Polyoxygenated Steroids of Marine Origin

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I. Introduction

Sterols of marine organisms have been found to be comprised of most complex mixtures. Over 200 new monohydroxysterols have so far been identified from marine sources, and evidence has been presented for the occurrence of over 70 sterols from a single marine species.¹ Sterol patterns in marine invertebrates reflect the complexity of sterols arising through food chains. The capability of further biochemical modification of dietary sterols makes the sterol mixtures even more complex. The symbiotic relationship between organisms also complicates the sterol compositions.²

In the last 20 years many sterols with unprecedented structures have been isolated from marine sources. For many years the known carbon skeleton of sterols ranged from C_{27} to C_{29} , and the carbon variation occurred exclusively in the side chain at C_{24} .³ After the discovery of the C26-sterols, first detected in 1970 from the mollusk *Placopecten Magellanicus*⁴ and later found widespread in marine invertebrates and also in a marine phytoplankton,⁵ thus suggesting that all the marine C26-sterols originate from phytoplankton, a number of "nonconventional" sterols have been reported. These include sterols having side chains modified by the apparent loss of carbon atoms or by the addition of extra carbon atoms at biogenetically unprecedented positions of a normal C₈ side chain, as well sterols with unusual nuclei. The discovery of novel sterols as major components of the extracts of marine organisms is not the rule, although there have been many exceptions, especially from sponges. Typical examples are aplysterol (9) and 24(28)-didehydroaplysterol (10), the first sterols found⁶ with a methyl group at C-26, which have been found as the major sterols of the sponges of the genus Aplysina (Verongia); calysterol (51), the major sterol (90% of the sterol mixture) of the sponge Calyxnicaensis;⁷ petrosterol (48), isolated from the sponge Petrosia ficiformis;^{8,9} strongylosterol (27), the sole sterol of Strongylophora durissima;¹⁰ and xestosterol (15) and sutinasterol (37), apparently the result of four biomethylation, isolated as the predominant sterols of Xestospongia muta¹¹ and a Xestospongia sp.,¹² respectively. A series of 19-nor- 5α -stanols and Anorstanols, both with conventional and nonconventional side chains, again from marine sponges, have been firstly isolated as the major sterol components from Axinella polypoides¹³ and A. verrucosa,¹⁴ respectively. One more interesting case of discovery of a novel sterol as major component of a mixture is dinosterol (11) from Gonyaulax tamarensis,¹⁵ which is the most characteristic sterol of dinoflagellates and possibly the biological precursor of the cyclopropyl sterol gorgosterol (45) found in zooxanthellae-containing hosts.¹⁶ Structures of the nonconventional side chains of marine monohydroxysterols are shown in Figure 1.

Several excellent reviews on the structures, biosynthesis and distribution of marine sterols are available.^{2,55,-62} A paper dedicated to the biosynthesis of marine sterols side chain is published in this issue. This review focuses on polyoxygenated steroids from marine organisms, which are a growing group of metabolites with potential interesting biological and pharmacological activities and is intended to be all inclusive with respect to its coverage. Polyhydroxysteroids have been found in algae, and virtually in every marine invertebrata phyla, *i.e.* Porifera, Coelenterata, Bryozoa, Mollusca, Echinodermata, Arthropoda, and Tunicata as well as in fish.

II. Polyoxygenated Steroids of Marine Algae

Polyoxygenated sterols have been occasionally isolated from red (Rhodophyta) and brown (Phoeophyta) algae, but have not yet been reported from the green (Chlorophyta) ones (Figure 2).

A. Red Algae

Dihydroxy steroids 57 and 58 along with 24,25epoxycholesterol 59 have been isolated as minor components of Asparagopsis armata⁶³ and Rissoella verruculosa;⁶⁴ 57 and 58 have been also isolated from Rhodymenia palmata⁶⁵ and the diol 58 has been found



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Luigi Minale was born in Tripoli, Libia, in 1936 and obtained his Laurea in Chemistry (1960) and Libera Docenza (1967) from the University of Naples. From 1961 to 1969 he was at the Institute of Organic Chemistry of the University of Naples with Professors R. A. Nicolaus and M. Piattelli. In 1969, he moved to the Institute for the Chemistry of Molecules of Biological Interest, C.N.R., Arco Felice and became Director of this institute in 1973 (until 1981) as well as, almost at the same time, Professor of Organic Chemistry at the University of Catania (1975). In the 1977 he moved to the University of Napoli Federico II, Faculty of Pharmacy, and became Director of Department of the Chemistry of Natural Products in 1984 (until 1990). He has been visiting Professor at the University of Aberdeen (Scotland, U.K.; 1973) and at the Tohoku University, Sendai (Japan; 1986). Professor Minale has published more than 200 papers in the chemistry of natural products and organic chemistry. He is member of the Advisory Board of the Gazzetta Chimica Italiana.

in Liagora distenta⁶⁶ and Scinaia furcellata.⁶⁶ It has been suggested that these side chain oxygenated sterols may be artifacts caused by autoxidation during the isolation process. Francisco *et al.*⁶³ have shown that 57 and 58 can indeed be produced by the autoxidation of desmosterol (cholesta-5,24-dien-3 β -ol); they are nevertheless present in the fresh red alga A. armata.

Two ecdysone-like sterols, pinnasterol (60) and acetylpinnasterol (61), both showing biological activity as molting hormones, have been isolated from Laurencia pinnata.⁶⁷ The structures were secured by X-ray crystallography of the acetylpinnasterol (61). In contrast with ecdysone and the many ecdysteroids isolated from arthropods and certain plant species, all having the 22R configuration, which is very important for high



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activity, the algal pinnasterol and acetylpinnasterol possess the 22S configuration. The same configuration has been found in $(20R, 22S)-5\alpha$ -cholesta-9(11),24(25)diene-3 β ,6 α ,20,22-tetrol, the aglycon of the asterosaponin protoreasteroside (497) isolated from the starfish Protoreaster nodosus and Pentaceraster alveolatus.⁶⁸

 5α -Cholestane-3,6-dione (62) has been reported from some red algae species,⁶⁹ and recently the novel 11α hydroxy- 5α -cholestane-3,6-dione (63), of potential utility as corticosteroid intermediate, has been isolated from *Acanthophora spicifera*.⁷⁰ Another diketosteroid, 5β cholest-3-ene-7,11-dione (64) was isolated from *Hypnea musciformis*.⁷¹ Recently 2α -oxa-2-oxo- 5α -hydroxy-3,4dinorcholestane (65), the first example of a ring A-dinorsteroid from a natural source, has been isolated from *Laurencia obtusa*⁷² as a minor component. The structure was confirmed by synthesis from 2-(hydroxymethylene)cholest-4-en-3-one and the 5α -hydroxy stereochemistry was determined by NMR experiments.

B. Brown Algae

The first dihydroxysteroid isolated from a marine source was saringosterol (66) identified by Ikekawa *et al.*⁷³ in two brown algae. Since then 66 and 24oxocholestane (67) have been isolated⁷⁴⁻⁷⁶ from a number of brown algae. Recently the presence of 66 and 67 along with 24,28-epoxyfucosterol (68) in the brown alga *Hizikia fusiformis* has been reported.⁷⁷ Their origin was considered as doubtful: they could have been *bona fide* constituents of the algae or be artifacts caused by oxidation during the isolation process. Knights⁷⁴ showed that the presence of saringosterol (66) and the 24-ketosterol 67 in the brown alga *Ascophyllum nodosum* were artifacts arising from autoxidation during the air-drying of the alga, the fresh material containing only fucosterol. An intriguing steroid diol with a novel side chain 69 was isolated from *Desmarestia aculeata*.⁷⁸ An unusual sterol with a 24-(vinyloxy) 23-ene side chain 70 was isolated from *Sargassum thumbergii*⁷⁹ as a minor component, the predominant sterol of the mixture being fucosterol. Artifactual formation of 70 appears very unlikely and the authors have suggested a biogenetically possible process involving the removal of some leaving group at C-23 of fucosterol 24,28-epoxide, with simultaneous cleavage of the C-24,28 bond.

Steroids containing a 3,5-dien-7-one and 3,5-diene 7α -hydroxycholestane nucleus were isolated from *Fucus* evanescens.⁷⁵ Ikekawa and colleagues have suggested that the 3,5-dien-7-ones are probably formed during the isolation procedure from the 3β -hydroxy-5-en-7-one counterparts.⁷⁵ We would note that the steroid Δ^5 - 3β , 7α - and Δ^5 - 3β , 7β -diols, along with the Δ^5 - 3β -hydroxy 7-ones, are well recognized autoxidation products of Δ^5 -sterols.⁸⁰ thus leaving the origin of the nuclear oxygenated sterols of *F. evanescens* as doubtful.

C. Microalgae

The Haptophyceae are microscopic unicellular algae, which are widely distributed in the ocean and often constitute a major proportion of marine phytoplankton. A novel sterol sulfate, with the dinosterol side chain, hymenosulfate, 71 has been recently isolated from the cultured marine haptophyte *Hymenomonas* sp.⁸¹ Sulfation is typical in the biosynthesis of secondary metabolites in many marine invertebrates, especially echinoderms, but hymenosulfate is the first report of a sterol sulfate from marine microalgae. This steroid has a potent SR (sarcoplasmic reticulum) Ca-releasing activity.

III. Polyoxygenated Steroids of Porifera (Sponges)

The sponges have yielded the most varied and biogenetically unprecedented array of sterols found among the invertebrate phyla. Most of the 200 new monohydroxysterols found in marine organisms have been isolated from the sponges.⁶² The uniqueness of sterols in the cell membranes of sponges is probably related to the presence of unusual fatty acids in their phospholipids.⁸²⁻⁸⁵

In addition to the monohydroxylated sterols, a number of polyoxygenated sterols have been isolated from sponges; most of them have appeared in the literature in the last two years.

Halistanol sulfate (72) isolated from Halichondria moorei⁸⁶ is the first example of the growing list of sulfated polyhydroxysteroids from sponges (Figure 3), which are very attractive because of their potential activity against HIV virus.^{87,88} Both the tert-butyl moiety at the end of side chain and the 2β , 3α , 6α trihydroxy functions in halistanol are biosynthetically intriguing. Another trisulfated steroid, sokotrosterol sulfate (73), has been isolated together with the previous 72 from two Halichondriidae species.⁸⁹ The side chain of 73 is unprecedented and involves a "normal" alkylation at C-24 of a standard C_8 side chain and the addition of two extra methyl groups at C-26 and one extra methyl group at C-25. Halistanol and sokotrosterol belong to an unusual class of steroids which possess side chains with quaternary carbons. The side chain of 73 is related

to that of mutasterol (26), and the authors⁸⁹ have proposed a biosynthesis via three consecutive SAM biomethylations of codisterol (24S-methyl-5,25-cholestadien-3 β -ol) or epicodisterol (24R). Recently in a continuing search for the biosynthesis of marine lipids. Djerassi and his group⁹⁰ have elucidated the biosvnthesis of mutasterol in the Caribbean sponge Xestospongia muta by feeding selected radioactive precursors and have shown that codisterol is efficiently transformed (10 times faster than its epimer) into mutasterol (26). Interestingly, the isolation of the 26-norsokotrasterol sulfate (74) with the side chain very similar with that of mutasterol (26), the difference being in the position of the double bond, from the sponge Trachyopsis halichondroides (family Halichondriidae) has been reported.⁹¹ Very recently Fusetani and his group⁹² reported the isolation of four more steroids (75-78) with the halistanol sulfate nucleus from the sponge Epipolasis sp. The unusual side chain of halistanol sulfate B (76) is unprecedented for naturally occurring sterols.⁶² A fifth new sulfate steroid isolated from the same sponge, halistanol sulfate E(79) is related to 72 by the introduction of an additional hydroxyl group at C-15.92 Two more steroids with the halistanol sulfate nucleus (80 and 81), showing anti-HIV activity, have been isolated from the sponge Pseudoaxinissa digitata (order Axinellida, family Axinellidae).93

Three novel sterol sulfates (82-84), sharing the same tetracyclic nucleus with a sulfate at C-3 and the 19methyl group oxidized to a carboxyl group, have been isolated from the sponge Toxadocia zumi.94 Antimicrobial, antifeedant, and cytotoxic properties were reported for these steroid sulfates, and the authors suggest that they might be in part responsible for the lack of fouling organisms on Toxadocia zumi.94 A 3βsulfoxy-4 β -hydroxypregnane (85) has been isolated from the sponge Stylopus australis.⁹⁵ A novel group of sterol sulfates (86-90) was discovered as antiviral substances in the sponge Petrosia weinbergi.^{87,88} The structure of the major weinbersterol disulfate A (86) was determined by a 2D INADEQUATE NMR experiment performed on a 150-mg sample in conjunction with its HMBC, HETCOR, COLOC, and COSY data, which established the complete connectivity for the molecule.⁸⁷ Weibersterol disulfate B (87) is related to A (86), but lacks the hydroxyl group at C-20 and a further hydroxyl group is located at C-18. Both compounds possess an unprecedented cyclopropane-containing side chain, showing one more examples for the diversity in the side-chain structures of sponge sterols, and exhibited in vitro activity against the feline leukemia virus. Weinbersterol disulfate A (86) also showed activity against the human immunodeficiency virus. Three minor sterol sulfates (ortho ester disulfates A-C, 88-90) in the same sponge were isolated, and their structures were found to have an ortho ester functionality.⁸⁸ They appear the first reported examples in the steroid class of this particular combination of functionalities. It is of interest to note that the trans diaxial disulfate $(2\beta, 3\alpha)$ and the trans AB ring juncture (5α) H) have been previously encountered in the 5α cholestane 2β , 3α , 26-triol sulfates (530) isolated from an echinoderm, the ophiuroid Ophiaracna incrassata.⁹⁶

An array of polyoxygenated sterols have been isolated from sponges of the genus *Dysidea* (Figure 4). The



1, Damiriana hawaiana¹⁷ (sponge)



5, occelasterol Pseudoptamilla occelata¹⁹ (anellid)



9, aplysterol Aplysina aerophoba⁶ (sponge)



13, Gonyaulax monilata²⁴ (dinoflagellate)



17, Eutreptia viridis²⁷ (euglenid)



21, stelliferasterol Jaspis stellifera³⁰ (sponge)



25, Sinularia ramulosa³³ (soft coral)



29, pulchrasterol Aciculites pulchra36 (sponge)



- 2, Placopecten magellanicus¹⁸ 3, Placopecten magellanicus¹⁸ 4, Placopecten magellanicus¹⁸ (scallop) (scallop)



6, dinoflagellates²⁰



10, Aplysina aerophoba⁶ (sponge)



14. ficisterol Petrosia ficiformis²⁵ (sponge)



18, Pseudoaxinissa sp.²⁸ (sponge)



22, isostelliferasterol Jaspis stellifera³⁰ (sponge)



26, mutasterol Xestospongia muta³⁴ (sponge)



30, Pseudoaxinissa sp.³⁷ (sponge)



7, Halichondria panice a^{21}

(sponge)

11. dinosterol

Gonyaulax tamarensis¹⁵

(scallop)

8, Pseudoaxinella lunacharta²² (sponge)



12, peridinosterol Peridinium foliaceum²³ (dinoflagellate)



16, verongulasterol Verongula cauliformis²⁶ (sponge)



20, Verongula cauliformis²⁹ (sponge)



24, siapalosterol Sinularia siapalosa³² (soft coral)



28, durissimasterol Strongylophora durissima 35 (sponge)



31, Pseudoaxinissa sp.³⁷ (sponge)

32, Strongylophora durissima³⁵ (sponge)



(dinoflagellate)

15, xestosterol Xestospongia muta¹¹ (sponge)



19, Pseudoaxinissa sp.²⁸ (sponge)



23, reimersterol Sinularia remei³¹ (soft coral)





(sponge)



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33, Strongylophora durissima³⁵ (sponge)



37, sutinasterol Xestospongia sp.¹² (sponge)



41, glaucasterol Sarcophytum glaucum⁴² (soft coral)



45, gorgosterol zooxanthellae46,47



49, Crysophyta sp.⁵⁰ (algae)



53, Calyx podatypa⁵²

(sponge)



34, xestospongesterol Strongylophora durissima³⁸ (sponge)



38, Halichondriidae sponges⁴⁰



42, Siphonoborgia sp.43 (soft coral)



46, Calyx nicaensis⁴⁸ (sponge)



50, hebesterol Petrosia hebes⁵¹ (sponge)



54, Calyx nicaensis53(sponge)

55, Calyx nicaensis⁵³ (sponge)

56, Callyspongia diffusa⁵⁴ (sponge)

Figure 1. Nonconventional side chains of marine monohydroxysterols. All compounds possess the cholest-5-ene tetracyclic nucleus except 12 and 13 (4α -methylcholestane nucleus), 14 (5α -cholestane nucleus), 17 (cholesta-5,7-diene nucleus), and 29 (cholest 7-ene nucleus).

first example of this group of steroids is 9α , 11α epoxycholest-7-ene- 3β , 5α , 6β , 19-tetrol 6-acetate (91) isolated from an unidentified species of Dysidea



35, Strongylophora durissima³⁹ (sponge)



39, Xestospongia sp.¹² (sponge)



43, Cystoseira sp.44 (algae)



47, nicasterol Calyx nicaensis⁴⁹ (sponge)



51, calysterol Calyx nicaensis⁷ (sponge)



36, Xestospongia sp.³⁹

(sponge)

40, Spheciospongia sp.⁴¹

(sponge)

48, petrosterol Petrosia ficiformis^{8,9} (sponge)



52, Calyx nicaensis 48 (sponge)







collected in Guam.⁹⁷ The 19-hydroxyl group was

previously found in one sterol from the soft coral

Lithophyton viridis⁹⁸ and later in many sterols from



Figure 2. Polyoxygenated sterols from marine algae.

Dysidea. The 9,11-epoxide is the most novel aspect of this sterol. The shape of the 11-H proton signal (doublet with J = 4.7 Hz) in the ¹H NMR of 91 was the key argument supporting the 9α , 11α stereochemistry, as the 3.5-Hz coupling between H-6 and H-7 was considered by the authors⁹⁷ more compatible with a 6β -acetoxy than with a 6α -acetoxy stereochemistry. The stereochemistry at C-6 has subsequently been revised by Fujimoto *et al.*⁹⁹ as the 6α -isomer through further NMR measurements involving the evaluation of the pyridine-induced shifts of the protons at C-4 of the corresponding 6-ol.

