

Avances en Química

ISSN: 1856-5301 clarez@ula.ve

Universidad de los Andes Venezuela

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Avances en Química, vol. 8, núm. 3, septiembre-diciembre, 2013, pp. 131-138
Universidad de los Andes
Mérida, Venezuela

Available in: http://www.redalyc.org/articulo.oa?id=93330145008



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www.saber.ula.ve/avancesenquimica Avances en Química, 8(3), 131-138 (2013)

Artículo científico



Naphthopyranones from Rhizomes of Paepalanthus diffissus

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Recibido: 04/11/2013 **Revisado**: 06/12/2013 **Aceptado**: 08/12/2013

Resumen

Del extracto en diclorometano obtenido de los rizomas de *Paepalanthus diffissus* Moldenke (Eriocaulaceae) fueron aisladas las naftopiranonas bioactivas (+)-semi-vioxantina [3] y vioxantina [8]. Estos compuestos fueron caracterizados en base a estudios espectroscópicos, incluyendo experimentos de RMN uni- y bi-dimensionales. La revisión de la literatura indica que estos metabolitos son encontrados frecuentemente en hongos y líquenes, pero son compuestos raros en plantas superiores. La presencia de (+)-semi-vioxantina en plantas con flores, es descrita aquí por primera vez.

Palabras clave: Eriocaulaceae; Paepalanthus; naftopiranonas; (+)-semi-vioxantina; vioxantina

Abstract

From a dichloromethane extract obtained of rhizomes of *Paepalanthus diffissus* Moldenke (Eriocaulaceae) were isolated the bioactive naphthopyranones (+)-semi-vioxanthin [3] and vioxanthin [8]. These compounds were characterized on the basis of spectroscopic studies, including 1D- and 2D-NMR experiments. The literature review indicated that these metabolites are frequently found in fungi and lichens, but they are rare compounds in higher plants. Presence of (+)-semi-vioxanthin in flowering plants is here described by first time.

Keywords: Eriocaulaceae; Paepalanthus; naphthopyranones; (+)-semi-vioxanthin; vioxanthin

Introduction

Eriocaulaceae is a small family of flowering plants, which contains about 1400 species grouped in 11 genera¹, although the most recent molecular studies tends to reduce the number of genera to ten². Most of the species included in this family are distributed in mountainous regions of South America, especially in the rocky savannas of Brazil and the õtepuisö (table mountains) of Guyana and Venezuela; only a few species assembled in five genera, extend their habitat into temperate regions of North America and Europe, tropical Africa and eastern Asia³. *Paepalanthus* represents the largest genus of the family, with over 450 species distributed disjunctly in neotropical South America, occidental Africa and Madagascar⁴.

The genus *Paepalanthus* is well documented as a good source of secondary metabolites such as flavonoids⁵⁻¹⁰, naphthopyranones⁹⁻¹⁴, naphthoquinones¹⁵ and caffeic acid derivatives⁶. Currently, some of these compounds possess a notable pharmacological interest due to their proven biological activity; for example, it has been reported that the naphthoquinone 5-methoxy-3,4-dehydroxanthomegnin [1],

isolated from *Paepalanthus latipes*, possess immune modulatory effects on nitric oxide production on LPS-stimulated macrophages¹⁶, in addition to cytotoxic, antitumoral, antioxidant and anti-*Helicobacter pylori* activity¹⁵⁻¹⁸. In the same way, the isocoumarin paepalantine [2], a metabolite present in different extracts of *Paepalanthus bromelioides* and *P. vellozioides*, exhibits a wide range of biological activities, including antioxidant^{19,20}, anti-inflammatory²¹, antimicrobial²²⁻²³ cytotoxic^{12,24}, genotoxic²⁵ and mutagenic^{12,26} activity. Also several flavonoids identified in *Paepalanthus ssp.* have shown to possess biological properties such as antioxidant potential⁸, mutagenicity⁹ or antimycobacterial activity¹⁰.

