Volatile Compounds from Marine Algae

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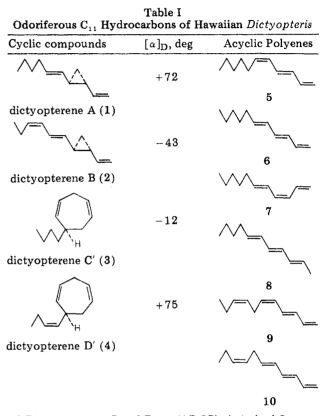
The fragrances of terrestrial plants have attracted the attention of man since antiquity. It is therefore not surprising that some of the earliest chemical investigations have been conducted on the odoriferous constituents. Organic chemists now immediately associate the odors of trees and shrubs with monoterpenes, of spices with phenols and their ethers, and of fruits and flowers with simple aliphatic esters.

The odors connected with marine plants are much less familar. Unlike the countless number of odoriferous terrestrial plants, relatively few marine plants possess odor. Prior to 1966, when we began our work on the odor of seaweed, only one volatile substance had been identified from wet, undecomposed seaweed and described in the literature. In 1935 Haas¹ had shown that dimethyl sulfide is spontaneously evolved from the red algae Polysiphonia fastigiata and P. nigrescens when the seaweeds are exposed to air. Bywood and Challenger² later found that dimethyl sulfide is similarly evolved from the green alga Enteromorpha intestinalis. Actually, the odor of dimethyl sulfide can be detected in many seaweeds immediately after crushing the plants. The formation of dimethyl sulfide results from an enzymatic decomposition of dimethyl- β -propiothetin,³ a metabolite of methionine⁴ that is fairly widespread in both benthic and planktonic marine plants.⁵

$CH_3SCH_2CH_2CHCO_2^- \rightarrow (CH_3)_2SCH_2CH_2CO_2^- \rightarrow (CH_3)_2S$ NH.⁴

The Odor of Brown Seaweed. From the steam distillate of dried Dictyopteris divaricata, a brown alga that grows on the coasts of Japan, Takaoka and Ando⁶ obtained an oil with a "beach odor" which has been shown by Irie and co-workers to be a mixture of sesquiterpenes of the cadinene type.⁷ A sesquiterpene, also of the cadinene type, has also been found to be the major hydrocarbon of D. zonarioiodes from Mexico.⁸

In striking contrast to the species of *Dictyopteris* that grow along the coasts of Japan and Lower California, the Hawaiian species D. plagiogramma or D. australis elaborates novel non-isoprenoid C₁₁ compounds in the essential oil (Table I) instead of sesquiterpenes.⁹ Each species grows abundantly on the sublittoral reef flats surrounding the Hawaiian Islands, and large amounts of the seaweed are deposited on the shores during the summer months by heavy surf. The odor of Dictyopteris can frequently be detected in the air around the beaches. Known as *limu lipoa* (seaweed gathered from the deep) in Hawaii, the freshly chopped seaweed



^a Dictyopterenes C and D are (1R,2S)-cis-1-vinyl-2-(trans-1'-hexenyl)cyclopropane and (1R,2S)-cis-1-vinyl-2-(trans-1, cis-3-hexadienyl)cyclopropane, respectively, the proposed precursors of dictyopterenes C' and D'.

is used as a condiment with raw fish and other foods.¹⁰

Two interesting dialkenylcyclopropanes, dictyopterenes A and B, are the major constituents of the essential oil, and the related hydrocarbons, dictyopterenes C' and D', are minor components. Their

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 (4) R. C. Greene, J. Biol. Chem., 237, 2251 (1962).
 (5) (a) F. Challenger, R. Bywood, P. Thomas, and B. J. Hayward, Arch. Biochem. Biophys., **69**, 514 (1957); (b) R. G. Ackman, C. S. Tocher, and J. McLachlan, J. Fish. Res. Board Can., **23**, 357 (1966).

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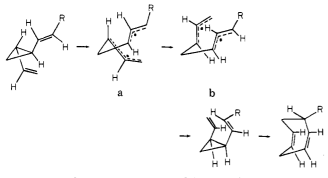
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Garden, Lawai, Kauai, Hawaii, 1974.

Richard E. Moore was born in San Francisco, Calif., in 1933. He received his B.S. and M.S. degree in chemistry from the University of San Francisco and his Ph.D. degree in organic chemistry from the University of California, Berkeley. After 3 years of postdoctoral work he joined the chemistry faculty at Hawali, where he is presently a professor. His research interests have been focused primarily on the chemistry of natural products, in particular those associated with marine plants.

structures were deduced from spectral and chemical data^{9e} and verified by synthesis.¹¹ Lemieux oxidation of either 1 or 2 gave (+)-trans-cyclopropanedicarboxylic acid, establishing the absolute configuration of the cyclopropane ring as R,R. Both 1 and 2 underwent stereoselective, thermally induced Cope rearrangements9c to 1,4-cycloheptadienes, and the products were identical, except for optical properties, with natural 3 and 4. To rationalize the stereoselectivity in the isomerization of 1 or 2, the allyl system in the intermediate diradial a¹² must rotate faster¹³ than the bulkier heptenyl (R = n-butyl) or heptadienyl (R =cis-1-butenyl) group to give the conformer b, which then recyclizes to the *cis*-1,2-dialkenylcyclopropane¹⁴ and rearranges by a conventional Cope rearrangement to the cvcloheptadiene. The absolute configurations of the



cycloheptadienes were secured by converting them to the known butylsuccinic acids.