Herbasterol (92), a polyhydroxylated 9,11-secosterol, is responsible for the ichthyotoxic and antimicrobial activities observed in the methanol extract of Dysidea herbacea.¹⁰⁰ The structure of **92** was elucidated by interpretation of spectral data and conversion, on either acid or basic treatment, into 19-norherbasterol (**93**). During the retroaldol reaction, the stereochemistry at C-6 undergoes inversion, and the authors¹⁰⁰ explained this phenomenon by assuming an A/B cis ring junction in herbasterol (**92**), the inversion occurring a C-10 during the retroaldol reaction. A further toxic polyoxygenated steroid **94** has been isolated from the Mediterranean Dysidea tupha.¹⁰¹ A group of eight new polyhydroxysterols, all showing a common 5 α -cholest-7-ene-2 α ,3 β ,5,6 β ,9 α ,11 α ,19-heptol framework **95** and various conventional side chains, have been isolated



Figure 3. Steroid sulfates from sponge.

from Dysidea etheria,¹⁰² collected in Bermuda. The 6β -hydroxy stereochemistry has been confirmed by evaluating the pyridine-induced shifts of the protons at C-4 according to the method suggested by Fujimoto et al.^{99,103} Specifically, the chemical shift of the 4β -proton at δ 3.12 requires the hydroxyl group at C-6 to be in the β -configuration. The major sterol of this group 95 with $R_1 = R_2 = Ac$ and the cholesterol side chain was active in the cytotoxic KB assay.¹⁰² Three minor polyhydroxylated sterols 96-98 in the same sponge were isolated and their structures were found to be analogous to those discovered earlier 95, but distinguished by a

 5β -skeleton,¹⁰⁴ which is a rare feature among sterols from sponges. Herbasterol (92) from Dysidea herbacea¹⁰⁰ and a series of coprostanols from Petrosia ficiformis¹⁰⁵ are the only other examples of 5β -sterols from Porifera. The authors¹⁰⁴ have suggested that both the $5\alpha,6\beta$ - and $5\beta,6\alpha$ -hydroxylated series cooccurring in the same sponge could arise from a $5\alpha,6\alpha$ -epoxide common precursor. More recently another group of polar sterols **99–102** have been discovered in a specimen of Dysidea herbacea,¹⁰⁶ collected near Massawa, Ethiopia. Strong support for the assignment of configurations at C-6 of the two epimers **100** and **101** was



Figure 4. Polyhydroxysteroids from sponges of the genus Dysidea.

derived from the comparison of the ¹H NMR data. The small coupling constant between H-6 and H-7 (<1 Hz) in 100, when compared with J of 5.5 Hz observed in the spectrum of 101, determined the configuration 6α -OAc in 100 and the 6β -OAc in 101. The NOE observed between 19-Me and H-6 in 100 reinforced these assignments. Extensive NOE measurements also yielded data that allow the stereochemistry of 102 to be determined.¹⁰⁶ Two other polyhydroxysteroids 103 and 104 were reported from the sponge *Dysidea fragilis*¹⁰⁷ collected in the Black Sea and their structures were found to be related to **95** earlier isolated from *D. etheria*,¹⁰² but lacking the 19-hydroxyl group. Several other polyoxygenated sterols were isolated from sponges (Figure 5). Three pregnane steroids 105-107 were isolated from *Haliclona rubens*.¹⁰⁸ They appeared to be the first example of pregnane derivatives isolated from marine sources. Two novel unusual pregnanes, agnatasterone A (108) and B (109), pregnatrienolones without much relationship on reference steroids, have recently been discovered in Axinella agnata¹⁰⁹ collected off Roscoff, France, and in Northern area of Ile de Batz. The pregnane structures of 105-109 immediately suggest steroid hormones, but the biogenesis and the role of these sponge-derived pregnane steroids have yet to be elucidated.

Sterols with the Δ^7 -3 β ,5 α ,6 β -triol nucleus of 99 have previously been described from Spongionella gracilis,¹¹⁰ Hippospongia communis,¹¹¹ Spongia officinalis,¹¹¹ and Ircinia variabilis.¹¹¹ Sterols with this nucleus have also been isolated from the bryozoan Myriapora truncata¹¹² and from the scallop Patinopecten yessoensis.¹¹³ Very recently the 24-ethylcholestane- 3β , 5α , 6β -triol and its 6-keto derivative have been isolated from the sponge Spirastrella incostans.¹¹⁴ The 6β-hydroxy stereochemistry in the Δ^7 derivatives was deduced from spectral data and supported by comparison with synthetic model compounds.¹¹³ Spongionella gracilis¹¹⁵ also gave Δ^7 -3 β ,6 α -dihydroxysterols (110), Spongia officinalis¹¹⁶ gave $\Delta^7 - 3\beta, 5\alpha, 6\beta, 9\alpha$ -tetrahydroxysterols (111) and Hippospongia communis^{117,118} has yielded 5.6-secosterols (112) with various conventional side chains. The Δ^7 -3 β .5 α .6 β -trihydroxysterols have been isolated from sponges which contain $\Delta^{5,7}$ -sterols as main sterol components and it has been suggested¹¹⁰ that they could arise from $\Delta^{5,7}$ -sterols. The 5,6-secosterols could be in turn derived from the cleavage of the C-5/ C-6 bond of the Δ^{7} -3 β ,5 α ,6 β -trihydroxysterols.¹¹⁷ We note that the polyoxygenated derivatives of ergosterol characterized by Δ^7 -3 β , 5 α , 6 β -triol and Δ^7 -3 β , 5 α , 6 β , 9 α tetrol partial structures were previously isolated from the terrestial fungus Polyporus versicolor.¹¹⁹

More recently a group of norsterols, incisterols (113 and conventional Δ^{22} side chains), with an unprecedented highly degraded skeleton have been isolated from the Mediterranean sponge *Dictyonella incisa*.¹²⁰ The authors have proposed that they could arise from $\Delta^{5,7}$ -sterols, through the formation of $3\beta,5\alpha,6\beta$ -triols, both of which are present in *D. incisa*, followed by a successive cleavage of the C-5/C-6 linkage to give 5,6secosteroids related to 112. Subsequent cleavage of the C-9/C-10 linkage would result in the removal of the entire ring A including the 19-methyl group. The incisterols, by the authors own admission, are probably artifacts of methanol incorporation, since methanol was used in the extraction. An endoperoxide or a keto acid are suggested as the possible natural products.¹²⁰

Recently a few more secosterols have been recovered from sponges. A new 9,11-secosterol 114 has been isolated from the Mediterranean sponge Spongia officinalis.¹²¹ 9,11-Secosterols were earlier found in a gorgonian¹²² and in a soft coral of the genus Sinularia^{123,124} and later in the sponge Dysidea herbacea.¹⁰⁰ A 8,9-secosteroid, jereisterol A (115), and an 8,14-secosteroid, jereisterol B (116), have been isolated from the Pacific sponge Jereicopsis graphidiophora.¹²⁵ Jereisterol A and B combine the unique 3β -methoxy group with a rare secostructure. Surprisingly this sponge lacks the usual 3β -hydroxysterols, but contains, 3β -methoxysteroids with normal Δ^5, Δ^0 and the rare $\Delta^{8,14}, \Delta^8$ and $\Delta^{7,9(11)}$ nuclei; some others have further oxygenation in the nucleus (117-121).¹²⁶

Anthosterones A (122) and B (123), which represent the first examples of a new type of ring A contraction in a steroid nucleus, have been isolated from the sponge *Anthoracuata graceae*.¹²⁷ The structure of anthosterone A (122) was verified by single-crystal X-ray diffraction analysis.¹²⁷ The authors suggested a possible biosynthesis of the anthosterone nucleus through a benzilic acid rearrangement of a 2,3-diketosteroid precursor as a ring contraction step. Δ^4 -3,6-Diketosteroids 124 were also isolated in the same sponge.¹²⁷ Steroidal Δ^4 -3,6-diketones have been then found in the sponge Geodia cydonium¹²⁸ and Cinachyra tarentina,¹²⁹ where they cooccur with the more common steroidal Δ^4 -3-ketones. $\Delta^{4,7}$ -3.6-Diketones 125 with the conventional 24-methyl-24-ethyl and cholesterol side chains were isolated from Raphidostila incisa.¹³⁰ where they cooccur with a major amount of 5,8-epidioxy- Δ^6 -sterols. Three unusual steroidal $\Delta^{5,8(9)}$ -3 β -hydroxy 7-ketones (126) have been isolated from the sponge Clathrina clathrus.¹³¹ Contignasterol (127), a highly oxygenated steroid with the "unnatural" 14β -proton configuration and cyclic hemiacetal functionality in its side chain. has been isolated from the sponge Petrosia contignata,¹³² collected in Papua New Guinea. The authors acknowledge than the 14β -epimers of a number of semisynthetic 15-ketosteroids were found to be more stable that the corresponding 14α -epimers, and they attempted to epimerize contignasterol (127) with base in order to determine the relative stability of its 14α and 14 β -epimers. Unfortunately all attempts failed. The extraction and chromatography conditions are unlikely to have caused epimerization, and the authors have assumed that 127 exists exclusively as the 14β epimer in the sponge. Thus, contignasterol (127) represents the first naturally occurring steroid with the 14 β -configuration, although steroids with a 14 β -hydroxyl group (e.g. digitoxin) are well known from nature. A few months later Shoji et al.¹³³ reported the isolation of xestobergsterol A (128) and B (129), which are the second examples of steroids with the 14β -H proton configuration, from the sponge Xestospongia bergquistia collected in Okinawa. The structures of xestobergsterol A (128) and B (129) are unique also in that they are pentacyclic steroids with the fifth ring linking C-16 to C-23. Extensive NOE measurements indicated a boat conformation of ring C. Both 128 and 129 strongly inhibited histamine release from rat mast cells induced by anti-IgE.

New $\Delta^{8(14)}-3\beta,7\alpha$ -dihydroxysterols (130) have been isolated from the Mediterranean Pellina semitubulosa.¹³⁴ Rare D-ring unsaturated steroid 3β .16 α -diols (131) have been isolated from the Mediterranean sponge Topsentia aurantiaca.¹³⁵ Sterols with a D-ring unsaturation are rare, having been until now discovered only in the sponge Homaxinella trachys¹³⁶ and from cultured marine dinoflagellates.¹³⁷ A series of Δ^5 -3 β ,7 β and Δ^5 -3 β ,7 α -diols, along with the 3 β -hydroxy Δ^5 -7ones, well-recognized autoxidation products of Δ^5 sterols, have been isolated from Haliclona oculata¹³⁸ from the bay of Funds, Canada, and Stelodoryx chlorophylla¹³⁹ from New Caledonia. They were accompanied by side chain oxygenated sterols, such as 24-oxo-, Δ^{22} -24-oxo-, $\Delta^{24(28)}$ -25-hydroxysteroids, ^{138,139} and short side chain ketones, such as the pregnane derived 20-ketones and the 26,27-bisnor Δ^{22} -24-ketones.¹³⁹ also found in a sponge of genus Hirtio¹⁴⁰ and in Damiriana hawaiana,¹⁷ Haliclona rubens,¹⁰⁸ and Psammaplysilla purpurea.¹⁴¹ Also these side chain oxygenated steroids could be of abiotic origin, deriving through oxidation.

In the echinoderms, sea cucumber and starfish are known to produce saponins. Besides these echinoderms, only a limited number of marine organisms have been shown to contain steroidal or triterpenoid glycosides. Recently, glycosides with cholestane- and но







115









117

сн₃о♥











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116



Figure 5. Polyoxygenated steroids from other sponge species.

lanostane-type aglycons have also been isolated from some sponges (Figure 6). Several norlanostane triterpenoid glycosides, sarasinosides A_1 , B_1 , C_1 (132) have been isolated from Asteropus sarasinosum.^{142,143} They are penta- (A_1, B_1) and tetraglycosides (C_1) containing two amino sugars (NAcGlc and NAcGal) each and have been reported to be ichthyotoxic and to inhibit cell division of fertilized starfish eggs.¹⁴² A 4-methyl steroidal diglycoside 133 has been isolated from Erylus lendenfeldi,¹⁴⁴ collected in the Red Sea. More recently two novel glycosides eryloside C (134) and D (135) have been isolated from a sponge of the genus $Erylus^{145}$ collected at depth of 500 m in south of New Caledonia. The novel lanostane derived aglycon of 134 and 135 features a rare 14-carboxyl group and a 24-methylene-25-methyl side chain. It appears closely related to penasterol, a lanostane-derived metabolite with the 14carboxy group and the cholestane side chain, isolated from the Okinawan sponge Penares sp.¹⁴⁶ Both penasterol and erylosides C and D may have biosynthetic implications concerning the synthesis of sterols in the sponges, in which the loss of the 14-methyl group could involve the intermediacy of a carboxylic acid rather than an aldehyde. A novel steroidal saponin, pachastrelloside A (136), has been isolated from the sponge Pachastrella sp.147 This glycoside inhibits cell division of fertilized starfish eggs.

IV. Polyoxygenated Steroids of Coelenterata

The phylum Coelenterata is subdivided into the classes Hydrozoa, Cubozoa, Scyphozoa, and Anthozoa. The latter class is subdivided into Hexacorallia and Octocorallia. Polyhydroxysteroids have been found in Hydrozoa and Anthozoa.

A. Hydrozoa

Marine hydroids are the simplest Cnidarians, generally characterized by an alternation of generation; they occur in most habitats from the shore to the deep sea. The sessile animals (polyps) grow as bushlike colonies on stones, shells, or seaweeds. Only a few reports on their chemical constituents, including sterols, have appeared, due largely to the difficulties of collection and identifying the biological material. Cholesterol is usually the predominant sterol in hydroids^{2,148} and polyhydroxysteroids have been discovered in species of the genus Eudendrium sp. (Figure 7). Cimino et al.¹⁴⁹ first found 3-oxocholest-4-ene-4,16 β ,18,22(R)tetrol 16,18-diacetate (137) in Mediterranean hydroids of the genus Eudendrium (sp. rameum, racemosus, and ramosum) as well in their predators, several species of nudibranchs (e.g. Hervia peregrina, Flabellina affinis, Coryphella lineata). Other C-18 oxygenated sterols 138-141, containing the rare 2α , 3α -diol functionality, have been isolated from Eudendrium glomeratum,¹⁵⁰⁻¹⁵² collected in the Bay of Napoli. The 2α -deoxy analog 142 has been isolated from the same organism.¹⁵²

B. Anthozoa

1. Hexacorallia

The subclass Hexacorallia comprises organisms such as sea anemones (order Actinaria) and zoanthids (order Zoanthidea), which are characterized by the production sugar chains:



136

Figure 6. Polyoxygenated steroid glycosides of sponges.

of very powerful toxins.¹⁵³ Sea anemone toxins are polypeptides or proteins whose amino acid sequences exhibit a high degree of homology and have become very useful tools for studying the voltage-dependent Na⁺ channel in nerve, cardiac, and muscle cells. Toxins isolated from several Palythoa spp. were the most toxic marine natural products before the discovery of maitotoxin,¹⁵⁴ one of the causative agent of ciguatera, a poisoning caused by ingestion of coral reef fishes. Palytoxin is uniquely distinct from other molecules with which organic chemists have previously dealt in terms of molecular size and of structural complexity; it has molecular weight of about 3000 Da but totally lacks repeating units, such as amino acid, sugar, and fatty acid residues, commonly found in molecules of that size.155

Publications on sterols from Hexacorallia are quite limited and report on the occurrence of conventional Δ^5 -sterols.² Recent papers from Pietra and colleagues¹⁵⁶⁻¹⁵⁸ account for the discovery of the zoanthid

Gerardia savaglia as an unexpected marine source of moulting hormones (Figure 8). The authors first reported the isolation of ecysterone (143),¹⁵⁶ and then of ecdysone (144) and ajugasterone C (145),¹⁵⁷ the latter an ecdysteroid previously thought to be an exclusive and rare plant product. All three compounds were obtained in large amounts. The fourth ecdysteroid isolated as minor component from Gerardia savaglia is the new gerardiasterone 146.158 Ecdysteroids have been identified, in trace amounts, in arthropods,¹⁵⁹ nematodes and anellids,¹⁶⁰ mollusks,¹⁶¹ and, in relative large amounts, in terrestrial plants.¹⁵⁹ The occurrence of ecdysteroids in such large amounts in the zoanthid G.savaglia poses a number of intriguing questions about their origin and role. The authors have noted that G. savaglia, after it had been kept for 15 months in an aquarium, still gave ecdysteroids in roughly the same large amounts as immediately after its collection, 157,158 and they suggested that ecdysteroids can be, at least in part, synthesized within the zoanthid from, possibly,





146, gerardiasterone

Figure 8. Ecdysteroids from the zoanthid Gerardia savaglia (Anthozoa, Hexacorallia).

was isolated from the gorgonian *Pseudopterogorgia* americana together with the known gorgosterol 45.¹²² The structure and the absolute configuration was established by X-ray analysis. Hippurin-1 (149), with a unique oxygenation pattern and a spiroketal ring structure on the side chain of a 24-methylcholestane skeleton, was isolated from *Isis hippuris*,¹⁶³ a common gorgonian found on the Great Barrier Reef. A singlecrystal X-ray analysis of the monoacetate secured the structure and relative stereochemistry of hippurin-1. ORD studies of the 11-keto and 3,11-diketo derivatives established the absolute configuration shown in 149, which is the normal configuration found in sterols. Two related steroids of further complex structural features in the same gorgonian were isolated, and their structures

Figure 7. Polyhydroxysteroids from hydroids of genus *Eudendrium*.

dietary cholesterols. The role of ecdysteroids as hormones in the zoanthid is ruled out because of the high concentration of the steroid, and a possible defensive role has been suggested. Finally, the authors conclude that it is likely that ecdysteroids will prove to be much more widely distributed in the marine environment than was thought.^{156,157} The finding that pinnasterol (60), an ecdysteroid-like sterol, is a constituent of the red alga Laurencia pinnata⁶⁷ might support the hypothesis.

2. Octocorallia

The subclass of Octocorallia comprises different orders among which Gorgonaceae and Alcyonacea are distinguished by their high content of polyhydroxylated sterols, which often are the major steroids. The structures of the polyoxygenated steroids from Gorgonaceae are shown in Figure 9.

 5α -Cholestane- 3β ,5,6 β ,9-tetrol (147) one of first polyhydroxysteroids isolated from marine species was obtained from *Pseudopterogorgia elisabethae*.¹⁶² After the isolation of 147 it was suggested⁵⁶ that a more detailed examination of the polar constituents of gorgonians will result in the isolation of others oxygenated sterols.

