Chemistry and Biology of Siderophores from Marine Microbes

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Abstract: Microbial siderophores are multidentate Fe(III) chelators used by microbes during siderophore-mediated assimilation. They possess high affinity and selectivity for Fe(III). Among them, marine siderophore-mediated microbial iron uptake allows marine microbes to proliferate and survive in the iron-deficient marine environments. Due to their unique iron(III)-chelating properties, delivery system, structural diversity, and therapeutic potential, marine microbial siderophores have great potential for further development of various drug conjugates for antibiotic-resistant bacteria therapy or as a target for inhibiting siderophore virulence factors to develop novel broad-spectrum antibiotics. This review covers siderophores derived from marine microbes.

Keywords: siderophores; marine microbes; synthesis; biosynthesis

1. Introduction

Marine microbial community diversity and chemodiversity lead to the generation of numerous biological secondary metabolites. These compounds increase the chances of finding valuable drug candidates [1–3]. Marine siderophores are a type of low molecular weight natural products, the functions of which are to facilitate the microbial acquisition of iron in the ocean environment [4–6]. Marine microbes typically require micromolar levels of iron for their growth, yet the concentration of iron in the surface water of the world ocean is only 0.01–2 nM [7]. In order to overcome this lack of iron, marine microbes have evolved siderophore-mediated delivery system to obtain iron. Siderophores coordinated to insoluble Fe(III) are first transported into microbial cells by membrane bound iron-siderophore receptors. Afterwards, iron is then released from siderophores, typical via reduction of Fe(III) to Fe(II) by microbe-mediated redox processes [8–10]. Based on the efficient siderophore-mediated iron acquisition mechanisms, marine siderophores may be developed as novel antimicrobials by covalently attaching clinical antibiotics to marine siderophores or novel iron chelator desferals [11–14]. Furthermore, marine siderophores have also exhibited various non-classical biological functions, for example, as agents which may interfere with quorum sensing regulation and swarming in bacteria, as mediators of mutualistic interactions, and as secreted signaling molecules regulating virulence factors of pathogens [15–17]. These wide range of activities exhibited their great potential in medicine. The present review gives a comprehensive overview of siderophores from marine microorganisms (Table 1).
Table 1. List of abbreviations.

| Abbreviation | Full Name                                      |
|--------------|------------------------------------------------|
| CAS          | Chrome azurol sulfonate                        |
| ED<sub>50</sub> | Median effective dose                           |
| IC<sub>50</sub> | Half maximal inhibition concentration            |
| NRPSs        | Nonribosomal peptide synthetases               |
| HSC          | N-hydroxy-N-succinylcadaverine                  |
| HSDs         | N-hydroxy-N-succinyl diamines                  |
| ArCP         | Aryl carrier protein                            |
| Cy           | Cyclisation                                     |
| PCP          | Peptidyl carrier protein                        |
| NIS          | NRPS-independent siderophore                    |
| hLys         | N<sup>6</sup>-hydroxy-l-lysine                  |
| ahLys        | N<sup>6</sup>-acetyl-N<sup>6</sup>-hydroxy-l-lysine |
| 3-DHS        | 3-dehydroshikimate                              |
| 3,4-DHBA     | 3,4-dihydroxybenzoic acid                       |
| 2,3-DHBA     | 2,3-dihydroxybenzoic acid                       |

2. Diversity of Siderophores from Marine Microorganisms

Recently, a large number of siderophores with diverse structures have been discovered in the marine microorganisms (Table 2). According to their functional groups and hydrophobicity, siderophores from marine microorganisms are divided into seven types, including hydroxamates (Figure 1), α-hydroxycarboxylates (Figure 2), catecholates (Figure 3), mixed hydroxamates/α-hydroxycarboxylates (Figure 4), mixed α-hydroxycarboxylates/catecholates (Figure 5), mixed hydroxamates/catecholates (Figure 6), and other types of siderophores (Figure 7) [18–21].

2.1. Hydroxamate-Type Siderophores

The distinct characteristic of some acyl peptidic hydroxamate-type siderophores is the presence of a nonpolar fatty acid tail. Many marine bacteria have been reported to produce large suites of acyl peptidic hydroxamate siderophores. Marine bacterium Marinobacter sp. DS40M6 produces the suit of marinobactin siderophores A–F (1–6) and HG (7) [22,23]. The conversion of exogenously added <sup>15</sup>N-labeled 1 to the bacterial culture suggests that bacteria produce 7 via the hydrolysis of the acyl amide bond of 1 rather than as a precursor to the compounds 1–6 or independent production. It is worth noting that compound 7 can still coordinate Fe(III) via three iron-coordinating moieties, as these moieties are not affected by deacylation. Marine bacterium Vibrio sp. R-10 is also reported to produce a suite of amphiphilic siderophores, amphibactins (8–17) [24]. Each amphibactin has the similar structure, including a peptidic headgroup composed of one serine residue and three ornithine residues but differs in the saturated, unsaturated, or hydroxylated acyl appendage ranging from C-14 to C-18. They are cell-associated siderophores. This association may play an important role in cell defense against siderophore diffusion in the marine environments. Moreover, several acyl peptidic hydroxamate siderophores, moanachelins (18–22) [25] from the marine bacterium, Vibrio sp. Ntl, and amphibactins U–V (23–24) [26] from two marine bacterial species (Synechococcus sp. PCC 7002 and Vibrio cyclitrophicus 1F53) have also been identified.