5-Methoxy-3,4-dehydroxanthomegnin [1]

Paepalantine [2]

In the light of the foregoing, as part of our continuing phytochemical studies on medicinal plants of Venezuela@s Andean, in this paper we describe the isolation of two naphthopyranones from rhizomes of *Paepalanthus diffissus* Moldenke, a species found commonly in Andean moors. These naphthopyranones were identified as

(+)-semi-vioxanthin [3] and vioxanthin [8], known antifungal antibiotics often isolated from fungi²⁷⁻²⁹ and lichens³⁰; presence of vioxanthin in flowering plants is up to now limited to a few species of the genus *Paepalanthus* ^{20,31}, but to the best of our knowledge, (+)-semi-vioxanthin has so far not been found in higher plants.

(+)-Semi-Vioxanthin [3]:
$$R_1=R_2=H$$

(+)-Semi-Vioxanthin 9-Acetate [4]: $R_1=Ac;\ R_2=H$

(+)-Semi-Vioxanthin, 9,10-Diacetate [5]: $R_1 = R_2 = Ac$

(+)-Semi-Vioxanthin 9-Methyl Ether [6]: $R_1 = CH_3$; $R_2 = H$

(+)-Semi-Vioxanthin 10-Methyl Ether [7]: $R_1 = H$; $R_2 = CH_3$

$$\begin{array}{c} \mathbf{H_{3}C_{H_{11}}} \\ \mathbf{H_{3}C_{H_{11}}} \\ \mathbf{O} \\ \mathbf{$$

Vioxanthin [8]

Materials and methods

General

Melting points were determined using a Fisher-Johns apparatus and they are uncorrected. Optical rotations were measured on a 60Hz-Steeg & Reuter G.m.b.H. polarimeter using CHCl₃ as solvent. UV spectra were obtained in a Perkin-Elmer spectrophotometer, Lambda 3B, using quartz cells with 1cm thick and methanol (Merck-Uvasol) as solvent. IR measurements were obtained on a Perkin-Elmer FT-1725X spectrophotometer as KBr pellets. 1D and 2D NMR spectra in CDCl₃ were acquired using a Bruker-Avance DRX-400 instrument, operating at 400 MHz for ¹H and 100 MHz for ¹³C. Mass spectra were recorded on a Hewlett-Packard Mass Spectrometer, model 5890 (70 eV). TLC were developed on 0.25 mm layers of silica gel PF 254 (Merck); spots were visualized using UV light (254 and 365 nm) and subsequently by spraying with a mixture v/v CH₃COOH-H₂O-H₂SO₄ (20:4:1) and then heating with air flow at 100°C for few minutes. VCC was performated with silica gel Merck 60 (63-200 m, 70-230 mesh). Size-exclusion chromatography columns were packed with Sigma Sephadex LH-20.

Plant material

Plant material (rhizomes) was collected at õPáramo de San José de Acequias, Municipio Campo Elías, Estado Mérida, Venezuelaö. Species was identified as *Paepalanthus diffissus* Moldeke by Eng. Juan Antonio Carmona Arzola,

Department of Pharmacognosy and Organic Medicaments, Faculty of Pharmacy and Bioanalysis, University of Los Andes (ULA); a *voucher specimen* (Amaro-Luis *et al.*, N° 2342) was deposited at the Herbario MERF of this faculty.

Extraction

Rhizomes of *Paepalanthus diffissus* (ca 1.80 Kg) were air-dried, ground and exhaustively extracted with hexane and then with dichloromethane in a soxhlet. The solutions obtained were filtered and concentrated *in vacuum* on a rotary evaporator, to afford respectively, 54.8 g and 60.4 g of crude extracts.

Isolation and identification of the constituents

The dichloromethane extract was preadsorbed on silica gel and chromatographed (VLC) over silica gel 60, eluting with hexane, dichloromethane and EtOAc in mixtures of increasing polarity. Fifty-four (54) fractions of 500 mL were collected, concentrated *in vacuum*, and combined according to the TLC characteristics to afford twelve major fractions (A-L).

