Review

Marine Indole Alkaloids—Isolation, Structure and Bioactivities

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Abstract: Indole alkaloids are heterocyclic natural products with extensive pharmacological activities. As an important source of lead compounds, many clinical drugs have been derived from natural indole compounds. Marine indole alkaloids, from unique marine environments with high pressure, high salt and low temperature, exhibit structural diversity with various bioactivities, which attracts the attention of drug researchers. This article is a continuation of the previous two comprehensive reviews and covers the literature on marine indole alkaloids published from 2015 to 2021, with 472 new or structure-revised compounds categorized by sources into marine microorganisms, invertebrates, and plant-derived. The structures and bioactivities demonstrated in this article will benefit the synthesis and pharmacological activity study for marine indole alkaloids on their way to clinical drugs.

Keywords: marine alkaloids; indoles; natural products; cytotoxicity; antibacterial; bioactivities

1. Introduction

Marine natural products have incomparable skeleton diversity and novelty relative to terrestrial source ones. They often exhibit superexcellent physiological activities and occupy an important position in today’s pharmaceutical industry as a continuously rich source of potential drugs [1–5]. The diversity of their structure enables them to have a broader range of pharmacological activities and action mechanisms, such as neuroprotection, analgesia, smoking cessation, antibacterial, antiviral, antitumor, anti-hypotension, and anti-hyperlipidemia [6].

The indole nucleus is one of the most crucial ring systems in nature. It has been termed a “privileged structure” in respect of pharmaceutical development. Viibryd (vilazodone, neurological disorders), decapetyl (tripirelmin, hormonal disorders), symdeko (tezacafitor and ivacaftor, genetic disorders), cialis (tadalafil, sexual health), cubicin (daptomycin, anti-bacterial), zepatier (elbasvir and grazoprevir, infectious diseases), tagrisso (osimertinib, oncology), sutent (sunitinib, oncology), zoladex (goserelin, oncology), alecensa (lectinib, oncology) and lupron (leuprolide, oncology) are all indole-containing top 200 small molecule pharmaceuticals by retail sales in 2018, which were summarized by Njarðarson Group (The University of Arizona, https://njardarson.lab.arizona.edu, 30 October 2021). Due to the high market occupancy and diverse physiological activities, indole alkaloids are now a research hotspot for pharmacologists. In recent years, pharmacological activities of indole alkaloids have been reviewed, including indole alkaloids with anti-diabetic activity [7], anti-malarial potential [8], anti-depression and anti-anxiety activity [9], antitumor and anti-drug-resistant cancer activity [10,11], and immune-regulatory activity [12]. This review focuses on marine indole alkaloids discovered since 2015, when the last comprehensive review, covering the time from 2003 to 2015, was reported by Netz and Opatz [13]. In this review, the newly isolated and structure-revised indole alkaloids from 2005 to 2021, 472 in total, are reported by the classification of sources. All the chemical structures are drawn in this review, and the bioactivities are discussed. The
general information of the cell lines mentioned in this review are listed in Table S1. The sources and bioactivities of all the reviewed marine indole alkaloids are summarized in Tables S2 and S3. The structures of these marine indole alkaloids were elucidated by various spectroscopic techniques. High-resolution mass spectrometer (HRMS) and 1D/2D nuclear magnetic resonance (1D/2D NMR) are the primary techniques for structure determination. Ultraviolet (UV) and infrared (IR) data are also used as auxiliary proofs. For compounds with chiral centers, the absolute configurations could be determined by specific rotation, electronic circular dichroism (ECD), X-ray single-crystal diffraction and Marfey’s method, etc. During the discovery of natural products, structures were mistaken especially for absolute configurations, which happened occasionally. Chemical total synthesis of the natural product and comparing the NMR spectroscopy between the synthetic product and the natural product is another precise but complex and expensive method for structure determination and revision.

2. Marine Microorganisms

2.1. Marine-Sourced Bacteria

Marine-sourced bacteria are one of the richest producers of bioactive natural products. There are 64 new indole metabolites isolated from marine-sourced bacteria, including 38 from sediment-sourced bacteria and 12 from sponge-sourced bacteria. If classified by the source of bacterial species, most of the indole alkaloids are found from actinomycetes.

2.1.1. Sediment-Sourced Bacteria

Isonasesazime B (1), an antimicrobial diketopiperazine dimer, was isolated from *Streptomyces* sp. SMA-1 by bioassay-guided separation (Figure 1). *Streptomyces* sp. SMA-1 was one of the 613 actinobacterial strains isolated from the sediments collected from the Yellow Sea, China [14]. Indolepyrazines A (2) and B (3) were isolated from *Acinetobacter* sp. ZZ1275, and they showed antimicrobial activities against methicillin-resistant *Staphylococcus aureus* (MRSA), *Escherichia coli* (E. coli), and *Candida albicans* with minimum inhibitory concentration (MIC) values of 12 µg/mL, 8–10 µg/mL, and 12–14 µg/mL, respectively. Indolepyrazine A (2) is the first indole-pyrazine-oxindole alkaloid, and both 2 and 3 are the first reported natural products isolated from marine-derived *Acinetobacter* species [15]. Streptopenylindoles A–C (4–6) were acquired from *Streptomyces* sp. ZZ820. Streptopenylindoles A and B were enantiomers that were separated by the preparation of Mosher’s method. No inhibiting activities of the streptopenylindoles were reported for the tested MRSA and *E. coli* [16]. 3-hydroxy-N-methyl-2-oxindole (7–8) were obtained from marine *Salinispora arenicola* strain from sediments of Brazil, and they showed no antibacterial activity against Gram-positive (*Enterococcus faecalis* and *Staphylococcus aureus*) and Gram-negative (*E. coli*) bacteria strains [17]. Two new chlorinated bisindole alkaloids, dionemycin (9) and 6-OMe-7”dichlorochromopyrrolic acid (10) were isolated from the deep-sea derived *Streptomyces* sp. SCSIO 11791. In vitro antibacterial and cytotoxic assays revealed that compound 9 shows anti-staphylococcal activity with a MIC range of 1–2 µg/mL against six clinic strains of MRSA isolated from human and pig. The cytotoxicity of the trichloro-bisindole 9 was evaluated on human cancer cell lines NCI-H460, MDA-MB-231, HCT-116, HepG-2, and noncancerous MCF10A with IC_{50} values ranging from 3.1 to 11.2 µM. Structure–activity relationship analysis of compounds 9, 10, and seven known analogs showed C-6” chlorine as an essential pharmacophore in their cytotoxic activities [18].

A total of 18 new indolocarbazole alkaloids (11–28) isolated from *Streptomyces* sp. DT-A61, A65, A68, A22, OUCMDZ-3118 and *lingchengensis* ULS14 were reported in a roll during 2018 and 2019 (Figure 2). Compounds 11–25 were evaluated for cytotoxic activity against PC3 cell line. Compound 20 showed the strongest cytotoxic activity against PC3 with an IC_{50} value of 0.15 µM, and the other indolocarbazoles exhibited moderate activities against the PC3 (IC_{50} = 0.8–41.3 µM). Compounds 11–25 were also tested for various enzyme inhibition activities of protein kinase C and bruton tyrosine kinase. Compound 12...
displayed significant and selective inhibition against ROCK2 and the other indolocarbazole also showed moderate inhibition activities to different kinases. Compound 26 showed moderate activity with IC\textsubscript{50} values of 0.91–1.84 μM for the tested protein kinases enzyme inhibition activities. Compound 27 was moderately effective against the A549 and MCF-7 cell lines with IC\textsubscript{50} values of 1.2–1.6 μM. The IC\textsubscript{50} of 28 against the HeLa cell line was 0.075 μg/mL [19–24].

![Figure 1. Chemical structures of 1–10.](image1)

![Figure 2. Cont.](image2)
Two new brominated bis-indole metabolites, 5-bromometagenediindole B (29), and 5-bromometagenediindole C (30) were separated under the guidance of LC-MS from the 25D7 clone derived *E. coli* fermentation broth, in which 5-bromoindole was added. 5-Bromometagenediindole B (29) demonstrated moderately cytotoxic activity against MCF-7, B16, CNE-2, BEL-7402, and HT-1080 tumor cell lines in vitro (Figure 3) [25]. 3,3′-bis-indole (31) were isolated from sediment-derived actinomycete *Nocardiopsis* sp. G057 as a natural product for the first time. Compound 31 exhibited antimicrobial activity against several strains of bacteria, and the yeast *Candida albicans* with values of MIC ranging from 64 to 256 µg/mL. Cytotoxic evaluation of compound 31 against four cancer cell lines (KB, LU-1, HepG-2, and MCF-7) indicated that 31 produced a weak inhibition against KB and LU cell lines (IC$_{50}$ = 12.5 and 25.6 µg/mL) [26]. 1-methyl-4-methylthio-β-carboline (32) was tracked by the GNPS MS$^2$ fragmentation pattern analysis tool and separated by a scale-up liquid culture of *Achromobacter spanius*. No bioactivity was reported [27]. Spiroindimicins E (33) and F (34) were identified by combined genomics-metabolomics profiling of marine *Streptomyces* sp. MP131-18, and demonstrated the potential of actinomycetes in combinatorial biosynthesis of secondary metabolites [28].
Taromycin B (35) was produced by heterologous expression of the activated taromycin biosynthetic gene clusters from marine actinomycete *Saccharomonospora* sp. CNQ-490. It showed potent activity against methicillin-resistant *Staphylococcus aureus* and vancomycin-resistant *Enterococcus faecium* (Figure 4) [29]. New xiamycin analogs (36–38) were isolated via genome mining of *Streptomyces xinghaiensis* NRRL B-24674T, and the bioactivity was not evaluated [30].