Additionally, several macrocyclic hydroxamate siderophores were discovered in the marine bacteria, such as alcaligin (25) [27], bisucaberin (26) [28], avaroferrin (27) [29], and putrebactin (28) [30]. Alcaligin (25) could promote Fe(III) uptake in the iron limited cells. It was originally discovered from Alcaligenes denitrificans isolated from sediments of lake Biwa, Japan [31]. Recently, it was also obtained from Alcaligenes eutrophus CH34 [27]. Bisucaberin (26) was isolated from marine bacterium Alteromonas haloplanktis SB-1123 [28]. It could inhibit the growth of both invasive micropapillary carcinoma and L-1210 leukemia cells with IC<sub>50</sub> of 12.7 and 9.7 µM, respectively, and sensitize tumor cells to macrophage-mediated cytolysis [32]. Studies of ecology showed that the increasing contents of bisucaberin (26) and acidification enhanced the solubilities and dissolution rates of ferric hydroxides. This suggested that siderophores generated by marine bacteria could grasp ferric...
hydroxides in aeolian particles to meet the growth of phytoplankton in the ocean environment. Thus, the production rate of marine siderophores may play a vital role in the survival of phytoplankton and biogeochemical cycle of iron in the sea environment [33]. As an open form of 26, a linear dimeric hydroxamate class siderophore, bisucaberin B (29) was isolated from the marine bacterium Tenacibaculum mesophilum collected in the Republic of Palau [34]. Its ferric ion-chelating activity was consistent with that of its macrocyclic counterpart 26. Moreover, the siderophore avaroferin (27) was recently identified in the marine bacterium Shewanella algae B516 by heterologous expression of deep-sea sediment metagenomics DNA [29]. It is a macrocyclic heterodimer of N-hydroxy-N-succinyl-putrescine and N-hydroxy-N-succinyl cadaverine that is shown to inhibit the swarming of Vibrio alginolyticus B522. Another new cyclic trihydroxamate compound, thalassosamide (30), was isolated from the marine-derived bacterium Thalassospira profundimaris [35] and showed moderate antibiotic activity against Escherichia coli and Pseudomonas aeruginosa with the same MIC value of 64 µg/mL in vitro. In vivo antibacterial assays showed thalassosamide (30) could reduce the infection of P. aeruginosa without toxicity compared with untreated control animals in the murine thigh P. aeruginosa infection model.

New bioactive siderophores, fradiamine A (31) and fradiamine B (32), were isolated from the Streptomyces fradiae MM456M-mf7 derived from the marine sediments of Calyptogena Community, Sagami Bay, Japan. They contain two alkyl amines asymmetrically bonded to citrate, rarely observed in siderophores. These molecules exhibited medium antibiotic activity against Clostridium difficile BAA-1382 of IC50 values of 32 and 8 µg/mL, respectively. However, their antimicrobial activities were cancelled dose-dependently under the presence of Fe(III) [36]. A novel albisporachelin siderophore (33) was obtained from iron-depleted culture broths of marine actinomycete Amycolatopsis albispora WP1. The study of iron-chelating ability suggested that its hydroxamate moieties were involved in iron binding, which was consistent with its structure [37]. Moreover, the variable siderophore metabolome of the marine actinomycete Salinispora tropica CNB-440 revealed the production of six known siderophores, desferrioxamine A1, A2, B, D1, D2, and E (34–39), and one novel siderophore desferrioxamine N (40) [38]. Among these, 36 has been developed as an iron chelator desferal for treating iron overload in man.

![Figure 1. Cont.](image_url)
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Figure 1. Hydroxamate-type siderophores from marine microbes.
2.2. α-Hydroxycarboxylates

Vibrio ferriferrin (41) was originally isolated from an enteropathogenic estuarine bacterium, *Vibrio parahaemolyticus*. Recently, it was also obtained from *Marinobacter* species collected from Pacific and Atlantic Oceans. 41 was considered as one of the weakest iron chelators of known marine siderophores according to the metal–ligand binding constant for 41–Fe(III) (10^{24.02(5)}). However, 41–Fe(III) complex was shown to be more sensitive to photolysis than other marine photoactive siderophores, such as aerobactin, petrobactin, marinobactins, or aquachelins. Taken together, 41 photodegradation may be an evolutionarily adaptive response by which the marine bacteria share photochemically produced soluble iron with their algal associates possibly through the exchange with algal-excreted metabolites [39].

![Figure 2. α-Hydroxycarboxylate-type siderophores from marine microbes.](image)

2.3. Catecholates

Structural elucidation revealed the discovery of a novel siderophore, nigribactin (42) derived from marine bacteria *Vibrio nigripulchritudo* [40]. It could enhance the expression of *spa* encoding a major surface bound virulence factor, Protein A by inducing *spa* transcription. However, another potent siderophile, enterobactin failed to affect the expression of *Staphylococcus aureus* virulence genes. This suggested the influence of 42 on *spa* expression might be independent from its siderophore activity. Vanchrobactin (43) from the marine bacterium *Vibrio anguillarum* is a dipeptide composed by arginine and serine residues linked to the 2,3-dihydroxybenzoyl moiety [41–43]. Its catecholate and salicylate groups are two potential bidentate coordination sites of Fe^{3+} [44]. The marine bacterium *Vibrio* sp. DS40M4 was reported to produce 43, a new triscatechol amide siderophore, trivanchrobactin (44), and a new related siderophore, divanchrobactin (45) [45]. Among them, 43 and 45 may be derived from the hydrolysis products of 44. All compounds were found to be non-cytotoxic against P388 murine leukemia cell lines. In addition, a novel triscatecholate siderophore, turnerbactin (46) was isolated from the shipworm endosymbiont *Teredinibacter turnerae* T7901 [46]. It is a trimer of *N*-(2,3-dihydroxybenzoic acid)-l-Orn-l-Ser with three monomeric units through the Ser ester linkages. Its structure was similar to the catecholate siderophore 43. They have a serine backbone, 2,3-dihydroxybenzoic acid functional moiety, a positively charged, and a hydrophilic spacer amino acid.

