# Naturally Occurring Diterpenoid Dimers: Source, Biosynthesis, Chemistry and Bioactivities

Authors

Affiliation

#### Li-Gen Lin, Carolina Oi Lam Ung, Zhe-Ling Feng, Li Huang, Hao Hu

State Key Laboratory of Quality Research in Chinese Medicine, Institute of Chinese Medical Sciences, University of Macau, Macau, PR China

Key words

- diterpenoid dimers
- structure
- source
- biosynthesis
- synthesis
- bioactivities

received	April 13, 2016
revised	July 14, 2016
accepted	August 1, 2016

Bibliography DOI http://dx.doi.org/ 10.1055/s-0042-114573 Published online August 19, 2016 Planta Med 2016; 82: 1309–1328 © Georg Thieme Verlag KG Stuttgart · New York · ISSN 0032-0943

#### Correspondence Dr. Li-Gen Lin

Dr. Li-Cen Lin State Key Laboratory of Quality Research in Chinese Medicine Institute of Chinese Medical Sciences University of Macau Avenida da Universidade Taipa 999078 Macau PR China Phone: + 85 3 88 22 80 41 Fax: + 85 3 28 84 13 58 ligenl@umac.mo

# Abstract

Diterpenoid dimers are rare in nature and mainly found in higher plants including the families Acanthaceae, Annonaceae, Asteraceae, Calceolariaceae, Chrysobalanaceae, Cupressaceae, Euphorbiaceae, Fabaceae, Lamiaceae, Liliaceae, Meliaceae, Rhizophoraceae, Taxaceae, Velloziaceae, and Zingiberaceae. In addition, a few diterpenoid dimers have been also reported from fungi (Psathyrellaceae), liverworts (Scapaniaceae), and a gorgonian (Gorgoniidae). They feature a wide variety of structures due to different core skeletons, linkage patterns, substituents, and configurations. Accordingly, diterpenoid dimers exhibit a broad range of bioactivities, including cytotoxic, anti-inflammatory, antimicrobial, antimalarial, and antifouling properties, which have attracted more and more research interests in the past decades. This review with 176 metabolites from 109 references provides a comprehensive and up-todate overview of the source, biosynthesis, structure, synthesis, and bioactivities of diterpenoid dimers.

# Abbreviations

<b>V</b>	
DGAT:	diacylglycerol O-acyltransferase
ECD:	electronic circular dichroism
LPS:	lipopolysaccharide
NO:	nitric oxide
TRAIL:	tumor necrosis factor-related apopto-
	sis-inducing ligand

# Introduction

Diterpenoids are a structurally diverse class of natural products, consisting of four isoprene units to form a 20-carbon backbone. Based on their core structures, diterpenoids can be classified into linear, macrocyclic, bicyclic, tricyclic, tetracyclic and pentacyclic types [1]. Naturally occurring diterpenoids are always found in a polyoxygenated form: i) with hydroxyl groups, which are often esterified by aliphatic or aromatic acids, or etherified by alcohols; ii) with formyl and carbonyl groups, which have often reacted with hydroxyl groups to form hemiacetal or acetal moieties; or iii) with carboxyl groups, which often form esters or lactones through reaction with alcohols. Moreover some diterpenoids have been found with opened and rearranged ring structures. Diterpenoids exhibit various biological activities such as cytotoxic, anti-microbial, and anti-inflammatory properties, and have been identified as active principles in some traditional medicines [2,3]. Due to their structural diversity and broad bioactivities, diterpenoids have attracted increasing research attentions, resulting in the identification of a growing number of compounds. Some of these compounds have been proven to be clinically effective. Taxol, for example, is an unusual diterpenoid discovered from Taxus brevifolia (Taxaceae), which inhibits the normal breakdown of microtubules during cell division, and is widely used in therapy against ovarian, breast, and lung cancer [4]. Salvinorin A, a diterpenoid isolated from Salvia divinorum, has psychoactive effects on humans, and has been used as an analgesic [5, 6].

Diterpenoid dimers are a rather uncommon subclass of diterpenoids, which are composed of two 20-carbon diterpenoid units linked through one or two C–C bond, ester bond, ether bond or a ring moiety. On the basis of a literature search in various databases, including Google Scholar, SciFinder, Web of Science, and Scopus, using "diterpenoid dimer" or "tetraterpenoid" as key words, 176 naturally occurring diterpenoid dimers were retrieved, with various bioactivities including cytotoxic, anti-inflammatory, antimicrobial, antimalarial, and antifouling activities (**• Table 1**). Due to their structural complexity and the wide range of bioactivities, naturally occurring diterpenoid dimers have attracted more and more research interests in the past decades. Furthermore, the development of purification and structural elucidation methods, especially preparative HPLC and high resolution NMR, makes it now possible to identify compounds at trace amounts. Chiral HPLC dramatically accelerated the isolation of stereoisomers, especially those with complex structures like diterpenoid dimers [7-9]. In recent years, ECD calculations have been widely used to determine the absolute configurations of diterpenoid dimers [7,9–11]. According to our literature review, around 31 publications related to diterpenoid dimers were found before 2000, and this number has increased to 88 in January 2016. However, up to now, no systematical review was carried out on this particular group of diterpenoids. The present review describes the biosynthesis, occurrence and structures of diterpenoid dimers identified up to date, summarizes their bioactivities, gives examples of total synthesis, and explores some research perspectives on diterpenoid dimers.