2.1.2. Sponge-Sourced Bacteria

Enhypyrizinones A and B (39 and 40), from a marine-derived myxobacterium *Enhygromyxa* sp., showed weak activity (MIC values > 128 µg/mL) against *E. coli*, methicillin-resistant *Staphylococcus aureus* (MRSA), and methicillin-sensitive *Staphylococcus aureus* (MSSA) (Figure 5) [31]. Investigation of the bioactive secondary metabolites of the sponge-
derived actinomycete Rubrobacter radiotolerans led to the isolation and characterization of another new naturally rare dimeric indole derivative (41). Compound 41 showed moderate antichlamydial activity with IC$_{50}$ values of 46.5–96.4 μM against different Chlamydia [32]. Rhodozeppinone (42), a new azepino-diindole alkaloid, was isolated and identified from the broth culture of Rhodococcus sp. UA13, which had been previously recovered from the Red Sea sponge Callyspongia aff. Implexa. Rhodozeppinone (42) exhibited significant antibacterial and antitrypanosomal activities against Staphylococcus aureus NCTC 8325 (IC$_{50}$ = 8.9 μg/mL) and Trypanosoma brucei brucei TC221 [IC$_{50}$ = 16.3 (48 h) and 11.8 (72 h) μg/mL], respectively [33]. Anthranoside C (43), discovered from actinomycete Streptomyces sp. CMN-62, which was originated from a marine sponge, could inhibit influenza H1N1 virus with an IC$_{50}$ value of 171 μM (ribavirin as positive control, IC$_{50}$ = 133 μM) [34]. Lysinibacillus fusiformis was one of the 48 sponge-associated microbes identified from Halichondria okadai by testing 720 kinds of culture conditions. Lysiformine (44) was isolated from Lysinibacillus fusiformis, and displayed cytotoxicity toward mouse leukemia P388 cells with an estimated IC$_{50}$ value of 10 μM [35]. Saccharomonosporine A (45), a novel brominated oxo-indole alkaloid and convolutamydine F (46) were isolated from Saccharomonospora sp. UR22 and Dietzia sp. UR66 co-culture. Compound 45 was a potential Pim-1 kinase inhibitor that mediate the tumor cell growth inhibitory effect with an IC$_{50}$ value of 0.3 ± 0.02 μM on Pim-1 kinase and significant antiproliferative activity against HT-29, (IC$_{50}$ = 3.6 μM) and HL-60, (IC$_{50}$ = 2.8 μM) [36]. Photopiperazines A–D (47–50) were isolated from sponge-derived actinomycete AJS-327. The cytotoxicity of photopiperazines A–D mixture was evaluated on four cancer cell lines. It showed 12,000-fold selective toxicity toward U87 and SKOV3 than MDA-MB-231 and HCT-116 cell lines with the IC$_{50}$ values of 0.41 nM and 0.75 nM, respectively [37].

![Chemical structures of 39–50.](image1)

**Figure 5.** Chemical structures of 39–50.
2.1.3. Miscellaneous

A new phenylamine-incorporated angucyclinone (51) was discovered from marine *Streptomyces* sp. PKU-MA00218 (Figure 6). Compound 51 was produced from a nonenzymatic conversion of the type II PKS-produced precursor. In addition, 18 new phenylamine-incorporated angucyclinone derivatives were prepared by the efficient nonenzymatic conversion under mild conditions. All 19 compounds showed different degrees of activity on nuclear factor erythroid 2-related factor 2 (Nrf2) transcription in HepG2 cells at 10 μM [38]. Sulfadixiamycins A–C (52–54) are a new class of antibiotics featuring sulfanilamide and dapsone substructures firstly reported from natural sources. They were discovered from recombinant *Streptomyces* species harboring the entire xiamycin biosynthesis gene cluster and exhibited moderate antimycobacterial activities and potent antibiotic activities even against multidrug-resistant bacteria [39].

![Chemical structures of 51–54.](image)

Figure 6. Chemical structures of 51–54.

Two psychrotolerant bacterial strains *Vibrio splendidus* T262 and *Arthrobacter psychrochitiniphilus* T406 were isolated from the gastrointestinal tract of a fish and the excrement of penguins near the South Orkney Islands in Antarctica. Seven new indole alkaloids, trisindolal (55), turbomycin C–F (56–59), 4-(1H-indol-3-yl-sulfanyl) phenol (60), and 2-(indol-3-ylmethyl)-indol-3-ylethanol (61) were obtained from T262 and T406 (Figure 7). Trisindolal (55) was active against the peronosporomycetes *Botrytis cinerea* and *Phytophthora infestans*, and showed pronounced potency and selectivity in a panel of 11 human tumor cell lines derived from 10 different tumor histotypes [40]. 6-Bromo-N-propionyltryptamine (62) were isolated and identified from a marine bacterium *Pseudoalteromonas rubra* strain isolated from seawater and exhibited weak 5-HT2A receptor antagonist activity (~10% inhibition, 10 μM) [41]. Another simple indole alkaloid (63) was isolated from the deep-sea-derived bacterium *Bacillus subterrneus* 11593 and displayed no anti-allergic bioactivity [42]. Compound 64 was
isolated from *Pseudovibrio denitrificans* strain isolated from seawater. It showed cytotoxic effect against L929 cells (EC$_{50}$ = 7 µM) and A549 cells (EC$_{50}$ = 8 µM) [43].

Figure 7. Chemical structures of 55–64.

2.2. Marine-Sourced Fungi

Marine fungi are important components of marine microorganisms, and they are the main source of marine natural products. Among them, Cephalosporin C is the brightest star molecule as the first marine antibiotic [44,45]. In this part, 257 new indole alkaloids were summarized, including 93 from sediment-derived fungi, 62 from coral-derived fungi, 19 from bivalve-mollusk-derived fungi, 20 from Mangrove-sourced fungi, 16 from marine alga endophytic fungi, and 20 from sponge-sourced fungi.

2.2.1. Sediment-Sourced Fungi

Cyclopiamides B–J (65–73), nine new oxindole alkaloids were isolated from the sediment-derived fungus *Penicillium commune* DFFSCS026 (−3563 m in the South China Sea) (Figure 8). Compounds 65–73 and positive control (ochratoxin A) displayed lethal activity on brine shrimp with LC$_{50}$ values of 25.2, 38.5, 14.1, 24.8, 25.6, 34.7, 16.4, 33.5, 26.7 and 6.2 mg/mL, respectively. Compounds 65–73 showed no cytotoxicity towards human carcinoma HepG-2 and HeLa cell lines at a concentration of 100 mg/mL, and no anti-influenza virus H1N1 activity under their largest concentration of non-toxic towards the tested MDCK cell [46].
Haenamindole (74) and citreoindole (75) were isolated from a South China Sea deep-sea fungus, *Penicillium citrinum* MF006 (Figure 9) [47]. The structures of the rare alkaloids 74 and 75 were revised based on detailed spectroscopic and C3 Marfey’s analysis [48,49].

Penicimutamides A–E (76–80) and a structure-revised asperverin (81), six new prenylated indole alkaloids, were isolated from a diethyl sulfate mutagenesis mutant of the marine-derived fungus *Penicillium purpurogenum* G59. Compound 76 and compounds 78–80 displayed less than 28.5% inhibition rates on human K562, HL-60, HeLa and BGC-823 cell lines at the concentration of 100 μg/mL, while compound 77 showed remarkable inhibition rates (77.3%–92.7% at 100 μg/mL) and the further determined IC50 values ranged from 20 to 52 μg mL−1 (Figure 10) [50,51].
Asperversiamides A–H (82–89) and structure-revised iso-notoamide B (90) were isolated from the marine-derived fungus *Aspergillus versicolor* (Figure 11). Asperversiamide G (88) exhibited a potent anti-inflammatory activity with an IC\textsubscript{50} value of 5.39 μM against iNOS (Figure 10) [52]. Eight new diketopiperazines (91–98) were isolated from a marine-derived fungus *Aspergillus versicolor* MF180151, and they showed no activity against the tested pathogens (*Candida albicans, Bacillus subtilis, Staphylococcus aureus, Methicillin-resistant S. aureus, Pseudomonas aeruginosa, and Bacillus Calmette-Guérin*) [53]. Roquefortine J (99) was founded in *Penicillium granulatum* MCCC 3A00475 and with an IC\textsubscript{50} value of 19.5 μM against HepG2 tumor cells [54].

Acremonpeptide D (100), together with Al(III)-acremonpeptide D (101) were obtained from the marine fungus *Acremonium persicinum* SCSIO 115 (Figure 12). In vitro bioassays revealed Al(III)-acremonpeptide D (101) as moderate antiviral agents for herpes simplex virus 1 with an EC\textsubscript{50} value of 14 μM (Figure 11) [55]. Cytoglobosins H (102) and I (103) were isolated from the deep-sea-derived fungus *Chaetomium globosum*, which was obtained from a deep-sea sediment sample (−2500 m depth) of the Indian Ocean. They showed weak to no cytotoxicity against MDA-MB-231, LNCaP and B16-F10 cell lines (IC\textsubscript{50} > 9 μM) [56].