(+)-Semi-vioxanthin [3]: From combined fraction E [22-26, eluted with hexane-CH₂Cl₂ (7:3)] precipitated an apple green solid residue (≅ 5.8 g), which was partially purified by flash chromatography (hexane-EtOAc 4:1); crystallization from mixtures EtOAc/hexane provided

pure yellow needles (\cong 353 mg) detected in TLC plates as a homogeneous green-yellow spot, m.p. = 192-193°C, [α]_D: +7.3° (CHCl₃). UV, λ_{max} (nm): 263, 379 (CH₃OH); 228, 243, 404 (CH₃OH + AlCl₃). IR (KBr), ν_{max} (cm⁻¹): 3384 (-OH), 2978-2850 (C-H), 1650 (C=O), 1582 (C=C), 1158 and 1124 (C-O), 844 and 598 (=C-H). ¹H NMR (Table 1). ¹³C NMR (Table 2). EI-MS: m/z (%) 275 (60.12) [M⁺ +1], 274 (42.31) [M⁺], 256 (47.31), 229 (27.72), 200 (15.81), 186 (23.18), 158 (16.03), 141 (22.20), 129 (41.63), 115 (49.82), 102 (33.64), 77 (32.95), 62 (22.31), 43 (37.75).

Vioxanthin [8]: Combined fraction J [42-46, eluted with CH₂Cl₂-EtOAc (7:3)] was chromatographed on a Sephadex-LH20 column using as eluent a mixture of hexane-CH₂Cl₂-CH₃OH (0.5:4:0.5) which allowed to obtain twenty six fractions. From fraction N° 6 precipitated a crystalline solid as pale yellow needles (10.2 mg.); m.p. = 196-198°C (descomposition); [α]_D: + 4.6° (CHCl₃). UV, λ_{max} (nm): 270, 387 (CH₃OH); 232, 276, 418 (CH₃OH + AlCl₃). IR (KBr), ν_{max} (cm⁻¹): 3398 (-OH), 2976-2848 (C-H), 1632 (C=O), 1584 (C=C), 1128 and 1092 (C-O), 856 and 568 (=C-H). ¹H NMR (Table 1). ¹³C NMR (Table 2). EI-MS: m/z (%) 548 (4.25) [M⁺+2], 546 (18.42) [M⁺], 531 (57.30), 517 (62.25).

Acetylation of (+)-semi-vioxanthin [3]

Compound [3] (210 mg) was dissolved in pyridine (14 mL) and treated with Ac_2O (35 mL) at room temperature overnight. Cold water was added to the reaction mixture and immediately it was extracted with CH_2Cl_2 . The organic layer was washed in successive stages with aqueous solutions of HCl (10% ν/ν), NaHCO₃ (2%) and water, dried on MgSO₄ and evaporated to yield a solid residue (142 mg) that showed in TLC two spots. Separation was carried out on a silica gel column using as eluent hexane- CH_2Cl_2 (1:4) to furnish compounds [4] (36 mg) and [5] (22 mg).

- (+)-Semi-vioxantin-9-monoacetate **[4]**: pale yellow flakes; m.p. = 186-188°C. IR (KBr), $v_{max.}$ (cm⁻¹): 3424 (-OH), 1770 (C=O), 1648 (C=O), 1623 (C=C), 1210 (C-O), 858 (=C-H). ¹H NMR (Table 1). ¹³C NMR (Table 2).
- (+)-Semi-vioxantin-9,10-diacetate **[5]**: pale yellow needles; m.p. > 200°C. IR (KBr), $\nu_{max.}$ (cm⁻¹): 1770 (C=O), 1716 (C=O), 1632 (C=O), 1580 (C=C), 1212 (C-O), 860 (=C-H). 1 H NMR (Table 1). 13 C NMR (Table 2).

