

NEW DITERPENOIDS FROM THE ROOTS OF *Croton hirtus* (EUPHORBIACEAE)

(Diterpenoid-Diterpenoid Terbaharu Daripada Akar *Croton hirtus* (Euphorbiaceae))

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Abstract

The root of *Croton hirtus* was collected at Tanjung Sepat, Banting for study of its chemical compounds. This sample was dried before being ground and extracted using the methanol solvent at room temperature for three days yielding 18.25 gram of crude extract. The crude extracts obtained were analysed using thin layer chromatography (TLC) then fractionated via vacuum column chromatography (VLC) and proceed the isolation using radial chromatography (RC) to get the pure compounds. The pure compounds obtained were elucidated by nuclear magnetic resonance (NMR), Fourier Transform Infrared (FT-IR) and mass spectroscopy to confirm their structures and from that analysis, the compounds were identified as two new compounds naming (-)-5,8-dihydroxyjatrophan-3-one (**1**) and (+)-14,16,17-trihydroxykauran-1-on (**2**) with total weight 4.9 mg and 4.5 mg respectively. This two compound were reported for this time on this genus.

Keywords: *Croton hirtus*, diterpenoid, euphorbiaceae

Abstrak

Akar *Croton hirtus* dikumpulkan di Tanjung Sepat, Banting untuk mengkaji sebatian kimianya. Sampel ini dikeringkan sebelum dikisar dan diekstrak dengan menggunakan pelarut metanol pada suhu bilik selama tiga hari dan ia menghasilkan 18.25 g ekstrak mentah. Ekstrak mentah yang diperoleh dianalisis dengan menggunakan kromatografi lapisan nipis (TLC) kemudian difraksikan melalui kromatografi cecair vakum (VLC) dan pengasingan diteruskan dengan menggunakan kromatografi radial (RC) untuk mendapatkan sebatian tulen. Sebatian tulen yang diperoleh ditentukan oleh resonan magnetik nuklear (RMN), inframerah transformasi Fourier (FTIR) dan spektroskopi jisim untuk mengesahkan struktur mereka dan dari analisis itu, sebatian telah dikenal pasti sebagai dua sebatian baru yang diberi nama (-)-5,8- dihidroksijatrophan-3-one (**1**) dan (+)-14,16,17-trihidroksikauran-1-on (**2**) dengan jumlah berat masing-masing sebanyak 4.9 mg dan 4.5 mg. Kedua-dua sebatian ini dilaporkan buat kali pertama bagi genus ini.

Kata kunci: *Croton hirtus*, diterpenoid, euphorbiaceae

Introduction

Croton hirtus, also known as *Croton glandulosus*, is a shrub belonging to genus *Croton* and family Euphorbiaceae. They were distributed widespread around the world as trees, shrubs and herbs with about 1300 species. *Croton* is widely found in Malaysia and in several areas of Indonesia except Kalimantan, Sulawesi and the Sunda Islands [1]. Extensive work has been carried out on various species of *Croton*, resulting in the isolation of mainly terpenoid class of compounds, namely sesquiterpenoids [2 – 5], diterpenoids [6 – 13], and triterpenoids [14 – 17]. Non-terpenoid was isolated such as alkaloids, flavonoids, lignans and megastigmane glycosides [18]. Work on Brazilian *C. hirtus* has mainly concentrated on diterpenes [19] and essential oils [20]. So, with this background information, the aim of this present study are to analyse and characterize two new diterpenoid that have been isolated from root of genus using the spectroscopy NMR (1D and 2D), FT-IR and mass spectroscopy.

Materials and Methods

Raw materials

The roots of *C. hirtus* were collected in Tanjung Sepat, Banting, Selangor, Malaysia and herbarium specimen (SM-461) was deposited in the Herbarium of Universiti Kebangsaan Malaysia.

