Bioactive Nitrogenous Secondary Metabolites from the Marine Sponge Genus Haliclona

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Abstract: Marine sponge genus Haliclona, one of the most prolific sources of natural products, contains over 600 species but only a small part of them had been classified and chemically investigated. On the basis of extensive literature search, this review firstly summarizes 112 nitrogenous secondary metabolites from classified and unclassified Haliclona sponges as well as from their symbiotic microorganisms. Most of these substances have only been found in Haliclona sponges, and display diverse bioactive properties with potential applications in new drug discovery.

Keywords: marine sponge; Haliclona; symbiotic microbe; secondary metabolite; alkaloid; bioactivity

1. Introduction

The marine environment is the largest treasure trove of creatures, including plants, animals and microorganisms. People never stop conducting chemical studies of marine organisms owing to their potential capability to produce bioactive secondary metabolites that are potential sources of leads for new drug development. So far over 34,000 articles about marine-derived natural products research have been published [1]. Marine sponges, which are immemorial organisms, are widely distributed around the world and comprise over ten thousand species, most of which live in the sea, while only one percent are freshwater sponges [2]. These creatures are a great source of natural products with a broad spectrum of biological properties. Numerous sponge-derived chemicals, especially alkaloid compounds, display pharmacological effects, such as didemnin B, cytarabine, trabectedin, vidarabine, etc.

The marine sponge genus Haliclona belongs to the family Chalinidae, order Haplosclerida, class Demospongiae [3]. Its skeleton is made up of grids of single spines or a network of spongy fibrous branches without an epidermal skeleton [2]. The Haliclona genus consists of over 600 species distributed throughout the world [3]. However, only a few dozen specimens collected from the Pacific Ocean [4], north Indian Ocean [5], Atlantic Ocean [6], and Mediterranean Sea [7] have been chemically investigated. These studies suggest that Haliclona sponges are some of the most prolific producers of bioactive secondary metabolites. On the basis of a extensive literature search using SciFinder, this review firstly summarizes all nitrogenous substances from the marine sponge genus Haliclona and its symbiotic microbes.

2. Bioactive Alkaloids from Haliclona Genus

As many as 103 alkaloid secondary metabolites have been isolated and characterized from Haliclona sponges since 1994. However, only 32 of them (compounds 1–32) were from seven classified Haliclona species including H. baeri, H. cymaeformis, H. densaspicula, H. exigua, H. koremella, H. nigra...
and *H. tulearensis*, and are respectively introduced in detail according to their biological sources. Other alkaloids (compounds 33-103) from unclassified *Haliclona* sponges are grouped into three types on basis of their chemical structures, namely 3-alkylpyridine, amide and depsipeptide, and miscellaneous alkaloids.

### 2.1. Haliclona Baeri

There is only one report on a chemical study of *Haliclona baeri* collected from the coast of Jongbrii Province (Thailand) [8]. One new nitrogenated compound maleimide-5-oxime (1) along with one benzoic derivative and two tetillapyrones was separated from this sponge (Figure 1). The follow-up bioassay tests suggested that compound 1 had weak cytotoxic activity against the human DAOY medulloblastoma cell line at 50 μg/mL [9].

![Figure 1. Chemical structure of compound 1 from Haliclona baeri](image)

### 2.2. Haliclona Cymaeformis

Fractionation of the ethanol extract of *Haliclona cymaeformis* collected from a Xuwen coral reef (Guangdong, China) using silica gel column chromatography led to isolation of eleven alkaloids, including one indole alkaloid 2, six nucleosides 3–8 and four sterols (Figure 2) [10]. Subsequently two pairs of 6-oxypurine regioisomers substituted at the 7 or 9 position (compounds 9–12) were purified from the same specimen (Figure 2) [11].