Using the chrome azurol sulfonate (CAS) assay-guided isolation, three new siderophores, dibenarthin (47), streptobactin (48), and tribenarthin (49) were isolated from the marine actinomycete *Streptomyces* sp. YM5-799 collected from Hokkaido in north Japan. Among these, the ED_{50} values of 47 and 48 were 117 µM and 156 µM, slightly stronger than that of the control, deferoxamine mesylate (ED_{50} = 195 µM), whereas 49 exhibited a weak binding with Fe(III) (ED_{50} = 937 µM). These siderophores might play a key role in the survival of marine-derived *Streptomyces* sp. YM5-799 under iron-limited conditions [47]. The marine *Penicillium bilaii* from Port Huon, Tasmania, Australia was also reported to produce a rare catechol-type siderophore, pistillarin (50) [48]. It showed potent iron chelation as well as antioxidant activity (free radical-scavenging activity), and significantly protected DNA against the damage of hydroxyl radicals generated by the Fenton reaction [49].
Figure 3. Cont.
Figure 3. Cont.
with two different fatty acid appendages. The ochrobactin–Fe(III) complexes are visible sunlight or UV light. Fe(II) is produced by the photoreaction of ochrobactin–Fe(III) \[56\].

Marinobacter DS40M3, isolated from the sample of seawater under iron limited and pure culture conditions led to the sediments below a depth of 1714 m, where sunlight would not penetrate. Thus, the photoreactivity of the Fe(III)-loihichelins is their photoreactivity. However, Halomonas LOB-5 is obtained from sediments below a depth of 1714 m, where sunlight would not penetrate. Thus, the photoreactivity of the Fe(III)-loihichelins may not be mainly responsible for Fe(III) uptake. Halomonas aquamarina DS40M3, isolated from the sample of seawater under iron limited and pure culture conditions led to the biosynthesis of aquachelin siderophores A–D (57–60), I (61), and J (62) \[22,23,52\]. Among them, 61 and 62 were more hydrophilic than 57–60. When H. aquamarina DS40M3 and Marinobacter sp. DS40M6 were co-cultured, a novel aquachelin siderophore HG (63) from hydrolysis of acyl aquachelins was produced. However, acyl amidase activity of Marinobacter sp. DS40M6 is predominantly constrained to the cellular membrane. Thus, the acyl aquachelins within the Marinobacter membrane are mainly responsible for aquachelin HG (63) production \[29\]. An amphiphilic siderophore imaqobactin (64) is detected in the arctic marine bacterium Variovorax sp. RKJM285 \[53\]. Its structure has been implicated in the photoreduction of Fe(III) to Fe(II). Reduction may occur through a charge transfer from ligand-to-metal leading to the cleavage of 64. Metal-binding capabilities showed that 64 was capable of forming stable adducts with Fe(III), Ga(III), and Al(III). Moreover, 64 also appeared to be reactive with gold, but more studies were required to determine whether this was a reductive process. Of particular note, Fe(III)–64 and Ga(III)–64 salts showed no antimicrobial activity. However, 64 showed moderate antimicrobial activity against Staphylococcus warneri, methicillin-resistant S. aureus, vancomycin-resistant Enterococcus and Proteus vulgaris with the IC\(_{50}\) values of 28, 35, 11, and 14 \(\mu\)M, respectively. It may be attributed to the ability of 64 to bind Fe(III) or Ga(III), depressing microbial proliferation via cellular iron depletion.

The planktonic marine Vibrio sp. DS40M5 was reported to produce a mixed hydroxamates/\(\alpha\)-Hydroxy carboxylates siderophore, aerobactin (65) \[54\]. Subsequently, other marine environmental strains of Vibrio also were reported to produce 65. It was a shuttle molecule recycled in each cycle of transport \[55\]. Recently, three 65-based amphiphilic siderophores, ochrobactins A–C (66–68) were discovered from the marine proteobacterium Ochrobactrum sp. SP18. They contain a citric backbone, two lysine residues and different fatty acids. This is the first example of aerobactin-derivative siderophores with two different fatty acid appendages. The ochrobactin–Fe(III) complexes are photoreactive under visible sunlight or UV light. Fe(II) is produced by the photoreaction of ochrobactin–Fe(III) \[56\]. Moreover, three siderophores, synechobactins (69–71) from the marine cyanobacteria Synechococcus sp. PCC 7002 were identified by a high throughput method which is combined liquid chromatography
inductively coupled plasma mass spectrometry with high resolution electrospray ionization mass spectrometry [57].

Figure 4. Cont.
Two photoreactive siderophores, petrobactin (72) and petrobactin sulfonate (73), are produced by marine bacterium *Marinobacter hydrocarbonoclasticus* [58–60]. They are composed of a citrate bis-spermidine skeleton, with two 3,4-dihydroxy-dihydroxybenzoyl groups providing four donor groups for Fe(III) binding. Fe(III) complexes with 72 and 73 are photoreactive under visible sunlight, regulated by the Fe(III)–citrate moiety. This reaction leads to decarboxylation and reduction of Fe(III) to Fe(II). 73 is also the first marine siderophore including a sulfonated 3,4-dihydroxy aromatic ring. Alterobactin A (74) and B (75) was originally obtained from *Alteromonas luteoviolacea* isolated by oligotrophic and coastal waters. Alterobactin A (74) showed an exceptionally high affinity for Fe(III) [61,62]. It contains two unusual amino acids: (3R,4S)-4,8-diamino-3-hydroxyoctanoic acid and l-threo-β-hydroxyaspartic acid, attaching to a catechol carboxylate. 74–Fe(III) complex is stable in solution with little change of absorbance. However, in the absence of coordinated Fe(III), the serine ester of 74 will be hydrolyzed to alterobactin B (75) siderophore. New siderophores, pseudoalterobactin A (76) and B (77) were obtained from marine bacterium *Pseudoalteromonas sp.* KP20-4 isolated from the Republic of Palau [63]. They contain a catechol and two β-hydroxy-Asp residues. Pseudoalterobactins showed strong activity by a CAS assay, which was comparable to that of desferrioxamine B and enterobactin. Both 76 and 77 exhibited ED_{50} value of 20 μM (the reduction of the absorbance at 630 nm of the CAS solution by 50%), while desferrioxamine B and enterobactin showed ED_{50} values of 500 μM and 60 μM under the same conditions, respectively.
Three unusual siderophores, lystabactins A–C (78–80), were obtained from the marine bacterium *Pseudoalteromonas* sp. S2B [64]. In order to further determine their stability by chelating Fe(III), the