**Figure 10.** Chemical structures of 76–81.

**Figure 11.** Chemical structures of 82–99.

Acremonpeptide D (100), together with Al(III)-acremonpeptide D (101) were obtained from the marine fungus *Acremonium persicinum* SCSIO 115 (Figure 12). In vitro bioassays revealed Al(III)-acremonpeptide D (101) as moderate antiviral agents for herpes simplex virus 1 with an EC\textsubscript{50} value of 14 μM (Figure 11) [55]. Cytoglobosins H (102) and I (103) were isolated from the deep-sea-derived fungus *Chaetomium globosum*, which was obtained from a deep-sea sediment sample (−2500 m depth) of the Indian Ocean. They showed weak to no cytotoxicity against MDA-MB-231, LNCaP and B16-F10 cell lines (IC\textsubscript{50} > 9 μM) [56].
Two alkaloids, fumigatosides E (104) and F (105) were isolated from deep-sea derived fungal Aspergillus fumigatus SCSIO 41012 (Figure 13). Compound 104 showed significant antifungal activity against Fusarium oxysporum f. sp. momordicae with MIC at 1.56 μg/mL, and compound 105 exhibited significant activity against A. baumanii with a MIC value...
of 6.25 µg/mL [57]. Penijanthes C and D (106 and 107) were isolated from the marine-derived fungus *Penicillium janthinellum*. They displayed significant anti-Vibrio activity (MIC values ranging from 3.1 to 50.0 µM) against three pathogenic *Vibrio* spp. [58]. Two new compounds 195,20-epoxy-18-oxotryprostatin A (108) and 20-hydroxy-18-oxotryprostatin A (109) were discovered from the marine-derived fungus *Aspergillus fumigatus* MF071 from the Bohai Sea sediment. For the limited amounts of 108 and 109, no activities have been evaluated yet [59]. Four indole diketopiperazine alkaloids aspechinulins A–D (111–113 and 110) were isolated from the sediment-derived fungus *Aspergillus* sp. FS445. Compounds 111–113 represented the first examples of indole diketopiperazine derivatives constructing a C-5 unit at 11-NH through an imide linkage. Compound 113 exhibited the most potent inhibitory activities against NO production with the IC$_{50}$ value of 20 µM, which was as effective as the positive control aminoguanidine (IC$_{50}$ = 23.7 µM) [60].

![Chemical structures of 104–113.](image)

Three new prenylated indole 2,5-diketopiperazine alkaloids (114–116), one new indole alkaloid (117), and six pairs of new spirocyclic alkaloid enantiomers eurotinoids A–C (118–123) and variecolortins A–C (124–129) were characterized from the sediment-derived fungus *Eurotium* sp. SCSIO F452 (Figure 14). Compound 116 and all the spirocyclic alkaloids 118–123 showed significant radical scavenging activities against DPPH with IC$_{50}$ values ranging from 3.7 to 24.9 µM. None of the alkaloids (114–129) exhibited obvious cytotoxicity against SF-268 and HepG2 cell lines. Interestingly, (+)-enantiomers (118, 120

![Chemical structures of 104–113.](image)
and 122) exhibited more potent activities than the corresponding (−)-enantiomers (119, 121 and 123). (−)-enantiomer 124 exhibited stronger antioxidative activity than (−)-enantiomer 125, while (+)-enantiomers (126 and 128) showed more potent cytotoxicities against SF-268 and HepG2 cell lines than (−)-enantiomers (127 and 129), which indicated that different enantiomers might result in different biological activities [61–63].

Graphiumins I (130) and J (131) were isolated from the culture broth of the marine-derived fungus Graphium sp. OPMF00224 (Figure 15). Compounds 130 and 131 inhibited yellow pigment production by MRSA with IC_{50} values of 63.5 and 76.5 µg/mL, respectively, without inhibiting its growth, even at 250 µg/mL [64]. Dichotocejpins A and C (132 and 133) were isolated from the culture of the deep-sea sediment-derived fungus Dichotomomyces cejpii FS110. Compound 132 exhibited excellent inhibitory activity against α-glucosidase with an IC_{50} of 138 µM [65]. Cristazine (134) was isolated from the mudflat-sediment-derived fungus Chaetomium cristatum. It displayed potent radical-scavenging activity against 2,2-diphenyl-1-picrylhydrazyl (DPPH), with IC_{50} values of 19 µM, and cytotoxic activity against HeLa cells, with an IC_{50} value of 0.5 µM [66]. Chetracins E and F (135 and 136) were isolated from the fungus Acrostalagmus luteoalbus HDN13-530 and showed potent

Figure 14. Chemical structures of 114–129.
cytotoxic effects on A549, HCT-116, K562, H1975 and HL-60 with IC$_{50}$ values ranging from 0.2 to 3.6 µM [67]. In the study of fungal and bacterial co-cultivation, a new indole alkaloid brevianamide X (137) was isolated from the Aspergillus fumigatus MR2012 fermentation [68].

Figure 15. Chemical structures of 130–137.

Five new prenylated indole alkaloids, 17-hydroxynotoamide D (138), 17-O-ethylnotoamide M (139), 10-O-acetylsclerotiamide (140), 10-O-ethylsclerotiamide (141) and 10-O-ethylnotoamide R (142) were isolated from the co-culture of marine-sediment-derived fungi Aspergillus sulphureus KMM 4640 and Isaria felina KMM 4639 (Figure 16). Compound 139 inhibited the colony formation of human prostate cancer cells 22Rv1 at non-cytotoxic concentration of 10 µM [69]. Two brevianamides (143 and 144) were isolated from the deep-sea-derived fungus Penicillium brevicompactum DFFSCS025. Both of them exhibited no cytotoxicity against HCT116 and no antibacterial or antifungal activities against Streptococcus mutans, S. sobrinus and Fusarium oxysporum f. sp. cubense Race 1 and Race 4. Compound 143 showed no antilarval activity in the larval settlement bioassay [70].

Figure 16. Cont.
Six new prenylated indole diketopiperazine alkaloids, asperthrins A–F (145–150), were isolated from the marine-derived fungus Aspergillus sp. YJ191021 (Figure 17). Compound 145 exhibited moderate antifungal and antibacterial activities against Vibrio anguillarum, Xanthomonas oryzae pv. Oryzicola, and Rhizoctonia solani with MIC values of 8, 12.5, and 25 µg/mL, respectively. Furthermore, 145 displayed notable anti-inflammatory activity with an IC50 value of 1.46 ± 0.21 µM in Propionibacterium acnes induced human monocyte cell line (THP-1) [71].

Seven new indole marine natural products (151–157) were isolated from four mangrove swamp-derived fungi (Figure 18). Trypilepyrazinol (151) was isolated from the fungus Penicillium sp. IMB17-046. Trypilepyrazinol exhibited inhibitory activities against...
HIV-1 and HCV with IC$_{50}$ values of 4.6 and 7.7 µM, respectively. It also showed antibacterial activities against Helicobacter pylori. (including the drug-sensitive strain G27 and the drug-resistant strain 159), but inactive against Gram-positive Staphylococcus aureus and Bacillus subtilis and Gram-negative Pseudomonas aeruginosa and Klebsiella pneumonia [72]. Four new prenylated indole alkaloids (152–155) were isolated from Penicillium sp. SCSIO041218 and inactive to the tested anti-allergic bioactivity on IgE-mediated rat mast RBL-2H3 cells [73]. A new prenylated indole alkaloid, named paraherquamide J (156), was isolated from another mangrove rhizosphere soil-derived fungus Penicillium janthinellum HKI-6. No activity was found for the tested antibacterial, topoisomerase I (topo I) inhibitory activities and lethality towards brine shrimp Artemia salina [74]. Raistrickindole A (157), a new indole diketopiperazine alkaloid, was isolated from the fungus Penicillium raistrickii IMB17-034. Compounds 157 showed anti-HCV activity with an EC$_{50}$ value of 5.7 µM in the in vitro inhibitory assay against the hepatitis C virus life cycle [75].

2.2.2. Coral-Sourced Fungi

Wen-Jian Lan and co-workers have conducted a great deal of research focused on the chemical diversity of fungi associated with soft coral. In recent years, nineteen marine indole alkaloids (158–176) have been identified by an amino acid-directed strategy, which is a method of feeding various amino acids to marine fungi (Figure 19). Utilizing this strategy, dichotomoomycetes cepii F31-1, scedapins A–E (161–164, 160) and scequinadolines A–G (165–171), scetryptoquivaline A (172), scequinadoline I (173), and scequinadoline J (174) were isolated from Scedosporium apiospermum F41-1 by the same amino acid-directed strategy. Another two new bisindole alkaloids, pseudoindoles A–B (175–176), were isolated from L-tryptophan, L-phenylalanine, L-methionine, and L-threonine fed Penicillium janthinellum F31-1. Among these compounds, scedapin C (163) and scequinadoline D (168) displayed significant antiviral activity against hepatitis C and scequinadoline J (174), and they promote triglyceride accumulation in 3T3-L1 cells [76–79].
scequinadoline D (168) displayed significant antiviral activity against hepatitis C and scequinadoline J (174), and they promote triglyceride accumulation in 3T3-L1 cells [76–79].

Figure 19. Chemical structures of 158–176.

Between the years 2015 and 2016, Wen-Jian Lan and co-workers also isolated five indole alkaloids (177–181) directly from coral-derived fungi, including Pseudellones A–D (177–180) from Pseudallescheria ellipsoidea F42-3 and 181 from Pseudallescheria boydii. No significant tested bioactivity was reported (Figure 20) [80–82].