Methylation of (+)-semi-vioxanthin [3]

Compound [3] (124 mg) was treated with excess ethereal CH_2N_2 and solution was left standing overnight in a refrigerator at 4°C. Evaporation of ether yielded a colorless oil that was chromatographed on a silica gel column eluted

- with hexane-CH₂Cl₂ (1:4), to obtain pure TLC compounds **[6]** (4 mg) and **[7]** (23 mg).
- (+)-Semi-vioxanthin-9- methyl ether **[6]**: yellow solid; m.p. = 142-144 °C. ¹H NMR (Table 1).
- (+)-Semivioxanthin-10-methyl ether [7]: colorless oil. IR (KBr), ν_{max} . (cm⁻¹): 3302 (-OH), 2848 (C-H), 1712 (C=O), 1642 and 1576 (C=C), 1123 (C-O). ¹H NMR (Table 1). ¹³C NMR (Table 2).

Results and Discussion

(+)-Semi-vioxanthin [3] was obtained as pale yellow needles [m.p. = 192-193 °C, $[\alpha]_D$: +7.3° (CHCl₃)]. The presence in its EIMS of an ion molecular peak at m/z: 274 in conjunction with NMR data, allowed to establish the molecular formula C₁₅H₁₄O₅. Its IR spectrum showed absorption bands of hydroxyl groups (3384 cm⁻¹), a carbonyl group (1650 cm⁻¹) and C-O (1158 and 1124 cm⁻¹) and aromatic C-H bonds (1582, 844 and 598 cm⁻¹). Its ultraviolet spectrum exhibited maxima at 263 and 379 nm. The ¹H NMR spectrum of [3] (Table 1) indicated the presence in the molecule of three aromatic protons, a methoxyl group, dos hydroxyl protons and six aliphatic hydrogens that constitute a methylene, an oxymethine and a secondary methyl group. At the same time its ¹³C NMR (Table 2) shows, apart of ten signals typical of aromatic carbons, a peak assignable to a carbony group and four sp³ aliphatic carbons signals.

Comparing and contrasting the above information with the data derived from the analysis of the 2D-NMR spectra it was possible to conclude that [3] is a naphthopyranone with a lactone moiety, similar to paepalantine [2]. In effect, the naphthalene unit was identified by the presence of two aromatic \tilde{o} meta \ddot{o} -coupled protons [doublets at δ_H 6.54 and $\delta_{\rm H}$ 6.51; J = 2.35 Hz (H-6 and H-8); HMQC: H-6 \leftrightarrow C-6 $(\delta_C 99.5, =\underline{CH}); H-8 \leftrightarrow C-8 (\delta_C 101.6; =\underline{CH}); HMBC: C-7 \leftrightarrow$ H-6 \leftrightarrow C-8 and C-6 \leftrightarrow H-8], which characterize a 1,2,3,5tetrasubstituted benzene ring [A] with two substituents identified as a hydroxyl on C-9 (acute singlet at $\delta_{\rm H}$ 9.45, -OH) and a methoxy group on C-7 [δ_H 3.87, s, (H-12)]; correlations in HMBC spectra (Fig 1) confirmed the ubication of these substituents [HMBC: C-8 ↔ OH ↔ C-9 $(\delta_C 158.6; =CH-O-)/C-9a \leftrightarrow H-8 \text{ and } H-12 \leftrightarrow C-7$ $(\delta_C \ 162.7; = CH-O-)$]. The other two A-ring substitute carbons [δ_C 140.5; =C< (C-5a) and δ_C 108.4; =C< (C-9a)] conform the link between the second condensed pentasubstituted bencene ring [B], which also possesses two quaternary carbons bridge $[\delta_C 133.2; =C < (C-4a)]$ and $\delta_{\rm C}$ 99.2; =C< (C-10a)] that configure the fusion with a third aliphatic cycle [C]. The fifth substituent is a chelated hydroxyl group $[\delta_H \ 13.74, \ (-O\underline{H})]$ located at C-10