The first unique 9,11-secocholestane system was 3β , 11-dihydroxy-9,11-secogorgost-5-en-9-one (148), which







149, R=OAc 152, R=H







150









1**6**0



WR¹

164, R=H or Me or Me, 26-nor

O'



Figure 9. Polyoxygenated steroids of Gorgonaceae (Anthozoa, Octocorallia).

were determined as 3α -acetoxy-11 β -hydroxy-24-methyl-22,25-epoxy-5 α -furostan-18,20 β -lactone (150) and 3α acetoxy-11 β ,18:18,20:22,25-triepoxy-5 α -furostan (151).¹⁶⁴ The eptacyclic steroid 150 was converted to the octacyclic one 151 on reduction with $LiAlH_4$ at reflux followed by acetylation. A minor compound, presumably the 18-epimer was also formed. The 18R-anomer, which requires less strained cis junction of the two fivemembered rings composed of the bicyclic acetal, is the most stable epimer, as confirmed by treatment of 151 with *p*-toluenesulfonic acid in THF at room temperature for 30 h, which showed no signs of epimerization.¹⁶⁴ The authors remark that naturally occurring 18substituted steroids are uncommon. They were only found in the aglycons of holothurins,¹⁶⁵ glycosides of sea cucumbers with an 18,20 lactone functionality on the lanostane skeleton, and in the highly active hormone aldosterone and in its metabolic products.¹⁶⁶ More recently C-18 oxygenated sterols have been isolated from hydroids (Figure 7), sponges of the genus Dysidea (Figure 4), gorgonians (Figure 9) and soft corals of the genus Sinularia (Figure 10). Several related desacetoxy hippurins (e.g. 152) have also been described;¹⁶⁷ five more hippurins closely related to 149 were found in speciments of Isis hippuris collected in the Andaman Islands, India.¹⁶⁸ The same species also yielded a tetrahydroxylated steroid, which was identified as gorgost-5-ene- 3β , 7α , 11α - 12β -tetrol 12-monoacetate (153).169

A pregnane derivative 154 has been isolated from the Mediterranean gorgonian Eunicella cavolini.¹⁷⁰ Four new metabolites, muricin-1 through muricin-4 (155-158) which are unique aminogalactose saponins containing pregnane-derived aglycons have been isolated from the Pacific gorgonian Muricea fruticosa.¹⁷¹ They were supposed to be responsible in reducing fouling on M. fruticosa, possibly through effective growth inhibition of the pennate diatom Phaeodactylum tricornutum. The ecologies and life hystories of M. fruticosa in comparison with those of M. californica, the two major Muricea in California, give support to the potential role of these compounds in contributing to the reduced fouling of M. fruticosa. M. californica, which is found growing together with the morphologically very similar M. fruticosa, was consistently overgrown with typical encrusting plants and animals such as red algae, hydroids, ectoprocts, zoanthids, ascidians, mollusks, and bryozoans. In contrast M. fruticosa was found virtually free of surface fouling organisms. While each contained mixtures of triglycerides, sterols, and fatty acids, only the extracts of \overline{M} . fruticosa were found to contain muricin-1 to muricin-4.¹⁷¹ Another pregnane glycoside, 3β -pregna-5,20-dienyl β -D-galactopyranoside (159) has been isolated from the gorgonian *Pseudoplexaura wagenaari*.¹⁷² The pregna-1,4,20-trien-3-one was previously isolated from an alcyonacean coral *Gersemia rubiformis*.¹⁷³

A reactive sterol, 3,16-dioxocholesta-4,14-diene-15,-20-diol (160) has been isolated from the Mediterranean gorgonian Leptogorgia sarmentosa.¹⁷⁴ On treatment with acetic anhydride and pyridine 160 suffers the loss of the entire side chain through a retro-aldol cleavage. Three polyoxygenated steroids, one being guggulsterol III (161) so far thought to be a plant product, the remaining 162 and 163 being closely related to 161, have been discovered in the same species, L. sarmentosa.¹⁷⁵ The unusual upfield resonance of C-17 in guggulsterol III (161) ($\delta_{\rm C}$ 60.1 ppm) when compared with its acetate derivative weakly downshifted at $\delta_{\rm C}$ 60.4 ppm and of its 16-epimer 163 at $\delta_{\rm C}$ 68.1 ppm is explained as due to the hydrogen linkage between the hydroxyl groups at C-20 and C-16 in 161. A series of five C-18 hydroxylated homologues (164) of guggulsterol III (161) have been isolated from the same organism, L. sarmentosa.¹⁷⁶

A novel secosterol, astrogorgiadiol (165) has been isolated from a gorgonian Astrogorgia sp.,¹⁷⁷ as an inhibitor of cell division in fertilized starfish eggs. The authors suggested that 165 is likely biosynthesized from cholestanol via dienol-phenol rearrangement and cleavage of the 9,10-bond. Two more secosterols, closely related to astrogorgiadiol (165) have been reported from the gorgonian Caligorgia sp.¹⁷⁸

Two novel glycosides named dimorphoside A (166) and B (167) have been isolated from the gorgonian *Anthoplexaura dimorpha*,¹⁷⁹ widely distributed along the southern coast of Japan, as the major active constituents (inhibition of the development of fertilized sea urchin eggs) of that organism. The oxidation of the C-19 methyl group to carboxylate is a rare feature, although related steroid sulfates have been isolated from a sponge.⁹⁴

Sinularia sp. (Alcyonacea) has given three new 9,-11-secosterols with the same 9,11-seconucleus as 148 and conventional 24-methylene, 24-methyl, and cholestanol side chains (168-170).^{123,124} The same species also yielded a very minor component, which proved to be the C-8 epimer of 3β ,11-dihydroxy-24-methylene-9,11-secocholest-5-en-9-one (171).¹²⁴ According to the authors it is unlikely that the less stable 8α -H epimer is an artifact of the isolation procedure. The fact that a group of 9,11-secosterols with a variety of side chains has been discovered suggest that Sinularia species probably cleaves ring C of exogenous sterols similar



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Figure 10. Polyoxygenated steroids from Alcyonacea (Anthozoa, Octocorallia) of genus Sinularia.

to the situation observed in some sponges, e.g. 5,6secosterols in $Hippospongia\ communis$,¹¹⁷ incisterols from $Dictyonella\ incisa^{120}$ and the ring A contraction of Achantella acuta sterols.¹⁸⁰ In addition to the 9,11-secosterols, the soft corals of the genus *Sinularia* have been shown to produce a series of new polyhydroxylated sterols (Figure 10). The common 3β , 5α , 6β -trihydroxy sterols were isolated from

Sinularia dissecta.¹⁸¹ A novel group of eight polyhydroxylated sterols, all of them possessing an 11α hydroxy substituent of potential utility as corticosteroid intermediates have been then isolated from the same specie.¹⁸² Seven of these compounds possessed identical $1\alpha, 3\beta, 11\alpha$ -trihydroxycholest-5-ene nuclei (172), but differed in the side chains, which, in addition to the conventional C_8 and C_9 side chains, include those of gorgosterol (45) and 23-demethylgorgosterol (44).¹⁸³ In continuation with the examination of the extracts of S. dissecta the same authors isolated a major group of five sterols with the same $1\alpha, 3\beta, 11\alpha$ -hydroxylation pattern as well as an acetylated group located at C-18. 173.¹⁸³ In addition the same organism also yielded the analogous 13-formyl, 174, and 13-carboxyl, 175, derivatives.¹⁸³ The corresponding 5α ,6-dihydro derivatives of the major 24-methylene side chain sterols have also be found 176-179. The isolation of three groups of C-18 functionalized sterols from the same organism is so far unprecedented. The authors remark that the fact that similar nuclei possessing diverse side chains are encountered suggests the existence of enzyme systems-either in the coral or in some symbiont-that introduce the 1α - and 11α -hydroxyl groups into a dietary precursor and that may also functionalize the C-18 angular methyl group.¹⁸³ Eventual isolation of the enzyme systems may provide a possible practical route to otherwise rare sterols-notably those with oxygenated C-11 and C-18 positions.

Two new cholestadienone derivatives, 180 and 181, which possess epimeric cyclic ketals of the C-22 side chain position involving the oxidation of the C-18 methyl group, have been isolated from an apparently undescribed species of *Sinularia* genus.¹⁸⁴ Accurate ¹H NMR studies indicated the conformations of the six-membered ketal rings to be a twist boat in 180 and a chair in 181. The variation in the ring conformation was then related to the stereochemistry of the ketal carbon C-22. Similar sterols were described from the gorgonian *Isis hippuris*.^{163,164}

Two novel polyhydroxysterols, numersterols A and B, have been isolated from the South China Sea Sinularia numerosa, and their structures were determined as 24-methylenecholestane- $1\alpha.3\beta.5\alpha.6\beta$ -tetrol (182) and 25-methylene-22-homocholestane- 1β , 3β , 5α triol (183).¹⁸⁵ The structure of numersterol A (182) with nuclear hydroxylation pattern differing from that encountered in many polyoxygenated steroids from soft coral (e.g. Sarcophyton glaucum, 186-188 Lobophytum pauciflorum,¹⁸⁹ and Sclerophytum sp.¹⁹⁰) only in the stereochemistry at C-1, was confirmed by X-ray analysis.¹⁸⁵ The side chain of numersterol B (183) is apparently the result of two biomethylations at C-27 of normal cholesterol side chain. The corresponding 6-keto derivative of numersterol A (182) has been recently isolated from Sinularia microclavata,¹⁹¹ one of the most typical species found in Indo-Pacific coral reefs, collected at Ishigaki Islands, Okinawa. Sinularia crispa from the East coast of Sri Lanka has been shown to contain a novel steroid glycoside (184), showing spermatostatic activity.¹⁹²

Figure 11 shows the structures of polyoxygenated steroids found in Alcyonaceans other than those found in the genus *Sinularia*.

A common feature of many polyhydroxysterols from soft corals is the 3β , 5α , 6β -trihydroxy moiety. The first of the soft coral sterols found to have this pattern was 185 isolated from Sarcophyton elegans,¹⁹³ then found as 25-hydroxy analog 186 also in Sclerophytum sp.¹⁹⁰ Sclerophytum sp. have been shown to contain also the rare (24S)-24-methylcholestane-3 β ,5 β ,6 α ,25-tetrol (187),¹⁹⁰ isomeric with the common 186 and differing from that in the stereochemistry at C-5 and C-6. On pyridinium chlorochromate (PCC) oxidation followed by dehydration with thionyl chloride in pyridine, both 186 and 187 afforded the same Δ^4 -3,6-diketo derivative.¹⁹⁰ The polyhydroxysteroid 188 was isolated from the same soft coral Sarcophyton elegans.¹⁹⁴ After that a series of 1β , 3β , 5α , 6β -tetrahydroxysteroids (189–194) have been isolated from Sarcophyton glaucum, 186-188 one of the most common species found in Indo-Pacific coral reefs, collected at Ishigaki Islands, Okinawa, and Lobophytum pauciflorum,¹⁸⁹ from the same Ishigaki Islands, and from a species of the genus Sclerophytum,¹⁹⁰ collected in the Andaman and Nicobar Seas of the Indian Ocean. Except for some of the 5β -sapogenols and for the Δ^5 -ruscogenin,¹⁹⁵ 1 β -hydroxysteroids are rare in nature. It is also to be noted that 194 is the first report on the isolation of a polyoxygenated androstane derivative from marine invertebrates.¹⁸⁷ Among the complex mixture of polyhydroxysteroids of S. glaucum Kobayashi and Mitsuhashi¹⁹⁶ also isolated a minor polyhydroxysteroid and determined the structure of (24S)-24-methyl-5 α -cholestane-3 β ,5,6 β ,25 ξ ,26-pentol (195). Hydroxylation at C-26 is common among polyhydroxysteroids from starfish¹⁹⁷ and the 24-methyl 25.-26-diol side chain of 195 have been found in a steroidal nonaol from the starfish Archaster typicus,¹⁹⁸ but never found before in Anthozoa. More recently new additions to the group of 1β , 3β , 5α , 6β -tetrahydroxysterols have been found in Sarcophyton subviride,¹⁹⁹ soft coral collected from the Katchal Islands of Andaman and Nicobar Coasts. These include the $\Delta^{25(26)}$ -analog of 192. the C-18 hydroxy analog of 189 (196), the 25-O-acetyl derivative of 195, and 197 with the gorgostane skeleton. A new 3β , 5α , 6β -trihydroxy steroid with further hydroxylation at C-22 and C-24 (198) have been isolated from the soft coral Asterospicularia randalli 200 collected at Guam Island. Hydroxylation at C-22 was also found in lobophytosterol (199), depresosterol (200), and three other closely related sterols, 5β , 6β -epoxysterols 201-203 from the Red Sea soft coral Lobophytum depressum.²⁰¹ The occurrence of C-28 oxygenated sterols, which may be intermediate in the demethylation pathway, might be of biogenetic interest.

The soft coral Minabea sp. contains sterol lactones of the withanolide class, three are new C₂₈ compounds, minabeolides 1-3 (204-206), and five are C₂₇ derivatives, minabeolides 4-8 (207-211).²⁰² Instead of the Δ^2 -1keto ring A, very common in plant withanolides, the marine-derived withanolides have a Δ^4 -3-keto ring A. Further sterols with the common 3β , 5α , 6β -trihydroxy moiety feature and a fourth hydroxyl group found in the 7β -position have been isolated in the soft coral Anthelia glauca²⁰³ (212) collected at Laing Island, Papua, New Guinea, and also as 7β -acetate analogs 213-215 in a species of the genus Xenia collected at Zamamijima, Okinawa.²⁰⁴ Gorgosta- 3β , 5α , 6β -triol has been also isolated from the same species.²⁰⁴ The mixture con-





Figure 11. Polyhydroxysteroids from Alcyonacea (Anthozoa, Octocorallia) other than genus Sinularia.

taining the above four sterols exhibited a growthinhibitory activity against B-16 melanoma cells.²⁰⁴

Litophyton viridis is the source of the rare 19oxygenated sterols 216-218.98,205 Compound 216 with the 7 β -acetoxy functionality appeared to be the first report of a naturally occurring 19-hydroxy steroid.⁹⁸ The 5 β ,6 β -epoxide 218 showed antileukemic activity (IC₅₀ 0.5 μ g/mL) against P₃₈₈ leukemia cells *in vitro*.²⁰⁵

The 19-hydroxylated sterols are very rare in nature only found in a sponge⁹⁷ and in the soft coral *L. viridis*^{98,205} and are of interest as a possible intermediate in the biosynthesis of 19-norsterols. Examination of less polar fractions of the extract of *L. viridis* led Bortolotto *et* $al.^{206}$ to isolate 4α -methyl- 3β , 8β -dihydroxy- 5α -ergost-24(28)-en-23-one (219). The structure was secured by X-ray of the *p*-bromobenzoate derivative. At that time the 8β -hydroxy function appeared unique in marine sterols; several polyhydroxysteroids have been later isolated from starfish and the majority of them possess the 8β -hydroxy substituent.¹⁹⁷

The 3,4,5,6-oxygenated pattern discovered in lobosterol (220), isolated first from Lobophytum pauciflo rum^{207} and later in species of the genus Sclerophytum.^{208,209} is unprecedented in natural sterols. This was also the first marine sterol reported to have an A/B cis ring fusion. The available amount of lobosterol was insufficient for a complete chemical elucidation of the structure, and this determination was obtained by X-ray diffraction analysis of the 4-O-p-bromobenzoate.²⁰⁷ In addition to lobosterol (220) another Sclerophytum sp. of soft coral collected off the Andaman and Nicobar coasts has been shown to contain two new polyhydroxysterols, and a mansterol (221) and nicobarsterol (222).²⁰⁹ Both compounds 221 and 222 bear an oxygenated C-21, which is rare in marine sterols except for those found in echinoderm ophiuroids and one starfish. The structure of and amansterol (221) was secured by X-ray analysis, thus confirming the conventional C-20 configuration. Nicobarsterol (222) is a new type of 9,11secosterol with the C-11 oxidized at a carboxyaldehyde level and forms a seven-membered hemiacetal ring with the C-21 hydroxy group. A sterol with the 9,11glycolated structure like 221 could likely be a precursor of 222. Lead tetraacetate treatment of andamansterol (221) gave a 9,11-seco derivative having the same sevenmembered hemiacetal ring as 222, including the 11Rconfiguration.²⁰⁹ Examination of another Sclerophytum sp. resulted in the isolation of a new steroid 223, which is the first example to have C-21 oxidized to carboxvlate.210

In a continuing investigation of the soft corals of the Andaman and Nicobar coasts of the Indian Ocean by Kobayashi and colleagues, three new polyhydroxysterol glycosides were isolated from one sample identified as Alcyonium sp.,²¹¹ and their structures were determined as 24-methylenecholest-5-ene- 3β , 16β -diol $3-O-\alpha$ -Lfucoside (224) and its 7 β - (225) and 7 α -hydroxy (226) derivatives. From the chemotaxonomic viewpoint it may be of interest that this is the second example of Alcyonium sp. containing steroidal glycosides, which are rare in soft corals. Pregnene-type steroidal glycosides were previously isolated from an Okinawan coralreef Alcyonium sp. by Kobayashi et al.²¹² Two of these glycosides, pregnediosides A and B, were characterized as 4-O- β -D-arabinopyranosyl (227) and 4-O- β -D-xylopyranosyl (228) of 3β , 4α -dihydroxy- 5α -pregn-20-ene, respectively, the remaining three being their monoacetates.²¹² The role of the Alcyonium sp. steroidal glycosides is still obscure.