Extraction and isolation

Dried and ground roots of *C. hirtus* (415 g) were extracted with methanol to give a dark green extract (18.25 g). The crude extract (18.25 g) was partitioned between chloroform and water (1:1) after which the chloroform extract was further extracted with 10% *n*-hexane and 90% methanol to give methanol sub-extract. The sub-extract (3.52 g) was fractionated by using silica gel vacuum liquid chromatography eluted with increasing polarity of *n*-hexane- ethyl acetate to give ten fractions. The eluents that showed the same profile on thin layer chromatography (TLC) were combined to give four fractions (I-IV). Purification of fraction II (0.2027 g) was carried out using radial chromatography (RC) with silica gel plate of 1 mm thickness eluted with 60:40 *n*-hexane-ethyl acetate in 5% polarity increment to yield compound 1 (4.9 mg). Purification of fraction III (0.3163 g) was also carried out in a similar manner eluting with 1:1 *n*-hexane-ethyl acetate producing sub-fractions III^a-III^f. Sub-fraction (III^c= 0.0221 g) was further purified using RC with silica gel plate of 1 mm, eluting with 30:70 *n*-hexane-ethyl acetate, which resulted in the isolation of compound 2 (4.5 mg).

Results and Discussion

(-)-5,8-dihydroxyjatrophane-3-one (1), $[\alpha]_D^{25}$ -102.50 (*c* 0.40, CHCl₃), was isolated as a pinkish oily material with molecular formula C₂₀H₃₀O₃, obtained by HR-Cl-MS $[M+1]^+$ 319.2276 *m/z*. The IR spectra exhibited absorptions for hydroxyl groups (3402 cm⁻¹), cyclic aliphatic group (2922 cm⁻¹), and α,β-unsaturated carbonyl group (1709 and 1647 cm⁻¹) (Figure 1).

The ¹³C spectrum indicated the presence of 20 carbons consisting of five methyls, three methylenes, seven methines and five quaternary carbons. The most highly deshielded signal at δ_C 199.7 (C-3) integrated for carbon attributed to a carbonyl group, three signals for methane carbon at δ_C 145.3 (C-1), 125.7 (C-7) and 119.4 (C-12) indicating the presence of olefinic carbons that are linked with olefinic-quaternary carbons at δ_C 134.6 (C-2), 143.7 (C-6) and 137.7 (C-13), while the other two shielded methine carbons at δ_C 77.4 (C-5) and 67.7 (C-8) belongs to two carbinol groups in the molecule, while the other three carbons at δ_C 25.5 (C- 9), 31.6 (C-11) and 39.9 (C-14) is indicative of the aliphatic component of the cyclic system, which also contains five methyl carbons from C-16 to C-20. The DBE value of C₂₀H₃₀O₃ is 6, indicating that the compound consists of four double bonds (three olefinics and one carbonyl) and two rings of aliphatic group. The ¹H-NMR spectrum showed 18 proton resonance signals representing 30 protons. Eighteen of the signals came from the three signals, three each from methyl group at H-16 (δ_H 1.95, s), H-17 (δ_H 1.72, s), and H-20 (δ_H 1.61, s) bonded to an olefinic carbon while the other two protons at H-18 (δ_H 1.03, s) and H-19 (δ_H 1.17, s) represent the two non- equivalent methyl protons. Furthermore, six signals of methylene protons and seven signals of methine protons are from a cyclic aliphatic or jatrophane skeleton as the main structure in the molecule (Figure 2).

