REVIEW

# **Bioactive compounds from marine actinomycetes**

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Abstract Actinomycetes are one of the most efficient groups of secondary metabolite producers and are very important from an industrial point of view. Among its various genera, Streptomyces, Saccharopolyspora, Amycolatopsis, Micromonospora and Actinoplanes are the major producers of commercially important biomolecules. Several species have been isolated and screened from the soil in the past decades. Consequently the chance of isolating a novel actinomycete strain from a terrestrial habitat, which would produce new biologically active metabolites, has reduced. The most relevant reason for discovering novel secondary metabolites is to circumvent the problem of resistant pathogens, which are no longer susceptible to the currently used drugs. Existence of actinomycetes has been reported in the hitherto untapped marine ecosystem. Marine actinomycetes are efficient producers of new secondary metabolites that show a range of biological activities including antibacterial, antifungal, anticancer, insecticidal and enzyme inhibition. Bioactive compounds from marine actinomycetes possess distinct chemical structures that may form the basis for synthesis of new drugs that could be used to combat resistant pathogens.

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#### Introduction

Microbial natural products are an important source of both existing and new drugs. Among the producers of commercially important metabolites, bacteria have proven to be a prolific source with a surprisingly small group of taxa accounting for the vast majority of compounds discovered till date [1]. Among these, Actinomycetes are the most economically and biotechnologically priceless prokaryotes. Representative genera of actinomycetes include Streptomyces, Actinomyces, Arthrobacter, Corynebacterium, Frankia, Micrococcus, Micromonospora and several others. Secondary metabolites produced by actinomycetes possess a wide range of biological activities [1–4]. The genus Streptomyces alone produces a large number of bioactive molecules [5-128]. It has an enormous biosynthetic potential that remains unchallenged without a potential competitor among other microbial groups. A large number of Streptomyces spp. have been isolated and screened from soil in the past several decades [129, 130]. Consequently the chances of isolating a novel Streptomyces strain from terrestrial habitats have diminished. Above 500 species of Streptomyces account for 70-80% of relevant secondary metabolites as shown in Table 1 [5-127], with small contributions from other genera, such as Saccharopolyspora, Amycolatopsis, Micromonospora and Actinoplanes. An important reason for discovering novel secondary metabolites is to circumvent the problem of resistant pathogens, which are no longer susceptible to the currently used drugs [131, 132]. The number of deaths due to these clever pathogenic organisms is on the rise. Secondary metabolites from marine

S. No.	Compound	Source	Activity
1.	Erythromycin [5]	Saccharopolyspora erythrae	Antibacterial
2.	Rhamnose [6]	Saccharopolyspora spinosa	Essential component of insect control agent compound spinosad
3.	Zorbamycin [7]	Streptomyces flavovirdis	Antitumor
4.	Kanamycin [8]	Streptomyces kanamyceticus 12-6	Antibacterial
5.	Kanglemycin C (K-C) [9]	Nocardia mediterranei var. kanglensis 1747-64	Immunosuppressive
6.	Rapamycin [10]	Streptomyces hygroscopicus	Antifungal
7.	Pandavir (nigericin) [11]	Streptomyces hygroscopicus	Affects ion transport and ATPase activity
8.	FK520 Ascomycin [12]	Streptomyces hygroscopicus var. ascomyceticus	Antifungal, immunosuppressive, neutrophic
9.	Himastatin [13]	Streptomyces hygroscopicus	Antitumor
10.	Jinggangmycin [14]	Streptomyces hygroscopicus	Antifungal
11.	Oxytetracycline [15]	Streptomyces rimosus	Antibacterial
12.	Amphotericin B [16]	Streptomyces nodosus	Antifungal
13.	Asukamycin [17]	Streptomyces nodosus subsp. asukaensis	Antibacterial
14.	Tylosin [18]	Streptomyces fradiae	Antibacterial
15.	Urdamycin A [19]	Streptomyces fradiae	Antitumor
16.	Fosfomysin [20]	Streptomyces fradiae	Antibacterial
17.	CE-108[21]	Streptomyces diastaticus	Antifungal
18.	Rimocidin [22]	Streptomyces diastaticus var. 108	Antifungal
19.	Shurimycins A and B [23]	Streptomyces hygroscopicus	Antibacterial, antifungal
20.	Chloramphenicol [24]	Streptomyces venezuelae	Antibacterial
21.	Rifamycin [25]	Amycolatopsis mediterranei U-32	Antibacterial
22.	Amythiamicins [26]	Amycolatopsis sp.	Antibacterial
23.	Cyclo (L-leucyl-l-prolyl) [27]	Streptomyces sp. KH614	Antileukemic, anti-VRE (vancomycin-resistant enterococci)
24.	Ipomicin [28]	Streptomyces ipomoeae group III	Antibacterial
25.	Streptomycin [29]	Streptomyces griseus	Antibacterial
26.	Valinomycin [30]	Streptomyces griseus	Mitochondrial toxin
27.	Griseorhodin [31]	Streptomyces griseus FCRC-57	Telomerase inhibitor
28.	Fredericamycin A [32]	Streptomyces griseus FCRC-48	Antitumor
29.	Capuramycin [33]	Streptomyces griseus SANK 60196	Antibacterial
30.	Frigocyclinone [34]	Streptomyces griseus strain NTK 97	Antibacterial
31.	Clorobiocin [35]	Streptomyces coelicolor	Inhibitor of bacterial gyrase
32.	Meilingmycin [36]	Streptomyces nanchangensis	Antiparasitic
33.	Nanchangmycin [36]	Streptomyces nanchangensis	Insecticidal
34.	Eremomycin [37, 38]	Amycolatopsis orientalis subsp. eremomycini	Antimicrobial
35.	Nikkomycins [39]	Streptomyces ansochromogenus	Antifungal
36.	Avilamycin A [40]	Streptomyces viridochromogenes Tu57	Antibacterial
37.	Tubelactomicin A [41]	Nocardia sp.	Antibacterial