Kingston and Fallis²¹³ have reported the isolation of two unusual C₂₆ steroids, 12β -hydroxy-24-norcholesta-1,4,22-trien-3-one (229) and its acetate 230 from the sea raspberry *Gersemia rubiformis*, an alcyonacean coral from the cold waters off Newfoundland and Labrador. A $\Delta^{1,4}$ -3-ketopregnane 231 was also found in the same organism.²¹⁴

Very few polyhydroxysteroids have been found in species of the class Anthozoa other those found in the order Gorgonaceae and Alcyonacea and are shown in Figure 12. Four new cytotoxic steroids, named stoloniferones were isolated from the stoloniferan Okinawan soft coral Clavularia viridis²¹⁵ (order Stolonifera). They possess the same 11α -hydroxy- 5β , 6β -epoxy 2-en-1-one steroidal skeleton, but differ in their side-chain structures. AX-ray crystallographic analysis of stonoliferone D (235) confirmed the structures and the relative stereochemistries derived from spectroscopic studies, and CD measurements of the 2,3-dihydrostoloniferone C (234) secured the absolute stereostructures. Stoloniferones exhibited growth inhibition against P₃₈₈ leukemia cells. Unusual 20-epicholanic acid derivatives 236-239 were isolated from the sea pen Ptilosarcus guerneyi²¹⁶ (order Pennatulaceae). Characteristic properties of these compounds are (a) the shorter retention time than those of their natural C-20 isomer $(20\alpha$ -H) on GLC analysis and (b) the C-21 methyl protons resonance at 0.1 ppm higher field in the NMR spectra than the 20α -H compounds. The isolated steroids, as methyl esters, were identical with the synthetic 20-epicholanic methyl esters.²¹⁷ The authors analyzed also the sea pen's free sterols and found them to possess the "normal" C-20 stereochemistry.²¹⁷ Thus the free sterols are not the immediate biosynthetic precursor of the 20-episteroids, and their origin is still an open question. Cholestane- $3\beta, 5\alpha, 6\beta$ -triol has been isolated from the sea pen Pteroides esperi²¹⁸ (order Pennatulaceae) and found to have steatotic and cytotoxic activity. A series of highly oxygenated sterols 240-246 have been isolated from Anthipathes subpinnata (order Antipatharia) commonly named black coral.^{219,220} The Δ^{5} -3 β .7 β .19hydroxylation pattern of 246 has been previously found in 216 from the soft coral Litophyton viridis.98 Interesting A. subpinnata produces both 18- and 19hydroxylated steroids; the hydroxylation at C-20 is rare in marine steroids except for the aglycons of the asterosaponins found in starfishes.¹⁹⁷

V. Polyoxygenated Steroids of Bryozoa

Bryozoa are minute filter feeders animals which contain biological active alkaloids and macrolides. Despite the considerable interest in anticancer activities of bryostatins from *Bugula neritina*,²²¹ the chemical studies of bryozoans are proceeding slowly, probably because of the difficulties experienced in collecting sufficient material for analysis. Only two papers on sterols of bryozoans have appeared in the literature, one of which describes the isolation of five polyhydroxysterols, all possessing a common Δ^7 - 3β , 5α , 6β trihydroxy steroidal skeleton and saturated and Δ^{22} unsaturated conventional C₈, C₉, and C₁₀ side chains from *Myriapora truncata*.¹¹² These sterols have also been found in a sponge¹¹⁰ and in a mollusk.¹¹³

VI. Polyoxygenated Steroids of Marine Mollusca

More species of Mollusca have been analyzed for their sterol content and composition than have those of any other phylum, in part because of their nutritional



Figure 12. Polyhydroxysteroids from Octocorallia (Anthozoa) other than Gorgonacea and Alcyonacea.

value.^{2,60,62} Even so, the occurrence of polyoxygenated sterols in mollusks (Figure 13) is relatively rare. The highly oxygenated steroid 137, isolated from three species of the hydroids of the genus Eudendrium, was also found in their predators, the nudibranchs (class Gastropoda) Hervia peregrina, Flabellina affinis, and Coryphella lineata.¹⁴⁹ Recent studies have shown that most dorid nudibranchs utilize organic metabolites obtained from their diets as chemical defences against predation.²²² The steroid 137 seems to be one of the nudibranch metabolites to which no defensive role is ascribed. Two cholanic acid derivatives 247 and 248, with significant antifeedant activity, were isolated from the dorid nudibranch Aldisa sanguinea cooperi,²²³ but were not found in the sponge Anthoarcuata graceae on which it feeds. This nudibranch is apparently obtaining inactive metabolites from its diet and chemically modifying them to produce an active antifeedant.²²³ Two new steroids, diaulusterols A (249) and B (250) were isolated from the skin extracts of the dorid nudibranch Diaulula sandiegensis and were identified by interpretation of spectral data.²²⁴ Diaulusterols are related to pinnasterol (60) and acetylpinnasterol (61). which were isolated from the red alga Laurencia pinnata.⁶⁷ All four steroids share structural features with ecdysones. The 2α , 3α -diol array of diaulusterols are not commonly encountered in naturally occurring steroids. They have only been isolated from the hydroid Eudendrium glomeratum.¹⁵⁰⁻¹⁵² A new cytotoxic epoxysterol 251, active against the L_{1210} cell line, was isolated from the marine gastropod Planaxis sulcatus.²²⁵ This is very similar to the 9,11 α -epoxysterol 91 isolated from a sponge of Dysidea sp. and therefore 251 is suspected to be the precursor of 91.²²⁵ Δ^7 -3 β ,5 α ,6 β -Trihydroxysteroids have also been found in the scallop Patinopecten yessoensis (class Bivalvia), including one, 252, with the rare 9α -hydroxylation.¹¹³ The occurrence of Δ^5 -3 β -hydroxy-7-ketosterols 253 in the prosobranch mollusk Patingera magellanica has been reported.226 The possible abiotic origin of Δ^5 -7-ketones, wellrecognized autooxidation products of Δ^5 -sterols, can be suspected. A series of highly degraded steroids.



Figure 13. Polyhydroxysteroids of marine Mollusca.

named aplykurodins, has been isolated from two ophistobranchs of the genus Aplysia (class Gastropoda). The 4β -hydroxy lactones 254 and 255 have been isolated from A. kurodai;²²⁷ the 4β -acetoxy derivative 256 and the 4-keto analog 257 have been then isolated from the Mediterranean A. fasciata.228 The relative stereostructure of aplykurodine A (255) was deduced by X-ray crystallography, whereas the absolute configuration was determined by application of CD data.227 The interconversion of the 1-9 δ -lactones, aplykurodines, and their isomeric 1-4 γ -lactones has been investigated^{227,228} and the authors concluded that the δ -lactones are the kinetic products, whereas the γ -lactones are the thermodynamic ones.²²⁸ Aplykurodin B (256) and aplykurodinone (257) which were isolated from the external parts of the body of the animals and showed icthyotoxicity and antifeedant activity (feeding deterrence) were considered by the authors to be the defense allomones of A. fasciata against predators.²²⁸

The endocrine system of steroid hormones in mollusks has been established, and many reports on the biosynthesis and metabolism of progesterone, estrogens, and androgens have been published.²²⁹ Mytilus edulis (class Bivalvia), the mussel, had been reported to contain 0.2 pg g⁻¹ by weight of a substance, which showed ecdysone activity.²⁸⁰ The probable role of ecdysteroids in relation to calcification of the shell in mollusk has been suggested.^{161,231}

VII. Polyoxygenated Steroids of Echinodermata

The phylum Echinodermata, which comprises about 6000 living species, is divided in five classes: Crinoidea

(sea lilies and feather stars), Holothuroidea (sea cucumbers or holothurians), Echinoidea (sea urchins), Asteroidea (sea stars or starfish), and Ophiuroidea (brittle stars).

Among the echinoderms, starfish and sea cucumbers invariably contain saponing, which are responsible of their general toxicity. Chemically, saponing derived from sea cucumbers are triterpenoid glycosides whereas those from starfish are steroidal glycosides. The presence of oligoglycosides in both Holothuroidea and Asteroidea classes gives support to the opinion that sea cucumbers and starfish are phylogenetically closely related. In starfish and sea cucumbers, Δ^7 -sterols, which are probably a consequence of the presence of haemolytic saponins, are also predominant, whereas the other three classes contain the usual Δ^5 -sterols.² Haemolysis is a consequence of the abstraction of membrane cholesterol by the saponins and saponins are known to show a much lower affinity for Δ^7 -sterols, thus the presence of Δ^7 -sterols might explain the apparent immunity of starfish and sea cucumbers to their own saponins.²³² Ophiuroids, which have received moderate attention by chemists as compared to the above classes, have been reported to contain a series of sulfated polyhydroxysteroids and two steroidal glycosides. In the past few years a large number of metabolites have been isolated from echinoderms, mainly steroidal glycosides and polyhydroxylated steroids from starfish and triterpene glycosides from sea cucumbers, with cytotoxic, antifungal, and antineoplastic activity. The interest in these compounds has resulted in a number of monographs entirely or in part devoted to this subject. We should mention the works by Hashimoto (1979),²³³ Burnell and ApSimon (1983),²³² Krebs (1986),¹⁵³ Minale, Riccio, Pizza, and Zollo (1986),²³⁴ Quinn (1988),²³⁵ Stonik and Elyakov (1988),¹⁶⁵ Habermehl and Krebs (1990),²³⁶ and Minale, Riccio, and Zollo (1993).^{197,237} The present paper focuses on free and sulfated polyhydroxysteroids and on their oligoglycosides occurring in starfish and ophiuroids.

A. Asteroidea (Starfish)

Most of the work on chemical constituents of starfish has been initially prompted by the discovery of their toxic saponins and interest in their biological properties. In the recent years the structural studies of these molecules has grown up rapidly, largely exceeding the biological studies on individual compounds and almost 250 steroidal constituents, which include free and sulfated polyhydroxysteroids and steroidal glycosides. have been isolated from ca.50 different starfish species, belonging to 14 families and representative of the three major orders (Paxillosida, Valvatida, and Forcipulata) of the class Asteroidea. According to their chemical structures, the steroidal glycosides from starfish have been subdivided into three main groups: the asterosaponins, which are sulfated steroidal penta- and hexaglycosides; the cyclic glycosides, so far only found in two species of the genus Echinaster; and the glycosides of polyhydroxysteroids, which, although unnoticed for a long time, are as widespread as asterosaponins among starfish.²³⁸ These molecules, which usually occur in minute amounts, consist of a polyhydroxylated steroidal aglycon linked to one or two sugar units and can be found in both sulfated and nonsulfated forms. Analysis of the polar extractives of the starfish Tremaster novaecaledoniae has recently led to the discovery of a new class of steroidal glycosides. in which the polyhydroxylated steroidal aglycon present also a phosphate conjugation to which the sugars are glycosidically attached.239

In the present paper we will discuss polyhydroxysteroids, glycosides of polyhydroxysteroids, asterosaponins, and cyclic glycosides, in that order.

1. Polyhydroxysteroids

Starfish appear as the richest source of polyhydroxysteroids. In contrast with other marine phyla, in which polyhydroxysteroids have been isolated from only a limited number of species sometime belonging to the same genus, polyhydroxysteroids are widespread in starfish, where they have been found, usually as complex mixtures, in almost all species examined. More than 80 polyhydroxysteroids from starfish have been reported so far 258-343. The large majority of them possess a $3\beta, 6\alpha$ (or β), $8, 15\alpha$ (or β), 16β -pentahydroxycholestane nucleus, sometimes with additional hydroxyl groups at one or more of positions $4\beta, 5\alpha, 7\alpha$ (or β) and occasionally 14 α . A 26-hydroxyl function with 25Sconfiguration is usually present in the side chain; less commonly the side chain is hydroxylated at C-24 with 24S-configuration or functionalized in various ways. All hydroxyl groups are disposed on one side of the tetracyclic nucleus inducing an amphiphilic character in the molecules. Up to nine hydroxyl groups have been found in the nonols 310-312, 315, and 316, that to the best of our knowledge represent the most highly



Figure 14. 3,21-Dihydroxysteroids 3,21-disulfates from the starfish Euretaster insignis.

hydroxylated sterols isolated from a natural source. Polyhydroxysteroids occur sometimes in sulfated forms, with the sulfate group located at position 3β , 6α , 15α , or 24, and rare examples of phosphated derivatives have been recently isolated from the deep-water starfish *Tremaster novaecaledoniae*.²⁴⁰

Euretaster insignis is a unique example among the many starfish species examined, it is apparently devoid of both common asterosaponins and glycosides of polyhydroxysteroids and also lacks the type of polyhydroxysteroids occurring in all other species. The analysis of its polar extract yielded instead the 3β ,21dihydroxysteroid 3,21-disulfates 258-263 (Figure 14) as a mixture resistant to further attempts of separation. After solvolysis to remove the sulfate groups, the mixture was fractionated by HPLC to afford pure desulfated 259 and two more fractions, still containing mixed dihydroxysteroids, that were resolved in the individual components by acetylation followed by argentic silica gel column chromatography.²⁴¹

Although the large majority of polyhydroxysteroids from starfish (Table 1) possess a 8β -hydroxyl group, the compounds 264–272 lack this distinctive feature. These simpler polyhydroxylated nuclei are easily identifiable by their characteristic ¹H NMR spectral data (Tables 2 and 3). The 3β , $5, 6\beta$ -trihydroxy moiety in 265 is a common element in marine polyhydroxysteroids and is encountered in a large number of polyhydroxysteroids from starfish. Besides the relatively low field shifts of the 19-methyl protons, the occurrence of a 5α -hydroxyl group in 3β -hydroxy- 5α -steroidal nuclei is well evidenced by an ca. 0.5 ppm downfield shift experienced by the 3α -H resonance. Hydroxylation of C-26 is a further common feature in the large majority of polyhydroxysteroids from starfish. ¹H and ¹³C NMR spectral data of epimeric 26-hydroxysteroids show such small differences²⁴⁸ that direct comparison of both stereoisomer is required for uniequivocal assignment of configuration at C-25. For instance in the 500-MHz ¹H NMR spectra of **265** and **269** the signals of the 27methyl protons are observed at δ 0.934 (25S-isomer) and 0.925 (25*R*-isomer) ppm, respectively and equally small differences are observed for carbon signals C-24, C-26, and C-27 in the ¹³C NMR spectra [δ_{C} 34.1, 68.4, and 17.3 for the 25S-isomer **265** and 34.7, 68.6, and 17.1 for the 25R-isomer **269**].²⁴⁰ Since it was observed that

Table 1. Polyhydroxysterols from Starfish

Basic Nuclear Hydroxylation Pattern: $3\beta, 6\beta, 15\alpha, 16\beta$



no.	side chain	R	R1	R ²	sources and references
264	мулон	н	н	Н	Hacelia attenuata ²⁴²
265	м. Дон	ОН	н	Н	Luidia maculata, ²⁴³ Solaster borealis, ²⁴⁴ Myxoderma platyacanthum ²⁴⁵
266	и Долгон	ОН	н	SO_3 -Na ⁺	Myxoderma platyacanthum, ²⁴⁵ Rosaster sp. ²⁴⁶
267	м. Дон	Н	ОН	Н	Luidia maculata ²⁴³
268	м. Дон	ОН	ОН	Н	Luidia maculata ²⁴³
269	и стран	OH	н	Н	Tremaster novaecaledoniae ²⁴⁰
270	и. Дон	ОН	н	SO_3 -Na ⁺	Tremaster novaecaledoniae ²⁴⁰
271	и стран	Н	н	Н	Sphaerodiscus placenta ²⁴⁷



272, from Tremaster novaecaledoniae 240

Basic Nuclear Hydroxylation Pattern: 3β , 6α , $8, 15\alpha$, 16β



no.	side chain	R	R1	sources and references
273	м. Д. Сон	Н	Н	Protoreaster nodosus, ²⁵⁰ Poraster superbus, ²⁵¹ Pentaceraster alveolatus, ²⁵² Asterina pectinifera, ^{253,254} Patiria pectinifera, ²⁵⁵ Patiria miniata, ²⁵⁸ Rosaster sp. ²⁴⁶
274	и тон	ОН	Н	Protoreaster nodosus, ²⁵⁷ Patiria moniata, ²⁵⁸ Pentaceraster alveolatus, ²⁵² Solaster borealis ²⁴⁴
275	И СТАТАТАТАТАТАТАТАТАТАТАТАТАТАТАТАТАТАТА	Н	ОН	as 273, Pycnopodia helianthoides, ²⁵⁸ Solaster borealis ²⁴⁴
276	И СТАТАЛАН	ОН	ОН	as 273, Pycnopodia helianthoides, ²⁵⁸ Solaster borealis ²⁴⁴
277	нородини он 6-O-sulfate	ОН	ОН	Oreaster reticulatus ²⁵⁹

no.	side chain	R	R1	sources and references
278	и стран	Н	Н	Protoreaster nodosus ²⁵⁷
279	и сн	ОН	н	Protoreaster nodosus ²⁵⁷
280	0503 ⁻ Na ⁺	Н	Н	Poraster superbus ²⁵¹
	*			

Basic Nuclear Hydroxylation Pattern: 3β , 6α ,8, 15β , 16β



no.	side chain	R	R1	\mathbb{R}^2	sources and references
281	м. Дон	Н	н	н	Halityle regularis, ²⁶⁴ Nardoa gomophia, ²⁸⁵ Pycnopodia helianthoides, ²⁵⁸ Dermasterias inbricata, ²⁶⁶ Astropecten scoparius ²⁶⁷
282	и стран	ОН	н	н	Halityle regularis ²⁸⁴
283		ОН	н	н	Coscinasterias tenuispina ²⁶⁸
	3.0.sunate				
284	и он	н	он	н	Pycnopodia helianthoides, ²⁵⁸ Asterina pectinifera, ²⁵⁴ Astropecten scoparius ²⁶⁷
285	и. Дон	н	н	ОН	Dermasterias imbricata ²⁸⁹
286	и стори	ОН	ОН	н	Asterina pectinifera, ^{253,254} Patiria miniata, ²⁵⁶ Solaster borealis, ²⁴⁴ Astropecten scoparius ²⁸⁷
287	Маралана он	он	он	н	Asterina pectinifera ²⁵³
	6-0-surrate				
288	И СТАТОН	Н	Н	Н	Dermasterias imbricata ²⁶⁶
289	и.	н	н	н	Dermasterias imbricata ²⁶⁶
290	M. CH	он	н	н	Culcita novaeguineae ²⁶⁹
291	алуулар он	н	н	ОН	Dermasterias imbricata ²⁶⁶
292	м. Дон	ОН	ОН	н	Patiria miniata, ²⁵⁶ Astropecten scoparius, ²⁶⁷ Solaster borealis ²⁴⁴
293	My CH	ОН	ОН	Н	Patiria miniata ²⁵⁶
	6-O-sulfate				



294 from Dermasterias imbricata²⁶⁶

Basic Nuclear Hydroxylation Pattern: 3β , 6β , $8, 15\alpha$, 16β



Basic Nuclear Hydroxylation Pattern: 38,68,8,158,168



26-O derivatization of a signal natural stereoisomer with chlorides of (R)-(+)- and (S)-(-)- α -methoxy- α -(trifluoromethyl)phenylacetic acid (MTPA, the Mosher's reagent) results in significant differences in the ¹H NMR spectra of the two diastereomeric derivatives,²⁴⁹ the assignment of configuration at C-25 has been systematically made by comparison of the diastereotopic 26H₂ signals in the spectra of the (R)-(+)- and (S)-(-)-MTPA esters.¹⁹⁷ As exemplified by 265 and 269, in the spectra of the MTPA esters of the 25S-isomer 265 the 26-methylene proton signals appear much closer in the spectrum of the (R)-(+)-MTPA ester (δ 4.21 ppm, bd) than in that of the (S)-(-)-MTPA derivative (δ 4.13-4.29 ppm, dd), while the reverse occurs for MTPA esters