**Figure 5.** Mixed α-hydroxycarboxylates/catecholates siderophores from marine microbes.
2.6. Mixed Hydroxamates/Catecholates

Three unusual siderophores, lystabactins A–C (78–80), were obtained from the marine bacterium *Pseudoalteromonas* sp. S2B [64]. In order to further determine their stability by chelating Fe(III), the pFe scale was used to compare with the iron-binding stability of siderophores. pFe was the negative logarithm of aqueous, free Fe(III) at fixed concentrations of siderophore, Fe(III), and acidification (pH). The tighter a given siderophore bound Fe(III), the higher the pFe. The pFe of compounds 78 and 79 were calculated to be 26.0 and 27.5, respectively. This suggested that their stability of iron-binding was stronger than the tris-hydroxamate siderophore, desferrioxamine B, while weaker than the tris-catecholate siderophore, enterobactin. Anguibactin (81) siderophore was found in the marine pathogen *V. anguillarum* or *Vibrio* sp. DS40M4 [42,45]. It was evaluated cytotoxic effects on the P388 murine leukemia cell lines and was shown to be cytotoxic (IC<sub>50</sub> < 15 μM). It was also identified as an important virulence factor.

![Figure 6. Mixed hydroxamates/catecholates siderophores from marine microbes.](image)

2.7. Other Types of Siderophores

Besides, several other types of siderophores were also discovered in the marine microorganisms. A novel siderophore, piscibactin (82), was isolated from the marine fish pathogens *Photobacterium damsela* subsp. *piscicida* and *V. anguillarum*. The inactivation of piscibactin system would lead to a severe loss of the virulence degree. Therefore, piscibactin had a great impact in the expression of virulence, the ability to piscibactin seemed to be sufficient to confer maximal virulence to pathogens [65–67]. Two new oxazole–thiazole siderophore compounds, tetroazolemycins A (83) and B (84), were isolated from marine *Streptomyces olivaceus* FXJ8.012 [68]. Their heavy metal ion-binding ability was evaluated by Cu<sup>2+</sup>, Zn<sup>2+</sup>,...
Fe\textsuperscript{3+}, Pb\textsuperscript{2+}, Cr\textsuperscript{3+}, and Mn\textsuperscript{2+}. The results indicated that 82 and 83 had affinity for Cu\textsuperscript{2+}, Zn\textsuperscript{2+}, and Fe\textsuperscript{3+} but not affinity for Pb\textsuperscript{2+}, Cr\textsuperscript{3+}, and Mn\textsuperscript{2+}. Further studies showed the free-state tetroazolemycins did not show antimicrobial activity. Their metal ion complexes were also inactive against pathogens E. coli, S. aureus, Mycobacterium gilvum, Bacillus subtilis, P. aeruginosa, Candida pseudorugosa, Candida albicans, Rhizoctonia solani, Aspergillus fumigatus, and Fusarium oxysporum, inactive against A/H1N1 influenza virus, and inactive against human lung adenocarcinoma cell line A549 and murine macrophage cell line P388D. However, their Zn\textsuperscript{2+} complexes showed weakly activity against pathogenic Klebsiella pneumoniae with MICs of 125–250 µg/mL and 125 µg/mL, respectively. Thus, it was speculated that the antimicrobial activity of oxazole/thiazole siderophores might be closely related to Zn\textsuperscript{2+}.

![Chemical structures](image)

**Figure 7.** Other types of siderophores from marine microbes.

### 3. Biosynthesis of Siderophores from Marine Microorganisms

To date, more than 80 siderophores have been isolated from marine microorganisms, however, only part of siderophore biosynthetic pathways have been reported. A large number of novel siderophore biosynthetic pathways still need to be further studied. Therefore, researchers should focus more on the biosynthesis of siderophores from marine microorganisms to obtain structurally diverse siderophores as drug candidates.

#### 3.1. NRPS-Mediated Siderophore Biosynthetic Pathway

Recently, it has become apparent that the biosynthesis of siderophores involves covalently tethered a series of intermediates to the multienzymes throughout the assembly process and is catalyzed by nonribosomal peptide synthetases (NRPSs). NRPSs are responsible for assembling structurally complex peptides from small building blocks such as carboxyl or amino acids. Each module is highly efficient dedicated to the activation, modification, and incorporation of one small building block into the compound and harbors all key enzymatic activities as specialized domains to catalyze the single chemical reaction (Table 3). The de novo study of biosynthesis showed EnzA–D (NRPSs) were mainly responsible for the biosynthesis of siderophores, bisucaberin B (29) and bisucaberin (26) (Scheme 1). EnzA and EnzB catalyzed the biosynthesis of intermediates from L-lysine to cadaverine, and cadaverine...
to N-hydroxy-cadaverine, respectively. EnzC activated the acylation of N-hydroxy-cadaverine with succinyl-CoA to produce N-hydroxy-N-succinylcadaverine (HSC) monomer. EnzD as the last step noncovalently bound to HSC and ATP to catalyze the oligomerization–macrocyclization reaction of HSC to form 29 or 26. Firstly, a HSC molecule and ATP noncovalently bound to the active site of EnzD. A second molecule of HSC also noncovalently bound to EnzD, which dissociated from the active site of EnzD. The activated carboxyl group of the first molecule of HSC underwent a base-promoted addition–elimination reaction with the protonated amino group of the second molecule of HSC to yield the linear dimer 29. The activated carboxyl group and protonated amino group of 29 could further undergo an intramolecular base-promoted condensation to generate 26 [34,69,70]. In the biosynthetic network of 27, it exhibits the similar scheme of biosynthesis with 26 and 29. Firstly, MbsA–C catalyze successive reactions, resulting in ornithine and lysine to the N-hydroxy-N-succinyl diamines (HSDs). MbsD then utilizes both N-hydroxy-N-succinyl-putrescine and N-hydroxy-N-succinyl cadaverine as immediate precursors and catalyzes both the oligomerization and final macrocyclization, leading to the biosynthesis of 27.