Table 1	(Commueu)		
S. No.	Compound	Source	Activity
38.	Benzanthrins A and B [42]	Nocardia lurida	Antibacterial
39.	Azureomycins A and B [43]	Pseudonocardia azurea nov. sp.	Antibacterial
40.	Nogalamycin [44]	Streptomyces nogalater	Antibacterial
41.	Aclacinomycin A (aclarubicin) [45]	Streptomyces galilaeus	Antitumor
42.	Cinerubin R [46]	Streptomyces eurythermus	Antibacterial
43.	Scopafungin [47]	<i>Streptomyces hygroscopicus</i> var. <i>enhygrus</i> var. nova UC-2397	Antifungal, antibacterial
44.	Spiramycin [48]	Streptomyces ambofaciens	Antibacterial
45.	Pristinamycin I [49]	Streptomyces pristinaespiralis	Antibacterial
46.	Lankacidin [50]	Streptomyces rochei	Antibacterial
47.	Lankamycin [50]	Streptomyces rochei	Antibacterial
48.	Actinomycin C [51]	Streptomyces chrysomallus	Antitumor
49.	Duanomycin [52]	Streptomyces sp.	Antitumor
50.	Midecamycin [53]	Streptomyces mycarofaciens	Antibacterial
51.	Avermectin [54]	Streptomyces avermitilis	Anthelminthic
52.	Oligomycin [55]	Streptomyces avermitilis	Cell growth inhibitor
53.	Resormycin [56]	Streptomyces platensis	Herbicidal, antifungal
54.	Ileumycin [57]	Streptomyces lavendulae	Antifungal
55.	Mitomycin C [58]	Streptomyces lavendulae	Antitumor
56.	Lomofungin [59]	Streptomyces lomodensis	Antifungal, antibacterial
57.	Kalafungin [60]	<i>Streptomyces tanashiensis</i> strain Kala UC5063	Antifungal, antibacterial, antiprotozoal
58.	Thiamycins [61]	Streptomyces michiganensis var. amylolyticus var. nova	Anthelminthic, antiprotozoal
59.	Axenomycins [62]	Streptomyces lisandri nov. sp.	Anthelminthic, antiprotozoal, antifungal
60.	Neihumicin [63]	Micromonospora neihuensis	Cytotoxic
61.	Fortimicin A (Astromicin) [64]	Micromonospora olivasterospora	Antibacterial
62.	Gentamicin [65]	Micromonospora purpurea var. violaceae	Antibacterial
63.	Tetracycline [66]	Streptomyces aureofaciens	Antibacterial
64.	Monomycin [67, 68]	Actinomyces circulatus var. monomycini	Antibacterial
65.	PC-766 B [69]	Nocardia brasiliensis	Antioxidant
66.	Medecamycin [70, 71]	Streptomyces mycarofaciens	Antibacterial
67.	Dunaimycins [72]	Streptomyces diastatochromogenes	Immunosuppressive, antimicrobial
68.	Novobiocin [73]	Streptomyces niveus	Antibacterial
69.	Carminomycin [74]	Actinomadura carminata	Antitumor
70.	Maduramycins [75]	Actinomadura rubra	Antibacterial
71.	MM461156 [76]	Actinomadura pelletieri	Antiviral, antibacterial
72.	Verucopeptin [77]	Actinomadura verrucosospora	Antitumor
73.	Saptomycins [78]	Streptomyces sp. HP 530	Antitumor, antimicrobial
74.	Oxaprapalines B, D, G [79]	Streptomyces sp. G324	Antitumor
75.	Lavendamycin [80]	Streptomyces lavendulae	Antitumor
76.	Chlorocarcins A, B, C [81]	Streptomyces lavendulae No. 314	Antitumor, antibacterial