![Figure 2. Chemical structures of compounds 2–12 from Haliclona cymaeformis.](image)

### 2.3. Haliclona Densaspicula

Two novel alkaloids with six hexacyclic diamines, densanins A (13) and B (14) (Figure 3), were found in methanol extract of the sponge *Haliclona densaspicula* from Keomun Island (Korea) and their absolute chemical structures were determined by 1D and 2D NMR spectral analysis and Mosher reactions. Biological evaluation indicated that compounds 13 and 14 possess potent inhibitory effects on lipopolysaccharide-induced nitric oxide production in BV2 microglial cells with IC50 values of 1.05 and 2.14 μM, respectively [4,12].
2.4. Haliclona Exigua

Chemical study of the sponge specimen Haliclona exigua collected from the coastal areas of India and Indonesia afforded nine alkaloid derivatives, including xestospongin D (15), araguspongins C-E (16–18), 3α-methylaragusponge C (19), neopetrocyclamines A (20) and B (21), papuamine (22) and haliconadidamine (23) (Figure 4) [5,13–17]. Compound 16 was the most common alkaloid from H. exigua and shown to have strong inhibitory activity against human lymphatic filarial parasite B, promastigote and intracellular amastigote forms of Leishmania donovani, and anti-fouling and competitive inhibition of NOS [5,15–17]. This compound was later purified from an Indonesian sponge Neopetrosia chaliniformis and could inhibit zebrafish embryos with a LD50 value of 4.3 μg/mL [18]. Compound 22 possessed remarkable cytotoxicity toward human glioblastoma cell line SF-295 with a GI50 value of 0.8 μM and 23 could control the transfer of MDA-MB-231 breast cancer cells [14]. Additionally, these alkaloids 22 and 23 obtained from an Okinawan sponge H. panicea and were found to inhibit the growth of Mycobacterium bovis BCG, M. intracellulare and M. smegmatis [19,20].

2.5. Haliclona Nigra

Fractionation of the aqueous extract of the marine sponge Haliclona nigra collected from northern coast of Papua New Guinea resulted in the discovery of two new hexapeptides, haligramides A (24) and B (25), together with waiakeamide (26) (Figure 5) [21]. Their chemical structures and configurations were elucidated by extensive NMR analyses and oxidative reactions.
2.6. Haliclona Tulearensis

Five new alkaloids, halitulin (27), halic isolorensin (29), isohaliclorensin (30), haliclorensin B (31) and haliclorensin C (32) together with isohalitulin (28) (Figure 6), were purified from the sponge Haliclona tulearensis collected from Sodwana Bay (South Africa) [22–24]. Compound 29 was a novel diamino derivative possessing an azacyclodecane ring, and exhibited strong cytotoxicity against P-388 mouse leukemia cells with an IC₅₀ value of 0.1 mg/mL [21]. In vitro biological evaluation results suggested that compounds 27, 29, 31 and 32 had significant cytotoxicity against P-388 with IC₅₀ values of no more than 0.1 μg/mL [23,24].

2.7. Other Haliclona Spps.

Until now, up to 71 nitrogenated compounds 33–103 have been discovered from unidentified marine sponge species of Haliclona. According to their chemical structures, these substances can be grouped into three classes, including 3-alkylpyridines, amides and depsipeptides, and miscellaneous alkaloids.

2.7.1. 3-Alkylpyridines

3-Alkylpyridine analogs with linear and cyclic frameworks such as navenones, halitoxins, niphatynes, niphatesines, haminols, viscosamine, etc [25] are the most common alkaloids isolated from the marine sponge genus Haliclona sponges. These substances possess pronounced biological activities.