Table 2.	¹ H NMR	Data of R	epresentative	Polyhy	droxysterols	from Starfis	h
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no.	hydroxylation pattern	3-H	4-H	6-H	7 β- Η	15-H	16-H	18-H ₈	19-H ₈	others
264	3β,6β,15α,16β	3. 5 7 m		3.76 br s		3.80 dd (10.5, 3.0)	4.00 dd (8.5, 3.0)	0.95 s	1.07 s	
265	3β,5,6β,15α,1 6 β	4.04 m		3.50 br s		3.76 dd (10.5, 3.0)	4.00 dd (8.5, 3.0)	0.94 s	1.21 s	
266	3β,5,6β,15α,16β 15-O-sulfate	4.08 m		3.50 br s		4.38 dd	4.33 dd	1.00 s	1.20 s	
26 7	3β,6β,7α,15α,16β	3.60 m		3.62 t (3.5)	3.91 t (3.5)	3.89 dd (10.5, 3.0)	4.04 dd (7.5, 3.0)	0.96 s	1.04 s	
268	3β,5,6β,7α,15α,16β	4.05 m		3.55 d (3.0)	4.02 t (3.0)	3.88 dd (10.5, 3.0)	4.04 dd (7.5, 3.0)	0.96 s	1.17 s	
273	3β,6α,8,15α,16β	3.59 m		3.66 dd (10.0, 3.0)	2.43 dd (12.0, 3.0)	4.06 dd (11.5, 2.5)	4.00 dd (8.5, 2.5)	1.15 s	1.03 s	
274	3 \$,4\$,6 a,8,15a,16\$	3. 46 m	4.28 br s	4.24 m	2.50 dd (12.0, 3.0)	4.09 dd (10.5, 3.0)	4.01 dd (7.5, 3.0)	1.15 s	1.21 s	
275	3 6,6 0,70,8,150,166	3 .58 m		3.81 m	3.81 m	4.16 dd (11.5, 2.5)	4.03 dd (8.5, 2.5)	1.15 s	1.03 s	
276	3 \$,4\$,6 \$\alpha,7\$\alpha,8,15\$\alpha,16\$	3 .46 m	4. 22 m	4.25 dd (11.5, 2.5)	3.88 d (2.5)	4.18 dd (10.5, 2.5)	4.03 dd (8.5, 2.5)	1.15 s	1.21 s	
2 77	$3\beta,4\beta,6\alpha,7\alpha,8,15\alpha,16\beta$ 6-O-sulfate	3.52 m	4.26 br s	5.03 dd (11.5, 2.5)	4.18 d (2.5)	4.18 dd (10.0, 2.5)	4.02 dd (7.0, 2.5)	1.15 s	1.30 s	
281	3 6,6 0,8,156,166	3.62 m		3.74 td (10.5, 4.0)	2.42 dd (12.5, 4.0)	4.40 dd (6.8, 5.6)	4.25 t (6.8)	1.27 s	1.02 s	
282	3 \$,4\$,6 \$,8,15\$,16\$	3.50 m	4.29 br s	4.22 td (10.5, 4.0)	2.49 dd (12.5, 4.0)	4.40 dd (6.8, 5.6)	4.25 t (6.8)	1.27 s	1.19 s	
284	3β,6α,7α,8,15β,16β	3.53 m		3.87 td (12.5, 2.5)	3.90 d (2.5)	4.50 dd (6.5, 5.5)	4.25 t (6.5)	1.28 s	1.02 s	14-H: 1.43 d (5.5)
285	3 6,6 0,8,140,156,166	3. 50 m		3.77 td (10.5, 4.0)	2.14 dd (12.5, 4.0)	4.00 d (6.5)	4. 37 t (6 .5)	1.35 s	1.04 s	
286	3β,4β,6α,7α,8,15β,16β	3.47 m	4.22 br s	4.32 td (11.0, 2.5)	3.98 d (2.5)	4.51 dd (6.5, 5.5)	4.26 t (6.5)	1.27 s	1.19 s	5-H: 1.52 dd (11,2, 3.0) 14-H: 1.42 d (5.5)
295 296 297 302	3 <i>β</i> ,6 <i>β</i> ,8,15 <i>α</i> ,16 <i>β</i> 3 <i>β</i> ,6 <i>β</i> ,7 <i>α</i> ,8,15 <i>α</i> ,16 <i>β</i> 3 <i>β</i> ,4 <i>β</i> ,6 <i>β</i> ,7 <i>α</i> ,8,15 <i>α</i> ,16 <i>β</i> 3 <i>β</i> ,6 <i>β</i> ,8,15 <i>β</i> ,16 <i>β</i>	3.43 m 3.50 m 3.60 m	4.10 br s	3.89 br s 3.72 dd (3.1, 2.9) 4.04 dd (3.1, 2.9) 3.60 dd (3.0, 2.9)	2.46 dd (15.0, 2.5) 3.87 d (3.1) 3.85 d (3.0) 4.03 d (3.0)	4.17 dd (11.2, 7.5) 4.21 dd (10.0, 2.5) 4.21 dd (10.0, 2.5) 4.50 dd (5.0, 6.2)	4.02 dd (7.5, 2.5) 4.000 dd (7.5, 2.5) 4.000 dd (7.5, 2.5) 4.25 t (6.2)	1.15 s 1.15 s 1.15 s 1.30 s	1.20 s 1.15 s 1.43 s 1.18 s	
303	36,46,66,8,156,166	3.55 m	4.08 br s	4.00 dd	4.02 d (3.0)	4.50 dd (5.0, 6.2)	4.23 t (6.2)	1.30 s	1.45 s	

^a Data (δ , ppm) were mostly obtained at 250 MHz from solution in CD₃OD and are referred to the central line of the CHD₂OD signal (δ 3.34 ppm).

Table 3. Selected ¹H NMR Data of Side Chains of Polyhydroxysterols from Starfish

	no.	21-H	22, 23-H's	$26 - H_2$	27-H ₈	28-H's	other
my Y	он 273	0.96 d (7.0)		3.46 dd (10.5, 6.0) ca. 3.32 ^b	0.94 d (7.0)		
	Он 288	1.06 d (6.5) 5.51 dt (16.0, 6.5)	5.60 dd (6.5, 16.0)	3.46 dd (10.5, 6.0) ca. 3.32 ^b	0. 92 d (7.0)		
May	289 `он	1.00 d (7.5)		3.41 dd (11.0, 7.5) 3.63 dd (11.0, 6.0)	1.10 d (7.5)	4.78 br s 4.85 br s	
	278 `он	0.95 d (7.0)		3.40 dd (10.5, 6.0) ca. 3.32 ^b	0.82 d (6.5)	0.84 d (6.5)	
	279 `он	1.04 (7.0)	5.45 m	3.41 dd (11.0, 6.0) 3.56 dd (11.0, 6.0)	0.91 d (7.0)	0 .9 9 d (7.0)	20-H: 2.55 m 24-H: 2.10 m
	293 Сон	1.07 (7.0)	5.54 m	3.42 d (11.0) 3.50 d (11.0)	1.10 s	1.01 d (7.0)	
۵۹۵ مخ لم	s [−] Nat ⁺ 280	0. 96 d (7.0)		0.89 d (7.0)	0.92 d (7.0)		29-H ₂ : 4.06 m (desulfated: 3.60 m)
" My Contraction of the second							,

^a Data (δ , ppm) were mostly obtained at 250 MHz from solution in CD₅OD and are referred to the central line of the CHD₂OD signal (δ 3.34 ppm). ^b Partially overlapping with solvent signal.

of the 25*R*-isomer 269, the resonance being closer in the (S)-(-)-MTPA ester (δ 4.19-4.23 ppm, dd) and more separated in the (R)-(+)-MTPA (δ 4.14-4.28 dd). It has also been observed that derivatization on ring D, as sulfate group at C-15 in 266 and 270, can affect the extent of the separation of the 26-methylene proton signals, thus it is always advisable to assign the configuration by direct comparison of both (R)-(+)-and (S)-(-)-MTPA derivatives.^{240,245}

Compounds 269, 270, and 272, all isolated from the deep-water starfish *Tremaster novaecaledoniae*, represent the only examples of 26-hydroxysteroids from starfish with 25R configuration. 272 is also the only one to exhibit a 3α -hydroxy- 5β -steroidal nucleus, as indicated by the low field signal of the angular methyl

carbon-19 at 26.1 ppm and by the multiplet at δ 3.54 ppm, having the typical shape for a 3β -hydroxymethine signal in a 5β -steroid.

Compounds 273, 275, and 276, with the basic 3β , 6α , 8, 15α , 16β -hydroxylation pattern, were the first polyhydroxysteroids to be isolated from a starfish, the Pacific Ocean species *Protoreaster nodosus*;²⁵⁰ they were later shown to be the most common polyhydroxysteroids from starfish. They exhibited moderate cytotoxic and anticancer activities.^{253,260} The structures of these first steroids were determined on the results of NMR and mass spectral analysis, chemical transformations and related spectroscopic data. Acetylation of 273 followed by oxidation with Jones reagent afforded 3β , 6α , 15α , 26-tetrakis(acetyloxy)-8-hydroxy- 5α -cholestan-16-



 $\label{eq:Figure 15.} Unusual polyhydroxysteroids with a side-chain methyl group oxidized to carboxyl group from the starfish Myxoderma platyacanthum.$

one. The elimination of the side chain with migration of one hydrogen (m/z 464, McLafferty rearrangement)followed by 18-methyl fission (m/z 449) in the mass spectrum was diagnostic of a 16-ketosteroid. Acetylation of 275 and 276 afforded likewise the corresponding tetraacetates $[3\beta, 6\alpha, 15\alpha, 26$ -tetrakis(acetyloxy)], which, on oxidation with Jones reagent, gave the 3β , 6α , 15α ,-26-tetrakis(acetyloxy)-7 α ,8-dihydroxy-5 α -cholestan-16one and 3β , 6α , 15α , 26-tetrakis(acetyloxy)- 7α , 8-dihydroxy- 5α -cholestane-4,16-dione, respectively. Treatment of 273, 275, and 276 with p-bromobenzovl chloride in pyridine led to the formation of the corresponding $3\beta.6\alpha.15\alpha.26$ -tetrakis(p-bromobenzoates) with CD curves displaying strong positive first and negative second Cotton effects ($\Delta_{\epsilon 252}$ +37.2/ $\Delta_{\epsilon 235}$ -20.0) in agreement with the expected positive chirality of the three major dibenzoates interactions $(3\beta/6\alpha, 3\beta/15\alpha, 6\alpha/15\alpha)$ in a cholestane skeleton with 5α -H absolute configuration. The 25S configuration was assigned, as described before, by ¹H NMR pattern of the 26- H_2 signals of the (R)-(+)and (S)-(-)-MTPA derivatives.249

Steroids 274, 278, and 279 were later isolated from the same starfish species, *Protoreaster nodosus*;²⁵⁷ has since been isolated also from other species. The stereochemistry of the side chains of 278 and 279 is suggested after the stereoselective synthesis of the 24methyl-26-hydroxysteroid models.²⁶¹ Compound 280 is a quite rare example of non-glycosylated C-29 polyhydroxysteroid, these compounds have been more commonly found as aglycons in several glycosides of polyhydroxysteroids (see section VII.A.2). The 24*R* configuration is suggested on the basis of the reported chemical shifts of the isopropyl methyls in both ¹H and ¹³C NMR spectra and comparison with model compounds.^{262,263}

Steroids 281-293 share the common 3β , 6α , $8, 15\beta$, 16β hydroxylation pattern, while 294 is a rare example of a 26-hydroxysteroid missing the 16β -hydroxyl function. The inversion of stereochemistry at C-15, with respect to the previous ones, is connected with a slight downfield shift of the 18-methyl proton signal and with major changes of the 7-, 15-, and 16-proton signals, whose shapes and chemical shifts become distinctive features of this structural subgroup. The cis-15,16-dihydroxy stereochemistry was supported by the formation of a 15,16-monoacetonide on treatment of the hexol 281 with Me_2CO and TsOH and by the formation of a 3,4:15,-16-bisacetonide on similar treatment of the heptol 282.264 The chemical shift differences of the 26methylene protons in the 26-O(R) - (+) and 26-O(S) - (-)(-)-MTPA esters (δ 4.26–4.31 dd and 4.20–4.41 dd, respectively) were used to establish the 25S configuration of 292.²⁵⁶ The 14 α -hydroxyl group in 285 and 291 induces substantial modifications in both ¹H and ¹³C NMR spectra vs their 14-deoxy analogs 281 and 289, most significant in the ¹³C NMR spectrum are the large γ -upfield shifts observed for the C-7 (-4.6 ppm), C-9 (-8.9 ppm), C-12 (-5.4 ppm), and C-17 (-10.0 ppm) signals. The absolute configuration of the side chain in 291 has been suggested by comparison of ¹H and ¹³C NMR data with those of the steroid 315 from Archaster tipicus.

The steroids 295-301 originate the structural group having the $3\beta,6\beta,8,15\alpha,16\beta$ -hydroxylation pattern. Most significant modifications of ¹H NMR signals vs their 6α -counterparts refer to the shape and chemical shift of the 6-H signal and to the 19-methyl proton signal, downfield shifted by ca. 0.17 ppm. Compounds 299-301 are very minor components of the Mediterranean starfish *Hacelia attenuata* and their structures with truncated side chains were mainly deduced by interpretation of NMR spectra. The occurrence of these truncated steroids may be of some interest as indicators of the capability of the starfish to oxidize dietary sterols.

Steroids 302-305, all isolated recently from the starfish Solaster borealis,²⁴⁴ represent a further and minor variation in the stereochemical disposition of the nuclear hydroxyl groups. Compounds 302 and 303 are epimeric with the previous 284, 286, 296 and 297 from which they differ alternatively for the stereochemistry at C-6 or at C-15. The relative *threo* configuration at C-24 and C-25 of 305 was assigned by comparison of the NMR spectral data of the side chain with those of synthetic model compounds, while the absolute (24R, 25S) configuration followed from derivatization with (R)-(+)-MTPA chloride and comparison of the ¹H NMR spectrum with those of the 26-O-(R)-(+)-MTPA esters of the two *threo* synthetic models (*i.e.* the 24R, 25S- and 24S, 25R-isomeric pair).

Polyhydroxysteroids 306-309 (Figure 15), isolated from the starfish Myxoderma platyacanthum²⁴⁵ together with 265 and 266, have the same $3\beta,5,6\beta,15\alpha$ tetrahydroxycholestane nucleus with different side chains: 26-acid; Δ^{22} -24-methyl 26-acid; Δ^{22} -27-nor-24methyl 26-acid; and 24-carboxymethyl. In 306 and 307 the carboxylic acid function is found as the amide derivative of taurine. They are examples of the structural variety of steroids cooccurring in the same organism and also constitute the first reported isolation from starfish of steroids with a methyl group oxidized to carboxyl. The absolute configuration of the side chain stereogenic centers in 307-309 was determined



Figure 16. Polyhydroxysteroids from the starfish Archaster typicus.

by spectral comparison with stereoisomeric $\Delta^{22(E)}$ -24methyl 26-acid,²⁶¹ 24-methyl-27-nor 26-acid,²⁴⁵ and 24carboxymethyl²⁶³ steroidal models obtained by stereoselective synthesis. Only very small differences were observed in the ¹H NMR spectra of the two epimeric 24-methyl-27-nor 26-acid synthetic models, mainly dealing with the shifts of the olefinic protons: double doublets centered at δ 5.32 with the internal lines coincident in the 24S-isomer and separated by 1 Hz in the 24R-isomer. The olefinic pattern in the spectrum of the natural **308** was coincident with that of the 24S- model, thus suggesting the reported absolute configuration.

Figure 16 illustrates the variety of highly hydroxylated steroids encountered in Archaster typicus;^{198,272} these steroids have been isolated in relatively large amounts as compared to the very limited fraction of steroidal glycosides in this starfish. Compounds 313 and 318 were shown to be moderately cytotoxic and to cause inhibition of growth of human lymphoma cells at a dose of 0.05 μ g/mL.²⁶⁰ The structures of the nonols 310–312, 315, and 316, the most highly hydroxylated









Figure 17. Polyoxygenated steroids from the starfish Styracaster caroli.

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steroids isolated from natural sources, were determined essentially upon accurate spectral analysis. Major

support in the elucidation of structures 310-312 came from 2D ¹H-¹³C NMR heterocorrelation experiments

Chart I









(HETCOR and COLOC) that, through the long-range correlations with quaternary carbons C-5 and C-14, allowed firm location of tertiary hydroxyl groups and consequent build up of the tetracyclic framework. Stereochemical arrangement of hydroxyl groups was deduced by NOEDS experiments showing throughspace interactions between 6-H and 19-Me. 9-H and 7-H, 15-H and 18-Me. Furthermore, on treatment with acetone and TsOH, compound 310 afforded a $3\beta.4\beta$: 14α , 15α -bisacetonide, thus supporting the 14α -hydroxy stereochemistry. The application of the Horeau's method of kinetic resolution to this bisacetonide allowed the assignment of the 24R configuration to C-24. The assignment of the stereochemistry at C-24 and C-25 in 315 and 316 required NMR spectral comparison with synthetic model compounds.¹⁹⁸ The structure 310, which has the remarkable feature of eight sequential hydroxyl groups protruding from the same side of the molecule, has been confirmed by a single-crystal X-ray diffraction study.²⁷³ The crystal packing of molecules was shown to be a consequence of the amphiphilic character; the hydroxyl groups develop an intricate and extensive network of hydrogen bonds connecting the

molecules through their hydrophilic moieties and giving rise to double layers, which interact through the hydrophobic surfaces.