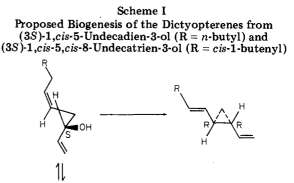
Several acyclic undecapolyenes, viz. three 1,3,5-undecatrienes (5-7), four 2,4,6-undecatrienes, of which only one (8) has been fully characterized, and two 1,-3,5,8-undecatetraenes (9, 10), are found with the dictyopterenes in the essential oil in moderate to trace amounts.^{9e} The compositions of the 1,3,5-undecatriene, 2,4,6-undecatriene, and 1,3,5,8-undecatetraene mixtures are essentially the same as the photostationary-state mixtures resulting from the *p*-benzoquinone photosensitized equilibrations of the trienes and tetraene in the presence of azulene as a quencher. For example, 5 and 6 are the predominant isomers of 1,3,5-undecatriene in both the essential oil and photostationary state mixture. From proton NMR evidence all four natural 2,4,6-undecatrienes are trans- Δ^4 isomers.^{9e} The same mixture of 2,4,6-undecatrienes is formed when 1,cis-3, cis-5-undecatriene¹⁵ is thermally isomerized and the

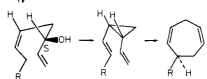
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(12) The preferential ring opening to intermediate a is suggested from circular dichroism and nuclear magnetic resonance studies of 1 and 2 at elevated temperatures: S.-Y. C. Lee, J. C. Craig, P. A. Kollman, R. E. Moore, J. A. Pettus, Jr., and S. Rothenberg, submitted for publication. The upper energy state conformation of a trans-dialkenylcyclopropane is gauche (dihedral angle <90°) and not s-cis as needed for formation of diradicals with cis-alkenyl radical groups. Cyclopentenes are not formed in the Cope rearrangements of 1 and 2 as the intermediate diradical has structure a.

(13) Internal rotational preference is shown in the thermal enantiomerization and diastereoisomerization of cis- and trans-1-cyano-2-isopropenylcyclopropane: W. von E. Doering and K. Sachdev, J. Am. Chem.

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(14) C. Ullenius, P. W. Ford, and J. E. Baldwin, J. Am. Chem. Soc.,
94, 5910 (1972); J. E. Baldwin and C. Ullenius, *ibid.*, 96, 1542 (1974).
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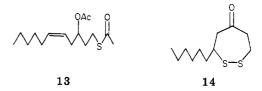
resulting cis-2, cis-4, trans-6-undecatriene is photoisomerized in the presence of *p*-benzoquinone and azulene.

Mass spectral analysis of the oil indicates that small amounts of compounds having the molecular compositions $C_{11}H_{14}O$ and $C_{11}H_{16}O$ are also present. Two odoriferous, dextrorotatory dihydrotropones related to dictyopterene C' have been isolated, and their structures have been deduced to be 11 and 12.¹⁶ Selenium



dioxide oxidation of 3 produced 11, indicating that C-6 in 11 has the R configuration. Catalytic hydrogenation of 12 led to a 3-butylcycloheptanone, which exhibited a CD curve similar to that of (-)-(3S)-methylcycloheptanone, showing that C-6 in 12 is also R^{17}

Less volatile organosulfur compounds such as 13 and 14 (absolute configurations undetermined) have been



isolated from the chloroform-methanol extract of the dried alga.^{9d,18} None of the sulfur compounds, however, exhibits a penetrating odor like the hydrocarbons and ketones. All of the compounds possess a C_{11} unit attached to sulfur with an oxygen substituent at C-3 and are obviously related to the hydrocarbons.

The most attractive precursors to both the hydrocarbons and sulfur compounds appear to be 1-undecen-3-ols.¹⁹ 1-Alken-3-ols are ubiquitous²⁰ minor constituents of animal and plant lipids and are me-

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(18) (a) P. Roller, K. Au, and R. E. Moore, Chem. Commun., 503 (1971); (b) R. E. Moore, ibid., 1168 (1971); (c) A. E. Asato and R. E. Moore, Tetrahedron Lett., 4941 (1973).

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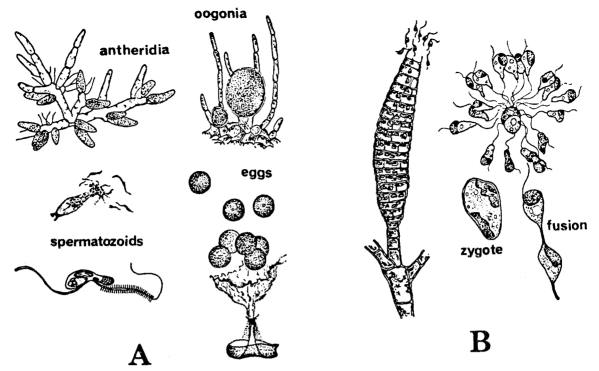


Figure 1. Sexual reproduction in (A) Fucus serratus and (B) Ectocarpus siliculosus (copyright 1972, Westdeutscher Verlag GmbH; reproduced, with permission, from *Rheinisch-Westfael*. Akad. Wiss., No. 217 (1972)).