Chemical study of a marine sponge Haliclona sp. from New Zealand led to isolation of halilocyclin C (33) and two new alkaloids dehydrohalilocyclins C (34) and F (35) (Figure 7), which were the first examples of cyclic 3-alkylpyridinium alkaloid (3-APA) monomerz with an unsaturated alkyl chain [26]. An anti-fouling mixture of poly 3-alkylpyridinium salts (36) as well as haminols (37-38) was firstly isolated from the methanol extract of Haliclona sp. collected in Terra Nova Bay, Ross Sea (Antarctica) (Figure 7) [27]. From an Indonesian sponge Haliclona sp. no. 95546, a new alkaloid, 3-dodecyl- pyridine (39) bearing a terminal cyano group, was purified and found to possess moderate in vitro cytotoxicity against tumor cell lines A549, MCF7 and Hela with the IC₅₀ values of 41.8, 48.4, 33.2 μM, respectively. Two new alkaloids 40–41 with dimeric and trimeric 3-APA moieties were isolated from the methanolic extract of the sponge Haliclona sp. collected in the Pacific coast of
Guatemala (Figure 7) [28]. Bioassay results indicated that compounds 40 and 41 had low cytotoxic effect on murine macrophage J774.A1 and fibrosarcoma WEHI-164 cell lines and human epithelial kidney HEK-293 with IC₅₀ > 20 μg/mL.

Figure 7. Chemical structures of compounds 33-70 from unclassified Haliclonasps.
Haliclocyclamines A–C (42–44), three new cyclic bis-1,3-APA derivatives, found together with five analogs cyclostellettamines A–C (45–47), E (48) and F (49), were separated from the EtOH extract of *Haliclona* sp. collected at Manado in Indonesia (Figure 7) [29, 30]. Compounds 44 and 49 could inhibit vaccinia H-1-related phosphatase (VHR) at 17–18 μM and compound 45 possessed cytotoxic effect on HeLa cells with an IC₅₀ value of 0.9 μg/mL and P388 cells with an IC₅₀ value of 1.06 μg/mL. A heterogeneous and inseparable mixture of five cyclic 3-APAs, cyclohaliclonamines A–E (50–54), was isolated from an Okinawa sponge *Haliclona* sp., which compounds 52–54 were the first tetrameric, pentameric and hexameric 3-APAs from natural sources (Figure 7) [25]. By solvent partition, Sephadex LH-20 gel permeation and HPLC, ten cyclic bis-1,3-APA derivatives (55–64) were separated from the MeOH extract of *Haliclona* sp. collected at the shore of Jeju Island (Korea) and characterized using combined NMR and FAB-MS/MS analyses (Figure 7) [31]. Compounds 56 and 62–64 had moderate antibacterial activities against *Staphylococcus aureus* with the same MIC values of 12.5 μg/mL, and compound 63 exhibited the highest cytotoxic against A549 cell-line with a LC₅₀ value of 14.7 μg/mL. Six tetracyclic alkaloids, haliclonacyclamines A–D (65–68) and halicyclamines A (69) and B (70) consisting of two 3-alkyl hexahydropyridine units were separated from *Haliclona* sp. grown on acroporid coral substrate on the Heron Island (Australia), and compounds 65 and 66 could inhibit the growth of leukemia P-388 cells with IC₅₀ values of 0.8 and 0.6 μg/mL, respectively (Figure 7) [32,33]. Halicyclamines A (69) derived from the Indonesian sponge *Haliclona* sp. 05A08 was shown to possess inhibitory effect on inosine-phosphate dehydrogenase at 1 μg/mL and anti-microbial activity against *Mycobacterium smegmatis* and *M. bovis* BCG with MIC values of 2.5 and 1.0 μg/mL, respectively [34]. Compound 70 showed cytotoxicity against HeLa cells with an IC₅₀ value of 12 μM and inhibitory effect on chymotrypsin and caspase with IC₅₀ values of 0.42 and 0.48 μM, respectively [35].