# **Biosynthesis**

The biosynthesis of diterpenoids has been widely investigated and involves different modes of cyclization of geranylgeranyl diphosphate (GGPP) [12]. To date, four different classes of synthases have been cloned with their functional proteins sequenced, including casbene synthase, ent-copalyl diphosphate synthase, taxadiene synthase, and abietadiene synthase. In contrast, the biosynthesis of diterpenoid dimers, especially the identification and characterization of the key dimerization processes, has been seldom studied. There are different ways in which diterpenoid dimers can be linked, including a rotatable or atropisomeric C-C bond, an ether C-O-C linkage, an ester bond, or a ring formed through homo- or hetero-Diels-Alder reaction. The incorporation of a malonic acid unit to form a diester linkage is observed in the diterpenoid dimers from the families Calceolariaceae and Asteraceae. Most of diterpenoid dimers arise as a result of a Diels-Alder condensation of two monomeric moieties. Yue and colleagues [7] proposed that aphadilactones A-D were formed from two molecules of nemoralisin-type diterpenoid through enzyme-catalyzed Diels-Alder reaction. Later, the total synthesis of these compounds using Diels-Alder cyclization further supported this hypothesis [13]. Until now some progresses have been achieved in the identification of natural Diels-Alderase [14]. Ichihara and colleagues [15, 16] isolated and purified an enzyme from Alternaria solani that catalyzes the [4+2] cycloaddition of prosolanapyrone III to the exo adduct solanapyrone A and endo adduct solanapyrone D. This enzyme was the first Diels-Alderase reported. In 2000, a study carried out by Vederas's group [17] identified a lovastatin nonaketide synthase, which catalyzes intramolecular Diels-Alder endo closure of (E,E,E)-(R)-6-methyl-dodecatri-2,8,10-enoic acid N-acetylcysteamine thioester to a bicyclic system. In a later study, Tanaka's group [18] reported for the first time the 1.70 Å resolution crystal structure of a natural Diels-Alderase, fungal macrophomate synthase, in complex with pyruvate. The authors also determined the active site of the enzyme as large and hydrophobic, with amino acid residues that can form hydrogen-bonds to the substrate 2-pyrone. Additionally, several artificial Diels-Alderases have been generated to catalyze different reactions [19-21]. However, to the best of our knowledge, no Another common dimerization way in the diterpenoid dimers is through a C–C linkage. Some studies [22,23] propose that this kind of linage is formed through Michael addition or aldol condensation. The formation of an ether bond and a dioxane ring is believed to occur through hemiacetal and acetal reactions. Despite many plausible biosynthetic pathways have been proposed, the nature and extent of enzymes involvement in the formation of diterpenoid dimers remain unclear. There has been no direct observation of enzymatic dimerization of the monomer units. The biosynthesis of diterpenoid dimers is worth further investigation as this promising progress has just started.

# Occurrence

Diterpenoid dimers are rare in nature, and they have been discovered in eighteen families of higher plants, fungi, liverworts and a gorgonian so far. The majority of diterpenoid dimers have been reported from higher plants including the families Acanthaceae (Andrographis), Annonaceae (Annona and Xylopia), Asteraceae (Baccharis), Calceolariaceae (Calceolaria), Chrysobalanaceae (Parinari) Cupressaceae (Calocedrus, Chamaecyparis, Cryptomeria, Cunninghamia, Juniperus, and Taiwania), Euphorbiaceae (Croton, Euphorbia, and Neoboutonia), Fabaceae (Caesalpinia, Cylicodiscus, and Erythrophleum), Lamiaceae (Ballota, Clerodendrum, Isodon, Plectranthus, Premna, Salvia, and Teucrium), Liliaceae (Fritillaria), Meliaceae (Aphanamixis), Rhizophoraceae (Ceriop), Taxaceae (Taxus and Torreya), Velloziaceae (Vellozia), and Zingiberaceae (Alpinia). They are widely distributed throughout the plants from roots, barks, stems, bulbs, leaves, seeds to fruits (O Table 1). Some diterpenoid dimers were found in fungi [Psathyrellaceae (Coprinus)], liverworts [Scapaniaceae (Scapania)] and a gorgonian [Gorgoniidae (Antillogorgia)].