![Scheme 1. The proposed biosynthesis of bisucaberin B (29) and bisucaberin (26).](image)

Vibrio ferrin (41) is synthesized by five biosynthetic enzymes PvsA-E (NRPSs). The biosynthetic pathway starting with enzyme PvsE. PvsE has a highly homologous with 2,6-diaminopimelate decarboxylase, indicating PvsE catalyzes decarboxylation from serine to ethanolamine. PvsB and PvsD form amide bond between L-alanine and 2-ketoglutaric acid, or pentanoic acid and ethanolamine. PvsA is perceived to form an ester bond between ethanolamine and citrate. The final product vibrio ferrin (41) is secreted by PvsC which is most likely to be a transporter involved in secreting the product from the cell [71].

The biosynthetic and regulatory elements of vanchrobactin (43) in marine bacteria *V. anguillarum* has been completely elucidated [42–44]. It proceeds through a NRPS-mediated pathway encoded by chromosomal genes to produce 43 (Scheme 2). VabB and VabE–F have been identified as three NRPSs. Isochorismate synthase (VabC) first started with the conversion of chorismate into isochorismate. Isochorismatase (VabB) and 2,3-dihydro-2,3-dihydroxybenzoate dehydrogenase (VabA) then catalyzed the biosynthesis of intermediates from isochorismate to 2,3-dihydro-2,3-dihydroxybenzoic acid (2,3-dihydro-2,3-DHBA), and (2,3-dihydro-2,3-DHBA) respectively to 2,3-DHBA. Once 2,3-DHBA was synthesized, VabE (NRPS) activated 2,3-DHBA to its acyl-adenylated intermediate in an ATP-dependent manner, and then VabE (NRPS) transferred this activated intermediate to VabB. VabB further bound with 2,3-DHBA through its ACP domain. VabF (NRPS) was involved in the last steps of 43 assembly. It was composed of two modules, including C-A1-PCP domain (condensation-adenylation-peptidyl-carrier-protein) in the N-terminal, and C-A2-PCP-TE domain (condensation-adenylation-peptidyl-carrier-protein-thioesterase) in the C-terminal. VabF A1 domain activated an arginine residue, and synthesized 2,3-dihydroxybenzoyl-arginine precursor through its PCP domain. VabF A2 domain activated a serine residue. The TE domain as the last domain was to release 43 into solution.
Mechanism of anguibactin (81) biosynthesis has been characterized in V. anguillarum (Scheme 3) [72]. A series of NRPSs are involved in V. anguillarum anguibactin (81) biosynthesis, such as AngB, AngD, AngE, AngM, AngN, and AngR. VabD and VabE are functional homologues of AngD and AngE, respectively. Each protein provides important functional domain(s) for anguibactin (81) biosynthesis. The adenylation (A) domains of VabE or likely of AngE activate 2,3-DHBA into 2,3-DHBA-AMP [73,74]. An aryl carrier protein (ArCP) domain of the C-terminal region of AngB is responsible for tethering 2,3-DHBA-AMP. The adenylation (A) domain of AngR is likely related to the activation of cysteine. It is worth noting that cyclisation (Cy) and peptidyl carrier protein (PCP) domains of AngR are not functional due to the replacement of the first aspartic acid by asparagine in the Cy domain, and the replacement of an essential serine by alanine in the PCP domain [75,76]. AngD or VabD is responsible for transferring the phosphopantetheinyl moiety to serine residues located in the ArCP domain of AngB and PCP domain of AngM, making them become active forms [74,77]. The PCP domain of AngM is used to tether the activated cysteine, and its C domain is likely to catalyze peptide bond formation between cysteine from AngM and 2,3-DHBA from AngB. AngN contains two Cy domains, Cy1 and Cy2 that could condense and cyclize 2,3-DHBA and cysteine to produce thiazoline [78,79]. The C domain of AngM could attach N-hydroxyhistamine to the dihydroxyphenyl–thiazoline–thioester to form anguibactin (81). AngT as the last step is to release anguibactin (81) from AngM [72].

Scheme 3. The proposed biosynthetic pathway of anguibactin (81).
Irp1-5 of *Photobacterium damselae* subsp. *piscicida* are mainly involved in the biosynthesis of siderophore piscibactin (82) (Scheme 4) [65,80]. Firstly, Irp5 (salicylate-activating enzyme) initiates piscibactin assembly, and then transfers the activated salicylate to aryl carrier protein domain of Irp2 (Iron-regulated protein 2). The ArCP domain of Irp2 is specific for cysteine activation. The activated cysteine is transferred to PCP1 and PCP2 simultaneously. Then, Irp2 catalyzes the continuous addition and cyclization of every cycle to form two thiazoline rings connected to a salicylate portion. Irp1 (PKS/NRPS multifunctional enzyme) is composed of 11 domains including a carboxyl-terminated thioesterase domain and three elongation modules. The ketosynthase domain in Irp1 loads a malonyl group which is linked to the ACP domain. The PCP3 domain of Irp1 catalyzes the condensation/heterocyclization of methyl-cysteine. Irp4 (thioesterase) hydrolyzes the thioester linkage. The final maturation of piscibactin (82) must be accomplished by Irp3 (reductase) which reduces the middle thiazoline ring to the thiazolidine.