tabolites of fatty acids. 1-Undecen-3-ols have not been detected in Dictyopteris and other seaweeds as yet. (3S)-cis-1,5-Octadien-3-ol (15), however, is a minor component of the essential oil of the red seaweed Chondrococcus hornemanni,²¹ but C₈ hydrocarbons related to the C_{11} hydrocarbons of *Dictyopteris* are not associated with it. If the hypothetical 1-undecen-3-ols have the S configuration at C-3, then concomitant dehydration and cyclization of (3S)-cis-1.5-undecadien-3-ol leads to either 1 or (1R.2S)-cis-1-vinyl-2-(trans-1'-hexenyl)cyclopropane (dictyopterene C), the latter rearranging spontaneously at biotic temperatures^{11a} to 3 (Scheme I). Similarly, (3S)-1,cis-5,cis-8-undecatriene-3-ol leads to either 2 or 4 via (1R,2S)-cis-1-vinyl-2-(trans-1,cis-3-hexadienyl)cyclopropane (dictyopterene D).

The Role of Volatile Hydrocarbons in the Sexual **Reproduction of Brown Seaweeds.** In the late 1940's Cook and his collaborators²² discovered that motile spermatozoa are attracted to mature eggs of Fucus serratus by a volatile substance secreted by the eggs. The same phenomenon was observed for F. vesiculosus. The chemotactic principle could be readily expelled from an aqueous extract of the eggs by a stream of inert gas and was completely recovered when the carrier gas was passed through a strongly cooled trap containing a small amount of frozen seawater. Interestingly, the active substance from F. serratus not only attracted the sperm of F. servatus but also attracted those of F. vesiculosus and F. spiralis. The nonspecificity of the pheromone indicated that the same chemotactic agent was produced by the closely related Fucales. The mechanism of this very striking phenomenon was

(21) F. X. Woolard, B. J. Burreson, and R. E. Moore, J. Chem. Soc., Chem. Commun., 486 (1975). concluded to be purely physical, as the chemotactic response was sensitive only to a gradient concentration of the pheromone. Chemotactic activity could be induced by a large number of simple organic compounds. Of the hydrocarbons that were tested, *n*-hexane corresponded closest in biological and physical behavior with the natural pheromone. An attempt to identify the chemotactic principle by mass spectrometry gave inconclusive results, and work was discontinued.

Some 20 years later Müller and Jaenicke²³ reisolated the volatile attractant from the eggs and oogonia of F. *serratus* and purified it by gas chromatography. Spectral and chemical examination of the resulting odoriferous oil showed that it was an isomer of 1,3,5octatriene which they named fucoserratene. From biogenetic considerations fucoserratene was predicted to be 1,*trans*-3,*cis*-5-octatriene (16),²⁴ and this was later proven by synthesis.²⁵ Compared with *n*-hexane, the sperm attractant had enormous chemotactic activity.

Earlier, Muller had studied chemotaxis in another brown seaweed *Ectocarpus siliculosus*.²⁶ Noting that the female gametes possessed odor whereas the isomorphic male gametes did not, he collected an odoriferous oil from female cultures and tested it for chemotactic activity. When Vaseline droplets impregnated with the pheromone were placed in the vicinity of active spermatozoids, these simulated eggs became surrounded by halos of male gametes. A collaborative effort with Jaenicke resulted in the first successful isolation and characterization of a sperm attractant from a marine alga.²⁷ The pheromone, ectocarpene (4), was a C₁₁

(23) D. G. Müller and L. Jaenicke, FEBS Lett., 30, 137 (1973).

(24) L. Jaenicke, D. G. Müller, and R. E. Moore, J. Am. Chem. Soc., 96, 3324 (1974).

(25) L. Jaenicke and K. Seferiadis, Chem. Ber., 108, 225 (1975).

(26) D. G. Müller, Planta, 81, 160 (1968).

(27) (a) D. G. Müller, L. Jaenicke, M. Donike, and T. Akintobi, Science, 171, 815 (1971);
 (b) L. Jaenicke, T. Akintobi, and D. G. Muller, Angew. Chem., Int. Ed. Engl., 10, 492 (1971).

^{(22) (}a) A. H. Cook, J. A. Elvidge, and I. Heilbron, Proc. R. Soc. London, Ser. B, 135, 293 (1948); (b) A. H. Cook, J. A. Elvidge, and R. Bentley, *ibid.*, 138, 97 (1951).

