# A REARRANGED CHAMIGRENE DERIVATIVE AND ITS POTENTIAL BIOGENETIC PRECURSOR FROM A NEW SPECIES OF THE MARINE RED ALGAL GENUS *LAURENCIA* (RHODOMELACEAE)

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(Received 30 July 1984)

Key Word Index—Laurencia, Rhodomelaceae, Rhodophyta, marine algae, halogenated chamigrene, sesquiterpenoids

Abstract—The structures of a novel rearranged sesquiterpenoid and a biogenetically-related chamigrene derivative have been determined by combined spectral and chemical methods. These sesquiterpenoids were components of an undescribed *Laurencia* species, and each was toxic toward the damselfish *Pomacentrus coeruleus* 

## INTRODUCTION

Over the past decade, red seaweeds of the genus Laurencia have become of considerable interest based upon their unprecedented synthesis of a wide diversity of halogenated sesquiterpenes, diterpenes, acetogenins and aromatic compounds [1, 2] In connection with our continuing investigations of the chemical adaptations of tropical marine algae, we recently encountered an apparently undescribed species of the red seaweed Laurencia along the east coast of central Florida We wish to report that this Laurencia species produced halogenated sesquiterpenoids mainly of the chamigrene class Described here, in detail, is the isolation and structure elucidation of two new sesquiterpenoids, 1 and 3 The alcohol 1 possesses a new rearranged carbon skeleton thus adding further biosynthetic capacity to this interesting genus of marine algae Alcohol 3, a typical bromochamigrene, possesses the structural features required to act as a biosynthetic precursor to 1

Although considerable chemical study has been directed toward the halogenated metabolites from Laurencia species, few studies have provided evidence of the biological significance of these molecules [3] We believe haloterpenoid and haloacetogenin synthesis represents a defensive adaptation against the abundant herbivore populations in tropical habitats Although still under investigation, we find the alcohols 1 and 3 to illustrate significant ichthyotoxicity toward reef fishes

# **RESULTS AND DISCUSSION**

The sesquiterpenoid alcohols 1 and 3 were isolated by conventional chromatographic methods from the CHCl<sub>3</sub>-MeOH (2 1) extract of the alcohol preserved algae Alcohol 1, the major metabolite (5% organic extract) analysed for  $C_{15}H_{22}O_2BrCl$  by its combined HRMS and  $^{13}C$  NMR (Table 1) spectral features Absorption at 3500 cm $^{-1}$  in the IR spectrum of 1 indicated the compound was an alcohol Indeed, treat-

ment of 1 under standard acetylation conditions yielded the monoacetate 2

Consideration of the unsaturation inherent in 1, taken in combination with  $^{13}$ C NMR data, showed that the molecule was bicarbocyclic The  $^{13}$ C NMR spectrum showed the presence of an exomethylene functionality [154 8 (s), 104 1 (t)], a trisubstituted epoxide [61 0 (s), 59 4 (d)], a secondary, bromine-bearing carbon [64 8 (d)], a quaternary chlorine-bearing carbon [70 1 (s)] and a secondary alcohol carbon [68 7 (d)]

At 360 MHz the majority of the proton bands in 1 could be inter-related by spin-decoupling (Table 1) In  $C_6D_6$  most resonances were even more resolved and complete assignments were made (Experimental) Three separate spin systems, C-1-C-2, C-4-C-5 and C-8-C-9-C-10 were readily discerned The substituents, their coupling constants (characteristic of cyclohexane systems) and the presence of the spirocarbon at C-6 [13 C 49 8 (s)] suggested that 1 possessed a spiro 5 5 undecane skeleton characterized in *Laurencia* species by the chamigrene skeleton [1] Indeed, this was clearly consistent with the well-known 13 C NMR features for the bis-equatorial

Table 1	<sup>1</sup> H and	13C NMR	assignments for	r sesquiterpenoids	1 and 3*
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C No	1			3		
	s m	J (Hz)	<sup>13</sup> C†	s m	J (Hz)	<sup>13</sup> C†
1	4 47 dd	126, 56	68 7 d	4 40 dd	13, 61	68 2 d
2	2 92 m 2 62 dd	11 2, 5 6	35 8 t	2 70 m 2 46 dd	134,61	40 5 t
3			70 1 s			70 3 s
4	471 dd	13 5, 4 2	64 8 d	4 87 dd	132, 55	64 0 d
5	295 m 154 m		46 3 t	2 70 m 2 12 dd	139, 133	49 4 t
6			49 8 s			50 6 s
7			61 0 s			62 4 s
8	295 m		59 4 d	292 m		61 0 d
9	1 54 m 1 22 m		43 0 t	2 54 m 2 54 m		39 1 t
10	2 35 m		27 1 d	496 dd	94,88	61 9 d
11	-		1548 s	_		44 1 s
12	5 00 s 4 89 s		104 1 t	1 18 s		198 q
13	1 03 s		18 4 q	1 16 s		25 6 q
14	1 69 s		25 5 a‡	1 58 s		28 3 q
15	1 77 s		27 1 q‡	1 75 s		29 1 q