# **Diterpenoid Dimers from Plants**

# **Family Acanthaceae**

As part of a search for cell differentiation inducers on mouse myeloid leukemia (M1), the aerial parts of *Andrographis paniculata* were chemically investigated and four labdanoid dimers, namely bisandrograpolides A–D (1–4) ( $\odot$  Fig. 1), were isolated [24]. All these compounds were deduced to be dimers linked via a C–C single bond between C-12 and C-15' of andrographolide derivatives but the respective configurations at C-12 and C-15' of these compounds remained undetermined. Bisandrograpolides A–C showed potent phagocytic and growth-inhibitory activities against M1 cells, while bisandrograpolide D showed no induction of phagocytosis, but exhibited growth-inhibitory effects.

# Family Annonaceae

Thirteen diterpenoid dimers have been isolated from plants of the family Annonaceae but only from the genera *Xylopia* and *Annona* (**• Fig. 2**).

Acutifloric acid (**5**) was isolated from the stem barks of *X. acutiflora* [25], and was identified as a dimer resulting from a Diels-Alder condensation between 15-oxo-kaur-16-en-19-oic and labda-8(17),13,15-trien-19-oic. Phytochemical studies of the green fruits of *X. amazonica* and stem barks and leaves of *X. frutescens* resulted in the identification of amazonins B and A (**6, 10**) and

# Table 1 Naturally occurring diterpenoid dimers<sup>a</sup>.

No.	Name	Molecular formula	Source	Part	Rei
Plants					
Acanth					
1	bisandrograpolide A	C <sub>40</sub> H <sub>56</sub> O <sub>8</sub>	Andrographis paniculata	aerial parts	[24
2	bisandrograpolide B	C <sub>40</sub> H <sub>56</sub> O <sub>8</sub>	Andrographis paniculata	aerial parts	[24
3	bisandrograpolide C	C <sub>40</sub> H <sub>56</sub> O <sub>8</sub>	Andrographis paniculata	aerial parts	[24
4	bisandrograpolide D	$C_{41}H_{60}O_{9}$	Andrographis paniculata	aerial parts	[24
Annona	aceae				
5	acutifloric acid	C <sub>40</sub> H <sub>60</sub> O <sub>3</sub>	Xylopia acutiflora	stem barks	[25
6	amazonin B	C <sub>41</sub> H <sub>60</sub> O <sub>5</sub>	Xylopia amazonica	fruits	[20
7	frutoic acid	$C_{40}H_{60}O_4$	Xylopia frutescens	stem barks	[2
8	emarginatine D	C <sub>42</sub> H <sub>62</sub> O <sub>5</sub>	Xylopia emarginata	stem barks	[2
9	emarginatine A	C <sub>42</sub> H <sub>62</sub> O <sub>5</sub>	Xylopia emarginata	branches	[2
0	amazonin A	C <sub>41</sub> H <sub>60</sub> O <sub>5</sub>	Xylopia amazonica	fruits	[2
11	emarginatine C	C <sub>42</sub> H <sub>62</sub> O <sub>5</sub>	Xylopia emarginata	stem barks	[2
2	emarginatine B	C <sub>42</sub> H <sub>62</sub> O <sub>5</sub>	Xylopia emarginata	branches	[2
3	ent-methylisoozate dimer	$C_{42}H_{64}O_4$	Xylopia aromatic	stem barks	[2
4	ent-13'-nor-13'-oxomethylisoozate dimer	C <sub>41</sub> H <sub>62</sub> O <sub>5</sub>	Xylopia aromatic	stem barks	[2
15	ent-13-epoximethylisoozate dimer	C <sub>42</sub> H <sub>64</sub> O <sub>5</sub>	Xylopia aromatic	stem barks	[2
16	annonebinide A	C <sub>40</sub> H <sub>64</sub> O <sub>3</sub>	Annona glabra	stem barks	[3
17	annoglabayin	C <sub>38</sub> H <sub>62</sub>	Annona glabra	fruits	[3
Asterac					
18	bacchalejin 1	$C_{43}H_{60}O_{6}$	Baccharis lejia	aerial parts	[3