A new cytochalasin, 6-O-methyl-chaetoglobosin Q (182), was isolated from the coral-associated fungus Chaetomium globosum C2FI7, and no bioactivity was reported (Figure 21) [83]. Three new indole diketopiperazine alkaloids, 11-methylnoechinulin E (183), variecolorin M (184) and (+)-variecolorin G (185) were isolated from a soft coral-associated epiphytic fungus Aspergillus sp. EGF 15-0-3. They all have no in vitro toxicity against NCI-H1975/GR cell line at the concentration of 50 µM [84]. Seven new deoxyisoaustamide derivatives (186–192) were isolated from the coral-derived fungus Penicillium dimorphosporum KMM
Compounds 189–191 revealed a statistical increase in PQ (paraquat)-treated Neuro-2a cell viability by 30–39% at a concentration of 1 μM [85].

Figure 20. Chemical structures of 177–181.

A new cytochalasin, 6-O-methyl-chaetoglobosin Q (182), was isolated from the coral-associated fungus Chaetomium globosum C2F17, and no bioactivity was reported (Figure 21) [83]. Three new indole diketopiperazine alkaloids, 11-methylneoechinulin E (183), variecolorin M (184) and (+)-variecolorin G (185) were isolated from a soft coral-associated epiphytic fungus Aspergillus sp. EGF 15-0-3. They all have no in vitro toxicity against NCI-H1975/GR cell line at the concentration of 50 μM [84]. Seven new deoxyisoaustamide derivatives (186–192) were isolated from the coral-derived fungus Penicillium dimorphosporum KMM 4689. Compounds 189–191 revealed a statistical increase in PQ (paraquat)-treated Neuro-2a cell viability by 30–39% at a concentration of 1 μM [85].

Figure 21. Chemical structures of 182–192.

Seventeen fumiquinazoline-type alkaloids, versiquinazolines A–Q (193–209), were isolated from the gorgonian-derived fungus Aspergillus versicolor LZD-14-1 and the structures of cottoquinazolines B, D and C (210–212) were revised to enantiomers (Figure 22). Com-
pounds 193, 194, 199, 203, 208 and 209 exhibited inhibitory activities against thioredoxin reductase (IC$_{50}$ values ranging from 12 to 20 µM) [86,87].

Aspergillipeptide E (213) was isolated from Aspergillus sp. SCSIO 41501 and 213 showed evident antiviral activity against herpes simplex virus type 1 (HSV-1) with an IC$_{50}$ value of 19.8 µM under the non-cytotoxic concentrations against a Vero cell line (Figure 23) [88]. Luteoride E (214) was isolated and identified from a coral-associated fungus Aspergillus terreus. Luteoride E inhibited no α-Glucosidase inhibitory activity and
with moderate inhibitory activity against LPS-induced NO production [89]. Aspergillspins A-B (215–216) were isolated from the marine gorgonian-derived fungus Aspergillus sp. SCSIO 41501. They exhibited no cytotoxicity activities against the tested HL-60, HepG2 and MCF-7 cell lines and no antibacterial activities against Bacillus subtilis and E. coli [90]. Three new cycloheptapeptides, asperversiamides A–C (217–219), were isolated from coral-derived fungus Aspergillus versicolor CHNSCLM-0063 under the guidance of molecular networking and 1H NMR. Asperversiamides A–C (217–219) exhibited potent inhibitory activity against M. marinum [91].

Figure 23. Chemical structures of 213–219.

2.2.3. Mollusk-Sourced Fungi

From the Penicillium sp. KFD28, an endophytic fungus in a bivalve mollusk Meretrix lusoria collected from Haikou Bay, eleven new indole-diterpenoids named penerpenes B–H and J (220–226 and 227) and epipaxilline (228) were isolated in Du-Qiang Luo and You-Xing Zhao groups (Figure 24). All the compounds showed potent to moderate inhibitory activity toward protein tyrosine phosphatase PTP1B with IC50 ranging from 1.7 to 31.8 µM. Compounds 220, 226 and 227 showed inhibitory activity toward protein tyrosine phosphatase TCPTP with IC50 values of 5.0, 4.5, 35 and 14.7 µM, respectively [92–94].

Seven new quinazoline-containing indole alkaloids named aspertoryadins A–G (229, 230, 232, 233, 231, 234 and 235) were isolated from the marine-derived fungus Aspergillus sp. HNMF114, which was separated from the bivalve mollusk Sanguinolaria chinensis (Figure 25). Compound 229 bears an aminosulfonyl group in the structure, which is rarely encountered in natural products. Compounds 234 and 235 exhibited quorum sensing inhibitory activity against Chromobacterium violaceum CV026 with MIC values of 32, 32 and 16 µg/well, respectively [95]. A continuous work by feeding tryptophan to the marine-derived fungus Aspergillus sp. HNMF114 was carried out, another three new quinazoline-containing indole alkaloids aspertoryadins H–J (236–238) were obtained. The biological activity of these compounds against the insect ryanodine receptor (RyR) was tested using HEK cells stably expressing RyR from Spodoptera frugiperda (sfRyR) or RyR1
from rabbit (rRyR1) and R-CEPIA1er. Alkaloids 236–238 only showed a weak activation effect on sfRyR, which reduced the [Ca^{2+}]_{ER} by less than 7% [96].
2.2.4. Mangrove-Sourced Fungi

One of Bin-Gui Wang group’s studies focusing on the exploration of structurally unique and biologically active natural products from the mangrove-derived fungi isolated ten new indole alkaloids (Figure 26). Penioxamide A (239), a new prenylated indole alkaloid possessing a piperidine moiety, was isolated and identified from Penicillium oxalicum EN-201, an endophytic fungus obtained from the inner tissue of the fresh leaves of marine mangrove plant Rhizophora stylosa. Compound 239 bore the rare anti-relative configuration in the bicyclo[2.2.2]diazaocostane ring and showed potent brine shrimp lethality with LD_{50} values of 5.6 µM [97]. Three new diketopiperazines, including spirobrocazines A–B (240–241), were characterized from the mangrove-derived Penicillium brocae MA-231. Both 240 and 241 possess a 6/5/6/5/6 cyclic system. Compound 240 showed moderate and nonselective antimicrobial activity against E. coli, S. aureus and Vibrio harveyi, with MIC values of 32.0, 16.0 and 64.0 µg/mL, respectively, whereas no bioactivity was reported for the dearomatized 241 [98]. Six new indole-diterpenes rhizovarins A–C, E and F (242–244, 245 and 246) were identified from M. irregularis QEN-189, an endophytic fungus isolated from the fresh inner tissue of the marine mangrove plant Rhizophora stylosa. Compounds 242 and 243 showed activities against human A549 and HL-60 cancer cell lines with IC_{50} values ranging from 5.0 to 11.5 µM, and compound 245 exhibited activity against the A-549 cancer cell line with an IC_{50} value of 9.2 µM [99].

Three novel chaetoglobosins, named penochalasins I–K (247–249), were isolated from the culture of Penicillium chrysogenum V11 (Figure 27). Compound 248 greatly inhibited C. gloeosporioides (MIC = 25.08 µM), showing an antifungal activity higher than carben-dazim. Compound 247 exhibited marked cytotoxicity against MDA-MB-435 and SGC-7901 cells (IC_{50} < 10 µM) [100]. Compound 249 displayed significant inhibitory activities against C. gloeosporioides and R. solani (MICs = 6.13, 12.26 µM, respectively), and it also exhibited potent cytotoxicity against MDA-MB-435, SGC-7901 and A549 cells (IC_{50} < 10 µM) [101]. Neosartoryadins A (250) and B (251), with a unique 6/6/6/5 quinazoline ring system connected directly to a 6/5/5 imidazooindolone ring, together with compounds 252 and 253 were isolated from the endophytic fungus Neosartorya udagawae HDN13-313. Compounds 250 and 251 exhibited inhibitory effects against influenza A virus (H1N1) with IC_{50} values of 66 µM and 58 µM by the cytopathic effect (CPE) inhibition assay, which were smaller than the positive control ribavirin with the IC_{50} of 94 µM [102]. Three new
indole diterpenes, penicilindoles A–C (254–256), were isolated from the mangrove-derived fungus *Eupenicillium* sp. HJ002. After evaluating cytotoxic and antibacterial activities in vitro, penicilindole A (254) showed cytotoxic activity against human A549 and HepG-2 cell lines with IC$_{50}$ values of 5.5 and 1.5 µM, respectively [103]. Two prenylated indole 3-carbaldehydes (257 and 258) were purified from mangrove-derived endophytic fungus *Eurotium chevalieri* KUFA 0006, and they significantly inhibited the biofilm production in *S. aureus* [104].

Figure 27. Chemical structures of 247–258.