<sup>\*1</sup>H assignments were made by spin-decoupling and  $^{13}$ C assignments were made by comparison of the data with suitable chamigrene models  $^{1}$ H spectra were recorded in CDCl<sub>3</sub> solution at 360 MHz with internal TMS as standard,  $^{13}$ C spectra were recorded in bz- $d_6$  solution at 50 MHz with TMS as internal standard, abbreviations m = multiplet, s = singlet, d = doublet, t = triplet, q = quartet

dihalide constellation commonly encountered in Laurencia chamigrenes [4] However, decoupling analysis clearly showed that the usual C-10 bromine component of halochamigrenes was absent Instead, it could clearly be seen, in  $C_6D_6$  solution, that an axial proton was present, coupled to a methylene pair at C-9 (J=139,15 Hz) but also to an equatorial methyl group (J=66 Hz) This same proton sharpened when the exomethylene proton at  $\delta 456$  was irradiated, thus placing the exomethylene component at C-11

The relative stereochemistries of the substituents, C-1, C-3, C-4 and C-7, C-8 were assigned based upon coupling constant analyses for 6-membered rings and upon NOE measurements The alcohol at C-1 and the bromine at C-4 were clearly equatorial since the corresponding protons showed large axial-axial couplings Fortuitously, a NOE experiment successfully related the epoxide stereochemistry to the overall configuration of the vicinal dihalidebearing ring Irradiation of the epoxide methyl group at  $\delta$  1 69 resulted in the expected enhancement of the epoxide proton at  $\delta$  2.95, but also enhanced the axial proton on the adjacent ring at C-4 This latter enhancement can only occur if the C-7 epoxide methyl (C-14) and the C-5-C-4 substituent at C-6 are both 'down' as arranged in structure 1 This experiment was fortuitous in that it successfully related the relative stereochemistries of the two carbon rings

The structure of alcohol 1 was thus assigned as the

rearranged chamigrene resulting, biogenetically, from an apparent methyl migration from C-11 to C-10 Rearrangements of this nature, fostered by solvolysis of the bromine atom at C-10, have been implicated in the biosynthesis of numerous rearranged carbon skeletons isolated from *Laurencia* species [5] Alcohol 1, however, represents the first example of this rearrangement within the chamigrene class of sesquiterpenoids

The unrearranged chamigrene alcohol 3 was also isolated from this extract but as a more minor component (2% organic extract). Alcohol 3 analysed for C<sub>15</sub>H<sub>23</sub>O<sub>2</sub>Br<sub>2</sub>Cl by HRMS and <sup>13</sup>C NMR. The spectral features of 3 were similar to 1 and highly analogous to several closely related *Laurencia* chamigrenes [6]. As in 1, acetylation yielded the monoacetate 4, and complete proton spin-decoupling analyses led to the complete assignment of alcohol 3 was thus based upon the complete analogy of the spectral data of 3 with several closely related chamigrenes [6]. No efforts were made to determine the absolute stereochemistries of these metabolites

In initial experiments designed to predict the biological functions of *Laurencia* metabolites the ichthyotoxicities of alcohols 1 and 3 were measured using methods already described [7] At concentrations of 15 µg/ml alcohols 1 and 3 were lethal to the damselfish *Pomacentrus coeruleus* within a 1 hr period

<sup>†</sup>Multiplicities determined by single frequency off-resonance decoupling techniques

<sup>‡</sup>Values may be interchanged

## **EXPERIMENTAL**

Extraction and isolation procedures Laurencia species, Smithsonian herbarium No JN-11540, was collected August 19, 1982 at Rio Mar near Vero Beach, Florida. The Florida plants resemble L flagellifera as reported from Brazil However, J N considers this may represent an undescribed species until further anatomical studies can be done comparing Florida and Brazil specimens. The algae were immediately preserved in EtOH solution, and next thoroughly extracted with CHCl3-MeOH (2 1) to yield 2.5 g crude condensed extract. The extract was chromatographed over silica gel using conventional methods and several fractions were combined (eluted with 20% EtOAc in isooctane) and further purified by silica preparative HPLC (same solvent). These procedures resulted in the isolation of alcohols 1 (125 mg, 5% ext.) and 3 (50 mg, 2% ext.) as light mobile oils