 Table 1 (Continued)

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S. No.	Compound	Source	Activity
77.	Mimosamycins [81]	Streptomyces lavendulae No. 314	Antibacterial
78.	Lavendomycin [82]	Streptomyces lavendulae	Antibacterial
79.	Sohbumycin [83]	Streptomyces sp. 82-85	Antitumor, antibacterial
80.	Furaquinocins C, D, E, F, G, H [84]	Streptomyces sp. KO 3988	Antitumor
81.	Arizonins A1 and B1 [85]	Actinoplanes arizonaensis sp. nov.	Antibacterial
82.	Coloradocin [86]	Actinoplanes coloradoensis sp. nov.	Antibacterial
83.	Teichomycins [87]	Actinoplanes teichomyceticus nov. sp.	Antibacterial
84.	Lipiarmycin [88]	Actinoplanes deccanensis nov. sp.	Antibacterial
85.	Candiplanecin [89]	<i>Ampullariella reguralis</i> subsp. <i>mannitophila</i> subsp. nov.	Antifungal
86.	Victomycin [90]	Streptosporangium violaceochromogenes nov. sp.	Antitumor, antibacterial
87.	Maggiemycin and anhydromaggiemycin [91]	Streptomyces sp.	Antitumor
88.	Gilvusmycin [92]	Streptomyces sp.	Antitumor
89.	Kazusamycin [93]	Streptomyces sp.	Antitumor
90.	Okicenone [94]	Streptomyces sp.	Antitumor
91.	Hydramycin [95]	Streptomyces violaceus	Antitumor
92.	Musacin C [96]	Streptomyces griseovirdis	Anthelminthic, antiviral
93.	Kanchanamycins [97]	Streptomyces olivaceus	Antifungal, antibacterial
94.	Elloramycin [98]	Streptomyces olivaceus	Antitumor
95.	Fattiviracin A1 [99]	Streptomyces microflavus	Antiviral
96.	FK 506 [100]	Streptomyces tsukubaensis	Antiviral
97.	Retamycin [101]	Streptomyces olindensis	Antitumor
98.	Manumycin [102]	Streptomyces parvulus	Antitumor, enzyme inhibitory
99.	Granaticin [103, 104]	Streptomyces thermoviolaceus	Antibacterial
100.	Pimaricin [105]	Streptomyces natalensis	Antifungal
101.	Virginiamycin M [106, 107]	Streptomyces virginae	Antibacterial
102.	Daptomycin (commercialized as Cubicin) [108]	Streptomyces roseosporus	Antibacterial
103.	Enduracidin [109]	Streptomyces fungicidicus B5477	Antibacterial
104.	Apramycin [110]	Streptomyces tenebrabrius UD2	Antibacterial
105.	Mithramycin [111]	Streptomyces argillaceus	Antitumor
106.	Blasticidin S [112]	Streptomyces griseochromogenes	Antifungal
107.	Leptomycin [113]	Streptomyces lividans	Antifungal, antitumor
108.	Landomycin E [114]	Streptomyces globisporus	Antitumor
109.	Phenalinolactones A-D [115]	Streptomyces sp.	Antibacterial
110.	Pipalamycin [116]	Sreptomyces sp.	Apoptosis inducer, antibacterial
111.	Biphenomycin A and B [117]	Streptomyces griseorubiginosus	Antibacterial
112.	Streptocidins A-D [118]	Streptomyces sp. Tu6071	Antibacterial
113.	Zelkovamycin [119]	Streptomyces sp. K96-0670	Antibacterial
114.	Methylsulfomycin I [120]	Streptomyces sp. RSP9	Antibacterial

S. No.	Compound	Source	Activity
115.	YM-216391 [121]	Streptomyces nobilis	Anticancer
116.	RP-1776 [122]	Streptomyces sp.	Inhibit binding of platelet derived growth factor to its receptor
117.	RS-22 A, B and C [123]	Streptomyces violaceusniger	Antifungal, antibacterial
118.	Vicenistatin [124]	Streptomyces sp. Tu6239	Antitumor
119.	Ripromycin [125]	Streptomyces sp.	Antibacterial, antitumor
120.	Vinylamycin [126]	Streptomyces sp.	Antibacterial
121.	Cephamycin C [127]	Streptomyces lactamdurans	Antibacterial

 Table 1 (Continued)

actinomycetes may form the basis for the synthesis of novel therapeutic drugs, which may be efficient to combat a range of resistant microbes [133, 134].