Compound	[3]	[4]	[5]	[6]	[7]	[8]*
H-3	4.73 (m)	4.71 (m)	4.74 (m)	4.70 (m)	4.62 (m)	4.75 (m)
H-4	2.96(m)	2.95 (m)	2.96 (m)	2.97(m)	2.97(m)	3.01 (m)
H-5	6.85(s)	6.93 (s)	7.44(s)	6.85 (s)	7.20(s)	6.95 (s)
Н-6	6.54 (d) J = 2.35	6.87 (d) J = 2.40	6.95 (d) J = 2.35	6.57 (d) J = 2.33	6.60 (d) J = 2.33	6.70 (s)
H-8	6.51 (d) J = 2.35	6.74 (d) J = 2.40	6.82 (d) J = 2.35	6.46 (d) J = 2.33	6.55 (d) J = 2.33	-
H-11	1.54 (d) J = 6.32	1.53 (d) J = 6.32	1.47 (<i>d</i>) $J = 6.20$	1.52 (d) J = 6.30	1.51(d) J = 6.30	1.56 (d) J = 6.04
H-12	3.87(s)	3.90(s)	3.89(s)	3.90(s)	3.87(s)	3.84 (s)
C-9 (O <u>H</u>)	9.45 (s)	-	-	-	9.85 (s)	9.70(s)
C-10 (O <u>H</u>)	13.74 (s)	12.90 (s)	-	13.20 (s)	-	13.79 (s)
H-1ø	-	-	-	3.99(s)	4.12(s)	-
H-2ø	-	2.38(s)	2.39(s)	-	-	-
H-2ö	-	-	2.46 (s)	-	-	-

Table 1: $^1\text{H-NMR}$ (CDCl3, 400 MHz) Chemical Shifts (δ_H)

^{*} Only shows δ_H of a monomer unit, since δ_H of second monomer unit are identical

Table 2. C-INVIX (CDCR, 100 MILZ) Chemical Simils (or	CDCl ₃ , 100 MHz) Chemical Shifts (δ_C)	100 MHz)	(CDCl ₃ .	¹³ C-NMR	Table 2 : 1
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Compound	[3]	[4]	[5]	[7]	[8]*
C-1	171.6 (s)	170.3 (s)	169.9 (s)	162.8 (s)	171,1 (s)
C-3	76,5 (<i>d</i>)	76.1 (<i>d</i>)	74.6 (<i>d</i>)	74.5 (d)	76,0 (<i>d</i>)
C-4	34,7 (t)	35.1 (t)	35.9 (t)	36.3 (t)	34,2 (t)
C-4a	133,2 (s)	134.3 (s)	136.1 (s)	136.1 (s)	132,3 (s)
C-5	116,1 (<i>d</i>)	115.6 (d)	114.4 (d)	120.8 (d)	115,6 (d)
C-5a	140,5 (s)	140.8 (s)	139.5 (s)	139.7 (s)	134,5 (s)
C-6	99,5 (d)	104.9 (d)	104.5 (d)	98.6 (<i>d</i>)	97,5 (s)
C-7	162,7 (s)	162.2 (s)	159.9 (s)	162.0 (s)	162,3 (s)
C-8	101,6 (d)	111.8 (d)	122,9 (<i>d</i>)	102.6 (d)	107,7(s)
C-9	158,6 (s)	150.0 (s)	148.0(s)	157.4 (s)	154,9 (s)
C-9a	108,4 (s)	112.0 (s)	116.5 (s)	112.6 (s)	108,0 (s)
C-10	163,0 (s)	161.0 (s)	160.1 (s)	161.6 (s)	160,9 (s)
C-10a	99,2 (s)	101.5 (s)	113.2 (s)	109.5 (s)	98,8 (s)
C-11	20,8 (q)	20. 8 (q)	20.7(q)	20.7(q)	20,2 (q)
C-12	55,5 (q)	55.7 (q)	55.8 (q)	55.5 (q)	55,4 (q)
C-1ø	-	171.2 (s)	169.2 (s)	64.4(q)	-
C-2ø	-	21.3 (q)	21.2 (q)	-	-
C-1ö	-	-	169.6 (s)	-	-
C-2ö	-	-	21.6 (q)	-	-

^{*} Only shows δ_C of a monomer unit, since δ_C of second monomer unit are identical