Type of Compound	Structure	Type of Compound	Structure
Haloforms	CHBr ₃ CHBr ₂ I CHBrI ₂ CHI ₃	Halogenated 2-acetoxypropanes	BrCH ₂ CH(OAc)CHBr ₂ Br ₂ CHCH(OAc)CHBr ₂
	CHBr ₂ Cl CHBrCll	Halogenated 1,2-epoxypropanes	$\operatorname{Br}CH-CHCHBr_2$
Dihalomethanes	$\begin{array}{c} \mathbf{CH}_{2}\mathbf{Br}_{2}\\ \mathbf{CH}_{2}\mathbf{Br}\\ \mathbf{CH}_{2}\mathbf{I}_{2} \end{array}$	1,1,3,3-Tetrahalopropenes	$Br_2C=CHCHBr_2$ $Br_2C=CHCHBrCl$ $Br_2C=CHCHCl,$
Carbon tetrahalides	CBr4		BrIC=CHCHBr ₂
Carbonyl dihalides		3,3-Dihaloacroleins	$Br_2C = CHCHO$
2-Haloethanols	ICH ₂ CH ₂ OH	Halogenated butenones	$Br_2C = CHCOCH_3^b$
1,2-Dihaloethanes	BrCH ₂ CH ₂ I		$Br_2C = CHCOCH_1Br^b$
Halogenated acetaldehydes	Br ₂ CHCHO		$Br_2C = CHCOCH_2I^b$
Halogenated acetones	CH ₃ COCH ₂ Br		$Br_2C = CHCOCHBr_2$
	CH ₃ COCH ₂ I		$Br_2C = CHCOCHBrCl^b$
	CH ₃ COCHBr ₂		$BrClC=CHCOCHBr_2^b$
	BrCH, COCH ₂ Br		$Cl_2C = CHCOCHBr_2^{b}$
	BrCH, COCH, I		BrClC=CHCOCHBrCl ^b
	CH,COCBr,		
	CH ₃ COCBr ₂ Cl		
	BrCH, COCHBr,		
	ICH,ĆOCHBr,		
	Br, CHCOCHBr,		
	Cl ₃ CCOCCl ₃		

 Table II

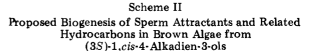
 Volatile Constituents of Hawaiian Asparagopsis taxiformis^a

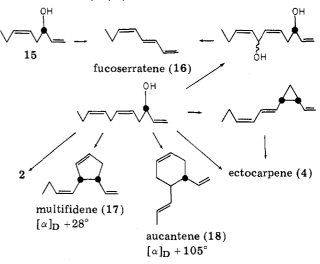
^a Collected from the reef between Diamond Head and Waikiki, Oahu. ^b The placement of the two olefinic halogens on C-4 is tentative.

hydrocarbon that proved to be identical in all respects, including optical properties, with the dictyopterene D' that my group had isolated from Hawaiian *Dictyopteris*.^{9c}

The studies of chemotaxis in *F. serratus* and *E. siliculosus* represented studies of two completely different fertilization processes. For the dioecious seaweed *F. serratus*, fertilization was oogamous: motile, biflagellated spermatozoids released by the antheridia of male plants fused with nonmotile, spherically shaped eggs released from the oogonia of female plants (Figure 1A). A C₈ hydrocarbon was produced by the eggs to attract the sperm to them. In the dioecious *E. siliculosus*, however, fertilization was isogamous and involved the fusion of motile, isomorphic male and female gametes (zoogametes) (Figure 1B). In this second case male gametes were attracted to female gametes by a C₁₁ hydrocarbon excreted by the females.

Müller then turned his attention to Cutleria multifida and a third type of fertilization process. In this dioecious alga fertilization was anisogamous: macroscopic, flagellated female gametes were fertilized by microscopic, flagellated male gametes. Again Muller observed that the females attracted the males with an odoriferous substance. Similar methodology was used to collect and purify the male-attracting oil of *Cutleria*, which consisted of three C_{11} hydrocarbons, only one of which was biologically active. From spectral and biogenetic arguments the sperm attractant multifidene was concluded to be a 3-(cis-1-butenyl)-4-vinylcyclopentene.²⁴ Synthesis has now shown that multifidene is the cis compound 17.²⁸ The two biologically inactive components of the male-attracting oil were also present in the essential oil of the male and asexual plants. One of these hydrocarbons was 4, whereas the other one, aucantene, was identified as trans-4-(trans-1-propenyl)-5-vinylcyclohexene (18).²⁴ The structure of au-





cantene has also been confirmed by synthesis²⁹ (see Scheme II).

Discovery of Haloforms in the Bonnemaisoniaceae. Asparagopsis taxiformis is an edible red seaweed that is highly favored in Hawaii for its strong aroma and flavor. Known locally as *limu kohu* (the supreme seaweed),¹⁰ the alga is a dioecious, gametophytic plant that alternates in its life cycle with a heteromorphic sporophyte, *Falkenbergia rufanolosa*. *A. taxiformis* is rich in iodine, but free molecular iodine is not present in the live plants. Its essential oil is composed of mainly bromine- and iodine-containing haloforms³⁰ and small to trace amounts of many other

⁽²⁹⁾ F.-J. Marner and L. Jaenicke, Chem. Ber., 108, 2202 (1975).
(30) B. J. Burreson, R. E. Moore, and P. Roller, Tetrahedron Lett., 473 (1975).

 Table III

 Halogenated Acids, Alcohols, and Amides in Extracts of

 Dried Hawaiian Asparagopsis taxiformis

-		
CH_2Cl_2 Extract ^a	H ₂ O Extract ^b	
1,1-Dihalo-2-propanols	Haloacetic acids	
1,3-Dihalo-2-propanols	Dihaloacetic acids	
1,1,3-Trihalo-2-propanols	Monohaloacrylic acids	
1,1,1,3-Tetrahalo-	3,3-Dihaloacrylic acids	
2-propanols		
1,1,3,3-Tetrahalo-	Trihaloacrylic acids	
2-propanols		
1,4,4-Trihalobut-3-	4,4-Dihalo-2-hydroxybut-	
en-2-ols	3-enoic acids	
1,1,4,4-Tetrahalobut-3-		
en-2-ols		
1,1,1,4,4-Pentahalobut-		
3-en-2-ols		
2,2-Dihaloacetamides		
•		

^a Br₂CHCH(OH)CH₂Br, Br₂CHCH(OH)CH₂Cl, Br₂-CHCH(OH)CHBr₂, Br₂C=CHCH(OH)CHBr₂, Br₂CHCON-H₂, BrICHCONH₂, and I₂CHCONH₂ are the major constituents. ^b Br₂CHCO₂H, BrICHCO₂H, I₂CHCO₂H, and Br₂C=CHCO₂H are the major constituents.

halogenated compounds (Table II).^{31,32} Related nonvolatile compounds are present in the methylene chloride³³ and aqueous extracts³⁴ of the lyophilized seaweed (Table III). Interestingly, none of these halogenated compounds exist in the asexual *F. rufa*nolosa.