Existence of cousins of terrestrial actinomycetes has been reported in the relatively untapped marine ecosystem. The immense diversity of this habitat along with it's underexploitation is the fundamental reason for attracting researchers towards it for discovering novel metabolite producers. Actinomycetes comprise about 10% of the bacteria colonizing marine aggregates and can be isolated from marine sediments [135]. Many actinomycete isolates from deep oceans contain non-ribosomal polyketide synthetase (NRPS) and polyketide synthetase (PKS) pathways, the hallmarks of secondary metabolite production [136]. There is an occurrence of distinct rare genera in the marine ecosystem as evidenced by the taxonomic description of the first marine actinomycete Rhodococcus marinonascens [137]. Actinomycetes have also been isolated from free swimming as well as sessile marine vertebrates and invertebrates [135]. Unusual actinomycetes belonging to *Micrococceae*, Dermatophilaceae and Gordoniaceae, have been isolated from sponges [133]. Tetrodotoxin-producing actinomycete has been isolated from puffer fish ovaries [138], the organism was found to be most closely related to Nocardiopsis dassonvillei.

Researchers are finding new genera from marine environments on a regular basis and discovering new metabolite producers never reported earlier. Actinomycete genera identified by cultural and molecular techniques from different marine ecological niches include Actinomadura, Actinosynnema, Amycolatopsis, Arthrobacter, Blastococcus, Brachybacterium, Corynebacterium, Dietzia, Frankia, Frigoribacterium, Geodermatophilus, Gordonia, Kitasatospora, Micromonospora, Micrococcus, Microbacterium, Mycobacterium, Nocardioides, Nocardiopsis, Nonomurea, Psuedonocardia. Rhodococcus. Saccharopolyspora, Salinispora, Serinicoccus, Solwaraspora, Streptomyces, Streptosporangium, Tsukamurella, Turicella, Verrucosispora and Williamsia [135]. Inspite of improvements being made in the cultural methods for the isolation of rare marine actinomycetes, many of these organisms still remain unculturable and have to be detected by using molecular techniques [139, 140]. Metagenomic methods are useful for characterizing microbes that cannot be cultivated and can also be used to isolate their genes [141].

#### Secondary metabolites from marine actinomycetes

Marine actinomycetes have proven to be efficient producers of new secondary metabolites as shown in Table 2 [142–182], which show a range of biological activities such as antifungal, antitumor, antibacterial, immunosuppressive, insecticidal and enzyme inhibition, to name a few.

Secondary metabolites produced by marine actinomycetes can be classified on the basis of their chemical structure as follows:

## 1. Terpenes and terpenoids

The most chemically diverse pool of secondary metabolites in nature is constituted by terpenes [183]. In 1956, novobiocin was isolated as the first antibiotic with a terpenoid side chain from *Streptomyces niveus* [184]. After this, the list of these compounds isolated from soil actinomycetes has increased as listed in Table 3 [185–193].

Terpenes are not only produced by the soil actinomycetes but also from the marine habitants as evidenced by the following compounds:

- I. Azamerone [142] is a meroterpenoid produced by a new marine bacterium related to the genus *Streptomyces*. It appears to be the first natural product with a phthalazione ring (Fig. 1).
- II. Three new pyrrolosesquiterpenes, glaciapyrroles A, B and C [143] are produced by a *Streptomyces* strain (NPSOO 8187). These compounds show antibacterial