**Scheme 4.** The proposed biosynthesis of piscibactin (82).

### 3.2. NRPS-Independent Siderophore Biosynthetic Pathway

NRPS-independent siderophore (NIS) synthetases are an emerging member of the family of new synthetases. They do not have the sequence or structural similarity to NRPSs. However, they also play an important role in the biosynthesis of siderophores. Many structurally diverse siderophores are biosynthesized by NIS synthetases (Table 3). NIS synthetases contain type A, type B, and type C based on sequence similarity criteria. Type A enzymes catalyze the condensation of amines or alcohols with a prochiral carboxyl group of citric acid. Type B enzymes catalyze the condensation of amines with γ-carboxyl group of α-ketoglutarate. Type C enzymes catalyze either the oligomerization and macrocyclisation of ω-amino-carboxylic acids, or an amine or alcohol with a monoamide derivative of citric acid. By selective gene knockout and reconstruction of *iuc* genes, aerobactin (65) biosynthetic pathway was deciphered. IucA–D are responsible for the biosynthesis of 65 (Scheme 5). They are highly stereoselective. IucB is confirmed to its l-lysine $N^6$ monoxygenase activity [81]. Similarly, IucD is also confirmed to its $N^6$-hydroxy-l-lysine (hLys) acetyltransferase activity [82]. IucB and IucD respectively catalyze the biosynthesis of intermediates from l-lysine to hLys, and hLys to $N^6$-acetyl-$N^6$-hydroxy-l-lysine (ahLys). IucA and IucC represent the archetypal Type A and Type C NIS synthetases, respectively [83]. IucA catalyzes the carboxymethyl group of citric acid to form a citryl-adenylate intermediate via the activation of ATP. IucC then consumes 3S,2'S-citryl-ahLys intermediate in a relatively modest catalytic efficiency to synthesize 65.
Based on the NRPS-independent assembly, the biosynthetic scheme of petrobactin (72) is developed (Scheme 6). AsbA (type A NIS synthetase) has been demonstrated to catalyze the ATP-dependent condensation of \(N^8\) of spermidine with citric acid to afford \(N^8\)-citryl-spermidine. Another NIS synthetase AsbB (type C NIS synthetase) catalyzes ATP-dependent condensation of \(N^8\)-citryl-spermidine or \(N^7\)-(3,4-dihydroxybenzoyl)-\(N^8\)-citryl-spermidine with spermidine. AsbF, a 3-dehydroshikimate (3-DHS) dehydratase, catalyzes the conversion of the bacterial metabolite 3-DHS to form 3,4-dihydroxybenzoic acid (3,4-DHBA). AsbC, a NRPS-like adenylyating enzyme, is responsible for catalyzing the ATP-dependent acylation of phosphopantetheine thiol of AsbD (a carrier protein) with 3,4-DHBA. AsbE as an acyltransferase is involved in the 3,4-dihydroxybenzoyl group transfer from the carrier protein AsbD to \(N^7\) and \(N^8\) of spermidine to afford \(N^7\)-(3,4-dihydroxybenzoyl)-spermidine and \(N^8\)-(3,4-dihydroxybenzoyl)-spermidine, respectively, suggesting that AsbE has relatively relaxed substrate specificity \[60,84,85\].

**Scheme 6.** The proposed biosynthetic pathway of petrobactin (72).

**4. Synthesis and Study of Siderophores from Marine Microorganisms**

Marine siderophores, as potentially valuable drug candidates, are attracting extensive attention of researchers. However, the content of siderophores from marine microbes is very low. Therefore, in order to further study their biological activities, mechanisms, absolute configurations, or structure–activity relationships, chemical syntheses of siderophores would be crucial.
4.1. Alcaligin

Alcaligin (25) had similar structural relationship to siderophores bisacaberen (26) and nocardamine. Bisacaberen (26) could inhibit the growth of both invasive micropapillary carcinoma and L-1210 leukemia cells with IC_{50} of 12.7 and 9.7 μM, respectively, and sensitize tumor cells to macrophage-mediated cytolysis [32]. However, this activity was absent in siderophore nocardamine. An investigation of the biological properties of 25 is particularly attractive. Therefore, Bergeron R.J. et al. [86] first performed the total synthesis of alcaligin (25) (Scheme 7). The starting point in the total synthesis of alcaligin was the regiospecific N-alkylation of ditosylate 1a providing monotosylate 2a. A second amino group was then coupled to C-1 of 2a by N-alkylation of trifluoroacetamide producing the diamide 3a. The primary amine 4a was produced by basic cleavage of 3a. Brief exposure of the N-(tert-butoxycarbonyl)amine 4a to trifluoroacetic acid led to the bis(benzyloxy)putrescine 5a. Addition of 2-[(tert-butoxycarbonyloxy)imino]-2-phenylacetonitrile to 5a resulted in the benzylxy amine 6a. 6a was next acylated with succinic anhydride to generate tert-butoxycarbonyl acid 7a. Subsequently, the condensation of 5a and 7a to produce the (benzylxy)amine 8a. The second succinate unit was coupled with 9a to generate tert-butoxycarbonyl acid 9a and 10a. Macrocyclic tetrabenzyl alcaligin 11a was obtained under the catalysis of diphenylphosphorazidate. Finally, the benzyl groups of 11a were removed to obtain alcaligin (25).