The haloforms are apparently formed by the classical haloform reaction of methyl or carboxymethyl ketones. Monohaloacetones, 1,1-dihaloacetones, and 1,1,1-trihaloacetones, which are intermediates in the conversion of acetone or acetoacetic acid to haloforms, are all found in the essential oil. Small amounts of 1,2-dihaloacetones, 1,1,2-trihaloacetones, 1,1,2,2-tetrahaloacetones, and even a hexahaloacetone are also present. Introduction of the halogens into the acetones is undoubtedly enzymatic,^{35,36} but the degradation of 1,1,-1-trihaloacetones into haloforms may not be. On standing in aqueous sodium bicarbonate solution, 1,-1,1-trihaloacetone such as 1,1,1-tribromoacetone and 1,1-dibromo-1-chloroacetone decompose slowly into haloforms and acetic acid.³¹

In the nonenzymatic bromination of acetone, the introduction of the first bromine is slow. Further bromination of bromoacetone in aqueous solution, however, is rapid and leads to 1,1,1-tribromoacetone. In glacial acetic acid, on the other hand, bromination of bromoacetone leads slowly to 1,1,3,3-tetrabromoacetone (19), as the reaction proceeds through an enol intermediate rather than through the enolate ion.

(31) B. J. Burreson, R. E. Moore, and P. P. Roller, J. Agric. Food Chem., 24, 856 (1976).

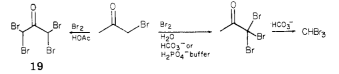
(32) While our work was in progress the isolation of several bromineand chlorine-containing acetones and butenones from *A. taxiformis* collected in the Gulf of California was reported: W. Fenical, *Tetrahedron Lett.*, 4463 (1974).

(33) F. X. Woolard, R. E. Moore, and P. P. Roller, *Tetrahedron*, 32, 2843 (1976).

(34) Presented by R. E. Moore, F. X. Woolard, and P. P. Roller, 172nd National Meeting of the American Chemical Society, San Francisco, Calif., August 1976, Abstract ORGN-083.

(35) For studies of enzymatic halogenation see: J. R. Beckwith, R. Clark, and L. P. Hager, J. Biol. Chem., 238, 3086 (1963); A. Taurog and E. M. Howells, *ibid.*, 241, 1329 (1966); J. A. Thomas, D. R. Morris, and L. P. Hager, *ibid.*, 245, 3129 (1970), and references therein.

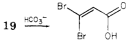
(36) Chloroperoxidase from the mold Caldariomyces fumago catalyzes the peroxidative chlorination of β -ketadipic acid to δ -chlorolevulinic acid and carbon dioxide: P. D. Shaw and L. P. Hager, J. Biol. Chem., 236, 1626 (1961).



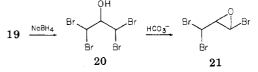
Further bromination of 19 occurs only in aqueous solution where it is brominated to pentabromoacetone, which decomposes into bromoform and dibromoacetic acid. Pentabromoacetone has not been detected in Hawaiian Asparagopsis, but dibromoacetic acid is a constituent of the aqueous extract of the seaweed.³¹

Most interesting is the presence of bromochloroiodomethane in Asparagopsis. The amount of this haloform is extremely small, and we have been unable to concentrate a sufficient quantity of this material to determine its optical properties. In the gas chromatograph it is very poorly resolved from the enormous amount of bromoform, the major constituent of the essential oil. All of the iodine-containing haloforms rapidly liberate iodine on exposure to air and light. This sensitivity suggests that the dihalomethanes, 1bromo-2-iodoethane, and carbonyl diiodide may be artifacts from photooxidation of iodoform, bromodiiodomethane, and dibromoiodomethane. The carbon tetrabromide, however, is probably formed by further bromination of bromoform in the seaweed.

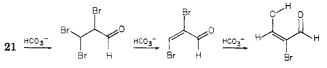
The 3,3-dihaloacrylic acids in Asparagopsis may be formed from either the haloform reaction of 4,4-dihalobutenones or from the Favorskii rearrangement of 1,1,3,3-tetrahaloacetones. When 19 is allowed to stand in aqueous sodium bicarbonate solution, it is converted cleanly into 3,3-dibromoacrylic acid.³⁴



The halogenated 2-propanols appear to be the precursors of the halogenated propenes and epoxypropanes. In nonenzymatic studies we have been unable to dehydrate 1,1,3,3-tetrabromopropan-2-ol (20) or to convert 2-acetoxy-1,1,3,3-tetrabromopropane to 1,1,3,3-tetrabromopropene under mild conditions. In aqueous bicarbonate solution, however, 20 readily cyclizes to *trans*-1,3,3-tribromoepoxypropane (21).³⁴