19	bacchalejin 2	C <sub>43</sub> H <sub>60</sub> O <sub>7</sub>	Baccharis lejia	aerial parts	[3
20	bacchalejin 3	C <sub>45</sub> H <sub>62</sub> O <sub>8</sub>	Baccharis lejia	aerial parts	[3]
21	bacchalejin 4	C <sub>47</sub> H <sub>64</sub> O <sub>10</sub>	Baccharis lejia	aerial parts	[3
Calceol	ariaceae				
22	foliosate	C <sub>43</sub> H <sub>64</sub> O <sub>4</sub>	Calceolaria foliosa	aerial parts	[3
23	glandulosate	$C_{43}H_{64}O_4$	Calceolaria glandulosa	aerial parts	[3
24	lepidate	C <sub>43</sub> H <sub>66</sub> O <sub>5</sub>	Calceolaria lepida	aerial parts	[3
25	polifosate	C <sub>43</sub> H <sub>64</sub> O <sub>4</sub>	Calceolaria polifolia	aerial parts	[3
26	petiolate	$C_{43}H_{64}O_4$	Calceolaria petioalaris	aerial parts	[3
27	bis-[ <i>ent</i> -9-epi-labda-8(17),(12Z),14-trien-19-yl] malonate	C <sub>43</sub> H <sub>64</sub> O <sub>4</sub>	Calceolaria densifolia	aerial parts	[3
Chrysol	balanaceae				
28	15-oxozoapatlin-13α-yl-10'α,16'α-dihydroxy-9'α-methyl-20'- nor-kauran-19'-oic acid γ-lactone-17'-aote	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Parinari campestris	leaves	[3
Cupress	saceae				
29	6-(abieta-6',8',11',13'-tetraenyl-12'-oxy)-7-methoxyabieta- 8,11,13-trien-12-ol	C <sub>41</sub> H <sub>58</sub> O <sub>3</sub>	Chamaecyparis obtusa	barks	[4
30	sugikurojin B	C <sub>41</sub> H <sub>58</sub> O <sub>3</sub>	Cryptomeria japonica	heartwood	[4
31	formosadimer A	C <sub>41</sub> H <sub>58</sub> O <sub>3</sub>	Calocedrus macrolepis	barks	[4
32	formosadimer B	C46H68O4	Calocedrus macrolepis	barks	[4
33	formosadimer C	C <sub>48</sub> H <sub>70</sub> O <sub>5</sub>	Calocedrus macrolepis	barks	[4
34	calocedimer C	C <sub>40</sub> H <sub>56</sub> O <sub>3</sub>	Calocedrus macrolepis	barks	[4
85	calocedimer D	C44H60O5	Calocedrus macrolepis	barks	[4
36	formosaninol	C <sub>40</sub> H <sub>56</sub> O <sub>4</sub>	Juniperus formosana	heartwood	[4
37	formosanin	C <sub>42</sub> H <sub>60</sub> O <sub>4</sub>	Juniperus formosana	heartwood	[4
38	sugikurojin C	C <sub>40</sub> H <sub>56</sub> O <sub>4</sub>	Cryptomeria japonica	heartwood	[4
39	obtusanol A	C <sub>40</sub> H <sub>56</sub> O <sub>4</sub>	Chamaecyparis obtusa	heartwood	[4
40	obtusanol B	$C_{40}H_{54}O_5$	Chamaecyparis obtusa	heartwood	[4
11	bicunningine A	$C_{40}H_{50}O_4$	Cunninghamia lanceolata	barks	[4
12	bicunningine B	C <sub>40</sub> H <sub>52</sub> O <sub>4</sub>	Cunninghamia lanceolata	barks	[4
3	taiwaniadduct B	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides	leaves	[4
14	taiwaniadduct C	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides	leaves	[4
15	taiwaniadduct D	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides	leaves	[4
16	taiwaniadduct E	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides	leaves	[4
	taiwaniadduct F	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides	leaves	[4
47					
	taiwaniadduct G	C <sub>40</sub> H <sub>56</sub> O <sub>7</sub>	Taiwania crypomerioides	leaves	[4
47 48 49	taiwaniadduct G taiwaniadduct H	C <sub>40</sub> H <sub>56</sub> O <sub>7</sub> C <sub>39</sub> H <sub>56</sub> O <sub>6</sub>	Taiwania crypomerioides Taiwania crypomerioides	leaves leaves	[4] [4]

