Review

Briarane Diterpenoids Isolated from Octocorals between 2014 and 2016

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Abstract: The structures, names, bioactivities, and references of 124 briarane-type natural products, including 66 new metabolites, isolated between 2014 and 2016 are summarized in this review article. All of the briarane diterpenoids mentioned in this review were isolated from octocorals, mainly from Briareum violacea, Dichotella gemmacea, Ellisella dollfusi, Junceella fragilis, Junceella gemmacea, and Pennatula aculeata. Some of these compounds exhibited potential biomedical activities, including anti-inflammatory activity, antibacterial activity, and cytotoxicity towards cancer cells.

Keywords: briarane; octocoral; Briareum; Dichotella; Ellisella; Junceella; Pennatula

1. Introduction

Following previous review articles from our research group focused on marine-origin briarane-type natural products [1–5], this review covers the literature from 2014 to January 2017, and describes 124 briarane-related diterpenoids (including 66 new metabolites), most of which are characterized by the presence of a γ-lactone moiety fused to a bicyclo[8.4.0] ring system, obtained from various octocorals (Figure 1), mainly Briareum violacea, Briareum spp. Dichotella gemmacea, Ellisella dollfusi, Junceella fragilis, Junceella gemmacea, and Pennatula aculeata. Many of these compounds exhibited interesting bioactivities in vitro, which might indicate a potential for use in biomedical applications. This survey of briarane-related compounds is presented taxonomically according to genus and species.
The structures of new briaranes and elastase release, with IC\(\Delta\) spectroscopic data (Figure 3) [16–20]. Briarenolide J (27) was established by chemical and spectroscopic methods, and its monobenzoyl derivative [15]. At a concentration of 10 \(\mu\)g/mL, briaranes and six known metabolites, solenolides A [10] and D (=briaexcavatolide E) [1,10–12], excavatolide, deacetylstylatulide, and asbestinum were isolated from the octocoral and collected from the waters of Taiwan [15]. The structures of new briaranes 1–10 were established by chemical and spectroscopic methods, and determination of the absolute configuration of briaviolide A (1) was completed by X-ray diffraction analysis of its monobenzoyl derivative [15]. At a concentration of 10 \(\mu\)g/mL, briaranes 5 and 9 were found to exert moderate inhibitory activities on elastase release (inhibition rate = 26.0% and 28.8%, respectively) and superoxide anion production (inhibition rate = 34.2% and 28.7%, respectively) by human neutrophils [15].

2.1. Briareae violacea (Family Briareidae)

The taxonomic position of octocorals affiliated with the genus Briareum (=Asbestia, Pachyclavularia, and Solenopodium) [6] has been found to be situated near the transition between Alcyonacea and Gorgonacea, in both taxonomic and chemical terms [6–8]. At a concentration of 10 \(\mu\)g/mL, briaranes and six known metabolites, solenolides A [10] and D (=briaexcavatolide E) [1,10–12], excavatolide, deacetylstylatulide, and asbestinum were isolated from the octocoral Briareum violacea, collected from the waters of Taiwan [15]. The structures of new briaranes 1–10 were established by chemical and spectroscopic methods, and determination of the absolute configuration of briaviolide A (1) was completed by X-ray diffraction analysis of its monobenzoyl derivative [15]. At a concentration of 10 \(\mu\)g/mL, briaranes 5 and 9 were found to exert moderate inhibitory activities on elastase release (inhibition rate = 26.0% and 28.8%, respectively) and superoxide anion production (inhibition rate = 34.2% and 28.7%, respectively) by human neutrophils [15].

![Figure 1](image1.png)

**Figure 1.** Possible biogenetic origin of briarane-type metabolites. The numbering system shown is that presently in use [1].

2.2. Briareum sp.

In continuing chemical studies of the constituents of an octocoral identified as Briareum sp. collected from the southern waters of Taiwan, 22 new briarane derivatives, briarenolides J–Y (11–26) and ZI–ZVI (27–32), were obtained, and their structures determined based on analysis of their spectroscopic data (Figure 3) [16–20]. Briarenolide J (11) was the first 12-chlorinated diterpenoid to be isolated from Briareum sp. [16]. The relationships between the \(^1\)H and \(^13\)C NMR chemical shifts of 2-hydroxybriaranes possessing a \(\Delta^{3,5(16)}\)-conjugated diene moiety or a \(\Delta^{3,5}\)-conjugated moiety have been summarized [18]. Briarane 11 has been shown to inhibit superoxide anion generation and elastase release, with IC\(_{50}\) values of 15.0 and 10.0 \(\mu\)M, respectively [16]. In macrophage cells,
briaranes 12–14, 17, 20–24, 26, 28, and 32 were found to reduce the level of iNOS to 23.7%, 31.7%, 49.6%, 58.4%, 57.4%, 53.5%, 41.9%, 47.3%, 50.1%, 54.3%, 47.2% and 55.7%, respectively, at a concentration of 10 µM [17–20]. Briaranes 15, 17, 21–24, and 26 were found to reduce the level of COX-2 to 53.9%, 59.1%, 59.3%, 26.1%, 35.6%, 58.1% and 55.4%, respectively, at a concentration of 10 µM [18,19].