(cross peaks in HMBC among OH and C-10, C-9a and C-10a); the chelation requires that the lactone carbony group in ring [C] is situated on C-1. The gross molecular structure is completed with the presence en ring [C] of a methylene group [$\delta_{\rm H}$ 2.96, m, (H-4); HMQC: H-4 \leftrightarrow C-4 ($\delta_{\rm C}$ 34.7; >CH₂)] adjacent to a oxymethine [$\delta_{\rm H}$ 4.73, m, (H-3); HMQC: H-3 \leftrightarrow C-3 ($\delta_{\rm C}$ 76.5; >CH-O-)] that supports to the

secondary methyl [δ_H 1.54; d, J = 6.32 Hz (H-11); HMQC: H-11 \leftrightarrow C-11 (δ_C 20.8; - CH₃)]. Many other correlations in HMBC spectrum (Fig. 1) ensure the lineal condensation of the rings A/B/C and, unequivocally, confirm the position of substituents in [B] and [C] rings [HMBC: C-6 \leftrightarrow H-5 \leftrightarrow C-9a \leftrightarrow H-6 \leftrightarrow C-5 \leftrightarrow H-4 \leftrightarrow C-3 \leftrightarrow H-11 \leftrightarrow C-4 \leftrightarrow H-5 \leftrightarrow C-4a \leftrightarrow H-4 \leftrightarrow C-10a \leftrightarrow H-5 \leftrightarrow C-5a].

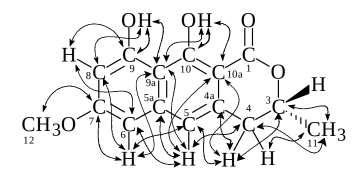
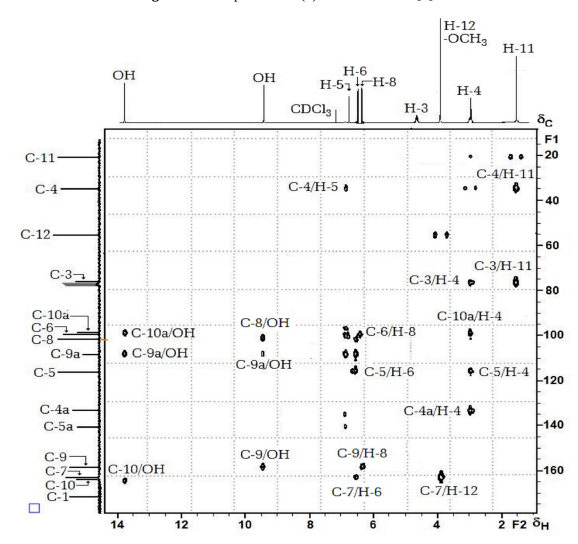


Fig. 1: HMBC Spectrum of (+)-Semivioxanthin [3]



The preceding analysis allows us to conclude that the structure of compound under study corresponds to 9,10dihydroxy-7-methoxy-3-methyl-1-oxo-1H-naphto [2,3-c] pyran. This structure possesses a single chiral center (C-3) whose possible configurations (R o S) characterizes two enantiomeric molecules. Both molecules have been previously described as natural products: levorotatory enantiomer, named (-)-semi-vioxanthin ($[\alpha]_D$: -10.6°), possesses configuration õSö in C-3 and it was obtained from cultures of Cryptosporiopsis abietina³², a coelomycete endophytic fungus isolated from Chamaecyparis obtusa; this enantiomer exhibited absicic activity against Hinoki cypress leaves and, in an inhibited antifungal test, spore germination Cladosporum herbarum. The dextrorotatory isomer, (+)-semi-vioxanthin (configuration R in C-3) is a rare natural compound first isolated from the fungus Penicillium citreo-viride²⁸ and subsequently also found in soil³³ and marine-derived fungi³⁴. Its properties as an antifungal antibiotic and as a tumor necrosis factor- α regulator have been documented³³⁻³⁴. Compound described in this study is dextrorotatory and consequently its configuration in C-3 is R with the secondary methyl group α -oriented; it is obvious that the same was clearly identified as (+)-semi-vioxanthin [3].