S. No.	Chemical group	Compound	Source	Activity
1.	Meroterpenoid	Azamerone [142]	Streptomyces sp.	None
2.	Pyrrolosesquiterpenes	Glaciapyrroles A, B and C [143]	<i>Streptomyces</i> sp. NPS008187	Antibacterial
3.	Amorphane sesquiterpenes [144]	$10\alpha$ , 15-dihydroxyamorph-4-en-3-one, $10\alpha$ , 11-dihydroxyamorph-4-ene and $5\alpha$ , $10\alpha$ , 11-trihydroxyamorphan-3-one [144]	Streptomyces sp. M491	None
4.	Sesquiterpene	Neomarinone [145]	Strain CNH-099	Cytotoxic
5.	Polyketide	Saliniketal A, saliniketal B [146, 147]	Salinispora arenicola	Anticancer
6.	Polyketide	Abyssomicin C [148]	Verrucosispora	Antibacterial
7.	Polyketide	SBR-22 [149]	Steptomyces psommoticus BT408	Antibacterial
8.	Polyketide	Daryamides [150]	Streptomyces sp. CNQ-085	Anticancer, antifungal
9.	Polyketide	Actinofuranones A and B [151]	Streptomyces sp.	Cytotoxic
10.	Peptide	Mechercharmycins [152]	Thermoactinomyces sp.	Antitumor
11.	Peptide	Thiocoraline [153]	Micromonospora	Anticancer, antibacterial
12.	Peptide	CyclomarinA [154]	Streptomyces sp.	Anti-inflammatory, antiviral
13.	Peptide	Piperazimycins [155]	Streptomyces sp.	Anticancer
14.	Peptide	Dehydroxynocardamine and desmethylenylnocardamine [156]	Streptomyces sp.	Enzyme sortase B inhibitor
15.	Peptide	Urukthapelstatin [157]	Mechercharimyces asporophorigenes YM11-542	Anticancer
16.	Peptide	Salinamides A and B [158]	Streptomyces sp.	Antibacterial, anti-inflammatory
17.	Caprolactone	R-10-methyl-6-undecanolide (6R,10S)- 10-methyl-6-dodeconolide [159]	Streptomyces sp. B6007	Phytotoxic, anticancer
18.	Butenolide	Butenolide [160]	Streptoverticillium luteoverticillatum	Anticancer
19.	Polycyclic xanthone	IB-00208 [161]	Actinomadura	Anticancer, antibacterial
20.	Piericidin	Piericidins C7 and C8 [162]	Streptomyces	Anticancer
21.	Quinone	Resistomycin [163]	Streptomyces corchorusii AUBN(1)/7	Antiviral
22.	Quinone	Tetracenomycin D [164]	Streptomyces corchorusii AUBN(1)/7	Anticancer, antibacterial
23.	Quinone	Resistoflavine [165, 166]	Streptomyces chibaensis AUBN(1)/7	Anticancer, antibacterial
24.	Quinone	Komodoquinone A [167]	Streptomyces sp. K53	Neuritogenic activity
25.	Quinone	Himalomycins A and B [168]	Streptomyces sp. B6921	Antibacterial
26.	Quinone	Helquinoline [169]	Janibacter limosus	Antibacterial
27.	Quinone	Chlorinated dihydroquinones [170]	CNQ-525	Anticancer, antibacterial
28.	Macrolide	Chalcomycin A [144]	Streptomyces sp. M491	None
29.	Macrolide	Arenicolide A [147, 171]	Salinispora arenicola	Antibacterial
30.	Macrolide	Marinomycins [172]	Marinispora	Anticancer, antibacterial

 Table 2
 Bioactive compounds produced by marine actinomycetes

S. No.	Chemical group	Compound	Source	Activity
31.	Alkaloid	K252c and arcyriaflavin A [173]	Z (2)0392	Anticancer
32.	Ester	Bonactin [174]	Streptomyces sp. BD21-2	Antibacterial, antifungal
33.	Manumycin derivatives	Chinikomycins A and B [175]	Streptomyces sp. M045	Anticancer
34.	Complex compounds	Trioxacarcins [176]	Streptomyces ochraceus and Streptomyces bottropensis	Anticancer, antimalarial
35.	Methylpyridine	Streptokordin [177]	Streptomyces sp. KORDI-3238	Anticancer
36.	Gamma lactam beta lactone	Salinosporamide A [147, 178]	Salinispora tropica	Anticancer
37.	Macrocyclic lactam	Aureoverticillactam [179]	Streptomyces aureoverticillaris	Anticancer
38.	Enzyme inhibitor	Alpha-amylase inhibitor [180]	<i>Streptomyces</i> <i>corchorusii</i> subsp. <i>rhodomarinus</i> subsp. nov	Enzyme Inhibition
39.	Enzyme inhibitor	Pyrostatins A and B [181]	Streptomyces sp. SA-3501	N-acetyl-beta- glucosaminidase inhibition
40.	Enzyme inhibitor	Pyrizinostatin [182]	Streptomyces sp. SA-2289	Pyroglutamyl peptidase inhibition

Table 2(Continued)

 Table 3
 Terpenes produced by soil actinomycetes

S. No.	Compound	Source	Activity
1.	Pentalenolactone I [185, 186]	Streptomyces filipinensis	Antibacterial, immunosuppressive
2.	Lavanduquinocin [185, 187]	Streptomyces viridochromogenes	Neuronal cell protection
3.	Napyradiomycins [185, 188]	Chiana rubra	Antibacterial
4.	Spirocardins A and B [185, 189]	Nocardia sp. SANK 64282	Antibacterial
5.	Benthocyanin A [185, 190]	Streptomyces prunicolor	Radical scavenger
6.	Benzastatin C [185, 191]	Streptomyces nitrosporeus	Antiviral
7.	Carquinostatin B [185, 192]	Streptomyces exfoliatus	Neuronal cell protection
8.	Moenomycin [185, 193]	Streptomyces bambergensis	Antibacterial

activities. Structures of glaciapyrroles A, B and C are shown in Fig. 2.

- III. Amorphane sesquiterpenes [144] (Fig. 3) namely  $10\alpha$ ,15-dihydroxyamorph-4-en-3-one,  $10\alpha$ ,11-dihydr -oxyamorph-4-ene and  $5\alpha$ ,10 $\alpha$ ,11-trihydroxyamor phan-3-one are produced by *Streptomyces* sp. M491. This is the first report of these sesquiterpenes from bacteria.
- IV. Neomarinone [145], a novel metabolite possessing a new sesquiterpene and polyketide-derived carbon skeleton and several derivatives of the marinone class of naphthoquinone antibiotics are produced by a taxonomically novel marine actinomycete (strain CNH-099). These bioactive molecules show moderate cytotoxicity towards human cancer cells.