There is a possibility that 21 is involved in the enzymatic biogenesis of 3,3-dibromoacrolein. Under nonenzymatic conditions, however, 21 slowly rearranges in aqueous bicarbonate and eliminates hydrogen bromide to form (*E*)-2,3-dibromoacrolein, a labile compound that is converted into the remarkably stable bromomalonodialdehyde on further standing in bicarbonate solution. Even though 21 is a constituent of the es-



sential oil, neither 2,3-dibromoacrolein nor bromomalonodialdehyde can be detected in the algal extracts. 34 3,3-Dihaloacroleins may serve as precursors to the 4,4-dihalobut-3-en-2-ols. A number of substrates such as acetate, malonate, haloacetate, dihaloacetate, and even haloform and methylene halide could provide the additional carbon atom.^{33,37} The 4,4-dihalobutenones, on the other hand, are probably formed by the Claisen condensation of a 3,3-dihaloacrylate with a one-carbon donor.

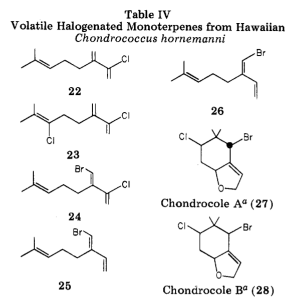
The presence of haloforms and other halogenated compounds in A. taxiformis suggests that this seaweed might be potentially dangerous to eat. In Hawaii the uncooked seaweed is frequently eaten with meats and fish. Since the odor and flavor are quite strong, however, relatively small quantities of the *limu* are added to foods. To the author's knowledge there has never been a case of illness attributed to the ingestion of *limu kohu*. Many of the compounds in Asparagopsis appear to be carcinogenic, and an epidemiological study might provide some valuable information. Although extensive data have been accumulated on the carcinogenicity of $CHCl_3$, CCl_4 , and other chlorine-containing compounds, relatively little is known about the bromine and iodine analogues.

Odoriferous halogenated compounds have also been isolated from other Bonnemaisoniaceae. Rinehart et al. have identified several bromine-containing 2-heptanones, primarily 1,1,3,3-tetrabromo-2-heptanone, and also 1-iodo-3,3-dibromo-2-heptanone, in the essential oil of *Bonnemaisonia hamifera*,³⁸ and very recently the research group at the Roche Institute of Marine Pharmacology in Sidney has found 3,4,4-tribromobutenone, 1,1,2-tribromoct-1-en-3-one, and six closely related compounds in *Ptilonia australasica*.³⁹

The Origin of Halocarbons in the Atmosphere and Water. Chloroform and carbon tetrachloride are constituents of the chlorocarbon complement of the upper atmosphere⁴⁰ and have been identified as trace contaminants of chlorine-treated drinking water.⁴¹ It is quite possible that the $CHCl_3$ and CCl_4 in both the air and purified water have algal origins and arise by reactions analogous to those in Asparagopsis for CHBr₃ and CBr_4 . To date we have not determined whether or not $CHCl_3$ and CCl_4 are produced by Hawaiian A. taxiformis. The presence of hexachloroacetone in the essential oil certainly suggests that they are. The Fenical and Goldberg groups, however, have recently shown that CHCl₃ and CCl₄, as well as bromodichloromethane, are constituents of A. armata from the Mediterranean.42

Methyl chloride, the dominant chlorocarbon in the atmosphere⁴³ and a possible regulator of the stratospheric ozone density, appears to have a marine origin and is not the result of direct industrial pollution.⁴⁴ In

- (43) E. P. Grimsrud and R. A. Rasmussen, Atmos. Environ., 9, 1010, 1014 (1975).
 - (44) J. E. Lovelock, Nature (London), 256, 193 (1975).



 a The optical rotation and absolute configuration has not been determined.

the sea its proposed precursor, methyl iodide, reacts nucleophilically with chloride ion and is converted into the more volatile methyl chloride, which then escapes into the air.⁴⁵ Methyl iodide is found in all ocean waters and may be a common extracellular metabolite of marine algae. The water in the vicinity of seaweeds such as the iodine-rich kelp *Laminaria digitata* contains a thousand times more methyl iodide than the open oceans, but its actual existence in this seaweed has not been proven. Methyl iodide, however, is produced by *Asparagopsis armata*.⁴²

Odoriferous Monoterpenes from Red Algae. Halogenated monoterpenes were first discovered in the digestive gland of the sea hare *Aplysia californica.*⁴⁶ Suspected to be of dietary origin, these compounds were eventually traced to species of *Plocamium* that are ingested by the mollusk.⁴⁷ Since then a fascinating array of novel compounds have been isolated from this genus of red seaweeds.⁴⁸ Most of the monoterpenes from *Plocamium*, however, are not very volatile and possess only faint odors.

Chondrococcus hornemanni, a red seaweed that grows in the subtropical and tropical Pacific Ocean in regions of heavy surf, has a strong turpentine-like odor that is manifested on crushing the plants. When inspected under a microscope, numerous oil droplets or possibly specialized cells can be detected throughout the thalli. These secondary structures are ruptured on freezing the seaweed, and the characteristic odor of the plant is liberated. Lyophilization of the seaweed removes the odoriferous constituent along with water, and the essential oil is recovered simply by extracting the condensate with methylene chloride. The essential oil

⁽³⁷⁾ H. Taguchi, H. Yamamoto, and H. Nozaki, J. Am. Chem. Soc., 96, 3020 (1974).