Identification of (+)-semi-vioxanthin [3] was also confirmed by obtaining some derivatives. Thus, on acetylation with Ac₂O/Py, compound [3] gave two acetyl derivatives, which were characterized by spectroscopic data (Table 1) as monoacetate [4] [IR, v_{max} : 1770cm⁻¹ (O=C-O-); ¹H NMR: substitution of OH singlet at δ_H 9.45, by a new 3H singlet at δ_H 2.38, s, $O=C-C\underline{H}_3$ (H-2 ϕ); ¹³C NMR: δ_C 171.2; $O=\underline{C}$ -O- (C-1 ϕ) and δ_C 21.3; O=C- $\underline{C}H_3$ (C-2ø)] and as diacetate [5] [IR, v_{max} : 1770 and 1716 cm⁻¹ (O=C-O-); ¹H NMR: substitution of both OH signals by two new singlets at $\delta_{\rm H}$ 2.39 and $\delta_{\rm H}$ 2.46; s, O=C-C $\underline{\rm H}_3$ (H-2ø and H-2ö); ^{13}C NMR: δ_{C} 169.2 and δ_{C} 169.6; O=<u>C</u>-O- (C-1ø and C-1ö) and δ_C 21.2 and δ_C 21.6; O=C- $\underline{C}H_3$ (C-2ø and C2ö)]. Treatment of 3 with CH₂N₂/ether gave two dimethyl ether derivatives: (+)-Semi-vioxanthin-9 methyl ether [6] $[\delta_H \ 3.99; \ s, \ -OCH_3 \ (H-1\emptyset)]$ and (+)-semivioxanthin-10methyl ether [7] [δ_H 4.12; s, -OC \underline{H}_3 (H-1 \emptyset) and δ_C 64.4; $O=C-CH_3(C-1\emptyset)$.

Vioxanthin **[8]:** yellow needles observed in TLC plates as a orange spot; m.p. = 196-198°C (descomposition); $[\alpha]_D$: + 4.6° (CHCl₃). Comparison of its ¹H NMR spectrum (Table 1) with that of (+)-semi-vioxanthin **[3]** revealed only two notable changes: The absence of singlet attributed to H-8 and the transformation of H-6 doublet in a sharp singlet $[\delta_H 6.70; s, (H-6/H-6\emptyset)]$; these changes indicate that C-8 is a

quaternary carbon (=C<) and that ring A is pentasubstituted. In accordance with the foregoing data, the ^{13}C NMR spectrum (Table 2) displays, in the DEPT-90, only two peaks assignable to aromatic methines [δ_{C} 115.6 and δ_{C} 97.5; =CH (C-5 and C-6)], and consequently, it is also observed a new peak typical of a quaternary sp² carbon [δ_{C} 107.7; =C< (C-8/8ø)]. The detection in its EI-MS of an ion molecular peak at m/z: 546 [M $^{+}$] allows us to conclude that this compound is a symmetric dimer of (+)-semi-vioxanthin [3], with a bridge C-8/C-8øbetween the two monomer units; the symmetry of the structure justifies the not duplicity of the NMR signals. The above analysis led to the structure [8], named in the literature as vioxanthin. This naphthopyranone has been previously isolated of several fungi²⁷⁻²⁹ and lichens³0 and also it has been found in other *Paepalanthus* species³1. Its wide range of biological activities is well documented in the literature 14,20,35,36 .

Acknowledgments

The authors are grateful to CDCHTA-ULA (C-1808-12-08-A) and to Venezuelan Ministry of Popular Power for Science, Technology and Innovation (MCTI), õScience Missionö Program (Grant Nº 2008000937), for financial support. Thank are also due to Eng. Juan Carmona Arzola, Department of Pharmacognosy and Organic Medicaments, Faculty of Pharmacy and Bioanalysis, University of Los Andes (ULA) for identification of plant material.

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