Chemistry, Faculty of Science, Universiti
- Lee CC. The sulphorhodamine (SRB) Assay and other332
- Approaches of Testing Plant Extracts and Derived Compounds333

# 327 CURRENT AUTHOR ADDRESSES

- mail: emrizal\_atiax@yahoo.com (Corresponding author)

- ijpt.iums.ac.ir | 5
- Teknologi Malaysia, 81310 Skudai, Johor Bahru, Malahysia. Email: farediah@kimia.fs.utm.my
- for Activities Related to Reputed Anticancer Activity. *Cancer* 334 H. M. Sirat, Department of Chemistry, Faculty of Science, Universiti Teknologi Malaysia, 81310 Skudai, Johor Bahru, Malahysia. Email: hasnah@kimia.fs.utm.my
- 336 mail: nashan@kimia.is.utm.my
  328E. Atiax, Department of Chemistry, Faculty of Science, Universiti
  329 Teknologi Malaysia, 81310 Skudai, Johor Bahru, Malahysia. E-337D. Arbain, Faculty of Pharmacy, University of Andalas, Padang, West Sumatra, Indonesia. E-mail: arbain\_d@yahoo.com