A group of 3β , 6α -disulfated steroids 319-322 has been more recently isolated from the "living fossile" species Tremaster novaecaledoniae collected at a depth of 530 m during exploration of the bathyal zone off New Caledonia.²⁴⁰ Furthermore, analysis of polar extracts led to the isolation of the steroid 323 and of a new group of glycosides of polyhydroxysteroids, the tremasterols A-C (459-461), all possessing both phosphate and sulfate conjugation.^{239,240} To the best of our knowledge they constitute the first reported isolation of phosphated steroids from a natural source. The presence of the phosphate group at C-6, at first indicated by the shape of the 6-proton signal, was confirmed by the proton noise-decoupled ¹³C NMR spectrum in which the C-5 and C-6 signals appeared as doublets due to the ³¹P-¹³C couplings through two and three linkages and finally by ³¹P NMR. Tremaster novaecaledoniae contains also the uncommon polyhydroxysteroids with 25R stereochemistry 269, 270, and 272.

A further 3β , 6α -disulfated steroid **324**, closely related to the aglycons of the asterosaponins also by the presence of an oxygenated function at C-23, has been isolated from the starfish *Aphelasterias japonica*.²⁷⁴

The very recent investigation of Styracaster caroli, a deep-water starfish collected between Thio and Lifou (New Caledonia) at a depth of 2000 m, revealed the occurrence of a very complex mixture of unprecedented polyhydroxysteroids (Figure 17) with unusual side chains.²⁷⁵ Compounds 327 and 331 show a unique 24ethyl-25-hydroxy-26-sulfoxycholestane skeleton, while compound 328 presents a Δ^{22} -23,24-dimethyl substitution pattern typical of dinosterol (11) and 329 is its 29-hydroxy derivative. An unusual oxygenation pattern is found in the C_{29} side chain of 332, with a sulfoxy group at C-28 and a hydroxyl group at C-29. Compounds 333-335 are further examples of polyhydroxysteroids possessing an amido function in the side chain, with a D-(-)-cysteinolic acid (2S-amino-3-hydroxypropanesulfonic acid) linked to a C-24 carboxyl group.

In a more limited number of cases polyhydroxysteroids with hydroxyl or sulfate groups at C-24 have been isolated, 336-343 (Chart I). The hydroxylation at C-24 is a structural feature commonly encountered in the aglycons of the many steroidal glycosides isolated from starfish and indeed 336 is the aglycon of the glycosides 351-359, while 338 and 339 have been found as the native aglycons in the glycosides 369-383 and 384-388, respectively. The unsaturated 341 is the aglycon of 391-393 and the 15-O-sulfated of 343 has been found in 434. The isomeric relationship at C-15 between 336 and 338 was confirmed by formation of an 8,15-phenylboronate in 338. The 24S configuration was suggested in 338 by ¹³C NMR data in comparison with those reported for 24(S)- and 24(R)-hydroxycholestanol. In fact differences observed between the spectra of the two epimers are very small^{277,278} and in the desulfated 340 the 24S configuration was later confirmed, upon derivatization with (R)-(+)- and (S)-(-)-MTPA, by the Mosher's method for determination of absolute configuration of chiral secondary carbinols.^{279,280} The steroid 341 was only isolated in admixture, with its 22,-23-dihydro congeners 338 resistant to further attempts of separation.

2. Glycosides of Polyhydroxysteroids

This group of steroidal glycosides from starfish shows a large degree of structural variability. Most of these compounds usually occur in minute amounts and are also widespread among starfish, having been found, usually as complex mixtures, in almost all the species investigated. They are composed of a polyhydroxylated steroidal aglycon and a carbohydrate portion mostly made up from one or two monosaccharide units (Figure 18) often linked to each other and glycosidically attached at C-3 or C-24 of the aglycon. Only very recently cytotoxic triglycosides (418, 419, and 421) were isolated from the New Caledonian species Fromia monilis.²⁸¹ They constitute the only examples of triglycosides among more than 100 different mono and diglycosides of polyhydroxysteroids isolated so far. The most common monosaccharides are D-xylopyranose, often methylated at position 2 and/or 4 and occasionally at position 3, and L-arabinose, found in its furanose form. Rare examples of xylofuranosides (371, 372, and

403), galactofuranosides (400, 441, and 444), fucofuranosides (389 and 390), and arabinopyranosides (417 and 420) have also been found.

The first representative of such compounds, the cytotoxic nodososide 431, was first isolated from Pacific Protoreaster nodosus²⁸² and later from other Valvatida species. This was followed by the structures of more than 100 different glycosides of polyhydroxysteroids. Structural variations originate from the hydroxylation pattern of the steroidal tetracyclic nucleus, the functionalization of the side chain, the presence of sulfate, and the nature and location of the saccharide moiety. Beside the invariable 3β -hydroxylation, hydroxyl groups are commonly found at the positions $6\alpha(\text{or }\beta)$, 8, $15\alpha(\text{or }\beta)$ β), and 24 of the aglycon with additional hydroxyl group-(s) at one or more of the positions 4β , 5α , 7α , and 16β . The glycosides of polyhydroxysteroids often occur as complex mixtures with free and sulfated polyhydroxysteroids and asterosaponins; for example the polar steroids from Coscinasterias tenuispina have been resolved into 19 constituents.²⁶⁸ Illustrative examples of the structural variety of steroidal glycosides cooccurring in the same organism are the 11 glycosides of polyhydroxysteroids isolated from the starfish Henricia laeviuscola,283 the eight steroidal diglycosides, the halitylosides, isolated from the starfish Halityle reg*ularis*²⁶⁴ along with two polyhydroxysteroids and four asterosaponins, and the 11 glycosides and five polyhydroxysteroids isolated during a recent reinvestigation of the polar extractives from the starfish Culcita novaeguineae.²⁶⁹

The structures of the currently known glycosides of polyhydroxysteroids are listed in Table 4 with their sources and references. Compounds 344–350 are uncommon examples of steroidal glycosides lacking the hydroxyl group at C-8 of the aglycon, a structural feature present in all other glycosides. Glycosides 351-363 are characterized by the steroidal aglycons with the basic 3β , 6α , 8, 15α -hydroxylation pattern, while in 364-368 the tetracyclic nucleus also bears a 16β -OH. The largest group of compounds presents aglycons with the basic 3β , 6α , 8, 15β -hydroxylation pattern, 369-402, with a single case of Δ^4 -unsaturation in 402. The (24S)-5 α cholestane- 3β , 6α , $8, 15\beta$, 24-pentol aglycon, first encountered in attenuatoside A II (369) from Hacelia attenuata, is the most common steroidal aglycon found in the glycosides from starfish. In the aglycons with the 3β , 6α , 8, 15β , 16β -hydroxylation pattern 403-422 the oxygenated C-28 and C-29 side chains prevail on the common 24-hydroxyl side chain only found in indicoside C (403). Compounds 423-436 constitute the group of glycosides whose aglycons possess the basic 3β , 6β , 8,- 15α -hydroxylation pattern; this group of compounds comprises several examples of those glycosides in which the two monosaccharides are not linked to each other but at positions 3 and 24 of the aglycon. Also in the group with 3β , 6β , 8, 15α , 16β -hydroxylation pattern 437-452 the C-28 and C-29 aglycons are predominant; moreover in the Δ^4 -unsaturated compounds 447-452 the xylosyl unit in constantly found linked at C-3 of the aglycon. The smallest group comprising compounds 453-455 has the basic 3β , 6β , 8, 15β -hydroxylation pattern.

Compounds 456–458 cannot be classified in any of the above structural groups; they contain aglycons with



Figure 18. Saccharide chains found in glycosides of polyhydroxysterols from starfish.

unsaturated tetracyclic nuclei, a quite unusual feature among glycosides of polyhydroxysteroids from starfish, only found in the above mentioned compounds with Δ^4 -steroidal skeleton.





name	no.	side chains	R	R1	R²	R ³	sources and references
	352	0-R ²	н	Н	н	A 3	Oreaster reticulatus, ²⁵⁰ Patiria miniata ²⁵⁶
miniatoside B	353	M	н	Н	Н	XA_1	Patiria miniata ²⁵⁶
crossasteroside B	354	Q-R ²	н	Н	Н	XX3	Crossaster papposus ²⁸¹
crossasteroside C	355		н	Н	Н	XA4	Crossaster papposus ²⁸⁷
borealoside A	356		н	н	н	XA3	So las ter borealis ²⁴⁴
borealoside B	357		Н	н	н	XA2	Solaster borealis ²⁴⁴
borealoside C	358		н	Н	Н	X	Solaster borealis ²⁴⁴
crossasteroside A	359		н	н	ОН	XX3	Crossaster papposus ²⁸⁸
crossasteroside D	360		н	Н	OH	XX2	Crossaster papposus ²⁸⁸
attenuatoside C	361	0.R ²	он	Н	н	\mathbf{A}_1	Hacelia attenuata ²⁸⁹
borealoside D	362	<u>O</u> -R ²	ОН	н	н	X,	Solaster borealis ²⁴⁴
6-epi-nodososide	363	φ. R ²	н	ОН	Н	XA5	Pentaceraster alveolatus ²⁵²

Basic Nuclear Hydroxylation Pattern: 3β , 6α , $8, 15\alpha$, 16β

		RO	он		н		
name	no.	side chains	R	R1	R²	R³	sources and references
	364	И ОН	X2	Н	Н		Poraster superbus ²⁵¹
	365	и он	X2	Н	OH		Poraster superbus ²⁵¹
	366	n , 1	Н	Н	Н	\mathbf{A}_1	Patiria pectinifera ²⁹⁰
miniatoside A	367	*	Н	н	Н	A4	Patiria miniata ²⁵⁶

name	no.	side chains	R	R1	\mathbb{R}^2	R ³	sources and references
asterosaponin P·2	368	⁰ ·ℝ ³	Н	OH	н	A ₂	Patiria pectinifera ²⁹¹

Basic Nuclear Hydroxylation Pattern: 3β , 6α , $8, 15\beta$



			R ¹	ōн			
name	no.	side chain	R	R1	\mathbb{R}^2	R ³	sources and references
attenuatoside A-II	369	*	Н	Н	Н	A ₁	Hacelia attenuata ²⁷¹
scoparioside A	370	۵-R ³	н	н	Н	A۶	Astropecten scoparius ²⁸⁷
scoparioside B	371	0.R ³	Н	н	Н	$\mathbf{X}\mathbf{f}_1$	Astropecten scoparius ²⁶⁷
indicoside B	372	<u>Ū</u> .R ³	н	Н	Н	$\mathbf{X}\mathbf{f}_2$	Astrospecten indicus ²⁹²
pycnopodioside A	373	Q.R ³	Н	н	н	X 1	Pycnopodia helianthoides ²⁵⁸
pycnopodioside B	374	^Q .R ³	So3-Na+	н	н	X 1	Pycnopodia helianthoides ²⁵⁸
pycnopodioside C	375	0-R ³	Н	н	н	Glc ₂	Pycnopodia helianthoides ²⁵⁶
luridoside A	376	0-R ³	Н	н	н	X ₅	Comasterias lurida ²⁹³
scoparioside C	377	0. R ³	Н	н	н	X ₆	Astropecten scoparius ²⁶⁷
attenuatoside A-I	378		Н	н	н	XA5	Hacelia attenuata ²⁹⁴
culcitoside C4	379	₩	Н	н	н	XA4	Culcita novaeguineae ²⁸⁹
halityloside E	380		н	н	Н	XA ₆	Halityle regularis, ²⁸⁴ Nardoa gomophia, ²⁸⁵ Sphaerodiscus placenta, ²⁴⁷ Culcita novaeguineae ²⁸⁶
glacialoside A	381	n,,,,,,,,,,,, ,,,,,,,,,,,,,,,,,,,,,,,,	X ₁	н	н	SO₃-Na+	Marthasterias glacialia ²⁹⁵
distolasteroside D_1	382	<u>□</u> .R ³	X ₁	н	н	X 1	Distolasterias nippon ²⁹⁶

name	no.	side chain	R	R1	R ²	R ³	sources and references
distolasteroside D ₃	383		S 1	Н	Н	Glc1	Distolasterias nippon ²⁹⁶
attenuatoside B-2	384	<u>0</u> . ℝ ³	н	он	н	A ₁	Hacelia attenuata ²⁸⁹
attenuatoside B-1	385	Q-R ³	н	он	н	XA5	Hacelia attenuata ²⁸⁹
culcitoside C5	386	₩,	н	ОН	Н	XA4	Culcita novaeguineae ²⁸⁹
halytiloside D (culcitoside C ₁)	387	₩	н	он	Н	XA ₆	Halityle regularis, ²⁶⁴ C. novaeguineae, ^{269,297} N. gomophia, N. novaecaledoniae, ²⁶⁵ Linkia guildingi ²⁹⁷
glacialoside B	388	₩	SO3-Na+	ОН	Н	X ₁	Marthasterias glacialis ²⁹⁵
imbricatoside B	389		н	н	ОН	QuiFuc <i>f</i>	Dermasterias imbricata ²⁶⁹
imbricatoside A	390	₽ , ₽ ⁰ . R ³	н	ОН	ОН	QuiFuc <i>f</i>	Dermasterias imbricata ²⁸⁹
luridoside B	3 91	⁰ . ℝ ³	н	н	Н	X ₅	Comasterias lurida ²⁹³
22-dehydrohalityloside E (poranoside A)	392	my ∼	н	н	Н	XA ₆	Sphaerodiscus placenta, ²⁴⁷ Porania pulvillus ²⁹⁸
distolasteroside D ₂	393	m <u>→</u> <u>·</u> ·	X 1	н	Н	X 1	Distolasterias nippon ²⁹⁶
22-dehydrohalytilodside D	394	₩	н	он	н	XA ₆	Sphaerodiscus placenta ²⁴⁷
coscinasteroside F	395	"	н	ОН	н	\mathbf{X}_1	Coscinasterias tenuispina ²⁸⁸
halityloside I (6-O-sulphate)	396	му О. R ³	н	он	н	XX4	Halityle regularis, ²⁶⁴ Nardoa gomophia ²⁶⁵
placentoside A	397	M	н	он	н	XX4	Sphaerodiscus placenta ²⁴⁷
culcitoside C ₈	398	R ³ . 0	н	ОН	Н	XA ₆	Coscinasterias tenuispina ²⁶⁸
culcitoside C7 6-sulfate	399	R ³ . 0	н	он	Н	XA ₆	Coscinasterias tenuispina ²⁸⁸
indicoside A	400		н	н	ОН	Galf	Astropecten indicus ²⁹⁹
gomophioside B	401		н	он	н	XA ₆	Gomophia watsoni ²⁷⁶



pisasteroside E 402 Pisaster giganteus³⁰⁰

Basic Nuclear Hydroxylation Pattern; 3β , 6α ,8, 15β , 16β

			Rđ		Т П П	J					
name	no.	side chain	R	R1	\mathbb{R}^2	R ³	sources and references				
indicoside C	403	″	Н	Н	Н	Xf ₂	Astropecten indicus ²⁹²				
coscinasteroside E	404	"уО- №	н	ОН	Н	\mathbf{X}_1	Coscinasterias tenuispina ²⁶⁸				
coscinasteroside D	405	и.,о. R3	Н	н	н	Glc2	Coscinasterias tenuispina ²⁶⁹				
coscinasteroside A	406	и.,О.,R ³	X 1	н	ОН	SO₃-Na+	Coscinasterias tenuispina ²⁶⁹				
culcitoside C_8	407	и. 	н	н	н	XX4	Culcita novaeguineae ²⁸⁹				
culcitoside C3	408	^{R³· 0, ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓}	н	н	н	XA ₆	Culcita novaeguineae ³⁰¹				
culcitoside C2	409	^{R³· 0, Ţ}	н	ОН	Н	XA ₆	Culcita novaeguineae ³⁰¹				
coscinasteroside C	410	₽ ^{3.} 0,	н	н	н	Glc3	Coscinasterias tenuispina ²⁸⁹				
pisasteroside A	411	^{₽³. 0, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,}	н	Н	Н	Glc2	Pisaster ochraceus and Pisaster brevispinus ³⁰²				
pisasteroside F	412	"	Н	н	н	Glc2	Pisaster giganteus ³⁰⁰				
halityloside B	413	°-R ³	н	Н	н	XX4	Halityle regularis, ²⁸⁴ Nardoa novaecaledoniae and N. gomophia, ²⁸⁵ Sphaerodiscus placenta, ²⁴⁷ Culcita novaeguineae ²⁶⁹				

name	no.	side chain	R	Rl	R²	R ⁸	sources and references
halityloside A	414	0. R ³	Н	ОН	Н	XX4	Halityle regularis, ²⁸⁴ Nardoa novaecaledoniae and N. gomophia, ²⁸⁵ Sphaerodiscus placenta, ²⁴⁷ Culcita novaeguineae ²⁸⁹
halityloside H	415	0-R ³	н	ОН	н	XA ₆	Halityle regularis, ²⁸⁴ N. gomophia ²⁸⁵
halityloside H 6-O-sulfate	416	0. R ³	н	он	Н	XA ₆	Halityle regularis, ²⁶⁴ N. gomophia ²⁶⁵
moniloside E	417	0-R ³	Н	он	н	AX	Fromia monilis ²⁸⁴
moniloside I	418	0- R ³	н	ОН	н	XXX ₁	Fromia monilis ²⁸⁴
moniloside G	419	0-R ³	н	ОН	н	XXX2	Fromia monilis ²⁸⁴
moniloside F	420	0. R ³	Н	ОН	Н	AX	Fromia monilis ²⁸⁴
moniloside H	421	^{0. R³}	н	он	н	XXX ₂	Fromia monilis ²⁸⁴
pisasteroside C	422	€. R ³	н	Н	н	X ₅	Pisaster brevispinus ³⁰²

Basic Nuclear Hydroxylation Pattern: $3\beta, 6\beta, 8, 15\alpha$

ſ	\sim	OH	, , , , , , , , , , , , , , , , , , ,
RO		ОН	•

name	no.	side chain	R	R1	\mathbb{R}^2	R ³	R4	sources and references
coscinasteroside B	423	Ø- R ⁴	Н	Н	H	SO₃-NA+	X ₁	Coscinasterias tenuispina, ²⁶⁹ Pycnopodia helianthoides ²⁵⁸
pisasteroside B	424	my	X ₁	Н	Н	н	SO3-Na+	Pisaster ochraceus ³⁰²
aphelasteroside A	425	₩	SO3-NA+	Н	H	н	X 1	Aphel as terias japonica ²¹⁴
5-deoxyisonodososide	426	۵. R ⁴	X ₂	Н	Н	Н	A 1	Acanthaster planci, ³⁰³ Choriaster granulatus ²⁸⁵