2.2.5. Alga-Sourced Fungi

Bin-Gui Wang’s group investigated the bioactive secondary metabolites of marine alga endophytic fungus. Thirteen new indole alkaloids were isolated, including four indolediketopiperazine alkaloids (259–262) from *Eurotium cristatum* EN-220, varioloid C (263) from *Paecilomyces variotii* EN-291, 4-epi-seco-shornephine A methyl ester (264) and 4-epi-seco-shornephine A carboxylic acid (265) from *Aspergillus alabamensis* EN-547 and three pairs of new N-methoxy-containing indolediketopiperazine enantiomers, acrozines A–C (266–271) from *Acrostalagmus luteoalbus* TK-43 (Figure 28). Compound 260 exhibited potent lethal activity against brine shrimp (LD$_{50}$ = 19.4 µg/mL) and weak nematicidal effect against *Panagrellus redivivus* (LD$_{50}$ = 110.3 µg/mL). Compound 263 exhibited cytotoxicity against A549, HCT116 and HepG-2 cell lines (2.5–6.4 µg/mL). Compounds 264–265 showed inhibitions against human pathogens *E. coli* and *M. luteus* and aquatic bacteria *Ed. ictaluri* and *V. alginolyticus* with MIC values ranging from 16 to 64 µg/mL [105–109]. 272 and 273 were
isolated from chemical-epigenetic cultures of *Aspergillus versicolor* OUCMDZ-2738 with 10 µM vorinostat (SAHA), and no antibacterial activity against the eight tested pathogenic microorganisms [108]. A new melatonin analog 6-hydroxy-N-acetyl-$\beta$-oxotryptamine (274) was isolated from the marine-derived fungus *Penicillium* sp. KMM 4672. It was not cytotoxic against neuroblastoma Neuro2a cells up to 100 µM and scavenged DPPH radicals by 48% at 100 µM. Compound 274 demonstrated increased cell viability in both 6-OHDA and PQ-induced neuronal cell damage models [110].

![Chemical structures of 259–274](image)

Figure 28. Chemical structures of 259–274.

2.2.6. Sponge-Sourced Fungi

Speradines B–D (275–277) were isolated from the sponge-derived fungus *Aspergillus flavus* MXH-X104 (Figure 29). Oxindoles 275–277 showed no activities on all the bioassay (cytotoxicities on P388, BEL-7402, A-549, Hela and HL-60 cells; inhibitory effects on H1N1 and HIV viruses, and antimicrobial activities on *Mycobacterium phlei*, *Staphylococcus aureus*, *Colibacillus sp.* and *Blastomyces albicans*) [111]. Methylthio-gliotoxin derivative 278 was firstly characteried from a sponge-derived fungus *Ascomycota Dichotomomyces cejpii* and the fungus was isolated from sponge *Callyspongia cf. C. flammea*. This study validates the anti-proliferative mechanisms of the newly isolated natural epipolythiodiketopiperazines via the inhibition of TNF-α-induced NF-κB activity [112]. Sartoryglabramide B (279) and fellutanine A (280) were isolated from sponge-associated fungus *Neosartorya glabra* KUFA 0702, and they exhibited no antibacterial and antifungal activities [113]. A new diketopiperazine dimer designated as SF5280-415 (281) was isolated from an ethyl acetate extract of the marine-derived fungus *Aspergillus* sp. SF-5280. Compound 281 showed inhibitory effects
against protein tyrosine phosphatase 1b (PTP1B) with an IC\textsubscript{50} value of 14.2 ± 0.7 \(\mu\)M [114]. Isopropylchaetominine (282) was isolated from fungus Aspergillus carneus using the OS-MAC (one strain many compounds) approach, and it showed potent cytotoxicity against the mouse lymphoma cell line L5178Y with IC\textsubscript{50} values of 0.4 \(\mu\)M [115]. Candidusin D (283) was isolated from the cultures of the marine sponge-associated fungus Aspergillus candidus KUFA 0062, and it was tested on various activities (antibacterial activity, biofilm formation inhibition activity and cytotoxic activity). Candidusin D showed cytotoxicity to all cell lines tested (HepG2, HT-29, HCT-116, A549, A375, MCF-7, U251 and T98G) except for T98G and HepG2 at the concentration of 100 \(\mu\)M [116]. Diketopiperazine dimer (284) was isolated from fungus Aspergillus violaceofuscus and it showed anti-inflammatory activity against IL-10 expression of the LPS-induced THP-1 cells with an inhibitory rate of 78.1% at a concentration of 10 \(\mu\)M [117]. 3-Hydroxysperadine A (285) was isolated from HMP-F28 induced extracellular alkalinization and \(H_2\)O\textsubscript{2} production in tobacco cell suspensions by a bioassay-guided fractionation and purification, and no activity was reported [118].

Aspergillamides C and D (286 and 287) were obtained from the marine sponge-derived fungus Aspergillus terreus SCSIO 41008 (Figure 30) [119]. Asterriquinones I–K (288–290), three new bis-indolyquinones, and asterriquinols G–I (291–293), three new bis-indolylbenzenoids, were isolated from the sponge-derived fungus Aspergillus sp. SCSIO 41,018. Asterriquinones I–K (288–290) displayed cytotoxic activities against K562, BEL-7042, SGC-7901, A549 and Hela cell lines [120]. A diketopiperazine–indole alkaloid fintiamin (294) was isolated from fungus Eurotium sp. It showed an affinity for the cannabinoid CB1 receptor at low micromolar concentrations [121].

![Figure 29](image-url)
2.2.7. Miscellaneous

Three new indole diterpenoids, 22-hydroxylshearinine F (295), 6-hydroxylpaspalinine (296) and 7-O-acetylmentol SB (297), were isolated from the sea-anemone-derived fungus Penicillium sp. AS-79. Only 296 exhibited activity against the aquatic pathogen Vibrio parahaemolyticus with a MIC value of 64.0 μg/mL, which was much bigger than the positive control chloromycetin with a MIC value of 0.5 μg/mL (Figure 31) [122]. Four new indole-diterpene alkaloids, asperindoles A–D (298–301), were isolated from the marine-derived fungus Aspergillus sp., associated with an unidentified colonial ascidian. Asperindole A (298) exhibited cytotoxic activity against PC-3, LNCaP and 22Rv1 with IC50 values of 69.4 μM, 47.8 μM and 4.86 μM, and induced apoptosis in 22Rv1 at the concentration of 0.3125 μM. Furthermore, 22Rv1 cells treated with asperindole A (298) for 48 h revealed...
an S-phase arrest [123]. Penicindopene A (302), a new indole diterpene, was isolated from the deep-sea fungus *Penicillium* sp. YPCMAC1. Compound 302 represented the first example of indole diterpene possessing a 3-hydroxyl-2-indolone moiety, and it exhibited moderate cytotoxicities against A549 and HeLa cell lines with IC\(_{50}\) values of 15.2 and 20.5 μM, respectively [124]. Misszrtine A (303) was isolated from marine sponge-derived fungus *Aspergillus* sp. SCSIO XWS03F03. Compound 303 represents the first example of N-isopentenyl tryptophan methyl ester with a phenylpropanoic amide arm, which exhibited a potent antagonistic activity on HL60 (IC\(_{50}\) = 3.1 μM) and LNCaP (IC\(_{50}\) = 4.9 μM) cell lines [125].

![Chemical structures of 294–303.](image)

Chaetoindolones A–D (304–307), 19-O-desmethylchaetogline A (308) and 20-O-desmethylchaetogline F (309) were produced by the marine fish-derived fungus *Chaetomium globosum* 1C51 through biotransformation (Figure 32). Alkaloids 304, 306, 308, and 309 showed antibacterial activities with MIC ranging from 8 to 128 μg/mL against *Xanthomonas oryzae pv. oryzae*, *Ralstonia solanacearum*, *Xanthomonas oryzae pv. oryzicola* and *Pseudomonas syringae pv. lachrymans*. Chaetoindolone A (304) was shown to inhibit the growth of the rice-pathogenic bacteria *Xanthomonas oryzae pv. oryzae* both in vitro and in vivo [126]. Three indole-diketopiperazines, spirotryprostatin G (310, an oxindole derivative), cyclotryprostatin F (311) and cyclotryprostatin G (312), were obtained by large-scale cultivation of the marine-derived fungus *Penicillium brasiliannum* HBU-136 with the aid of genomic analysis. Compound 310 displayed selective cytotoxicities against the HL-60 cell line with an IC\(_{50}\) value of 6.0 μM, whereas compounds 311 and 312 exhibited activities against the MCF-7 cell line with the IC\(_{50}\) values of 7.6 and 10.8 μM, respectively. However, all of the metabolites
appeared to be inactive in antibacterial and antifungal assays (MIC > 25 µM) [127]. Chemical investigation of secondary metabolites from the marine-derived fungus *Aspergillus austroafricanus* Y32-2 resulted in isolating two new prenylated indole alkaloid homodimers, di-6-hydroxydeoxybrevianamide E (313) and dinotoamide J (314). Each compound was evaluated for proangiogenic, anti-inflammatory effects in zebrafish model and cytotoxicity for HepG-2 human liver carcinoma cells. As a result, compound 314 exhibited proangiogenic activity in a PTK787-induced vascular injury zebrafish model in a dose-dependent manner [128].

![Chemical structures of 304-314.](image)

Asperginine (315), an alkaloid possessing a rare skeleton, was isolated from the cultural broth of the marine fungus *Aspergillus* sp., and it has no cytotoxicity against prostate cancer PC3 and human HCT-116 (Figure 33) [129]. Compounds 316–318 were isolated from the culture broth of a marine gut fungus *Aspergillus* sp. DX4H, and only showed weak inhibitory activity at 20 µg/mL against PC3 cell line [130]. Two new dioxopiperazine alkaloids (319 and 320) were isolated from Antarctic marine-derived *Aspergillus* sp. SF-5976. Compound 320 decreased PGE2 production in RAW 264.7 and BV2 cells, and 319 only...
showed inhibitory effects in BV2 cells and the same situation on LPS-stimulated NO production in RAW 264.7 and BV2 cells [131]. Quellenin (321) was isolated from deep-sea fungus Aspergillus sp. YK-76. It showed weak inhibition against the growth of S. parasitica with inhibition zones of 19.9 mm at the dosage of 200 µg/disc [132].