![Scheme 7. Chemical synthesis of alcaligin (25).](image)

4.2. Vanchrobactin

In the course of the studies into vanchrobactin (43), its stereochemistry could not be elucidated by spectroscopic analysis. In order to determine the absolute configuration of vanchrobactin (43), Soengas R.G. et al. [87] performed the total synthesis of vanchrobactin (43) (Scheme 8). 2,3-dihydroxybenzoic acid (2,3-DHBA) was first protected by benzyl bromide to generate tert-butoxycarbonyl acid (2,3-DHBA) was first protected by benzyl bromide to generate tert-butoxycarbonyl acid 2b. Subsequent saponification with barium hydroxide gave 2,3-dibenzoylbenzoic acid 2b. On the other hand, N^3-Cbz-{ornithine-OMe 3b was obtained from the Cbz analogue of o-ornithine copper complex. Coupling of 3b with 2,3-dibenzoylbenzoic acid 2b gave compound 4b. Deprotection of the benzyl groups in 5b, introduction of the guanidine functionality to obtain 6b and acidic hydrolysis produced vanchrobactin (43). Moreover, in order to further deduce structure–activity relationships of vanchrobactin (43), several 43 analogues were also synthesized by a similar strategy of the total synthesis of vanchrobactin (43) [88]. The results suggest the aromatic ring in catecholate siderophores is essential for the binding of the outer membrane receptors. The lack of stereochemistry of the amino acid scaffold will affect the siderophore activity in V. anguillarum. Moreover, microbes are not always selective in the use of siderophores, they can utilize several different siderophores to transport Fe(III) into microbial cells, even siderophores
with very different structures. The relatively low specificity of siderophore utilization may facilitate the design of drugs against different pathogens.

4.3. Petrobactin

Pandey R.K. et al. [89] performed efficient total synthesis of petrobactin (72) via antimony triethoxide mediated coupling (Scheme 9). It not only proved the structure of 72, but also provided sufficient yield for subsequent use in the therapeutic study. First, the hydroxyl group of N-dibenzyl-aminoalcohol 1c was mesyl protected to provide compound 2c. Nucleophilic displacement of O-Ms with amine 3c yielded 4c. Debenzylation of 4c under H2-Pd/C conditions to yield 5c. On the other hand, esterification of 6c under reflux conditions was performed, followed by benzyl protection of catechol unit to yield 8c. Ester–amide exchange was performed with amine 5c and acid 8c to produce 9c. The N-Boc group was removed from 9c to obtain amine 10c. Coupling of 10c and 12c was carried out using trimethylamine to yield protected petrobactin 12c. Finally, 12c was debenzylated to yield petrobactin (72). Moreover, sidechain-modified petrobactin derivatives were also synthesized for the study of iron binding, the results suggest both acylation and alkylation of the spermidine sidechain basically do not affect iron binding properties of the siderophore [90].

Scheme 8. Chemical synthesis of vanchrobactin (43).

Scheme 9. Chemical synthesis of petrobactin (72).
5. Conclusions and Perspectives

Siderophores from marine microbes are an important class of secondary metabolites that have intrigued microbiologists, chemists, and biochemists due to their structural diversity and potential applications in diseases and medicinal agents. However, with widely stated estimate that less than 1% marine microbes have been brought culture [91], the great challenge includes the development of new methods to bring more marine microbes into culture. Moreover, further detailed understanding of chemical synthesis and biosynthesis of marine siderophores will provide valuable insight for their study of biological activities, and elucidation of mechanism. In total, marine siderophore research will continue stimulating a multidisciplinary research effort that will facilitate their potential applications in medicine and health care.