Rearranged chamigrene 1 and acetate 2 The rearranged chamigrene alcohol 1 showed  $[\alpha]_D + 67^\circ$  (c 0.5, CHCl<sub>3</sub>) and analyzed for C<sub>15</sub>H<sub>22</sub>O<sub>2</sub>BrCl by HRMS, for [M-OH]<sup>+</sup> calc 331 0462, obsd 331 0451, for [M - HCl] + calc 312 0724, obsd 312 0699 The alcohol showed the following spectral features not reported in the text Table 1 IR (CHCl<sub>3</sub>) 3500, 2950, 1620, 1450, 1375, 1100, 1040 cm $^{-1}$ ,  $^{1}$ H NMR (360 MHz,  $C_{6}D_{6}$ )  $\delta$  3 98, 1H, dd, J = 130, 57 (C-1), 289 1H, t, J = 130 (C-2), 237, 1H, dd, J= 130, 57 (C-2'), 491, 1H, dd, J = <math>135, 42 Hz (C-4), 251, 1H, bs(C-8), 1 84, 1H, m (C-9), 1 00, 1H, ddd, J = 139, 139, 15 Hz (C-9)9'), 206, 1H, ddd, J = 139, 66, 15 Hz (C-10), 456, 1H, d, J= 11 Hz (C-12), 454, 1H, s (C-12'), 073, 3H, d, J = 66 Hz (C-13), 1 52, 3H, s (C-14), 1 56, 3H, s (C-15) Acetate 2 Acetylation of 1 (5 mg) was performed with a 20% molar excess of acetic anhydride in pyridine at room temp for 24 hr Ether extraction of the reaction mixture followed by water, acid and NaHCO3 wash of the combined ether extracts and reduction in vacuo yielded the monoacetate 2 which was not further purified Acetate 2 showed the following spectral features  $[\alpha]_D^{25} + 61^\circ$  (c 0 6, CHCl<sub>3</sub>), HRMS for  $[M-CHO]^+$  calc 361 0568, obsd 361 0536, IR (CHCl<sub>3</sub>) 2960, 1730, 1450, 1360, 1260 cm<sup>-1</sup> <sup>1</sup>H NMR (360 MHz, CDCl<sub>3</sub>)  $\delta$  5 50 (1H, dd, J = 10 7, 7 2 Hz), 4 72 (1H, s), 471(1H, s), 306(1H, dd, J = 141, 41 Hz), 298(1H, s), 276(2H, s)mult), 2 30 (1H, mult), 2 23 (1H, mult), 2 05 (3H, s), 1 81 (3H, s), 1 67 (3H, s), 1 60 (2H, mult), 1 41 (1H, t, J = 13 Hz), 1 00 (3H, d, J $= 66 \, \text{Hz}$ 

Chamigrene alcohol 3 and acetate 4 The chamigrene alcohol 3 showed  $\begin{bmatrix} \alpha \end{bmatrix}_D + 88^\circ$  (c 1 5, CHCl<sub>3</sub>) and analyzed for  $C_{15}H_{23}O_2Br_2Cl$  by HRMS calc for  $[M-Br]^+$  349 0570 obsd

349 0573 Alcohol 3 showed the following IR absorptions (CHCl<sub>3</sub>): 3500, 2960, 1450, 1390, and 1210 cm<sup>-1</sup> Acetate 4 The alcohol 3 (5 mg) was acetylated in a fashion identical as with 1 to yield the monoacetate 4 which was not purified further The acetate showed the following spectral features IR (CHCl<sub>3</sub>): 2950, 1730, 1450, 1360, 1260 cm<sup>-1</sup>, <sup>1</sup>H NMR (CDCl<sub>3</sub>)  $\delta$  5 47 (1H, dd, J = 121, 6 1 Hz), 4 83 (1H, dd, J = 132, 5 4 Hz), 4 14 (1H, dd, J = 108, 7 7 Hz), 2 94 (1H, d, J = 46 Hz), 2 69 (2H, mult), 2 54 (3H, mult), 2 11 (3H, s), 1 79 (3H, s), 1 60 (3H, s), 1 18 (3H, s), 1 15 (3H, s)

Acknowledgements—We wish to thank Dr James N Norris, Smithsonian Institution, Washington, DC, for the collection and taxonomic assessment of this Laurencia species. This is contribution #142 of the Marine Botany Project at the Smithsonian Marine Station at Link Port High resolution mass spectra were provided by the NIH Mass Spectrometry Resource Center UC San Francisco. The Center is supported by NIH grant RR01614 (A L Burlingame, principal investigator). This research is a result of financial support from NOAA, Office of Sea Grant, Department of Commerce under grant No NA80AA-D-00120. The US Government is authorized to produce and distribute reprints for governmental purposes, notwithstanding any copyright notation that may appear hereon.

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