# 2. Polyketides

- I. Saliniketal A (Fig. 4) and saliniketal B [146, 147], produced by *Salinispora arenicola*, are inhibitors of ornithine decarboxylase biosynthesis. Inhibition of ornithine decarboxylase production is an important strategy in the control of cancer since high levels of this enzyme lead to uncontrolled proliferation of cells. The Saliniketals are partly related in structure to the rifamycins.
- II. Abyssomicin C [148] (Fig. 5) is a polycyclic polyketide produced by *Verrucosispora*. It targets p-aminobenzoate (PABA) biosynthesis and therefore inhibits folic acid biosynthesis at an early stage as compared to the well-known synthetic sulpha drugs.

The abyssomicins are the first known bacterial secondary metabolites that can inhibit the biosynthesis of PABA. Targeting PABA production is an attractive strategy for arresting microbial growth since PABA directly leads to the production of folic acid, which is a precursor of purine biosynthesis. Humans lack this pathway; therefore the strategy will not be harmful to humans. Abyssomicin C shows antibacterial activity against gram-positive bacteria as well as clinical isolates of multiple resistant and vancomycin-resis-



Fig. 1 Azamerone

tant *Staphylococcus aureus*. Abyssomicin C and its analogues thus have a high potential to be developed as antibacterial agents against drug-resistant pathogens.

- III. A marine inhabitant known as *Streptomyces psom-moticus* produces antibiotic SBR-22 [149]. It shows antibacterial activity against methicillin-resistant *Staphylococcus aureus*.
- IV. Daryamides [150] (Fig. 6) are cytotoxic polyketides isolated from culture broth of a *Streptomyces* strain, CNQ-085. These bioactive compounds show weak to moderate cytotoxicity against the human colon carcinoma cell line HCT-116 and very weak antifungal activities against *Candida albicans*.
- V. Actinofuranones A and B [151] (Fig. 7) are isolated from the fermentation broth of a marine bacterium related to *Streptomyces* genus. Actinofuranones A and B show weak *in vitro* cytotoxicity against mouse splenocyte T-cells and macrophages.
- 3. Peptides
- I. Mechercharmycins [152] are new bioactive compounds obtained from marine-derived *Thermoactinomyces* sp. YM3-251. The cyclic structure of mechercharmycin A



Fig. 2 Glaciapyrroles A, B, C





5a, 10a,-11-trihydroxyamorphan-3-one

10a, 11-dihydroxyamorph-4-ene



10a-15-dihydroxyamorph-4-en-3-one

Fig. 3 Amorphane sesquiterpenes



Fig. 4 Saliniketal A







Fig. 6 Daryamides A, B and C



Fig. 7 Actinofuranones A and B

(Fig. 8) is essential for its strong antitumor activity, since the related compound mechercharmycin B (Fig. 9) does not show such an activity.

- II. Thiocoraline [153] is a new depsipeptide isolated from *Micromonospora*. It shows potent cytotoxicity against P-388, A-549 and MEL cell lines, and also a strong antimicrobial activity against gram-positive microorganisms. This compound binds to supercoiled DNA and inhibits RNA synthesis.
- III. Cyclomarins A-C [154] (Fig. 10) are cyclic peptides produced by a *Streptomyces* sp. They show anti-inflammatory and antiviral activities.
- IV. Piperazimycins [155] (Fig. 11) are cytotoxic hexadepsipeptides isolated from the fermentation broth of a

*Streptomyces* sp. Piperazimycin A exhibits potent *in vitro* cytotoxicity against multiple tumor cell lines.

- V. Two cyclic peptides dehydroxynocardamine [156] and desmethylenylnocardamine [156] along with nocardamine have been isolated from a *Streptomyces* sp. which has been obtained from an unidentified marine sponge. These new compounds exhibit weak inhibition against the enzyme sortase B.
- VI. Urukthapelstatin A [157] is a novel cyclic peptide produced by the thermoactinomycete bacterium *Mechercharimyces asporophorigenes* YM11-542. It inhibits the growth of human lung cancer A54 cells and shows cytotoxicity against a range of human cancer cell lines.



Fig. 8 Mechercharmycin A



Fig. 9 Mechercharmycin B





Fig. 10 Cyclomarin A

Fig. 11 Piperazimycins

VII. Salinamides [158] A and B are bicyclic depsipeptides produced by a *Streptomyces* sp., CNB-091, isolated from jelly fish *Cassiopeia xamachana*. These metabolites are useful as antibiotic and anti-inflammatory agents.