# Table 1 Continued

No.	Name	Molecular	Source	Part	Ref.
		formula			
Euphorl	piaceae				
51	crotoeurin A	C <sub>38</sub> H <sub>36</sub> O <sub>10</sub>	Croton euryphyllus	twigs and leaves	[49]
52	crotonkinensin C	C <sub>40</sub> H <sub>62</sub> O <sub>8</sub>	Croton tonkinensis	leaves	[50]
53	crotonkinensin D	C <sub>44</sub> H <sub>66</sub> O <sub>10</sub>	Croton tonkinensis	leaves	[50]
54	yuexiandajisu D	C <sub>38</sub> H <sub>48</sub> O <sub>4</sub>	Euphorbia ebracteolata	roots	[51]
55	langduin C	C <sub>40</sub> H <sub>50</sub> O <sub>10</sub>	Euphorbia fischeriana	roots	[52]
56	bisyinshanic acid A	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Euphorbia yinshanica	roots	[53]
57	bisyinshanic acid B	C <sub>40</sub> H <sub>58</sub> O <sub>5</sub>	Euphorbia yinshanica	roots	[53]
58	neoboutomannin	$C_{32}H_{26}O_6$	Neoboutonia mannii	barks	[54]
Fabacea		032.12000		build	[9.]
59	cyclodione	C <sub>40</sub> H <sub>56</sub> O <sub>4</sub>	Cylicodiscus gabunensis	barks	[55]
60	erythrophlesin A	C <sub>41</sub> H <sub>58</sub> O <sub>10</sub>	Erythrophleum succirubrum	leaves	[56]
61	erythrophlesin B	$C_{43}H_{60}O_{12}$	Erythrophleum succirubrum	leaves	[56]
62	erythrophlesin C	C <sub>43</sub> H <sub>63</sub> NO <sub>10</sub>	Erythrophleum succirubrum	leaves	[56]
63	erythrophlesin D	C <sub>45</sub> H <sub>65</sub> NO <sub>12</sub>	Erythrophleum succirubrum	leaves	[56]
64	erythrophlesin E	C <sub>44</sub> H <sub>63</sub> NO <sub>12</sub>	Erythrophleum fordii	leaves	[57]
65	erythrophlesin F	C <sub>42</sub> H <sub>61</sub> NO <sub>10</sub>	Erythrophleum fordii	leaves	[57]
66	erythrophlesin G	C <sub>44</sub> H <sub>63</sub> NO <sub>12</sub>	Erythrophleum fordii	leaves	[57]
67	erythrophlesin H	C <sub>45</sub> H <sub>65</sub> NO <sub>10</sub>	Erythrophleum fordii	barks	[58]
68	erythrophlesin I	C <sub>45</sub> H <sub>65</sub> NO <sub>11</sub>	Erythrophleum fordii	barks	[58]
69	caesanine D	$C_{42}H_{55}NO_7$	Caesalpinia sappan	seeds	[11]
Lamiace		C42H55NO7	Cuesulpinia suppan	seeds	[11]
70	persianone	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Ballota aucheri	aerial parts	[60]
70	inermes A	C <sub>52</sub> H <sub>74</sub> O <sub>19</sub>	Clerodendrum inerme	leaves	[61]
72	inermes B	C <sub>53</sub> H <sub>76</sub> O <sub>20</sub>	Clerodendrum inerme	leaves	[61]
72	trichotomone	C <sub>53</sub> H <sub>76</sub> O <sub>20</sub> C <sub>40</sub> H <sub>48</sub> O <sub>9</sub>	Clerodendrum trichotomum	roots	[10]
73	maoecrystal M		Isodon eriocalyx	leaves	[62]
74	bistenuifolin L	C <sub>48</sub> H <sub>64</sub> O <sub>16</sub>	,		[62]
	bistenuifolin M	C <sub>52</sub> H <sub>68</sub> O <sub>18</sub>	Isodon tenuifolius	aerial parts	
76		C <sub>50</sub> H <sub>66</sub> O <sub>16</sub>	Isodon tenuifolius	aerial parts	[64]
77	bisjaponin A	C <sub>40</sub> H <sub>52</sub> O <sub>12</sub>	Isodon japonicus	aerial parts	[65]
78	bisjaponin B	C <sub>40</sub> H <sub>54</sub> O <sub>12</sub>	Isodon japonicus	aerial parts	[65]
79	lushanrubescensin J	C <sub>40</sub> H <sub>52</sub> O <sub>12</sub>	Isodon rubescens	leaves	[66]