### 2.7.2. Amides and Depsipeptides

Three novel amides, 2-palmitamidoethane sulfonic acid (71), N¹-(2-aminoethyl)-N²-isopentylphthalamide (72) and N¹-isobutyl-N²-tridecylphthalamide (73), were obtained from a *Haliclona* sp. sponge collected off the coast of Hainan Island (China) (Figure 8) [36,37]. Haliclonin A (74), a new macrocyclic diamide from the Korean-derived sponge *Haliclona* sp. exhibited moderate antibacterial activity with a MIC value of 6.25 μg/mL against *Bacillus subtilis*, and cytotoxicity against the K562 leukemia cell line with an IC₅₀ of 15.9 μg/mL (Figure 8) [38]. Chemical analysis of a Rottnest Island-derived sponge *Haliclona* sp. afforded two new olefinic amides, salicylihalamides A (75) and B (76) (Figure 8) [39]. Chemical investigation of a *Haliclona* sp. sponge specimen from the Vanuatu Islands afforded three new amides, haliclamide (77), halipeptins A (78) and B (79) (Figure 8) [40,41]. The structures of compounds 78 and 79 were later revised and corrected by total chemical synthesis [42]. Compound 77 exhibited in vitro anti-tumor activity against human bronchopulmonary non-small-cell-lung-carcinoma line NSCLC-N6 with an IC₅₀ value of 4.0 μg/mL while 78 displays potent anti-inflammatory effect. One new depsipeptide, kendarimide A (80), was isolated and characterized from a Sulawesi Island-derived *Haliclona* sp. Sponge (Figure 8) [43]. A MTT assay suggested that this chemical had 87% growth inhibition on multi-drug resistance (MDR) cell line KB-C2 cells in the presence of 0.1 mg/mL colchicine. Two cyclic hexapeptides, waiakeamide (81) and its sulfone derivative 82, and five cyclic heptapeptides, the haliclonamides A–E (83–87) (Figure 8), were sequentially purified from a Palau-derived *Haliclona* sp. sponge [44–46]. Bioassay results showed that these peptides had potent antifouling activity at the concentration of 100 ppm, except for compounds 84 and 86.
Figure 8. Chemical structures of compounds 71-87 from unclassified *Haliclona* sps.
2.7.3. Miscellaneous Alkaloids

A new cytotoxic polycyclic alkaloid njaoamine I (88) containing a quinoline system and a known cytotoxic compound njaoamine G (89) were detected in the methanol extract of a Haliclona (Reniera) sp. sponge collected from Okuza Island (Tanzania) (Figure 9) [47]. Two isoquinoline alkaloids, 1-hydroxymethyl-7-methoxyisoquinolin-6-ol (90) and mimosamycin (91) (Figure 9), were purified from a Haliclona sp. sponge collected at Jessie Beazley Reef (Philippines) [48]. Interestingly, compound 91 was also produced by the marine sponge Cribrochalina and had strong cytotoxic effect on human tumor cell lines LOX, OVCAR-3 and HeLa cells with IC50 values of 10, 10, 2.6 μg/mL, respectively [48, 49]. Manzamines A (92) and Y (93) (Figure 9), two unusual alkaloids with β-carboline and isoquinoline skeletons and a 13-element dense, N-containing polycyclic structure unit, were isolated and characterized from two specimens of the sponge Haliclona sp., which were respectively collected from Manzamo and Iriomote Island [50,51]. Compound 92 could strongly inhibit the growth of mouse P-388 cells with an IC50 value of 0.07 μg/mL while 93 showed weak cytotoxicity on KB cells (IC50 = 7.3 μg/mL).
Figure 9. Chemical structures of compounds 88–103 from unclassified Haliclona sps.