## 4. Caprolactones

Two new caprolactones R-10-methyl-6-undecanolide and (6R,10S)-10-methyl-6-dodeconolide [159] are produced by a marine *Streptomyces* sp. isolate B6007. These caprolactones show a moderate phytotoxicity and low cytotoxocity against cancer cells.

# 5. Butenolides

*Streptoverticillium luteoverticillatum* produces four butenolides [160]. These butenolides show cytotoxicity against the murine lymphoma P388 and human leukemia K562 cell lines. This is the first report of isolation of butenolides from the marine ecosystem, which possess cytotoxic activity.

## 6. Polycyclic xanthones

IB-00208 [161] is a polycyclic xanthone isolated from the culture of *Actinomadura*. This compound possesses cytotoxicity against tumor cell lines and bactericidal activity against gram-positive bacteria.

#### 7. Piericidins

Piericidins C7 and C8 [162] show selective cytotoxicity against rat glia cells transformed with the adenovirus EIA gene and neuro-2a mouse neuroblastoma cells. These compounds are produced by a marine *Streptomyces* sp.

#### 8. Quinones

- Resistomycin [163] (Fig. 12), an antibiotic related to quinones, is produced by *Streptomyces corchorusii* AUBN(1)/7. This is an inhibitor of HIV-1 protease.
- II. Tetracenomycin D [164] (Fig. 13) is an anthraquinone antibiotic also produced by *Streptomyces corchorusii* AUBN(1)/7. It shows cytotoxicity against cell line HMO2 (gastric adenocarcinoma) and HepG2 (hepatic carcinoma) and possesses weak antibacterial activities against gram-positive and gram-negative bacteria.
- III. Resistoflavine [165, 166] (Fig. 14) is produced by Streptomyces chibaensis AUBN(1)/7. It shows cytotoxicity against cell line HMO2 (gastric adenocarcinoma) and HepG2 (hepatic carcinoma) and possess-



Fig. 12 Resistomycin



Fig. 13 Tetracenomycin D



R = H



es weak antibacterial activities against gram-positive and gram-negative bacteria.

- IV. Komodoquinone A [167] (Fig. 15) is a neuritogenic anthracycline isolated from the fermentation broth of a marine *Streptomyces* sp. K53. It induces cell differentiation in the neuroblastoma cell line, Neuro2A and arrests cell cycle at the G1 phase.
- V. Himalomycins A and B [168] (Fig. 16) are two new quinone antibiotics from a *Streptomyces* isolate, B6921. Himalomycins exhibit strong antibacterial activity against *Bacillus subtilis, Streptomyces*



Fig. 15 Komodoquinone A



Fig. 16 Himalomycins A and B

viridochromogenes, Staphylococcus aureus and Escherichia coli.

- VI. Helquinolines [169] (Fig. 17) are new tetrahydroquinoline antibiotic isolated from culture broth of *Janibacter limosus*. Helquinoline shows moderate activity against *Bacillus subtilis*, *Streptomyces virdochromogenes* Tu57 and *Staphylococcus aureus*.
- VII. CNQ-525 is a member of a new genus (tentatively called MAR4) within the family Streptomycetaceae, which produces three novel chlorinated dihydroquinones [170]. These compounds possess new carbon skeletons but are related to several previously reported metabolites of the napyradiomycin class. The metabolites possess significant antibiotic properties and cytotoxicity against cancer cells.
- 9. Macrolides
- I. *Streptomyces* sp. M491 is a marine actinobacterium that produces a macrolide antibiotic named Chalcomycin A [144] (Fig. 18) and also some terpenes.
- II. Some strains of *Salinispora arenicola* produce a series of macrolides exemplified by Arenicolide [147, 171]



Fig. 17 Helquinoline

(Fig. 19). These possess weak antibacterial activities against drug-resistant bacteria.

- III. Marinomycins [172] (Fig. 20) are polyene-like macrolides. A marine *Marinispora* produces these compounds, which are potent antitumor antibiotics with moderate activities against selected human tumors and drug-resistant bacterial pathogens. Marinomycin A inhibits the growth of human pathogenic bacteria such as methicillin-resistant *Staphylococcus aureus* and vancomycin-resistant *Enterococcus faecium*. These polyenes are highly photoreactive and undergo isomerization even at room light because of which their use in clinics as potential drugs has been discontinued. In spite of being polyenes, marinomycins however, do not shows antifungal activities typically associated with other polyene antibiotics.
- 10. Alkaloids

Two indolocarbazole alkaloids, K252c [173] (Fig. 21) and Arcyriaflavin A [173] (Fig. 22) are produced by a marine actinomycete Z(2)0392. Both of these alkaloids possess moderate cytotoxicity against the K562 cell line and induce apoptosis. This is the first report of the significant apoptosis inducing effect of indolocarbazole alkaloids against K562 cancer cells.