80	bisrubescensin C biexcisusin B	C <sub>40</sub> H <sub>56</sub> O <sub>12</sub>	Isodon rubescens Isodon excisus	leaves	[67]
81		C <sub>48</sub> H <sub>68</sub> O <sub>14</sub>		aerial parts	[68]
82	biexcisusin C	C <sub>48</sub> H <sub>68</sub> O <sub>16</sub>	Isodon excisus Isodon excisus	aerial parts	[68]
83 84	biexcisusin D biexcisusin E	C <sub>48</sub> H <sub>66</sub> O <sub>16</sub>		aerial parts	[68]
	bistenuifolin A	C <sub>48</sub> H <sub>66</sub> O <sub>16</sub>	Isodon excisus Isodon tenuifolius	aerial parts	[68]
85		C <sub>52</sub> H <sub>68</sub> O <sub>18</sub>	1	aerial parts	[64]
86	bistenuifolin B bistenuifolin C	C <sub>52</sub> H <sub>68</sub> O <sub>18</sub>	Isodon tenuifolius	aerial parts	[64]
87		C <sub>48</sub> H <sub>64</sub> O <sub>16</sub>	Isodon tenuifolius	aerial parts	[64]
88	bistenuifolin D	C <sub>50</sub> H <sub>66</sub> O <sub>16</sub>	Isodon tenuifolius	aerial parts	[64]
89	bistenuifolin E	C <sub>48</sub> H <sub>62</sub> O <sub>15</sub>	Isodon tenuifolius Isodon tenuifolius	aerial parts	[64]
90	bistenuifolin F xindongnin M	C <sub>48</sub> H <sub>68</sub> O <sub>16</sub>	Isodon rubescens	aerial parts	[64]
91	xindongnin N	C <sub>48</sub> H <sub>70</sub> O <sub>15</sub>	Isodon rubescens	leaves leaves	[69]
92	5	C <sub>48</sub> H <sub>68</sub> O <sub>15</sub>	Isodon rubescens		[69]
93	xindongnin O	C <sub>48</sub> H <sub>68</sub> O <sub>15</sub>	Isodon rubescens	leaves	[69]
94	bisrubescensin B	C <sub>40</sub> H <sub>58</sub> O <sub>13</sub>	Isodon excisus	leaves	[67]
95	biexcisusin A	C <sub>48</sub> H <sub>70</sub> O <sub>16</sub>		aerial parts	[68]
96	bispseurata F	C <sub>44</sub> H <sub>60</sub> O <sub>14</sub>	Isodon pharicus	aerial parts	[23]
97	bistenuifolin G	C <sub>48</sub> H <sub>66</sub> O <sub>17</sub>	Isodon tenuifolius	aerial parts	[64]
98	bistenuifolin H	C <sub>49</sub> H <sub>68</sub> O <sub>17</sub>	Isodon tenuifolius	aerial parts	[64]
99	bistenuifolin I	C <sub>49</sub> H <sub>68</sub> O <sub>17</sub>	Isodon tenuifolius	aerial parts	[64]
100	bistenuifolin J	C <sub>45</sub> H <sub>64</sub> O <sub>15</sub>	Isodon tenuifolius	aerial parts	[64]
101	bistenuifolin K	C <sub>47</sub> H <sub>66</sub> O <sub>16</sub>	Isodon tenuifolius	aerial parts	[64]
102	bisrubescensin A	C <sub>43</sub> H <sub>60</sub> O <sub>13</sub>	Isodon rubescens	leaves	[67]
103	rubescensin M	C <sub>40</sub> H <sub>58</sub> O <sub>9</sub>	Isodon rubescens	leaves	[70]
104	hebeiabinin E	C <sub>40</sub> H <sub>60</sub> O <sub>11</sub>	Isodon rubescens	leaves	[71]
105	hebeiabinin F	C <sub>40</sub> H <sub>56</sub> O <sub>9</sub>	Isodon rubescens	leaves	[71]
106	hispidanin A	C <sub>42</sub> H <sub>56</sub> O <sub>6</sub>	Isodon hispida	rhizomes	[72]
107	hispidanin B	C <sub>42</sub> H <sub>56</sub> O <sub>6</sub>	Isodon hispida	rhizomes	[72]
108	hispidanin C	C <sub>42</sub> H <sub>56</sub> O <sub>7</sub>	Isodon hispida	rhizomes	[72]
109	hispidanin D	C <sub>42</sub> H <sub>56</sub> O <sub>7</sub>	Isodon hispida	rhizomes	[72]
110	grandidone A	C <sub>40</sub> H <sub>48</sub> O <sub>9</sub>	Plectranthus grandidentatus	whole plants	[73]
					cont.