⁽³⁸⁾ J. F. Siuda, G. R. Van Blaricom, P. D. Shaw, R. D. Johnson, R. H. White, L. P. Hager, and R. L. Rinehart, Jr., J. Am. Chem. Soc., 97, 937 (1975).

⁽³⁹⁾ Paper presented by R. Kazlauskas, P. T. Murphy, R. Quinn, and R. J. Wells at the 10th International Symposium on Chemistry of Natural Products, IUPAC, Dunedin, New Zealand, August 1976.

⁽⁴⁰⁾ J. E. Lovelock, R. J. Maggs, and R. J. Wade, Nature (London), 241, 194 (1973).

⁽⁴¹⁾ Chem. Eng. News, 54 (16), 7 (1976).

⁽⁴²⁾ O. McConnell and W. Fenical, private communication; C.-W. Su and E. D. Goldberg, private communication.

⁽⁴⁵⁾ The residence time of methyl iodide in seawater is comparable to its rate of reaction with chloride ion: O. C. Zafiriou, J. Mar. Res., 33, 75 (1975).

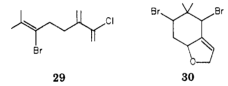
 ^{(46) (}a) D. J. Faulkner, M. O. Stallard, J. Fayos, and J. Clardy, J. Am.
 Chem. Soc., 95, 3413 (1973); (b) D. J. Faulkner and M. O. Stallard,
 Tetrahedron Lett., 1171 (1973).

⁽⁴⁷⁾ M. O. Stallard and D. J. Faulkner, Comp. Biochem. Physiol. B, 49, 25 (1974).

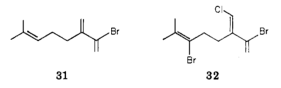
 ^{(48) (}a) J. S. Mynderse, D. J. Faulkner, J. Finer, and J. Clardy, Tetrahedron Lett., 2175 (1975); (b) J. S. Mynderse and D. J. Faulkner, Tetrahedron, 31, 1963 (1975); (c) P. Crews and E. Kho, J. Org. Chem., 40, 2568 (1975).

of subtropical C. hornemanni is composed of mostly myrcenes that are mono- and disubstituted with bromine and/or chlorine.⁴⁹ Usually bromine is found on the C-3 methylene and chlorine is attached to the C-2 and C-6 carbons of 7-methyl-3-methylene-1.6-octadiene (myrcene). For example, in a variety found at Black Point, Oahu, 22 and 23 are the most abundant volatile compounds, whereas in another variety collected at the Halona Blowhole, Oahu, 24 and 25 are the major constituents of the essential oil (Table IV). In addition to the myrcenes, the essential oil of the latter alga also contains a large amount of a cyclic monoterpene, chondrocole A. a dimethylhexahydrobenzofuran having the structure 27. Chondrocole A is only a minor volatile constituent of the alga from Black Point, but accompanies the isomeric chondrocole B (28) in that seaweed.^{49b} The chlorine and bromine of chondrocoles A and B are in positions analogous with the placement of a chlorine on C-6 and a bromine on the C-3 methylene of myrcene.

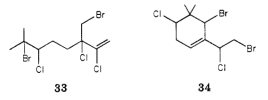
Chondrococcus also produces halogenated myrcenes with bromines at C-2 and C-6 and a chlorine on the C-3 methylene. Although we have not encountered any of these compounds in the essential oils of Hawaiian *Chondrococcus*, the ethereal extract of the lyophilized seaweed from Black Point curiously contains a 6bromomyrcene, 29, along with a related cyclic com-



pound, chondrocole C.⁵⁰ Ichikawa et al. have also reported that **31** and **32** are minor constituents of C. hornemanni from the Amami Island coasts of Japan.^{49a}



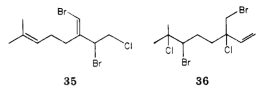
Several halogenated dihydro- and tetrahydromyrcenes are present in the ethereal extracts of the dried seaweeds. The *Chondrococcus* from Black Point elaborates mainly **33**, with smaller amounts of three closely related compounds. All four compounds display a Markownikoff addition of BrCl to the C-3 methylene, and two of these show that BrCl has been added to an anti-Markownikoff manner to the Δ^6 double bond. None of the major polyhydromyrcenes have BrCl added to the vinyl group; however, a related cyclic compound, **34**, has been found with this feature.⁵⁰ The variety



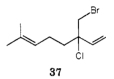
(49) (a) N. Ichikawa, Y. Naya, and S. Enomoto, *Chem. Lett.*, 1333 (1974);
(b) B. J. Burreson, F. X. Woolard, and R. E. Moore, *Tetrahedron Lett.*, 2155 (1975).

(50) B. J. Burreson, F. X. Woolard, and R. E. Moore, Chem. Lett., 1111 (1975).

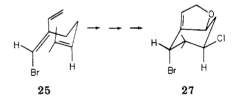
from the Halona Blowhole, on the other hand, elaborates 35 and 36 as the major di- and tetrahydromyrenes.