Four antifungal amino alcohols, halaminols A-D (94 – 97) (Figure 9), were purified from a Haliclona sp. sponge grown on the Great Barrier Reef and their relative configurations were deduced from the NMR characteristics of oxazolidinone derivatives and absolute configurations were determined by their MPA esters [52]. Two new uncommon amino ketones, (6Z,9Z,12Z,15Z)-1-{(2-phenyl-ethyl)amino}octadeca-6,9,12,15-tetraen-3-one (98) and (6Z,9Z,12Z,15Z)-1-(diethylamino)octadeca-6,9,12,15-tetraen-3-one (99) (Figure 9), were separated from an unclassified Haliclona sponge collected from Weizhou Island (Guangci, China) [53]. Chemical study of the EtOH extract of a Haliclona sp. sample collected from Iriomote Island (Japan) afforded two new halichondriamine derivatives, halichondriamine C (100) and 1-epi-halichondriamine C (101) along with papuamine (22) and halichondriamine (23) (Figure 9) [19]. Compounds 100 and 101 could inhibit the growth of Mycobacterium bovis BCG with the same MIC values of 0.5 μg/mL and M. intracellulare with MIC values of 1.0 and 0.5 μg/mL, respectively. Additionally, two purine derivatives, 1,3-dimethylpurine (102) and 1,3-dimethyl-6-imino (103) (Figure 9), were separated from a Haliclona sp. sponge grown on Hainan Island (China) [54].

3. Bioactive Alkaloids from Haliclona-Derived Microbes

Marine sponges are important hosts for a large community of microorganisms, which are shown to be great producers of secondary metabolites [55]. However, only eight alkaloids 104–112 have been separated from Haliclona sponge-derived microbes until now (Figure 10). Chemical analysis of the ethyl acetate extract of the strain Bacillus megaterium LC3CS2 symbiont of the sponge Haliclona oculata collected from Son Cha Peninsula (Vietnam) afforded three anti-microbial agents: 7,7-bis(3-indolyl)-p-cresol (104), cyclo-(Pro-Leu) (105) and cyclo-(Pro-Val) (106) (Figure 10) [56]. These chemicals had antimicrobial activities against Vibrio vulnificus, V. parahaemolyticus and Trichophyton mentagrophytes with MIC values ranging from 0.05 to 5.0 μg/mL. Compound 104, formerly obtained from a marine sponge Hyatell-derived microbe Vibrio sp. was shown to inhibit the growth of Bacillus cereus and Micrococcus luteus with MIC values of 0.5 and 0.005 μg/mL, respectively [56,57]. Alantrypinone (107) along with lovastatin, methyl ester of lactone ring-opened monacolin K, terrein, territrems B and ergosterol was separated from a F62 fungal strain associated with the sponge Haliclona simulans collected from the South China Sea (Figure 10) [58]. Screening of symbiotic strains from the marine sponge Haliclona sp. collected from the sea shore of Tateyama city (Japan) led to the discovery of four new Streptomyces strains [59]. Later chemical investigation of strains GE-23 GE-26 and SC-24 afforded five new alkaloids JBIR-30, -34, -35, -39 and -40 (108–112) (Figure 10). However, none of these compounds had potent cytotoxic effects on human cervical carcinoma HeLa cells and malignant pleural mesothelioma ACC-MESO-1 cells.
4. Conclusions

In summary, 112 nitrogenous secondary metabolites have been isolated and characterized from the marine sponge genus *Haliclona* and its derived microbes till now. Only five alkaloids (compounds 16, 22, 23, 91 and 104) were separated from other organisms. Therefore, this indicates that *Haliclona* sponges are some of the most prolific sources of exclusive bioactive alkaloids despite the fact only a handful of classified species had been chemically investigated. It is well-known that marine organisms have served as a primary source of bioactive natural products during the past several decades. Nowadays, however, a rapid decrease in the speed of discovery of new compounds from Nature strongly necessitates new research strategies and approaches. Microorganisms are ubiquitous in the ocean owing to their stronger adaptability. During long co-evolution with marine sponges, symbiotic microbes maybe play important physiological and ecological roles in promoting host growth and increasing the resistance to predators and omnivores by excreting toxic metabolites. Therefore, more efforts should be made to explore and identify unknown *Haliclona* sponges and their derived symbiotic microbes and to carry out chemical studies for the discovery of novel therapeutical agents.

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