Figure 3. Structures of briarenolides J–Y (11–26) and ZI–ZVI (27–32).
3. Gorgonacea

3.1. *Dichotella gemmacea* (Family Ellisellidae)

In 2014, Zhang et al. reported the isolation of seven new briarane derivatives, which were named gemmacolides AS–AY (33–39) (Figure 4), along with 10 known analogues, gemmacolides L [21], X (=dichotellide T) [22,23], AH, AJ, AO, AQ [24], junceellolides C and D [25], junceellin (=junceellin A) [25–32], and frajunolide K [33], from the South China Sea gorgonian coral, *D. gemmacea* [34]. Structural determination of new briaranes 33–39 was conducted using spectroscopic methods, and their absolute configurations were established based on the results of electronic circular dichroism (ECD) experiments [34]. Briarane 37 was found to exert a cytotoxic effect towards MG-63 (human osteosarcoma) cells, with an IC₅₀ value of 7.2 μM [34].

![Figure 4. Structures of gemmacolides AS–AY (33–39).](image)

A new briarane, dichotellide V (40) (Figure 5), along with four known briarane analogues, gemmacolide N [35], dichotellide J [23], junceellin A (=junceellin) [25–32], and junceellolide A [25], were isolated from *Dichotella gemmacea*, collected from Meishan Island, Hainan Province, China [36]. The structure of new briarane 40 was determined by spectroscopic methods, and none of the above compounds exhibited a cytotoxic effect on A549 (human epithelial lung carcinoma), BGC823 (human gastric cancer), H1975 (human non-small cell lung cancer), HeLa (human cervix adenocarcinoma), MCF7 (human mammary gland adenocarcinoma), or U-937 (human histiocytic lymphoma) tumor cells [36].

![Figure 5. Structure of dichotellide V (40).](image)
Eight known briaranes, junceellolide D [25], (+)-11β,20β-epoxyjuncceellolide D, (−)-11β,20β-epoxy-4-deacetoxyjuncceellolide D [30,37], junceol A [38], juncins H and K [39,40], praelolide [25,29–32,41,42], and juncceillin (=juncceillin A) [25–32–32] were obtained from *D. gemmacea*, collected from Meishan Island, Hainan Province, China in April 2009 [43]. Junceellolide D and praelolide showed antifouling activity against the settlement of larvae of barnacle *Balanus amphitrite* with EC50 values of 14.5 and 16.7 μM, respectively. Junceellolide D, (−)-11β,20β-epoxy-4-deacetoxyjuncceellolide D, juncin H, and praelolide exhibited lethality towards brine shrimp *Artemia salina* with lethal ratios of 90%, 85%, 60% and 75% at a concentration of 50 μg/mL [43].

In addition, seven new briaranes, gemmacolides AZ–BF (41–47) (Figure 6), and eight known analogues, dichotellides M and O [23], gemmacolide C [44], juncins P [45] and ZI [46], junceellolides D [25] and K [37], and (−)-4-deacetyljunceellolide D [30], were obtained from *D. gemmacea*, collected from the South China Sea [47]. The structures of new briaranes 41–47 were determined by spectroscopic methods. Briaranes 41–44, 46, 47, and dichotellide O, showed cytotoxicity towards A549 cells, with IC50 values of 28.3, 24.7, 34.1, 26.8, 25.8, 13.7 and 25.5 μM, respectively. Briaranes 42, 44, 46, 47, and dichotellide O, showed cytotoxicity towards MG-63 cells, with IC50 values of 15.8, 11.4, 30.6, 34.8 and 36.8 μM, respectively. Briarane 44 and dichotellide O exhibited antibacterial activity against the Gram-negative bacterium *Escherichia coli*, while dichotellide O demonstrated actitity against the Gram-positive bacterium *Bacillus megaterium* [47].