In 36 BrCl has added in a Markownikoff fashion to the Δ^6 double bond but in 35 has added anti-Markownikoff to the vinyl group.⁵⁰ Interestingly *C. hornemanni* from tropical Sri Lanka (Ceylon) does not produce halogenated myrcenes. In the extract of the seaweed, however, the major halogenated compound is 37.⁵¹



Biogenetically the halogenated myrcenes are probably formed by the enzymatic addition of BrCl to myrcene (38), a minor constituent of the essential oil,^{49a} followed by a dehydrohalogenation (e.g., $38 \rightarrow 37 \rightarrow 25$ or 26). In the nonenzymatic addition of halogen to myrcene, the first equivalent adds almost exclusively to the Δ^6 double bond, but monohalomyrcenes substituted at C-6 have not been found in *Chondrococcus*. Chondrocole A may be formed from 25. Similarly chondrocoles B and C could arise from 26.^{49b}



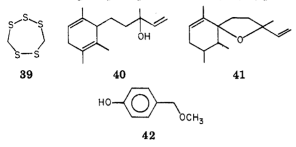
Future Directions. Actually, only a few benthic marine algae have been examined in any detail for volatile compounds. The isolation of 15 from Chondrococcus hornemanni suggests that 1-undecen-3-ols and C_8 hydrocarbons related to the dictyopterenes may eventually be found in seaweeds. The isolation of numerous chlorine-, bromine-, and iodine-containing compounds from Asparagopsis moves one to wonder whether organofluorine compounds might exist in some marine plants. Ethylene, an important regulator in the growth and development of terrestrial plants,⁵² has not been demonstrated in marine plants but may be produced concomitantly with dimethyl sulfide. The sulfur odor of seaweeds may not always be due to dimethyl sulfide. Some seaweeds are rich sources of sulfur and sulfur-containing lipids; Wratten and Faulkner,⁵³ for example, have recently identified lenthionine (39) and other cyclic polysulfides and their oxidation products in the odoriferous red alga Chondria californica. A few species of Laurencia, a genus of red seaweeds that has been a very rich source of interesting halogenated C_{15} compounds,⁵⁴ have a faint odor attributed mostly to

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unhalogenated sesquiterpenoids.55 In a study of Hawaiian L. nidifica, Erickson and her collaborators have described the isolation and structure elucidation of an interesting sesquiterpenoid alcohol, 40,56 a possible



(55) J. Cardellina, B. J. Burreson, P. P. Roller, and R. E. Moore, unpublished work.

link to dactyloxene B (41), isolated from the sea hare Aplysia dactylomela.⁵⁷ Several species of benthic algae are currently being investigated in my laboratory. In one case we have found that p-(methoxymethyl)phenol (42) is responsible for the foul odor of the red alga Martensia fragilis.58

It is a great pleasure to acknowledge the efforts of my collaborators, especially B. J. Burreson, J. A. Pettus, Jr., P. P. Roller, and F. X. Woolard, whose work is discussed here. I am very grateful to the National Science Foundation and to the Donors of the Petroleum Research Fund, administered by the American Chemical Society, for support of this research.

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Spin-Labeled Nucleic Acids

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Numerous spectroscopic techniques have been used to study the relationships which exist between the structure and the function of nucleic acid molecules. Some of the significant structure-function problems in the macromolecular nucleic acid field are: (a) helix-coil transition, (b) unfolding of the tertiary structure by temperature or chemical agents, (c) binding of metal ions such as magnesium, (d) interaction and intercalation of drugs, causing perturbations of the native structure, (e) recognition site on the transfer RNA (tRNA) by the corresponding aminoacyl-synthetase, and (f) interaction of specific proteins with RNA in the ribosomes.

Certain aspects of these problems have been illuminated by spin-labeling studies.

Nitroxide spin-labels are stable synthetic organic free radicals that can interact either covalently or noncovalently with biological macromolecules of interest. Much of the organic chemistry of nitroxides is presented elsewhere.¹⁻⁷ Spin-labels can provide information concerning structure and conformational changes because they are particularly sensitive to molecular motion and orientation, as well as to electric and magnetic environments. All these features can be found in biological systems, and they can be detected and monitored by using an electron paramagnetic resonance (EPR) spectrometer.

The spin-label technique or method has been applied to a variety of biological systems, but polynucleotides have received little attention despite publication of an initial study in 1967.8 Compared to other biosystems, nucleic acids present a particular challenge because they contain essentially only four types of monomers or bases (G, C, A, U, and/or T), which most of the time are either stacked or hydrogen-bonded to one another. Thus, specific spin-labeling of a particular site or base becomes almost impossible!

Because of this difficulty, most specific labeling studies of nucleic acids have concerned transfer RNAs. in which some rare bases are selectively reactive toward certain spin-label reagents. This Account treats this topic in depth and gives less exhaustive attention to investigations of DNA and RNA molecules. Spin-labeled mononucleotides were reviewed recently by Gaffney⁹ and are not covered here. Numerous articles or books treat the general topic of biological applications of spin-labeling.¹⁰⁻¹⁹

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Hermann Dugas received his B.Sc. (1964) from the Universite de Montreal and his Ph.D. (1967) in organic synthesis from the University of New Brunswick. After 1 year of postdoctoral work at New Brunswick and 2 years at the National Research Council of Canada in Ottawa, he joined the staff of the Departement de Chimle of the Université de Montréal in 1970. Professor Dugas's research interests relate to the study of structure and function of biological systems using NMR and EPR techniques.