# Table 1 Continued

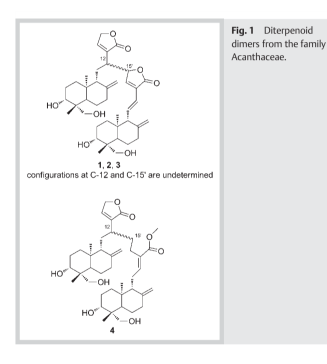
No.	Name	Molecular formula	Source	Part	Ref.
111	premnalatifolin A	C <sub>40</sub> H <sub>50</sub> O <sub>6</sub>	Premna latifolia	stem barks	[74]
112	obtusinone D	C <sub>40</sub> H <sub>52</sub> O <sub>6</sub>	Premna obtusifolia	roots	[75]
113	obtusinone E	C <sub>40</sub> H <sub>52</sub> O <sub>6</sub>	Premna obtusifolia	roots	[75]
114	hongencaotone	C <sub>40</sub> H <sub>50</sub> O <sub>5</sub>	Salvia prionitis	roots	[76]
115	bisprioterone A	C <sub>40</sub> H <sub>48</sub> O <sub>4</sub>	Salvia prionitis	roots	[77]
116	bisprioterone B	C <sub>39</sub> H <sub>46</sub> O <sub>5</sub>	Salvia prionitis	roots	[77]
117	bisprioterone C	C <sub>38</sub> H <sub>42</sub> O <sub>6</sub>	Salvia prionitis	roots	[77]
118	rosmanoyl carnosate	C <sub>40</sub> H <sub>48</sub> O <sub>7</sub>	Salvia canariensis	flowers	[78]
119	salviwardin A	C <sub>40</sub> H <sub>54</sub> O <sub>4</sub>	Salvia wardii	roots	[79]
120	salviwardin B	C <sub>40</sub> H <sub>52</sub> O <sub>4</sub>	Salvia wardii	roots	[79]
121	salvialeriafone	C <sub>39</sub> H <sub>48</sub> O <sub>7</sub>	Salvia leriaefolia	whole plants	[80]
122	salvialeriicone	C <sub>40</sub> H <sub>50</sub> O <sub>6</sub>	Salvia leriaefolia	whole plants	[83]
123	broussonetone A	C <sub>40</sub> H <sub>60</sub> O <sub>4</sub>	Salvia broussonetii	roots	[84]
124	broussonetone B	C <sub>40</sub> H <sub>60</sub> O <sub>4</sub>	Salvia broussonetii	roots	[84]
125	epirosmanol ester of 12-O-methyl carnosic acid	C <sub>41</sub> H <sub>54</sub> O <sub>8</sub>	Salvia officinalis	stems and leaves	[85]
126	-	C <sub>40</sub> H <sub>50</sub> O <sub>7</sub>	Salvia wagneriana	aerial parts	[86]
127	-	C <sub>40</sub> H <sub>42</sub> O <sub>9</sub>	Salvia wagneriana	aerial parts	[86]
128	biteuvisone A	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Teucrium viscidum	whole plants	[87]
129	biteuvisone B	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Teucrium viscidum	whole plants	[87]
Liliaceae		C401152O8	reachann visciadhn	whole plants	[07]
130	fritillebin A	C <sub>42</sub> H <sub>66</sub> O <sub>5</sub>	Fritillaria ebeiensis	bulbs	[88]
130	fritillebin B	C <sub>44</sub> H <sub>68</sub> O <sub>7</sub>	Fritillaria ebeiensis	bulbs	[88]
131	fritillebin C	C <sub>40</sub> H <sub>64</sub> O <sub>3</sub>	Fritillaria ebeiensis	bulbs	[88]
	fritillebin D				
133		C <sub>40</sub> H <sub>64</sub> O <sub>3</sub>	Fritillaria ebeiensis	bulbs	[89]
134	fritillebinide A	C <sub>40</sub> H <sub>64</sub> O <sub>2</sub>	Fritillaria ebeiensis	bulbs	[22]
135	fritillebinide B	C <sub>42</sub> H <sub>66</sub> O <sub>4</sub>	Fritillaria ebeiensis	bulbs	[90]
136	fritillebinide C	C <sub>42</sub> H <sub>66</sub> O <sub>4</sub>	Fritillaria ebeiensis	bulbs	[91]
137	fritillebinide D	C <sub>44</sub> H <sub>68</sub> O <sub>6</sub>	Fritillaria ebeiensis	bulbs	[92]
138	fritillebinide E	C <sub>44</sub> H <sub>68</sub> O <sub>6</sub>	Fritillaria ebeiensis	bulbs	[92]
Meliacea	le				
139	aphadilactone A	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[7]
140	aphadilactone B	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[7]
141	aphadilactone C	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[7]
142	aphadilactone D	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[7]
143	aphanamene C	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
144	aphanamene D	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
145	aphanamene E	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
146	aphanamene F	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
147	aphanamene K	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
148	aphanamene L	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
149	aphanamene M	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
150	aphanamene N	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
151	aphanamene G	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
152	aphanamene H	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
152	aphanamene I	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub> C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
	aphanamene				
154	i j	C <sub>40</sub> H <sub>54</sub> O <sub>8</sub>	Aphanamixis grandifolia Aphanamixis grandifolia	root barks	[8]
155	aphanamene O	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	1 5 1	root barks	[8]
156	aphanamene P	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[8]
157	aphanamene B	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	root barks	[95]
158	aphanamene A	C <sub>40</sub> H <sub>54</sub> O <sub>7</sub>	Aphanamixis grandifolia	root barks	[95]
159	aphadilactone I	C <sub>40</sub> H <sub>54</sub> O <sub>7</sub>	Aphanamixis grandifolia	leaves	[9]
160	aphadilactone E	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[9]
161	aphadilactone F	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[9]
162	aphadilactone G	C <sub>40</sub> H <sub>52</sub> O <sub>8</sub>	Aphanamixis grandifolia	leaves	[9]
Rhizopho	oraceae				
163	tagalsin I	C <sub>40</sub> H <sub>60</sub> O <sub>2</sub>	Ceriop tagal	stems and twigs	[96]
164	tagalsin J	C <sub>40</sub> H <sub>58</sub> O <sub>3</sub>	Ceriop tagal	stems and twigs	[96]
165	tagalsin L	C <sub>40</sub> H <sub>60</sub> O <sub>3</sub>	Ceriop tagal	roots	[97]
166	tagalsin M	C <sub>40</sub> H <sub>58</sub> O <sub>2</sub>	Ceriop tagal	roots	[97]
167	tagalsin N	C <sub>40</sub> H <sub>58</sub> O <sub>2</sub>	Ceriop tagal	roots	[97]
168	8(14)-enyl-pimar-2'(3')-en-4'(18')-en-15'(16')-en-dolabr-	C <sub>40</sub> H <sub>58</sub> O <sub>2</sub>	Ceriop tagal	roots	[98]
		10 30 - 2	,		1 -1

This document was downloaded for personal use only. Unauthorized distribution is strictly prohibited.