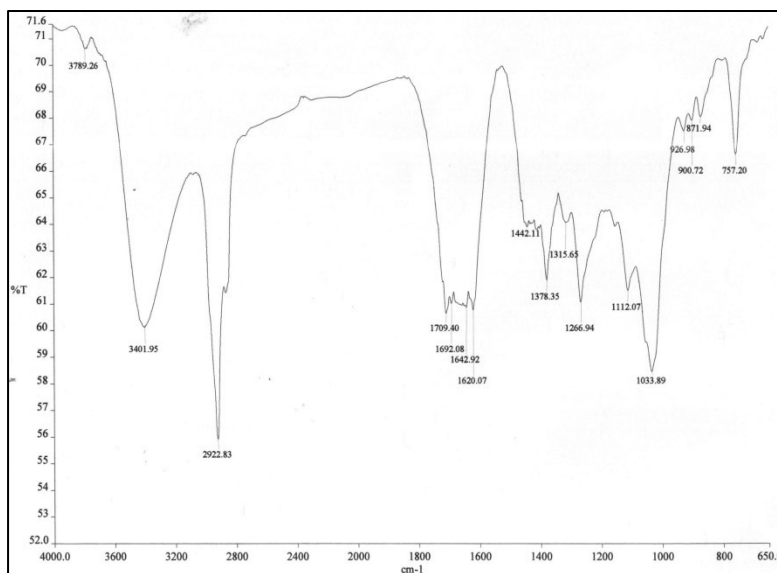


Figure 1. FTIR spectrum of (-)-5,8-dihydroxyjatrophan-3-one

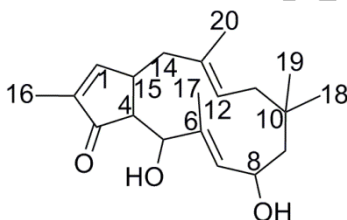


Figure 2. Chemical structure of (-)-5,8-dihydroxyjatrophan-3-one

The structure was confirmed by using 2D NMR. The HMBC experiment showed H-16 in correlation with C-1 and C-3, confirming the position of this methyl group bonded to C-2 at the cyclopentane ring and also C-20 with methylene (C-14) and olefinic-methine (C-12), confirming C-20 as the methyl carbon attached to the quaternary carbon at C-13. Meanwhile, non-equivalent dimethyl protons at H-18 and H-19 show 2J interaction with each other and the quaternary carbon at C-10. Each of the carbinol protons at H-5 and H-8 correlated with the carbonyl group at C-3 and the olefinic methine at C-7 respectively (Figure 3).

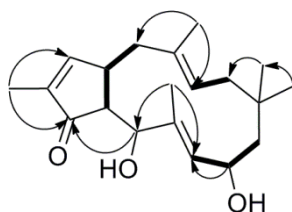


Figure 3. Significant ^1H - ^1H COSY (→) and HMBC (→) correlations of (-)-5,8-dihydroxyjatrophan-3-one

(+)-14,16,17-trihydroxykauran-1-on (**2**), $[\alpha]_D^{25} +30.00$ (c 0.40, CHCl_3) is a white needle crystal. The IR spectrum showed characteristic absorptions for hydroxyl (3244 cm^{-1}), and carbonyl (1696 cm^{-1}) (Figure 4).

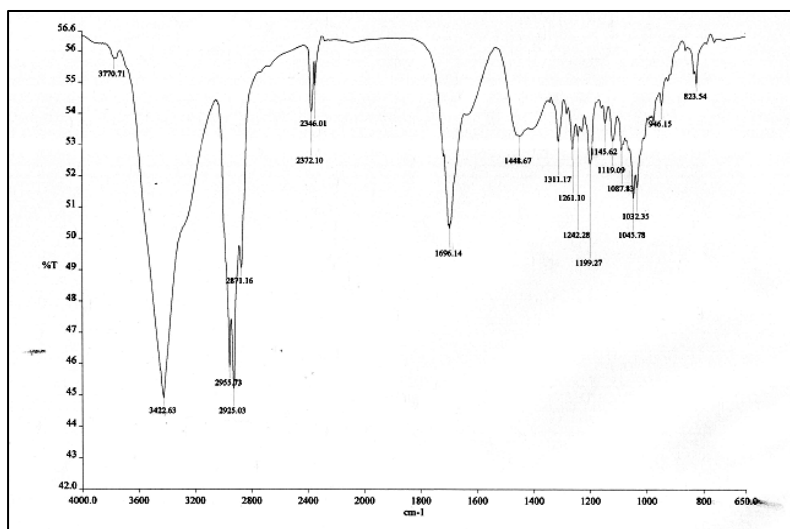


Figure 4. FTIR spectrum of (+)-14,16,17-trihydroxykauran-1-one

This compound contains 20 carbons observed from the ^{13}C NMR spectra, which shows that the compound is made up of three methyls, eight methylenes, four methines and five quaternary carbons (Table 1). The DBE value of five indicates that this compound have four cyclic systems with only one double bond. The highly deshielded carbon at δ_{C} 212.4 is a carbonyl. Meanwhile, three carbinol carbons appear δ_{C} at 76.5 (C-14), 89.5 (C-16) and 65.7 (C-17). Overlapping carbon signals with a significantly high intensity at δ_{C} 17.5-59.4 suggest that there is a long chain of cyclic aliphatic carbons. The appearance of a towering signal of protons indicates the presence of a series of long chain methylene groups forming the alkyl of the cyclic aliphatic group along with three hydroxyl groups and three methyls at the terminal of the chain. To determine methylene groups in the alkyl part, mass spectrometer was used. The MS spectral data shows that molecular weight of this compound is 336 m/z ($[\text{M}^+ + \text{H} - \text{H}_2\text{O}] = 319.2259$). Based on the discussion, this compound is identified as (+)-14,16,17-trihydroxykauran-1-one (Figure 5) with molecular formula of $\text{C}_{20}\text{H}_{32}\text{O}_4$.