Figure 33. Chemical structures of 315–321.

3. Marine Invertebrates
3.1. Sponges

Sponges are the simplest multicellular animals in the world, and they settle across the bottom of the sea with more than 10,000 species. There are 133 new indole related compounds isolated from invertebrates recently, and 109 are from 33 species of sponges. The marine sponge of the genus Hyrtios has been recognized as a rich source of unique bioactive products. In recent years, six references containing seven indole alkaloids have been reported, including hyrtinadines C (322) and D (323), ishigadine A (324), 5, 6-dibromoindole-3-carboxaldehyde (325), hyrtiodoline A (326), 3,4-dihydrohyrtiosulawesine (327) and bromoindole alkaloid (328) (Figure 34). Various activities were evaluated for these alkaloids. Compound 322 showed antifungal activity against A. niger (IC$_{50}$ = 32 µg/mL), while 323 displayed antibacterial activity against E. coli (MIC = 16 µg/mL) and B. subtilis (MIC = 16 µg/mL). Compound 324 exhibited cytotoxicity against L1210 murine leukemia cells (IC$_{50}$ = 3.3 µg/mL) in vitro. Compound 326 has the most potent antitrypanosomal activity, with an IC$_{50}$ value of 7.48 µM after 72 h. Compound 327 displayed potent inhibitory activities against isocitrate lyase (IC$_{50}$ = 92.9 µM) from Candida albicans. Compound 328 exhibited weak cytotoxicity against HCT-116, MCF-7 and HepG-2 (IC$_{50}$ > 100 µM), and it also inhibits the growth of S. aureus and E. coli at the concentration of 3 mg/mL with the inhibition zones of 14 and 21 mm, respectively [133–138].
Seventeen indole alkaloids (329–345) were isolated from the sponge Fascaplysinopsis reticulata collected in Mayotte and Xisha Islands (Figure 35). Fourteen of them were new oxygenated aplysinopsin-type enantiomers, (+)- and (−)-oxoaplysinopsins A–G (329–342), and the others were 6,6′-bis-(debromo)-gelliusine F (343), 6-bromo-8,1′-dihydro-isoplysin A (344) and 5,6-dibromo-8,1′-dihydro-isoplysin A (345). Compounds 331 and 332 showed tyrosine phosphatase 1B inhibition activity stronger than the positive control of acarbose. Compounds 333 and 334 exhibited cytotoxicity against the Hela cell line. Compounds 344 and 345 displayed antimicrobial activities towards Vibrio natrigens with MIC values of 0.01 and 1 μg/mL, respectively [140,141].

Figure 34. Chemical structures of 322–328.

Figure 35. Chemical structures of 329–345.

Sabrin R.M. Ibrahim group isolated ingenines C–F (346–349) from the Indonesian sponge Acanthostrongylophora ingens (Figure 36). Ingenine C (346) and D (347) were evaluated for their cytotoxic activity towards MCF-7, A549 and HCT-116 cell lines. It is noteworthy that compounds 346 and 347 exhibited cytotoxic activities against MCF-7 and HCT-116 with IC50 values of 4.33 and 6.05 and 2.90 and 3.35 μM, respectively. Ingenine E (348) exhibited cytotoxic activity against MCF-7, HCT-116 and A549 cell lines with IC50 values of 3.5, 0.67 and 2.15 μg/mL. Ingenine F (349) exhibited cytotoxic activity toward MCF-7, HCT-116 and A549 cell lines with IC50 values of 2.82, 1.00 and 2.37 μM, respectively [142–144]. Five new manzamine alkaloids (350–354) were isolated from an Indonesian Acanthostrongylophora sp. sponge. They exhibited weak cytotoxicity against A549 and K562 (LD50 = 4.6–12 μM), and moderate antibacterial activity to six bacteria (MIC > 1.6 ng/mL). Compounds 352–354 showed mild inhibition against the enzyme isocitrate lyase [145].

A novel pyridinium, tricepyridinium (355), and a novel benzoxazine–indole hybrid (356, racemic mixture) were obtained from the culture of an E. coli clone incorporating metagenomic libraries from the marine sponge Discodermia calyx. Compound 356 was speculated to be formed through a nonenzymatic process during the isolation procedure. The synthesized tricepyridinium bromide showed antimicrobial activity against Bacillus cereus, MSSA and Candida albicans with MIC values of 0.78, 1.56 and 12.5 μg/mL, but not against E. coli. In addition, tricepyridinium bromide had cytotoxicity to P388 cells with an IC50 value of 3.5 μM [146].
The synthesized tricepyridinium bromide showed antimicrobial activity against E. coli. Speculated to be formed through a nonenzymatic process during the isolation procedure. Metagenomic libraries from the marine sponge Discodermia calyx were subjected to an unbiased phenotypic assay on hONS cells as a model of Parkinson's disease followed by cluster analysis. It is noteworthy that compounds 346 and 347 exhibited cytotoxic activity towards MCF-7, A549 and HCT-116 cell lines. Two new indole alkaloids (350–354) were isolated from an Indonesian Acanthostrongylophora sp. sponge. They exhibited weak cytotoxicity against A549 and K562 (LD50 = 4.6–12 \( \mu \)g/mL). Ingenines C–F (346–349) from the Indonesian sponge Acanthostrongylophora ingens (Figure 36). Ingenine C (346) and D (347) were evaluated for their cytotoxic activity towards MCF-7, A549 and HCT-116 cell lines. It is noteworthy that compounds 346 and 347 exhibited cytotoxic activities against MCF-7 and HCT-116 with IC\(_{50}\) values of 4.33 and 6.05 and 2.90 and 3.35 \( \mu \)M, respectively. Ingenine E (348) exhibited cytotoxic activity against MCF-7, HCT-116 and A549 cell lines with IC\(_{50}\) values of 3.5, 0.67 and 2.15 \( \mu \)g/mL. Ingenine F (349) exhibited cytotoxic activity toward MCF-7, HCT-116 and A549 cell lines with IC\(_{50}\) values of 2.82, 1.00 and 2.37 \( \mu \)M, respectively [141–143]. Five new manzamine alkaloids (350–354) were isolated from an Indonesian Acanthostrongylophora sp. sponge. They exhibited weak cytotoxicity against A549 and K562 (LD50 = 4.6–12 \( \mu \)M), and moderate antibacterial activity to six bacteria (MIC > 1.6 \( \mu \)g/mL). Compounds 352–354 showed mild inhibition against the enzyme isocitrate lyase [144].

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Figure 35. Chemical structures of 329–345.

Sabrin R.M. Ibrahim group isolated ingenines C–F (346–349) from the Indonesian sponge Acanthostrongylophora ingens (Figure 36). Ingenine C (346) and D (347) were evaluated for their cytotoxic activity towards MCF-7, A549 and HCT-116 cell lines. It is noteworthy that compounds 346 and 347 exhibited cytotoxic activities against MCF-7 and HCT-116 with IC\(_{50}\) values of 4.33 and 6.05 and 2.90 and 3.35 \( \mu \)M, respectively. Ingenine E (348) exhibited cytotoxic activity against MCF-7, HCT-116 and A549 cell lines with IC\(_{50}\) values of 3.5, 0.67 and 2.15 \( \mu \)g/mL. Ingenine F (349) exhibited cytotoxic activity toward MCF-7, HCT-116 and A549 cell lines with IC\(_{50}\) values of 2.82, 1.00 and 2.37 \( \mu \)M, respectively [141–143]. Five new manzamine alkaloids (350–354) were isolated from an Indonesian Acanthostrongylophora sp. sponge. They exhibited weak cytotoxicity against A549 and K562 (LD50 = 4.6–12 \( \mu \)M), and moderate antibacterial activity to six bacteria (MIC > 1.6 \( \mu \)g/mL). Compounds 352–354 showed mild inhibition against the enzyme isocitrate lyase [144].

A novel pyridinium, tricepyridinium (355), and a novel benzoxazine–indole hybrid (356, racemic mixture) were obtained from the culture of an E. coli clone incorporating metagenomic libraries from the marine sponge Discodermia calyx. Compound 356 was speculated to be formed through a nonenzymatic process during the isolation procedure. The synthesized tricepyridinium bromide showed antimicrobial activity against Bacillus cereus, MSSA and Candida albicans with MIC values of 0.78, 1.56 and 12.5 \( \mu \)g/mL, but not against E. coli. In addition, tricepyridinium bromide had cytotoxicity to P388 cells with an IC\(_{50}\) value of 0.53 \( \pm \) 0.07 \( \mu \)g/mL. Compound 356 exhibited no antibacterial activity against the tested Bacillus cereus [145,146].