Table 4 (Continued)

name	no.	side chain	R	\mathbb{R}^1	\mathbb{R}^2	\mathbb{R}^3	R4	sources and references
laeviuscoloside I forbeside I	427	₩	X4	ОН	Н	Н	н	Henricia laeviuscola, ²⁸³ Asterias forbesi ³⁰⁴
granulatoside A	428	<u>0</u> -R ⁴	X2	ОН	Н	Н	A 1	Choriaster granulatus, ²⁸⁵ Thromidia catalai ³⁰⁵
laeviuscoloside G	429	<u>0</u> -R ⁴	X4	ОН	Н	Н	A 1	Henricia laeviuscola, ²⁸³ Asterias forbesi ³⁰⁴
isonodososide	430	<u>Q</u> . R ⁴	\mathbf{X}_2	н	ОН	Н	A 1	Acanthaster planci ³⁰³
nodososide	431	<u>0</u> . R ⁴	Н	н	ОН	Н	XA5	Protoreaster nodosus, ²⁸² Pentaceraster alveolatus, ⁶⁸ A. planci, Linkia Laevigata ³⁰⁶
echinasteroside B_2	432	<u>0</u> R ⁴	н	ОН	н	н	XA5	Echinaster sepositus ³⁰⁷
echinasteroside B ₁	433	<u>0</u> . R ⁴	н	он	н	COCH ₃	XA5	Eschinaster sepositus ³⁰⁷
scoparioside D	434	<u>₽</u> . R ⁴	Н	н	н	SO3-NA+	\mathbf{X}_1	Astropecten scoparius ²⁶⁷
laeviuscoloside H	435	<u>₽</u> - R ⁴	X4	он	н	Н	Н	Henricia laeviuscola ²⁸³
forbeside K	436	"	X2	он	н	н	\mathbf{A}_1	Asterias forbesi ³⁰⁴

Basic Nuclear Hydroxylation Pattern: $3\beta,6\beta,8,15\alpha,16\beta$



			R10⊦	1			
name	no.	side chain	R	R1	\mathbb{R}^2	R ³	sources and references
laeviuscoloside B	437		X4	Н	SO3-Na+		Henricia laeviuscola ²⁸³
	438	И СТАТИТАТИ ОН	\mathbf{X}_2	Н	Н		Poraster superbus ²⁵¹
thromidioside	439	ион	\mathbf{X}_2	ОН	н		Thromidia catalai ³⁰⁵
laeviuscoloside C	440	он т и.,	\mathbf{X}_2	Н	SO₃-NA+		Henricia laeviuscola ²⁸³
crossasteroside P_1	441		н	н	н	XGalf	Crossaster papposus ³⁰⁶
attenuatoside S-I	442		Н	н	SO₃-NA+	\mathbf{X}_1	Hacelia attenuata ³⁰⁹

Polyoxygenated Steroids of Marine Origin

Table 4 (Continued)

name	no.	side chain	R	R1	\mathbb{R}^2	R³	sources and references
laeviuscoloside E	443	0. R ³	X ₂	Н	SO3-NA+	Н	Henricia laeviuscola ²⁸³
crossasteroside P_2	444	0·R ³	Н	ОН	Н	XGal <i>f</i>	Crossaster papposus ³⁰⁸
attenuatoside S-II	445	0.R ³	Н	н	SO₃-NA+	X 1	Hacelia attenuata ³⁰⁹
attenuatoside S-III	446	M., O. R ³	н	Н	SO₃-NA+	X 1	Hacelia attenuata ³⁰⁹

Basic Nuclear Hydroxylation Patterns: $\Delta^{4}-3\beta,6\beta,8,15\alpha$ and $\Delta^{4}-3\beta,6\beta,8,15\alpha,16\beta$

		RC				1
name	no.	side chain	R	R1	R ²	sources and references
pisasteroside D	447		X ₁	Н	Н	Pisaster giganteus ³⁰⁰
aphelasteroside B	448	₩	X 1	SO₃-Na+	ОН	Aphelasterias japonica ²¹⁴
echinasteroside A	449	и устанон	X2	SO3-Na+	ОН	Echinaster sepositus, ³¹⁰ Henricia laeviuscola ²⁸³
laeviuscoloside D	450	M. J. Coh	X2	SO₃-Na+	ОН	Henricia laeviuscola ²⁸³
echinasteroside B	451		X ₂	SO₃-Na+	он	Eschinaster sepositus, ³¹⁰ Henricia laeviuscola ²⁸³
forbeside L	452	и. устори	X2	Н	ОН	Asteri as forbesi ³⁰⁴

Basic Nuclear Hydroxylation Pattern: 38,68,8,158





Further recent additions are the phosphated steroids tremasterols A-C (459-461), isolated from the deepwater species *Tremaster novaecaledoniae*.

3. Asterosaponins

The term asterosaponin was originally coined to generally designate toxic steroidal saponins occurring in starfish, by analogy with the term holothurin used for triterpenoid saponins of holothuroids. Following the elucidation of first structures and the subsequent discovery of different structural type of steroidal glycosides from starfish, the term has been used to indicate those highest molecular weight compounds unified by a number of structural analogies involving both the aglycons and the oligosaccharide chains. The aglycons are $\Delta^{9(11)}$ -3 β , 6α -dihydroxysteroids bearing a sulfate group at C-3 and often a 23-oxo function. The oligosaccharide chains, commonly made up by five or six sugar units, are always glycosidically linked at C-6. A close resemblance is also evident in this saccharide portions: sugars are in their pyranose form with β -anometric configurations (α for L-arabinose) and linked with a constant pattern of interglycosidic linkages. A

branching point is always located on the second monosaccharide (usually xylose or quinovose) starting from the aglycon and a terminal quinovose is always found 2-linked to the branched sugar. The more common sugars are D-fucose, D-quinovose, D-xylose, D-galactose, and D-glucose. Other less common monosaccharides are D-6-deoxy-xylo-hex-4-ulose and L-arabinose. The single exception to the general pattern of interglycosidic linkages is represented by the recently isolated santiagoside (488),³¹¹ with a 4-substituted glucose unit linked to the aglycon, instead of the 3-substituted unit present in all asterosaponins. We would note that the ¹³C NMR data of santiagoside (488) are very close to those assigned to related asterosaponins [e.g. marthasteroide C (489)], including the unusually downfield shifted signal at 91.0 ppm, which is a distinctive feature of ¹³C NMR spectra of all asterosaponins and has been assigned to the glycosylated carbon-3 of the monosaccharide unit (β -glucopyranosyl or β -quinovopyranosyl) directly attached to the aglycon.

Also asterosaponins are widespread among starfish, having been found in the majority of species examined. The toxic properties of starfish have been known for many years, but it was only in 1960 that Hashimoto

Figure 19. Steroidal aglycons found in astereosaponins from starfish.

and Yasumoto recognized that the toxicity is associated with compounds similar to plant saponins.³¹² Asterosaponins also possess ichtyotoxicity, 312,313 and toxicity has also been noted toward anellids, mollusks, arthropods and vertebrates.³¹⁴ Because of their general toxicity, it is probable that saponins act primarily as chemical defense agents, rejecting infectious aquatic fungi, protists, parasites, and predators. Makie et al. found that asterosaponins are the substances eliciting the escape response in some species of molluscs when in presence of starfishes.³¹⁵ In addition, these oligoglycosides, in some species of starfish, participate in reproduction processes. Ikegami et al. identified saponins as the spawning inhibitors in the ovaries of Asterias amurensis.³¹⁶ Fujimoto et al. observed that three steroidal saponins, designated Co-ARIS I, II, and III, isolated from the egg jelly of the starfish Asterias amurensis, are essential for inducing the acrosome reaction.³¹⁷ Starfish extracts and purified saponin fractions have shown a variety of pharmacological activities: hemolytic activity;^{314,318} in vitro cytotoxicity toward tumor cells;³¹⁹⁻³²¹ antiviral activity;³²² blockage of neuromuscular transmission in mammals;³²³ and antiinflammatory, analgesic, and hypotensive activities.³²⁴ Starfish saponins are also known to inhibit development of fertilized sea urchin eggs.^{318,325} A more recent study of the biological activity of representative saponins and related steroids from starfish confirmed a high incidence of cytotoxicity and inhibition of Grampositive bacteria but only weak antiviral activity and no inhibition of Gram-negative bacteria.²⁶⁰

Asterosaponins, as do all these polyhydroxysteroids and glycosides from starfish, also occur as complex mixtures that are fairly difficult to separate in the individual components. This has probably been the main reason for which most of the initial papers were concerned with the analysis of aglycons obtained by acid hydrolysis of partially purified saponin mixtures. This procedure resulted in the production of several artifacts, such as asterone (462),^{326,327} an artifact obtained by retro-aldol cleavage of the genuine thornasterol A sulfate (463),³²⁸ which has been the most widely reported steroid obtained by acid hydrolysis of asterosaponins. $\Delta^{20(22)}$ - and $\Delta^{17(20)}$ -steroids, along with a rearranged $\Delta^{13(14)}$ -17-methyl-18-norsteroid, certainly artifacts generated during acid hydrolysis, have also been reported.^{329,330} An excellent review of previous work on the aglycons can be found in the work of Burnell and ApSimon.²³²

The first complete structure of an asterosaponin, thornasteroside A (464), was reported in 1978 by

Table 5. Asterosaponins from Starfish*

Name	#	oligosaccharide chain	sources and references
Asterosaponins contair	ning tho	masterol A, aglycone A	
thornasteroside A	464	Fuc ¹⁻² →Gal ¹⁻⁴ →Xyl ¹⁻³ →Qui ¹⁻⁶ A	Acanthaster planci ³³¹ , Asterias amuren- sis ³³² , Luidia maculata ³³³ , Ophidiaster ophidianus ³³⁴ , Linkia laevigata ³³⁵ , Pro- toreaster nodosus and Pentaceraster alveo- latus ⁶⁸ , Halityle regularis ³³⁶ , Nordoa go- mophia ²⁶⁵ , Coscinasterias tenuispina ²⁶⁸ , Thromidia catalai ³⁰⁵ , Pisaster ochraceus and Pisaster brevispinus ³⁰³ , Pisaster gigan- teus ³⁰⁰ , Pycnopodia helianthoides ²⁵⁸
glycoside B ₂ = forbeside B	465	Qu i→Ga l4 Xy l3 Qu i6	Asterias amurensis ^{284, 337} , Asterias forbesi ³³⁸
acanthaglycoside C	466	$Fuc \xrightarrow{1-2} G[c \xrightarrow{1-4} Xy] \xrightarrow{1-3} Qui \xrightarrow{1-6} A$	Astropecten latespinosus ³³⁹ , Acanthaster planci340, Asterina pectinifera ³⁴¹ , Patiria miniata ³⁴²
maculatoside = luidiaglycoside B	467	Fuc	Luidia maculata ³³³ , Linkia laevigata ³³⁵
ophidianoside F	468	$\operatorname{Fuc} \xrightarrow{1-2} Xy \xrightarrow{1-4} Xy \xrightarrow{1-3} Qu \xrightarrow{1-6} A$	Ophidiaster ophidianus ³³⁴ , Linkia laevigata ³³⁵ , Thromidia catalai ³⁰⁵
regularoside B	469	Fuc ¹⁻² / _→ Fuc ¹⁻⁴ →Xyl ¹⁻³ / _→ Qui ¹⁻⁶ A ∫ 1-2 Qui	Halityle regularis ³³⁶ , Coscinasterias tenuispina ²⁶⁸ , Thromidia catalai ³⁰⁵
lævigatoside	470	Fuc ¹⁻² Ara ¹⁻⁴ →Qui ¹⁻³ →Qui ¹⁻⁶ A	Linkia laevigata ³³⁵
pectinioside A	471	Fuc → GIc → Qui ↓ 1-3 ↓ Qui ↓ 1-6 ↓ 1-2 Qui	Asterina pectinifera ³⁴³
ovarian asterosaponin I = Co Aris I = forbeside C	472	Fuc $\frac{1-2}{1-2}$ Fuc $\frac{1-4}{2}$ Qu i Qu i	Asterias amurensis ³ 17, 284, 344, 345 _, Asterias forbesi ³⁴⁶ , Asterias vulgaris ³⁴⁶
marthasteroside A ₂ = luidiaglycoside A	473	$Fuc \xrightarrow{1-3} Fuc \xrightarrow{1-2} Qu i \xrightarrow{1-4} Xy I \xrightarrow{1-3} Qu i \xrightarrow{1-6} A$	Luidia maculata ^{333, 347} , Marthasterias glacialis ^{348, 349}

Name	#	oligosaccharide chain	sources and references
marthasteroside A ₁	474	Fuc → Fuc → Ga I → Xy I → Qu i → Qu i → A	Marthasterias glacialis ^{348, 349} , Astro- pecten latespinosus ³³⁹ , Linkia laeviga- ta ³³⁵ , Nardoa gomophia ²⁶⁵ , Pisaster brevispinus ³⁰³ , Achantaster planci ³⁴⁰ , Astropecten scoparius ²⁶⁷
versicoside A = forbeside A	475	$Ga I \xrightarrow{1-3} Fuc \xrightarrow{1-2} Ga I \xrightarrow{1-4} Xy I \xrightarrow{1-3} Qu i \xrightarrow{1-6} A$ $\uparrow 1-2$ $Qu i$	Asterias amurensis versicolor ³³² , Pisaster ochraceus and Pisaster brevispinus ³⁰³ , Pisaster giganteus ³⁰⁰ , Asterias forbesi ³³⁸
pectinioside F	476	Ga I → GI C → Xy I → Qu i → A ↓ 1 - 2 Fuc Qu i	Asterias pectinifera ³²¹
pectinioside E	477	$Fuc \xrightarrow{1-4} Glc \xrightarrow{1-4} Qul \xrightarrow{1-3} Qul \xrightarrow{1-6} A$ $\uparrow 1-2 \qquad \uparrow 1-2$ $Xyl \qquad Qul$	>>
pectinioside G	478	Ara ¹⁻⁴ →GIc ¹⁻⁴ →Qui ¹⁻³ →Qui ¹⁻⁶ A	Asterina pectinifera ²⁵⁴ , Patiria miniata ³⁴²
forbeside H	479	Qui → XyI → Qui → A	Asterias forbesi ³⁵⁰
forbeside G	480	$Qu i \xrightarrow{1-2} Qu i \xrightarrow{CH_3} OH$	»>
forbeside F	481	$Fuc \xrightarrow{1-4} Qui$	>>
myxodermoside A	482	Ga I → Xy I → Qu i → A	Myxoderma platyacanthum ²⁴⁵
Asterosaponins containi	ng 24,25-	dehydro-thornasterol A, aglycone B	

achantaglycoside A 483 $Fuc \xrightarrow{1-2} Qu i \xrightarrow{1-4} Xy | \xrightarrow{1-3} Qu i \xrightarrow{1-6} B A chantaster planci 351$ $\downarrow 1-2 \\ Qu i$ achantaglycoside B 484 $Fuc \xrightarrow{1-2} G | c \xrightarrow{1-4} Xy | \xrightarrow{1-3} Qu | \xrightarrow{1-6} B A chantaster planci 340$ $\downarrow 1-2 \\ Qu i$ achantaglycoside D 485 $Fuc \xrightarrow{1-2} G | c \xrightarrow{1-4} Xy | \xrightarrow{1-3} Qu | \xrightarrow{1-6} B A chantaster planci 340$ $\downarrow 1-2 \\ Qu i$ $Fuc \xrightarrow{1-2} G | c \xrightarrow{1-4} Xy | \xrightarrow{1-3} Qu | \xrightarrow{1-6} B A chantaster planci 340$

Name	#	oligosaccharide chain	sources and references
Asterosaponins conta	uning marthaste	erone, aglycone C	
marthasteroside B	486	$Fuc \xrightarrow{1-2} Fuc \xrightarrow{1-4} Qui \xrightarrow{1-3} Glc \xrightarrow{1-6} C$ $\int_{1-2}^{1-2} Qui$	- Marthasterias glacialis ^{348, 349} , Luidia maculata ³³³ , Coscinasterias tenuispina ²⁶⁸
luidiaglycoside C	487	$\operatorname{Fuc} \xrightarrow{1-2} \operatorname{Qu} i \xrightarrow{1-4} \operatorname{Qu} i \xrightarrow{1-3} \operatorname{Glc} \xrightarrow{1-6} \operatorname{C} \overset{1}{ }_{1-2} \overset{1}{ }_{2-2} \overset{1}{ }$	Luidia maculata ³⁵²

Asterosaponins containing 24,25-dihydromarthasterone, aglycone D

santiagoside	488	Fuc <u>¹⁻⁴</u> Qui <u>¹⁻⁴</u> Gic <u>¹⁻⁶</u> D	Neosmilaster georgianus ³¹¹
marthasteroside C	489	$Fuc \xrightarrow{1-2} Fuc \xrightarrow{1-4} Qui \xrightarrow{1-3} Gic \xrightarrow{1-6} D$	Marthasterias glacialis ^{348, 349} , Luidia maculata ³³³ , Coscinasterias tenuispina ²⁶⁸
luidiaglycoside D	490	$Fuc \xrightarrow{1-2} Qu \xrightarrow{1-4} Qu \xrightarrow{1-3} G c \xrightarrow{1-6} D$ $\downarrow 1-2$ $Qu i$	Luidia maculata ³⁵²