Table 1. ^1H and ^{13}C NMR data (500 and 125 MHz, CDCl_3) for compounds **1** and **2**

Position	^{13}C (ppm)		^1H (ΣH , <i>mult.</i> , J in Hz)	
	1	2	1	2
1	145.3	212.4	6.30 (1H, <i>dd</i> , $J_1 = 9.5$ Hz, $J_2 = 1.1$ Hz)	-
2	134.6	37.8	-	2.40 (1H, <i>s</i>) 2.38 (1H, <i>d</i> , $J = 3.5$ Hz)
3	199.7	41.4	-	1.47 (1H, <i>m</i>) 1.19 (1H, <i>d</i> , $J = 4.0$ Hz)
4	28.1	33.2	1.53 (1H, <i>d</i> , $J = 6.2$ Hz)	-
5	77.4	53.2	4.13 (1H, <i>d</i> , $J = 6.2$ Hz)	1.27 (1H, <i>dd</i> , $J_1 = 10.2$ Hz, $J_2 = 6.9$ Hz)
6	143.7	40.2	-	1.87 (1H, <i>m</i>) 1.12 (1H, <i>dd</i> , $J_1 = 12.8$ Hz, $J_2 = 3.3$ Hz)

Table 1 (cont'd). ^1H and ^{13}C NMR data (500 and 125 MHz, CDCl_3) for compounds **1** and **2**

Position	^{13}C (ppm)		^1H (ΣH , <i>mult.</i> , <i>J</i> in Hz)	
	1	2	1	2
7	125.7	41.1	5.20 (1H, <i>dd</i> , $J_1 = 8.0$ Hz, $J_2 = 1.1$ Hz)	2.09 (1H, <i>dd</i> , $J_1 = 11.6$ Hz, $J_2 = 3.5$ Hz) 1.90 (1H, <i>m</i>)
8	67.7	56.0	5.26 (1H, <i>dd</i> , $J_1 = 8.0$ Hz, $J_2 = 6.6$ Hz)	-
9	25.5	59.4	2.09 (1H, <i>m</i>) 0.78 (1H, <i>m</i>)	1.86 (1H, <i>m</i>)
10	29.8	36.9	-	-
11	31.6	18.2	2.31 (1H, <i>m</i>) 2.25 (1H, <i>m</i>)	1.66 (1H, <i>dd</i> , $J_1 = 11.6$ Hz, $J_2 = 3.5$ Hz) 1.50 (1H, <i>m</i>)
12	119.4	43.4	4.70 (1H, <i>ddd</i> , $J_1 = 9.5$ Hz, $J_2 = 4.0$ Hz, $J_3 = 1.1$ Hz)	2.14 (1H, <i>d</i> , $J = 11.6$ Hz) 1.57 (1H, <i>m</i>)
13	137.7	41.5	-	2.61 (1H, <i>t</i> , $J = 6.5$ Hz)
14	39.9	76.5	2.23 (1H, <i>m</i>) 1.70 (1H, <i>m</i>)	4.48 (1H, <i>s</i>)
15	35.8	46.4	1.19 (1H, <i>d</i> , $J = 9.5$ Hz)	2.04 (1H, <i>d</i> , $J = 12.0$ Hz) 1.69 (1H, <i>m</i>)
16	12.1	89.5	1.95 (3H, <i>d</i> , $J = 1.1$ Hz)	-
17	10.3	65.7	1.72 (3H, <i>d</i> , $J = 1.5$ Hz)	3.83 (1H, <i>d</i> , $J = 12.0$ Hz) 3.64 (1H, <i>d</i> , $J = 12.0$ Hz)
18	16.1	21.3	1.03 (3H, <i>s</i>)	0.85 (3H, <i>s</i>)
19	29.3	33.3	1.17 (3H, <i>s</i>)	0.87 (3H, <i>s</i>)
20	15.4	17.5	1.61 (3H, <i>d</i> , $J = 1.1$ Hz)	1.25 (3H, <i>s</i>)