11. Esters

Bonactin [174] (Fig. 23) is an antimicrobial ester. Bonactin displays antimicrobial activity against gram-positive and gram-negative bacteria as well as against several fungi. Bonactin is produced by *Streptomyces* sp. BD21-2.

12. Chinikomycins

Chinikomycins A (Fig. 24) and B [175] are chlorinecontaining aromatic manumycin derivatives. They exhibit antitumor activity against different human cancer cell lines,



Fig. 18 Chalcomycin A



Fig. 19 Arenicolide A



Fig. 20 Marinomycin A

but are inactive as antiviral, antimicrobial and phytotoxic agents. These compounds are produced by *Streptomyces* sp. isolate MO45.

#### 13. Trioxacarcins

Trioxacarcins [176] (Fig. 25) are complex compounds showing high antibacterial activity against gram-positive and gram-negative bacteria, and some of them show high antitumor and antimalarial activities as well. Trioxacarcin A also exhibits antifungal activities. Trioxacarcin A, B and C are obtained from *Streptomyces ochraceus* and *Strep*-



Fig. 21 K252c



Figs. 22 Arcyriaflavin A

*tomyces bottropensis*. Some of these compounds possess extremely high antiplasmodial activity, which is comparable to that shown by artemisinin, the most active compound against the pathogen of malaria. The producers of trioxacarcins also biosynthesize the related metabolite, gutingimycin.

# 14. Methylpyridine

Streptokordin [177] a new cytotoxic compound of the methylpyridine class is isolated from the cultural broth of *Streptomyces* sp. KORDI-3238. It exhibits significant cytotoxicity against several human cancer cell lines but shows no growth inhibition against various microorganisms, including bacteria and fungi.



Fig. 23 Bonactin



Fig. 24 Chinikomycin A

 $\beta$ -lactam compounds which contain a four-membered  $\beta$ -lactam ring. The structure of  $\beta$ -lactam second ring allows these compounds to be classified into penicillins, cephalosporins, clavams, carbapenes and monobactans [194]. Most  $\beta$ -lactam compounds inhibit bacterial cell wall synthesis but others behave as  $\beta$ -lactamase inhibitors (e.g. cluvalanic acid) and even as antifungal agents (e.g. some clavams) [194], however salinosporamide A and aureoverticillactam show cytotoxicity against cancer cells.

16. Enzyme inhibitors

Some of the enzymes inhibitors reported from marine actinomycetes include:

I. Alpha amylase inhibitor from *Streptomyces corchorusii* subsp. *rhodomarinus*. subsp. nov [180].



#### Fig. 25 Trioxacarcins

- 15. Lactams
- I. Salinosporamide A [147, 178] (Fig. 26) is produced by *Salinispora tropica* which is found in oceanic sediments. Salinosporamide A is a potent proteasome inhibitor used as an anticancer agent that has entered phase I of the human clinical trials for the treatment of multiple myeloma. It inhibits proteasome activity by covalently modifying the active site threonine residues of the 20S proteasome.
- II. Aureoverticillactam, a novel 22-atom macrocyclic lactam [179] is isolated from *Streptomyces aureoverticillaris*. It shows cytotoxicity against various tumor cell lines.

Salinosporamide A and aureoverticillactam are lactams from marine actinomycetes. These are distinct from



Fig. 26 Salinosporamide A

- II. Pyrostatins A and B are inhibitors of n-acetyl-beta-glucosaminidase, produced by *Streptomyces* sp. SA-3501 [181].
- III. Pyrizinostatin is an inhibitor of pyroglutamyl peptidase, isolated from culture of *Streptomyces* sp. SA-2289 [182].

## Conclusion

Secondary metabolites produced from marine actinomycetes have distinct chemical structures, which may form the basis for the synthesis of new drugs. Salinispora alone produces a wide range of metabolites having different biological activities [146, 147, 171, 178]. Enrichment and selective isolation methods can also be used to isolate rare actinomycetes from marine ecological niches having the potential to biosynthesize novel bioactive compounds [140, 195-197]. A great hurdle however, in the search of these actinomycetes is that more than 90% of the organisms remain uncultivable under laboratory conditions. To explore the genomic diversity of the marine ecosystem and estimate their biosynthetic capability, the techniques of metagenomics can be used. Turbomycin is one of the first antibiotics to be discovered by metagenomics [198]. Isolation of long-chain acyltyrosine antibiotics from metagenomic libraries has also been reported [199]. Genes encoding enzymes responsible for the synthesis of secondary metabolites, are usually clustered on a contiguous piece of DNA. For expression of a single antibiotic there is a need for a large size DNA, which is a major challenge when DNA is isolated from soil, having high concentrations of humus and heavy metals as contaminants [200, 201]. But large insert metagenomic libraries can be prepared from marine samples with ease. By designing a suitable vector, which can accommodate large size inserts, it is possible to isolate novel bioactive compounds from marine unculturable actinomycetes [200, 201].

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