Figure 36. Cont.
Two new indole alkaloids (357 and 358) were obtained from *Spongia* sp. collected by SCUBA in the South Sea of Korea (Figure 37). They did not display any significant inhibitory activity on farnesoid X-activated receptor (FXR) up to 100 μM, and they were not cytotoxic to CV-1 cells up to 200 μM by MTT assay [147]. 1-(1H-indol-3-yl)propan-2-ol (359) was isolated from the Red Sea sponges *Haliclona* sp. and showed weak cytotoxic activities against the tested HepG-2, Daoy and HeLa by MTT assay [148]. Two bisindole alkaloids tethered by a guanidino ethylthiopyrazine moiety, dragmacidins G (360) and H (361), were isolated from *Lipastrotethya* sp. marine sponge. Dragmacidin G (360), and dragmacidin H (361), showed cytotoxicity against HeLa cells with IC50 values of 4.2 and 4.6 μM, respectively [149]. Chemical investigation of a specimen of *Jaspis splendens* collected from the Great Barrier Reef resulted in the isolation of a new bisindole alkaloid, splendamide (362), and 6-bromo-1H-indole-3-carboximidamide (363) are reported for the first time as naturally occurring metabolites. They were subjected to an unbiased phenotypic assay on hONS cells as a model of Parkinson’s disease followed by cluster analysis of cytological effects and showed similar biological activity in cluster B. under a Pearson’s correlation of 0.91 [150]. A new acrylic jasplakinolide congener (364) and another structure-revised acyclic derivative (365) were isolated from the Indonesian marine sponge *Jaspis splendens*, and the jasplakinolides inhibited the growth of mouse lymphoma (L5178Y) cells in vitro with IC50 values in the low micromolar to the nanomolar range [151]. A new cyclic peptide, jamaicensamide A (366), composed of six amino acids, including thiazole-homologated amino acid, was isolated from the Bahamian sponge *Plakina jamaicensis* collected from Plana Cay, and no bioactivity have been evaluated due to the insufficient quantities [152]. A novel brominated marine indole (367) was isolated from the boreal sponge *Geodia barretti* collected off the Norwegian coast. Compound 367 was inactive (IC50 > 690 μM) on electric eel AChE even with a structural resemblance with other known natural AChE inhibitors and showed somewhat higher inhibitory potential towards BChE (IC50 = 222 μM) [153]. Geobarrettin A–C (368–370) were isolated from the sub-Arctic sponge *Geodia barrette* by UPLC-qTOF-MS-based dereplication study. Both 369 and 370 reduced DC secretion of IL-12p40, but 370 concomitantly increased IL-10 production. Maturing DCs treated with 369 or 370 before co-culturing with allogeneic CD4+ T cells decreased T cell secretion of IFN-γ, indicating a reduction in Th1 differentiation [154].

Antibacterial-guided fractionation of an extract from a deep-water *Topsentia* sp. marine sponge led to the isolation of two new indole alkaloids, tulongcin A (371) and dihydrospongotone C (372) (Figure 38). Antibacterial, anti-HIV activity and cytotoxicity were evaluated for compounds 371 and 372. They showed strong antibacterial effects toward *S. aureus* with 1.2 and 3.7 μg/mL MICs. However, only weak to no inhibition toward *E. coli* at the maximum concentration tested (100 μg/mL) was reported. Both compounds inhibited HIV infection in HIV infectivity assays against the CCR5-tropic primary isolate YU2 and the CXCR4-tropic strain HxB2 with the IC50 values ranging from 2.7 to 4.5 μM. They were
inactive (IC_{50} > 10 \mu M) in cytotoxicity assays against a monkey kidney cell line (BSC-1) and a human colorectal tumor cell line (HCT-116) [155]. A new brominated indole 6-Br-8-keto-conicamin A (373) was identified from Haplosclerida sponge, and it showed moderate cytotoxic activity against the PANC-1 tumor cell line with the IC_{50} value of 1.5 \mu M [156]. Two new brominated bisindole alkaloids, dragmacidins I (374) and J (375), were isolated from the Tanzanian sponge Dragmacidon sp. They showed low micromolar cytostatic activity against A549, HT-29 and MDA-MB-231. The mechanism of the action was investigated through different molecular biology experiments, which indicated that these two dragmacidins act via the inhibition of Ser-Thr PPs [157]. Six new cyclopenta[g]indole natural products, trans-herbindole A (376) and trikentramides E–I (377–381), were isolated from the sponge Trikentrion flabelliforme, and there is no bioactivity reported [158].

![Figure 37. Chemical structures of 357–370.](image-url)
The first chemical investigation of the subtidal sponge *Spongosorites calcicola* led to the discovery of two new bisindole alkaloids of the topsentin family (382 and 383), and they showed very weak or no cytotoxic activity against the Hela cell line (Figure 39) [159]. Six new bisindoles, (Z)-coscinamide D (384), (E)-coscinamide D (385) and lamellomorphamides A−D (386–389) were isolated from a rare New Zealand deep-sea sponge, *Lamellomorpha strongylata*. Compounds 385, 386, 389 showed weak activity against MRSA at the concentration of 20 μM (14.9−18.2% inhibition) [160].

**Figure 38. Chemical structures of 371–381.**

**Figure 39. Chemical structures of 382–389.**

Guitarrins A–E (390–394), the first natural 5-azaindoles, and aluminumguitarrin A (395), the first aluminum-containing compound from marine invertebrates, were isolated
from the sponge *Guitarra fimbriata* (Figure 40). Guitarrin C (392) inhibited alkaline phosphatase from the marine bacterium *Cobetia marina* with an IC$_{50}$ value of 2.0 µM, being a natural inhibitor of alkaline phosphatase [161]. Two brominated oxindole alkaloids (396 and 397) were isolated from sponge *Callyspongia siphonella* with LC-HRESIMS-assisted dereplication and bioactivity-guided isolation. The sponge was collected from Hurghada along the Red Sea Coast. Oxindoles 396 and 397 exhibited diverse pharmacological activities, including antibacterial activity, biofilm inhibitory activity, antitrypanosomal activity and antitumor activity. They inhibited the growth of *Staphylococcus aureus* (MIC = 8 and 4 µg/mL), *Bacillus subtilis* (MIC = 16 and 4 µg/mL), *Pseudomonas aeruginosa* (49.32% and 41.76% inhibition at the concentration of 128 µg/mL), and *T. brucei* (13.47 and 10.27 µM for 72 h). In addition, they showed good cytotoxic effect toward HT-29, OVCAR-3 and MM.1S with IC$_{50}$ values ranging from 7 to 12 µM through non-programmed cell necrosis [162]. A naturally new alkaloid (398) was isolated from *Gelliodes* sp. collected in Vietnam, and showed no cytotoxicity against Hela, MCF-7 and A549 cell lines [163]. Myrindole A (399), a bis-indole alkaloid, was isolated from the deep-sea sponge *Myrmekioderma* sp. Myrindole A inhibits the growth of *E. coli* and *Bacillus subtilis* with MIC values of 37.5 and 18.5 µM, respectively [164]. The structures of a series of incorrectly reported sponge-derived dibrominated indole alkaloids, echinosulfone A (400) and the echinosulfonic acids A–D (401–404) were corrected [165–167]. Another two papers have also disclosed identical structure revisions for these dibrominated indole alkaloids (400–404) [168,169].

A bis-indole (405) and an alkynyl indole alkaloid (406) were isolated from the sponge *Plakortis* sp. collected from Zampa in Okinawa (Figure 41). The bis-indole was inactive...
against both P388 and B16 cells even at 100 µg/mL, while 406 showed cytotoxicity against P388 at 1 µg/mL (IC₅₀ = 0.6 µg/mL) and B16 cells at 100 µg/mL [170]. Zamamidine D (407) was isolated from an Okinawan Amphimedon sp. marine sponge, and it exhibited obvious antibacterial activity against the eight tested strains (Escherichia coli, Staphylococcus aureus, Micrococcus luteus, Aspergillus niger, Trichophyton mentagrophytes, Candida albicans and Cryptococcus neoformans) with IC₅₀ values ranging from 2 to 32 µg/mL [171]. An extract of the marine sponge Damiria sp., which represents an understudied genus, provided two novel alkaloids named damirines A (408) and B (409). Compound 408 showed selective cytotoxic properties toward six different cell lines in the NCI-60 cancer screen [172]. Makaluvamine W (410) was isolated from the Tongan sponge Strongylodesma tongaensis. Compound 410 was inactive to the tested HL-60 cell line and confirmed the requirement of an intact iminoquinone functionality required by these metabolites to be bioactive [173]. In the study of developing a metric-based prioritization approach by exact LC-HRMS, 411 and 412 were isolated in a case study from a sponge collected from a reef on the island of Tavarua, Fiji Islands. No activity was evaluated for 411 and 412 [174].

Five dibromoindole alkaloids (413–417) were isolated from sponge Narrabeena nigra collected around the Futuna Islands (Figure 42). They reduced the TBHP-induced cell death, which demonstrated their potential in neuroprotection, and showed almost no cytotoxic effect up to 10 µM on human neuroblastoma SH-SY5Y and microglia BV2 cells [175].

New indolo-imidazole alkaloids trachycladindoles H–M (418–423) were isolated from a deep-water Great Australian Bight sponge, Geodia sp. CMB-01063 (Figure 43). The trachycladindoles H–M did not exhibit growth inhibitory activity against the E. coli, Bacillus subtilis, Candida albicans human colorectal (SW620) or lung (NCI-H460) carcinoma cells [176].

A highly modified hexapeptide friomaramide (424) was isolated from the Antarctic sponge Inflatella coelosphaeroides, and it blocks more than 90% of Plasmodium falciparum liver-stage parasite development at 6.1 µM (Figure 44) [177]. Halicylindramides F–H (425–427) were isolated from a Petrosia sp. marine sponge collected off the shore of Youngdeok-Gun,
East Sea, Republic of Korea. Halicylindramides F (425) showed human farnesoid X receptor (hFXR) antagonistic activity, but it did not bind directly to hFXR [178].