Asterosaponins containing 24-nor-thornasterol A, aglycone E

ophidianoside B	491	$Fuc \xrightarrow{1\cdot 2} Ga \stackrel{1\cdot 4}{\longrightarrow} Xy \stackrel{1\cdot 3}{\longrightarrow} Qu \stackrel{1\cdot 6}{\longrightarrow} E$ $ \stackrel{\uparrow}{\underset{Qu i}{\stackrel{1\cdot 2}{\longrightarrow}}} Qu \stackrel{1\cdot 6}{\longrightarrow} E$	Ophidiaster ophidianusand Hacelia attenuata ³³⁴
ophidianoside C	492	$Fuc \xrightarrow{1-2} Xy \stackrel{1-4}{\longrightarrow} Xy \stackrel{1-3}{\longrightarrow} Qu \stackrel{1-6}{\longrightarrow} E$ $ \stackrel{\uparrow}{1}_{1-2} Qu \stackrel{1}{i}$ Qu i	~~

asterosaponins containing 22,23-epoxy-aglycone F

tenuispinoside A 493
$$Fuc \xrightarrow{1-2} Ga | \xrightarrow{1-4} Xy | \xrightarrow{1-3} Qu | \xrightarrow{1-6} F$$

$$Coscinasterias tenuispina268$$

$$fuc \xrightarrow{1-2} Qu | \xrightarrow{1-2} Qu | \xrightarrow{1-6} F$$

$$Vy | \xrightarrow{1-3} Qu | \xrightarrow{1-6} F$$

$$Vy | \xrightarrow{1-6} F$$

$$S = \frac{1-2}{Qu | 1}$$

Name_	#	oligosaccharide chain	sources and references	
asterosaponin con	ntaining 22,23-epoxy	-24-nor-aglycone G		
asteroside B	496	Qu i $\xrightarrow{1-2}$ Ga I $\xrightarrow{1-4}$ Xy I $\xrightarrow{1-3}$ Qu i $\xrightarrow{1-6}$ G \uparrow 1-2 Qu i	- Asterias amurensis ²⁸⁴	

asterosaponins containing the less common aglycones H, I, J, K and L

protoreasteroside	497	$Fuc \xrightarrow{1-2} Qu i \xrightarrow{1-4} Xy I \xrightarrow{1-3} Qu i \xrightarrow{1-6} H$ $\uparrow 1-2$ Qu i	Protoreaster nodosus and Pentaceraster alveolatus ⁶⁸
ovarian asterosaponin-4 = Co-Aris III	498	$\begin{array}{c} \operatorname{Qu} i \xrightarrow{1-2} \operatorname{Gl} c \xrightarrow{1-4} XyI \xrightarrow{1-3} \operatorname{Qu} I \xrightarrow{1-6} I \\ & \uparrow 1-2 \\ \operatorname{Qu} i \end{array}$	Asterias amurensis ^{317, 284, 353}
solasteroside A	499	Fuc ¹⁻² / _→ Fuc ¹⁻⁴ →Xyl ¹⁻³ / _→ Qui ¹⁻⁶ / ¹ / ₁₋₂ Qui	Solaster borealis ²⁴⁴
Co-Aris II	500	Fuc $\frac{1-2}{1-2}$ Fuc $\frac{1-4}{1-2}$ Qu i	Asterias amurensis ³¹⁷
tenuispinoside C	501	$Fuc \xrightarrow{1-2} Gal \xrightarrow{1-4} Xyl \xrightarrow{1-3} Qul \xrightarrow{1-6} K$ $ \int 1-2 Qul$	Coscinasterias tenuispina ²⁶⁸
asteroside D	502	$\begin{array}{c} \operatorname{Qu} i \xrightarrow{1-2} \operatorname{Gl} c \xrightarrow{1-4} \operatorname{Xy} I \xrightarrow{1-3} \operatorname{Qu} i \xrightarrow{1-6} L \\ & \uparrow 1-2 \\ \operatorname{Qu} i \end{array}$	Asterias amurensis ²⁸⁴

asterosaponins containing (24R)-24-methylthornasterol A, aglycone M

versicoside C
= thornasteroside B
steroside C
versicoside B
steroside B

$$503$$

Fuc $\frac{1\cdot2}{\rightarrow}$ Ga | $\frac{1\cdot4}{\rightarrow}$ Xy | $\frac{1\cdot3}{\rightarrow}$ Qu | $\frac{1\cdot6}{M}$
Asterias amurensis [cf.] versicolor 345,
Coscinasterias tenuispina²⁶⁸
Asterias amurensis²⁸⁴
 $\int_{1\cdot2}^{1\cdot2}$ Qu | $\frac{1\cdot6}{M}$
Asterias amurensis²⁸⁴
 $\int_{1\cdot2}^{1\cdot2}$ Qu | $\frac{1\cdot6}{M}$
versicoside B
 505
Ga | $\frac{1\cdot3}{\rightarrow}$ Fuc $\frac{1\cdot2}{\rightarrow}$ Ga | $\frac{1\cdot4}{\rightarrow}$ Xy | $\frac{1\cdot3}{\rightarrow}$ Qu | $\frac{1\cdot6}{M}$
Asterias amurensis²⁸⁴
 $\int_{1\cdot2}^{1\cdot2}$ Qu | $\frac{1\cdot6}{M}$
Asterias amurensis [cf.] versicolor³⁴⁵

Name	#	oligosaccharide chain s	sources and references
asterosaponin co	ntaining (24S)-24	4-methylthornasterol A, aglycone N	
acanthaglycoside	e F 506	Fuc ¹⁻³ →Fuc ¹⁻² →Gal ¹⁻⁴ →Xyl ¹⁻³ →Qui 1-2 Qui	Acanthaster planci ³⁴⁰
asterosaponins co	ontaining (24S)-2	2,23-epoxy-24-methyl aglycone O	
regularoside A	507	$Fuc \xrightarrow{1-2} Qu \stackrel{1-4}{\longrightarrow} Qu \stackrel{1}{\longrightarrow} Qu \stackrel{1}{\longrightarrow} Glc$	Halityle regularis ³³⁶
henricioside A	508	Ara —→Gic → Xyi → Qui 1-2 Qui	1-6 Henricia laeviuscola ²⁸³
pectinioside B	509	Gal —→GIc —→Xyl —→Qui	Asterina pectinifera ³⁴³
patirioside A	510	Fuc	Patiria miniata ³⁴²

asterosaponin containing (24S)-ethylthornasterol A, aglycone P

pectinioside C 511
$$\operatorname{Fuc} \xrightarrow{1-3} \operatorname{Fuc} \xrightarrow{1-2} \operatorname{Glc} \xrightarrow{1-4} \operatorname{Xyl} \xrightarrow{1-3} \operatorname{Qul} \xrightarrow{1-6} \operatorname{P} Asterina pectinifera^{341}$$

^a Qui = D-quinovose. Fuc = D-fucose. Xyl = D-xylose. Ara = L-arabinose. Gal = D-galactose. Glc = D-glucose. All monosaccharides are in their pyranose forms and the glycosidic linkages are β (α for L-arabinose).

Kitagawa and Kobayashi.³³¹ Thornasteroside A (464) is the most widely distributed asterosaponin, having

been isolated from 15 species representative of the three major orders of Asteroidea, and it is also a good example of the general structure of pentaglycosides, the more common components among asterosaponins. Thornasterol A sulfate (463), the aglycon of thornasteroside A (464), is by far the more common aglycon, but a number of asterosaponins containing aglycons with different functionalities in the side chain have also been isolated (Figure 19). The aglycon K of tenuispinoside C (501) is the only one having a different functionalization of the tetracyclic nucleus, with an extra 12α hydroxyl group. The structures of all reported asterosaponins are listed in Table 5 with sources and references.

A further minor constituent of Asterias forbesi, forbeside E (512), was shown to be the 6-O-quinovopyranosyl 4'-sulfate of (20R)-5 α -pregn-9(11)-ene-3 β , 6α ,-20-triol 3-sulfate,³⁵⁴ an aglycon also isolated as hydrolysis product of saponins from Asterias vulgaris.³⁵⁵ The corresponding asterone 462 analog, cheliferoside LI, has been reported from Lethasterias nanimensis chelifera.³⁵⁶

4. Cyclic Steroidal Glycosides

Toxic saponing of a completely different structural type have been discovered in two species of the genus Echinaster (Figure 20). They have a number of unusual features when compared to the more common asterosaponins: there is no sulfate group and charge is due to a glucuronic acid unit in the saccharide moiety. the Δ^7 -3 β ,6 β -dihydroxysteroidal nucleus is unprecedented, and, the most remarkable feature, the trisaccharide chain is cyclized between C-3 and C-6 of the aglycon, giving rise to a macrocyclic ring reminiscent of a crown ether. Sepositoside A (513), the major saponin from the Mediterranean starfish Echinaster sepositus,³⁵⁷ is accompanied by smaller amounts of three related saponins 514-516, which differ only in the structure of the side chain of the aglycons, all having a 22,23-epoxy functionality.³⁵⁸ The key step during structural study of sepositoside A (513) was the mild acid hydrolysis, which, by cleavage of the allylic ether linkage, gave rise to the opening of the macrocyclic ring

and formation of the UV-active glycoside, 517. A further representative of this class of glycosides was isolated from a Pacific starfish belonging to the same genus, *E. luzonicus*, and accordingly named luzonicoside A (518).³⁵⁹ Sepositoside A (513) is moderately toxic ($LD_{50} = 43 \text{ mg kg}^{-1}$ by ip injection in mouse) and showed cytotoxic activity toward bovine turbinate cells up to a level of 1 μ g mL^{-1.260} Both sepositoside A (513) and luzonicoside A (518) were slightly effective in the inhibition of cell division of fertilized sea urchin eggs (*ca.* 30% inhibition at 10⁻⁵ M) and showed antifungal activity.

B. Ophiuroidea (Brittle Stars)

Until a few years ago brittle stars (ophiuroids) had only received moderate attention as compared to the echinoderm classes of asteroids and holothuroids and only sporadic papers dealing with their sterols content had appeared in the literature. Quite recently we had the occasion to investigate a number of ophiuroid species from which we could isolate several sulfated polyhydroxysteroids and two steroidal glycosides (Figure 21).

The steroid glycosides, longicaudoside A (519) and B (520) have been isolated from the Mediterranean Ophioderma longicaudum.³⁶⁰ Both have a 5 α -cholestane-3 α ,6 β ,12 β ,21-tetrol aglycon bearing a sulfate group at C-3 and a monosaccharide residue at C-12 (β -Dxylopranosyl in 519 and β -D-glucopyranosyl in 520). In contrast with the hydroxylation at C-26, commonly encountered among polyhydroxysteroids from starfish, the polar steroids from ophiuroids are characterized by the hydroxylation at C-21, only found among

Figure 20. Cyclic steroidal glycosides from starfish of the genus Echinaster.

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QSO₃ Na*

OSO3 Na*

он

Figure 21. Polyhydroxysteroids from ophiuroids. starfish metabolites, in steroids from *Euretaster in*signis.²⁴¹ ¹³C NMR spectroscopy proved to be the better tool for differentiation between the two hydroxylated side chains. Noticeable differences are observed in the methyl and hydroxymethyl resonances: the C-21 hydroxylated side chain exhibits the typical signals for C-26 and C-27 at 23.0 and 23.1 ppm and the CH₂OH signal at 63.4 ppm, while the C-26 hydroxylated side

chain exhibits methyl carbon signals at 17.3 and 19.2 ppm and the CH₂OH signal at 68.4 ppm. The narrow signal at a rather low field (δ 4.72 ppm, $W_{1/2} = 7$ Hz) in the ¹H NMR spectra of both 519 and 520, shifted to δ 4.10 in the spectra of their desulfated derivatives, was indicative of a 3α -sulfato- 5α -stanol structure. The alternative 3β -sulfato- 5β -stanol structure could be eliminated mainly because of the chemical shift of C-19

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at 15.3 ppm. Ophioderma longicaudum also furnished unusual sulfated polyhydroxysterols together with a mixture of common Δ^5 -3 β -hydroxysterols.³⁶¹ The more polar sulfated sterol was characterized as 58-cholestane- 3α , 4α , 11β , 12β , 21-pentol 3, 21-disulfate (521), featuring the cis-A/B ring fusion, later also found in the polar steroids from the further species analyzed.^{96,362} A second group of components consisted in a mixture of disulfated 3α , 21-dihydroxysteroids 522-525, whose composition was established after solvolytic removal of the sulfate groups.³⁶¹ Additional sulfated polyhydroxysteroids have been then isolated recently from three further species of ophiuroids, Ophiocoma dentata, Ophiarthrum elegans, and Ophiorachna incrassata, all collected off Noumeà (New Caledonia).96 The major compound in all three species has been shown to be 5β -cholestane- 3α , 4α , 11β , 21-tetrol 3, 21-disulfate (526). Two minor components of Ophiocoma dentata possess the same nucleus as 526 but differ in the side chain, 527 and 528. Along with minor amounts of 528 and major amounts of the disulfated tetraol 526, Ophiarthrum elegans also contains the 11-keto derivative 529. Ophiorachna incrassata also contains major amounts of the steroid disulfate 526 along with minor amounts of the two more polar steroids 530 and 531, bearing three sulfate groups, which represent the first occurrence of 26-hydroxylation in ophiuroids. Major support for the presence of both 25S- and 25R-epimers is found in the ¹³C NMR spectra, where every side-chain carbon signal appears split into two peaks separated by 0.04-0.2 ppm. The pentol disulfate 521, the tetrol disulfate 526, and the Δ^5 -steroidal sulfates 523 and 525 have moderate cytotoxic activity.²⁶⁰

The investigation of the Pacific ophiuroid Ophiolepis superba, collected at Okinawa, Japan, has led to the isolation of seven sulfated polyhydroxysteroids (532-538), all with 3α ,21-disulfoxy- 4α -hydroxy substituents and the A/B cis ring junction.³⁶² The stereochemistry at C-24 and C-25 in 533 and 535 have been determined after the synthesis of model compounds.^{261,363} The $3\alpha.4\alpha.5\beta$ -trihydroxycholestane structure of 538 was derived from analysis of ¹H and ¹³C NMR data and confirmed by the synthesis of 5β -cholestane- 3α , 4α , 5triol and the alternative 5α -cholestane- 3β , 4β , 5-triol models. A comparison of the ¹H and ¹³C NMR spectra of 538 with those of its desulfated derivative allowed the placement of the sulfoxy group at C-3. The introduction of an additional hydroxyl group at C-2 β in the 3α -sulfoxy- 4α -hydroxy- 5β -steroidal nucleus, as in structures 536 and 537, leads to the appearance in the ¹H NMR spectrum of a signal at δ 4.03 overlapping with the 3-H signal. When the spectrum of the desulfated 536 was measured, the three hydroxymethine protons appeared as isolated signals at δ 3.81 (m), 3.23 (dd, J = 9.5, 3.5 Hz) and 3.94 (t, J = 3.5 Hz) ppm. Decoupling experiments proved that they were located in a sequential arrangement and allowed the inference of a 2β , 3α , 4α -trihydroxy- 5β -steroidal structure. The 24S configuration was assigned to 537 after the stereoselective synthesis of (24R, 22E)- and (24S, 22E)-24- $(hydroxymethyl)cholesta-5,22-dien-3\beta-ol.^{364}$

Stonik and Elyakov reported to have isolated cholest-5-ene- 3α , 4β , 21-triol 3-sulfate (539) from Ophiura sarsi.³⁶⁵

More recent additions to the sulfated polyhydroxysteroids from ophiuroids are a new 5 β -steroid, 3 β ,21disulfate 540 with a unique 4α ,9 α -ether bridge, and a minor related steroid 541 from Ophiomastix annulosa.³⁶⁶ O. annulosa also contained the more common tetrol 526 and its keto derivative 529.

VIII. Polyoxygenated Steroids of Arthropoda

In arthropods, the molting process is under the control of a group of polyhydroxysteroid hormones referred as ecdysones. Although much of the interest of ecdysone biochemistry has centered on Insecta, the role of these steroids in crustaceans has also received considerable attention. The structures of ecdysones reported in crustaceans are shown in Figure 22. In 1960, the

Figure 22. Structures of ecdysones found in Crustacea.

548

Figure 23. Polyhydroxysteroids from Tunicata.

552, R=H

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presence of a substance which showed ecdysone (542)activity was detected in crabs by Karlson and Skinner.367 In 1966, crustecdysone (543) was first isolated from the crayfish, Jasus lalandei, by Horn et al.368 and subsequently 2-deoxycrustecdysone (544)369 was isolated. The 20-hydroxyecdysteroids, callinecdysone A (inokosterone) (545) and callinecdysone B (makisterone A) (546) were isolated from Calinectes sapidus.³⁷⁰ Ecdysteroids metabolism in crustaceans has been reviewed by Goad^{2,229} and Ikekawa.⁶⁰

IX. Polyoxygenated Steroids of Tunicates

Several species of ascidians have been analyzed for their sterol content and were shown to contain typical mixtures of sterols with cholesterol as the major component.² Coprostanols along with Δ^4 -3-ketosteroids and 4α -methylsterols have been described from Ascidia nigra.³⁷¹ 5 β -Stanols have later been found in Halocynthia papillosa, Microcosmus sulcatus, Microcosmus savignyi, and Styela plicata.³⁷² The tunicates Phallusia

561, **∆**⁴ R=H Figure 24. Polyoxygenated steroids from fish.

mamillata and Ciona intestinalis were reported to contain 5,8-endoperoxides of several $\Delta^{5,7,9(11)}$ -sterols (547)³⁷³ and the 24-hydroperoxy-24-vinylcholesterol (548) along with its corresponding 24-hydroxy derivative.³⁷⁴ this latter previously found in an alga (Figure 23).

X. Polyoxygenated Steroids from Vertebrates (Fish)

The bile of vertebrates characteristically contains bile salts, and in most animals these are largely steroidal C_{24} carboxylic acid derivatives. In the case of some fish the bile salts include C₂₇ alcohols present as their sodium sulfate esters.

Scymnol (549, Figure 24) is the bile alcohol from shark bile and the first of such alcohols to be isolated.³⁷⁵ The history of the structure determination of this alcohol has been reviewed by Scheuer.³⁷⁶ The final structure 549 was deduced by two groups, Cross³⁷⁷ and Briggs and Haslewood.³⁷⁸ The two bile alcohols myxinol (550) and deoxymyxinol (551) were both isolated from hagfish. For myxinol, the first of the two to be isolated, ³⁷⁹ the 5 β -structure was initially proposed, but later work established the 5α -structure 550.³⁸⁰ Deoxymyxinol was then assigned the structure 551 by the same research group.³⁸¹

Certain fish have a self-defense mechanism consisting of the secretion of toxic substances that repel their predators. Among them are the Red Sea Moses sole Pardachirus marmoratus and its congener, the peacock sole Pardachirus pavoninus, which repel sharks by emission of their toxic secretion at the moment when they are about to be bitten. The chemical nature of the shark repellents has been shown by Tachibana. Nakanishi, and co-workers to consist of mixture of the peptidic paradaxins and two group of steroid glycosides. pavonins 1-6 (552-557) from P. pavoninus, 382,383 and mosesins 1-5 (558-562) from P. marmoratus.³⁸⁴ The chemical structure of both pavonins and mosesins has been established by spectroscopic studies and chemical conversions. Mosesin 4 (560) has been successfully obtained by synthesis.385 The biological activity of these toxins is believed to be related to their surfactant properties, that in the case of glycosides arises from a hydrophobic "top" and a hydrophilic "bottom" region. The respective roles of the steroidal glycosides and of the peptidic paradaxins in defense mechanism still remain obscure. It has been suggested that pavonins and mosesins are repellents that act on the shark's olfactory sense, whereas the proteinaceus toxin is possibly an antifeedant that acts on its gustatory sense. A review on chemical defense in fishes, which includes the fascinating history of the shark repellents from Pardachirus sp. has been published by Tachibana.³⁸⁶

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