![Figure 6. Gemmacolides AZ–BF (41–47).](image)

3.2. *Ellisella dollfusi* (Family Ellisellidae)

Zhou and coworkers isolated seven briaranes, including two new compounds, dollfusilins A (48) and B (49) (Figure 7), along with five known analogues, brianthein W [14,48,49], funicolide E [49], 9-deacetyl briareolide H [14,50,51], 9-deacetylstylatulide lactone [14], and umbraculolide A [29,52], from the organic extract of gorgonian coral *Ellisella dollfusi*, collected from the Xisha Sea area of the South China Sea [53]. The structures of new briaranes 48 and 49 were determined through comprehensive analysis of spectroscopic data. Brianthein W exhibited an effect of delayed hatching and notochord growth malformation toxicity towards zebrafish *Danio rerio* embryos with IC50 values of 30.6 and 18.9 μg/mL in 48 h, respectively. Funicolide E displayed egg coagulation and delayed hatching toxicity towards zebrafish embryos, with EC50 values of 33.6 μg/mL (24 h) and 29.8 μg/mL, respectively [53].
3.3. *Juncturella fragilis* (Family Ellisellidae)

Gorgonian corals belonging to the genus *Juncturella* have also been found to be major sources of briarane-related natural diterpenoids [54,55]. The gorgonian *J. fragilis*, collected from the South China Sea, was found to contain 12 new briaranes, fragilisinins A–L (50–61) [56] (Figure 8), along with seven known analogues, (+)-juncellolide A [30], juncellolide B [25], juncelol A [38], junceellonoid D [57,58], fragilide C [59], and frajunolides A [60] and E [33]. The structures of new briaranes 50–61 were determined by spectroscopic methods. Briaranes 58–61 were the first iodine-containing briarane derivatives to be isolated. The absolute configuration of briarane 50 was confirmed by single-crystal X-ray diffraction data [56]. Briaranes 54, 55, 59, (+)-juncellolide A, and junceellonoid D showed potent antifouling activities against the settlement of barnacle *Balanus amphitrite* larvae, with EC_{50} values of 14.0, 12.6, 11.9, 5.6, and 10.0 µM (LC_{50}/EC_{50} = >13, >14.5, >11.5, >33.3, >20), respectively [56].

![Figure 7. Structures of dollfusilins A (48) and B (49).](image)

![Figure 8. Structures of fragilisinins A–L (50–61).](image)

3.4. *Juncturella gemmacea* (Family Ellisellidae)

Four new briaranes, junceellolides M–P (62–65) (Figure 9) [61], along with seven known briaranes, junceellolides A–D [25], juncelin A [25–32], praelolide [25,29–32,41,42], and juncin ZI [46], were isolated from the gorgonian *J. gemmacea*, collected from the South China Sea [61]. The structures, including the absolute configurations, of new briaranes 62–65, were deduced on
the basis of spectroscopic analyses, particularly electronic circular dichroism (ECD) experiments, and from biogenetic correlations among briaranes 62–65.

![Images of chemical structures](image1)

**Figure 9.** Structures of junceellolides M–P (62–65).

3.5. *Junceella* sp. (*Family Ellisellidae*)

Three known briaranes, junceellin (=junceellin A) [25–32], praelolide [25,29–32,41,42], and junceellolide A [25], were claimed to have been obtained from a gorgonian coral *Junceella* sp., collected off the Vietnam Thu Island in May 2010 [62]. In the antimicrobial activity test, junceellin and praelolide exhibited weak antibacterial activity against the bacterium *Vibrio parahaemolyticus*. Junceellolide A was also found to display weak antibacterial activity against the bacterium *Candida albicans* [62].

4. *Pennatulacea*

*Pennatula aculeata* (*Family Pennatulidae*)

Investigation of the chemical constituents of *P. aculeata*, collected from Dinawan Island in Sabah, Malaysia, afforded novel briarane 2-acetoxyverecynarmin C (66) [63] (Figure 10). The structure of the new briarane 66 was elucidated by analysis of spectroscopic data, and this compound showed moderate inhibitory activity towards COX-1 and COX-2, with IC$_{50}$ values of 44.3 and 47.3 μM, respectively. The 2-acetoxy group in 66 was found to be located on the α-face, relative to Me-15 and H-10, which is a rare occurrence in briarane-related analogues.

![Images of chemical structure](image2)

**Figure 10.** Structure of 2-acetoxyverecynarmin C (66).

5. Conclusions

Since briarein A, the first briarane-type natural product, was prepared from the Caribbean octocoral *Briareum asbestinum* in 1977 [9], over 600 briarane-type diterpenoids have been isolated from a wide variety of marine life to date. A large portion of these natural compounds has been prepared from soft corals belonging to the orders Alcyonacea and Gorgonacea. Compounds of this type of diterpenoid have been demonstrated to possess various bioactivities in vitro, such as anti-inflammatory activity, antibacterial activity, and cytotoxicity towards cancer cells. For example, one of the compounds of this type, excavatolide B [13], has been proven to possess extensive biomedical bioactivities, such as anti-inflammatory, analgesic, the attenuation of rheumatoid arthritis activities, anticancer, and the modulation of the electrophysiological characteristics and calcium homeostasis in
atrial myocytes [64–67]. Due to the structural diversity and biomedical bioactivities, there has been little synthetic work on briarane analogues [68,69]. It is interesting to note that all briaranes reported as having been isolated between 2014 and 2016 were all collected from octocorals distributed in the Indo-Pacific Ocean, particularly from the South China Sea.

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