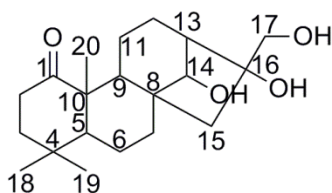


Figure 5. Chemical structure of (+)-14,16,17-trihydroxykauran-1-one

The structure of the compound was further confirmed using 2D NMR experiments including HMBC and COSY. From the HMBC experiments, the position of carbonyl carbon (C-1) was proven by correlations of C-2, C-5 and C-20 toward C-1. The proton H-14 in correlation with C-9 confirmed the position of this carbinol group bonded to C-8 and C-13 as branches to the other two carbinol groups at C-16 and C-17. COSY spectrum shows correlations between four proton methylene (H-2, H-3, H-6, H-7, H-11 and H-12) that forms the unsubstituted aliphatic component of the main ring skeleton in the molecule (Figure 6).

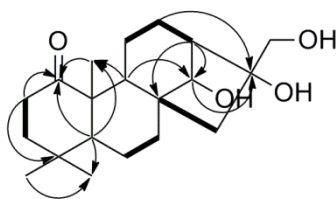


Figure 6. Significant ¹H-¹H COSY (-) and HMBC (→) correlations of (+)-14,16,17-trihydroxykauran-1-on

Chemotaxonomic significance

The genus *Croton* is the second largest genus of Euphorbiaceae with over 1300 species, and over 100 species from this genus have been studied phytochemically, resulting in the isolation and identification of more than 200 various types of diterpenoids [21,22] comprising of different skeleton types such as clerodane (most common), kaurane, cembranoid, halimane, isopimarane, labdane, phorbol and trachylobane [23].

In this study, a new jatrophane and kaurane-type diterpenoid, (-)-5,8-dihydroxyjatrophane-3-one (**1**) and (+)-14,16,17-trihydroxykauran-1-on (**2**) respectively, was obtained from the roots of *C. hirtus*. The occurrence of diterpenoids in Euphorbiaceae is not unusual. Compounds with jatrophane-type skeleton was isolated from the genus *Jatropha* for the first time, and widely identified in the other genera (Euphorbiaceae) as well. The roots and rhizomes of several *Jatropha* species was reported to contain jatrophane, hydroxyjatrophane A-C, 9 β -hydroxyisabellione, 9 β -13 α -hydroxyisabellione, citlaltirone, jatrophene, and japodagrone [24 – 32]. Meanwhile, four jatrophane polyesters were isolated from the aerial parts of the East Asian weed *Euphorbia mongolica* Prokh [33].

In 2010, Hegazy et al. [34] identified the new guyonianin E and F from the aerial parts of *Euphorbia guyoniana*. Meanwhile, euphodendrophane A (Euph A) and euphodendrophane B (Euph B) were isolated from the jatrophane fraction of *Euphorbia dendroidea* L. [35]. Since only these two genera were reported to biologically synthesize diterpenoid, thus chemotaxonomically, it can be assumed that the genus *Croton* is related to genera *Jatropha* and *Euphorbia*. Despite the large number of kaurane-type diterpenoids isolated from this genus, some species from other families also show the presence of this diterpenoid subclass. Four ent-kaurane diterpenoid glycosides, cussivantosides A–D were isolated with other known compounds from the dried leaves of *Cussonia vantsilana* Baker (Araliaceae) [36]. Other ent-kaurane derivatives such as ent-16-kauren-19-oic acid (kaurenoic acid) methyl ester, grandiflorenic acid methyl ester, ent-16-kauren-19-al, 16 α ,17-epoxy-15 α -angeloyloxykauran-19-oic acid methyl. Grandiflorenic acid and 15- α -angeloyloxy-16,17- α -epoxy- ent-16-kauren-19-oic acid were present only in the Asteraceae plant *Smilax sonchifolius* and *Smilax siegesbeckii* [37].

Conclusion

Two diterpenoids; (-)-5,8-dihydroxyjatrophane-3-one (**1**) and (+)-14,16,17-trihydroxykauran-1-on (**2**) were successfully isolated from the methanolic extract of *Croton hirtus*. To the best of our knowledge, this is the first report for both compounds from this species. Both of compounds can be used to carry out biological activities because this species was known as many useful in traditional medicine.

Acknowledgement

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