| Compd. | Name                  | PubChem CID | PubChem Database                      | Ref.        |
|--------|-----------------------|-------------|---------------------------------------|-------------|
| 1      | Marinobactin A        | –           | –                                     | [22]        |
| 2      | Marinobactin B        | –           | –                                     | [22]        |
| 3      | Marinobactin C        | –           | –                                     | [22]        |
| 4      | Marinobactin D        | –           | –                                     | [22]        |
| 5      | Marinobactin E        | –           | –                                     | [22]        |
| 6      | Marinobactin F        | –           | –                                     | [22]        |
| 7      | Marinobactin HG       | –           | –                                     | [23]        |
| 8      | Amphibactin           | –           | –                                     | [24]        |
| 9      | Amphibactin           | –           | –                                     | [24]        |
| 10     | Amphibactin           | –           | –                                     | [24]        |
| 11     | Amphibactin           | –           | –                                     | [24]        |
| 12     | Amphibactin           | –           | –                                     | [24]        |
| 13     | Amphibactin           | –           | –                                     | [24]        |
| 14     | Amphibactin           | –           | –                                     | [24]        |
| 15     | Amphibactin           | –           | –                                     | [24]        |
| 16     | Amphibactin           | –           | –                                     | [24]        |
| 17     | Amphibactin           | –           | –                                     | [24]        |
| 18     | Moanachelin           | –           | –                                     | [25]        |
| 19     | Moanachelin           | –           | –                                     | [25]        |
| 20     | Moanachelin           | –           | –                                     | [25]        |
| 21     | Moanachelin           | 122223347   | [25]                                  |             |
| 22     | Moanachelin           | –           | –                                     | [25]        |
| 23     | Amphibactin U         | –           | –                                     | [26]        |
| 24     | Amphibactin V         | –           | –                                     | [26]        |
| 25     | Alcaligin             | –           | –                                     | [27]        |
| 26     | Bisucaberin           | –           | –                                     | [28]        |
| 27     | Avaroferrin           | –           | –                                     | [29]        |
| 28     | Putrebacitin          | –           | –                                     | [30]        |
| 29     | Bisucaberin B         | –           | –                                     | [34]        |
| 30     | Thalassosamide        | –           | –                                     | [35]        |
| 31     | Fradiamine A          | 129008905   | [26]                                  |             |
| 32     | Fradiamine B          | 60151746    | [26]                                  |             |
| 33     | Albisporachelin       | –           | –                                     | [37]        |
| 34     | Desferrioxamine A₁    | –           | –                                     | [38]        |
| 35     | Desferrioxamine A₂    | –           | –                                     | [38]        |
| 36     | Desferrioxamine B     | –           | –                                     | [38]        |
| 37     | Desferrioxamine D₁    | –           | –                                     | [38]        |
| 38     | Desferrioxamine D₂    | –           | –                                     | [38]        |
| 39     | Desferrioxamine E     | –           | –                                     | [38]        |
| 40     | Desferrioxamine N     | –           | –                                     | [38]        |
| Compd. | Name          | PubChem CID | PubChem Database                          | Ref.  |
|-------|---------------|-------------|-------------------------------------------|-------|
| 41    | Vibrioferrin  | 11102119    | https://pubchem.ncbi.nlm.nih.gov/compound/11102119 | [39]  |
| 42    | Nigribactin   | –           | –                                         | [40]  |
| 43    | Vanchrobactin | –           | –                                         | [45]  |
| 44    | Trivanchrobactin | –         | –                                         | [45]  |
| 45    | Divanchrobactin| –           | –                                         | [45]  |
| 46    | Turnerbactin  | –           | –                                         | [46]  |
| 47    | Dibenarthin   | –           | –                                         | [47]  |
| 48    | Streptobactin | –           | –                                         | [47]  |
| 49    | Tribenarthin  | –           | –                                         | [47]  |
| 50    | Pistillarin   | –           | –                                         | [48]  |
| 51    | Loihichelin A | 101476230   | https://pubchem.ncbi.nlm.nih.gov/compound/101476230 | [51]  |
| 52    | Loihichelin B | 101476231   | https://pubchem.ncbi.nlm.nih.gov/compound/101476231 | [51]  |
| 53    | Loihichelin C | 101476232   | https://pubchem.ncbi.nlm.nih.gov/compound/101476232 | [51]  |
| 54    | Loihichelin D | 101476233   | https://pubchem.ncbi.nlm.nih.gov/compound/101476233 | [51]  |
| 55    | Loihichelin E | 101476234   | https://pubchem.ncbi.nlm.nih.gov/compound/101476234 | [51]  |
| 56    | Loihichelin F | 101476235   | https://pubchem.ncbi.nlm.nih.gov/compound/101476235 | [51]  |
| 57    | Aquachelin A  | –           | –                                         | [22]  |
| 58    | Aquachelin B  | –           | –                                         | [22]  |
| 59    | Aquachelin C  | –           | –                                         | [22]  |
| 60    | Aquachelin D  | –           | –                                         | [22]  |
| 61    | Aquachelin I  | –           | –                                         | [23]  |
| 62    | Aquachelin J  | –           | –                                         | [52]  |
| 63    | Aquachelin HG | –           | –                                         | [29]  |
| 64    | Imaqobactin   | –           | –                                         | [53]  |
| 65    | Aerobactin    | –           | –                                         | [54]  |
| 66    | Ochrobactin A | –           | –                                         | [56]  |
| 67    | Ochrobactin B | –           | –                                         | [56]  |
| 68    | Ochrobactin C | –           | –                                         | [56]  |
| 69    | Synechobactin | 122377042   | https://pubchem.ncbi.nlm.nih.gov/compound/122377042 | [57]  |
| 70    | Synechobactin | 122377043   | https://pubchem.ncbi.nlm.nih.gov/compound/122377043 | [57]  |
| 71    | Synechobactin | 122377044   | https://pubchem.ncbi.nlm.nih.gov/compound/122377044 | [57]  |
| 72    | Petrobactin   | –           | –                                         | [58]  |
| 73    | Petrobactin sulfonate | –       | –                                         | [59]  |
| 74    | Alterobactin A| –           | –                                         | [61]  |
| 75    | Alterobactin B| 101775921   | https://pubchem.ncbi.nlm.nih.gov/compound/101775921 | [62]  |
| 76    | Pseudoalterobactin A| 11434714 | https://pubchem.ncbi.nlm.nih.gov/compound/11434714 | [63]  |
| 77    | Pseudoalterobactin B| 11788080 | https://pubchem.ncbi.nlm.nih.gov/compound/11788080 | [63]  |
| 78    | Lystabactin A | –           | –                                         | [64]  |
| 79    | Lystabactin B | –           | –                                         | [64]  |
| 80    | Lystabactin C | –           | –                                         | [64]  |
| 81    | Anguibactin   | –           | –                                         | [42,45]|
| 82    | piscibactin   | 136754132   | https://pubchem.ncbi.nlm.nih.gov/compound/136754132 | [65–67]|
| 83    | Tetroazolemycin A| –         | –                                         | [68]  |
| 84    | Tetroazolemycin B| –         | –                                         | [68]  |
Table 3. The EC numbers of enzymes.

| Enzymes | EC Numbers |
|---------|------------|
| EnzA    | 4.1.1.7    |
| VabC    | 3.1.22.4   |
| VabD    | 2.7.8.7    |
| VabF    | 3.2.1.55   |
| Irp1    | 2.3.2.27   |
| Irp2    | 2.3.2.27   |
| Irp3    | 2.3.2.27   |
| Irp4    | 2.3.2.27   |
| AngB    | 3.3.2.1    |
| AngE    | 1.14.14.148|
| VabD    | 2.7.8.7    |
| IucA    | 6.3.2.38   |
| IucC    | 6.3.2.39   |
| IucD    | 1.14.13.59 |
| AsbF    | 4.2.1.118  |

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