Microsclerodermins B and J (428 and 429) were reported by Faulkner and co-workers and Li and co-workers from marine sponge Microscleroderma (Figure 45) [179,180]. The configuration of the C44 stereocenter of microscleroderm B was confirmed by total synthesis of dehydromicoscleroderm B, which was revised from 44S to 44R. The same method was applied with microsclerodermins J, and the configuration was revised to be 44R [181]. The structure of topsentin C (430) was revised by an efficient total synthesis [182].
A series of new indole-oxazole-pyrrole natural products breitfussins C–H (428–430) were isolated from marine hydrozoan Thuiaria sp. marine sponge collected off the shore of Young-Gun, East Sea, Republic of Korea. Halicylindramides F (425) and 2-amino-1,5-dihydro-5-(1H-indol-3-ylmethyl)-4H-imidazol-4-one (426) were isolated from the sea anemone Heteractis aurora, and no activity was reported (Figure 46) [183]. Two new natural products, bis-6-bromogramine (427) and 6-bromogramine (428), were isolated from the marine hydroid Abietinaria abietina. They activate NF-κB-dependent transcriptional activity in JB6 Cl 41 NF-κB cells at 1.6 μM [184]. A series of new indole-oxazole-pyrrole natural products breitfussins C–H (438, 435–437, 439, 440), along with breitfussin A and B, were isolated from marine hydrozoan Thuiaria breitfussi. The hydrozoan was collected from Bjørnøya, Svalbard (79.0293 N, 20.8574 E, at 48 m depth). Cytotoxic activity and kinase inhibition profiles of breitfussins C–F were evaluated against seven malignant cell lines, one non-malignant cell line and 13 kinases. Then, the pharmacological potential of breitfussin C was extended to the activities evaluation of

**Figure 44.** Chemical structures of 424–427.

**Figure 45.** Chemical structures of 428–430.

### 3.2. Cnidarians

Two new indole alkaloids 2-amino-1,5-dihydro-5-(1H-indol-3-ylmethyl)-4H-imidazol-4-one (431) and 2-amino-5-[(6-bromo-1H-indol-3-yl)methyl]-3,5-dihydro-3-methyl-4H-imidazol-4-one (432) were isolated from the sea anemone Heteractis aurora, and no activity was reported (Figure 46) [183]. Two new natural products, bis-6-bromogramine (433) and 6-bromogramine (434), were isolated from the marine hydroid Abietinaria abietina. They activate NF-κB-dependent transcriptional activity in JB6 Cl 41 NF-κB cells at 1.6 μM [184].
88 cancer cell lines and 468 kinases. The results showed excellent inhibition and selectivity against MDA-MB-468 with the IC₅₀ value of 340 nM and against the PIM1 and DRAK1 kinases with IC₅₀ and Kᵋ values of 200 and 390 nM, respectively. Further studies on potential off-target effects, toxicological effects, as well as relevant in vitro ADME, displayed the potential of breitfussins for selective kinase inhibitor development [185].

Figure 46. Chemical structures of 431–440.

### 3.3. Bryozoans, Tunicates and Molluscs

Three new halogenated, hexacyclic indoleline-imidazole alkaloids, securamines H–J (441–443), were isolated from Arctic bryozoan Securiflustra securifrons (Figure 47). Compounds 441 and 442 showed cytotoxicity against the human cancer cell lines A2058 (skin), HT-29 (colon), and MCF-7 (breast), as well as against nonmalignant human MRC-5 lung fibroblasts with IC₅₀ values ranging from 1.4 ± 0.1 to 5.3 ± 1.1 µM. The cytotoxicity of 441 was further evaluated and found to be time-dependent [186]. A new compound, 2,6-dibromo-N-methylgramine (444), was discovered in bryozoan Amathia verticillata, and no activity was reported [187]. 2,5-dibromo-1-methyl-1H-indole-3-carbaldehyde (445) was isolated from the bryozoan A. lamourouxi, which was collected from rock pools of Woolgoolga, Australia. Compound 447 was inactive for the tested antiplasmodial activity and HEK293 cytotoxicity at 40 µM [188]. Terminoflustrindoles (TFIns) B and C (444 and 445) were isolated from the bryozoan Terminoflustra membranaceatruncata, and they exhibit no antimicrobial activity against various microorganisms tested [189]. A new indole alkaloid 448 was isolated from a colonial marine tunicate, Didemnum sp., collected by SCUBA near Haeguengang, Korea. Compound 448 showed no pharmacological potential as an antibacterial agent and FXR antagonist [190]. Stolonines A and C (449 and 450) were isolated from a marine tunicate Cnemidocarpa stolonifera. An immunofluorescence assay on PC3 cells indicated that compounds 449 and 450 increased cell size, induced mitochondrial texture elongation, and caused apoptosis in PC3 cells [191]. Compounds 451 and 452 were reported in a case study of the metrics-based prioritization approach development. They were isolated from the pink mottled tunicate collected from Tavarua, Fiji Islands [174]. Orbicularisine (453), a novel spiro-indolofuranone fused to a thiazine skeleton, was isolated from gills of the mollusk Codakia orbicularis. Compound 453 was inactive against Enterococcus faecalis, Streptococcus pneumonia, Klebsiella pneumonia, E. coli, and Pseudomonas aeruginosa. Inhibition assays against a panel of kinases including Hs_CDK2/CyclinA, Hs_CDK5/p25, Hs_CDK9/CyclinT, Hs_RIPK3, Hs_Haspin, Hs_AuroraB, Ld_TLK, Hs_Pim1, Ssc_GSK3a/b, LmCK1, and Rn_Dyrk1A showed residual activities of more than 60% for 16 µM/mL.
Finally, the treatment of HCT-116 colon cancer cells and U87-MG glioblastoma cancer cells with concentrations up to 100 µM showed no activity [192]. Cesplamide E (454) was isolated from the Taiwanese soft coral Cespitularia taeniata, and it exhibited cytotoxicity against MCF-7, Daoy and Hela cancer cells with IC₅₀ of 17.5, 22.3, and 24.7 µM, respectively [193].

![Chemical structures](image)

Figure 47. Chemical structures of 441–454.

4. Marine Plants

4.1. Cyanobacteria

Two proline-rich cyclic peptides (455 and 456) were isolated from marine cyanobacterium Sympleca sp., collected from Minna Island, Japan and Bintan Island, Indonesia (Figure 48). Compound 456 possessed cytotoxicity against the MOLT4 and AML2 cancer cell lines with IC₅₀ values of 4.8 and 8.2 µM, respectively [194,195].
4.2. Red Algae

Eleven new tetrahalogenated indoles (457–467) were isolated from the red alga *Rhodophyllis membranacea*, collected from Moa Point, New Zealand (Figure 49). Compounds 457 and 459–461 showed no antifungal activity against wild-type *Saccharomyces cerevisiae* (baker’s yeast) and cytotoxicity against HL-60 promyelocytic leukemia cell line with IC$_{50}$ values higher than 10 µM [196]. Compounds 468–470 were isolated and identified from the red algae *Laurencia similis*, and 468 showed potent antibacterial activity against seven bacterial strains with MIC values ranging from 2 to 8 µg/mL [197].
4.3. Mangrove

Chemical investigation of the leaves and stems of the Chinese mangrove Acanthus ilicifolius Linn. led to the isolation and structure elucidation of one new pyrido[1,2-a]indole alkaloid named acanthiline A (471), and no bioactivity was reported (Figure 50) [198]. Marine alkaloid 472 was reported as 2-methylimidazo[1,5-b]isoquinoline-1,3,5(2H)-trione, and it was revised to be 473 by a total synthesis [199,200].

![Figure 50. Chemical structure of 471–473.](image)

5. Conclusions

In this article, we reviewed 472 indole alkaloids discovered from marine organisms with a vast of bioactivities during the year from 2015 to 2021. The alkaloids were grouped according to the sources, divided into marine microbes, invertebrates, and plants. Marine microbes are the main source of natural marine products, which is the case for indole alkaloids. A total of 321 new indole metabolites were isolated from marine microorganisms, including 64 from marine-sourced bacteria and 257 from marine-sourced fungi. Sponges have been abundant and stable sources of marine natural products over the years. Among the indole alkaloids discovered from marine invertebrates, sponge-derived make up the vast majority, with 109 out of 133 in total. Marine plants only contributed 18 indole compounds isolated from cyanobacteria, red algae and mangrove.

Due to the insufficient amount of compounds isolated, the bioactivity determination of natural products has always been a significant challenge. Although a number of top chemists are devoted to moving natural products synthesis from the laboratory to the factory, the total synthesis of some complex marine natural products remains a challenge, and not to mention industrialization. However, marine microbial fermentation has the characteristics of non-destruction of ecological resources, relatively low cost, and good reproducibility. Mass fermentation assisted by genetic engineering transformation can better realize industrial production. In addition, there are many kinds of marine microorganisms, which are inexhaustible sources for marine drugs development. At present, most marine indole alkaloids were evaluated for their antitumor and antimicrobial activities. This is not only because this class of compounds is more likely to exhibit such activity, but also because antitumor and antimicrobial activity measurements are generally easier to perform. It can be summarized from this review that marine indole alkaloids have rich skeleton structures and various pharmacological activities. How to transfer the chemical diversity to pharmacological activity diversity is another challenge. Therefore, it is expected that a more general and practical pharmacological activity assay should be developed and applied to the early drug development process.

Compounds with indole moiety typically have significant pharmacological activity. We hope that this review will promote the development of marine indole alkaloids in medicinal chemistry and pharmacology in terms of the extent of drug discovery.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/md19120658/s1, Table S1: General information of the cell lines; Table S2: Summary of the marine indole alkaloids isolated from marine microorganisms; Table S3: Summary of the marine indole alkaloids isolated from marine invertebrates and plants.
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