#### Table 1 Continued

No.	Name	Molecular formula	Source	Part	Ref.		
Taxaceae							
169	grandione	C <sub>40</sub> H <sub>56</sub> O <sub>6</sub>	Torreya grandis	stems	[99]		
170	diabietane ether	C <sub>41</sub> H <sub>60</sub> O <sub>5</sub>	Taxus cuspidata	needles	[100]		
Velloziaceae							
171	bismagdalenic acid	C <sub>40</sub> H <sub>60</sub> O <sub>4</sub>	Vellozia magdalenae	whole plants	[101]		
Zingiberaceae							
172	pahangensin C	C <sub>40</sub> H <sub>58</sub> O <sub>6</sub>	Alpinia pahangensis	rhizomes	[102]		
Fungi, liverworts and gorgonian							
Psathyre	ellaceae						
173	radianspene M	C <sub>40</sub> H <sub>50</sub> O <sub>6</sub>	Coprinus radians	fruiting body	[103]		
Scapaniaceae							
174	scapaundulin A	C <sub>32</sub> H <sub>48</sub> O <sub>6</sub>	Scapania undulata	whole plants	[104]		
175	scapaundulin B	C <sub>40</sub> H <sub>64</sub> O <sub>6</sub>	Scapania undulata	whole plants	[104]		
Gorgoniidae							
176	bisersolanolide	C <sub>40</sub> H <sub>48</sub> O <sub>10</sub>	Pseudopterogorgia bipinnata	whole animals	[105]		

<sup>a</sup>The diterpenoid dimers are grouped according to biological sources. For plants, families and genera are listed in alphabetical order.



frutoic acid (7), respectively [26,27]. In other studies, emarginatines A–D (**8**, **9**, **11**, **12**) were isolated from the branches and stem barks of *X*. *emarginata* [26,28]. These dimers are composed of kauranoid and labdanoid units, and could be considered as taxonomic markers of the genus *Xylopia*. Three diterpenoid dimers linked through a six-membered ring via a Diels-Alder condensation of two labdanoid units, namely *ent*-methylisoozate dimer (**13**), *ent*-13'-nor-13'-oxomethylisoozate dimer (**14**) and *ent*-13epoximethylisoozate dimer (**15**), were isolated from the stem barks of *X. aromatica* [29]. They were the first labdanoid dimers identified from the family Annonaceae.

Annonebinide A (**16**) was identified from the stems of *A. glabra* and determined to be a dimer with two *ent*-kauranoid units linked via an ester bond between C-17 and C-17' [30]. Annoglabayin (**17**), a kauranoid dimer, was isolated from *A. glabra* with its structure determined on the basis of spectroscopic analysis

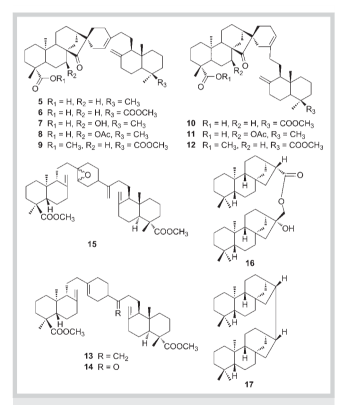


Fig. 2 Diterpenoid dimers from the family Annonaceae.

[31]. Annoglabayin has a unique C–C linkage between two nor*ent*-kauranoid units.

### Family Asteraceae

Four clerodanoid dimers linked via a C-18 malonate ester, bacchalejins 1–4 (**18–21**) (**• Fig. 3**), were reported from the aerial parts of *Baccharis lejia* [32]. These compounds represent the first and the only examples of diterpenoid dimers from the family Asteraceae.

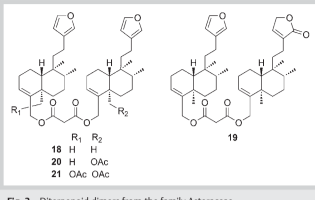


Fig. 3 Diterpenoid dimers from the family Asteraceae.

### Family Calceolariaceae

In the family Calceolariaceae, six diterpenoid dimers have been identified, all of which were from the genus *Calceolaria* (**• Fig. 4**). They are linked through C-17, C-18 or C-19 by a malonic acid unit. Three dimers composed of two isopimarane-type units, foliosate (**22**), glandulosate (**23**), and lepidate (**24**), were isolated from the aerial parts of *C. foliosa, C. glandulosa,* and *C. lepida,* respectively [33–35]. Additionally, polifosate (**25**) and petiolate (**26**) were identified from *C. polifolia* and *C. petioalaris,* respectively, which are composed of two pimarane-type units [36, 37]. Interestingly, bis-[*ent*-9-epi-labda-8(17),(12Z),14-trien-19-yl] malonate (**27**), a dimer composed of two labdane-type units, was also isolated from *C. densifolia* [38].

# Family Chrysobalanaceae

A kauranoid dimer, 15-oxozoapatlin-13 $\alpha$ -yl-10' $\alpha$ ,16' $\alpha$ -dihydroxy-9' $\alpha$ -methyl-20'-nor-kauran-19'-oic acid  $\gamma$ -lactone-17'aote (**28**) (**• Fig. 4**), was isolated from the leaves of *Parinari campestris* and identified on the basis of 2D NMR and ESI-MS [39]. It is the only diterpenoid dimer reported from the family Chrysobalanaceae.

# **Family Cupressaceae**

Plants from the family Cupressaceae are rich in abietane-type diterpenoids [3]. At present, twenty-two diterpenoid dimers have been identified from this family (**• Fig. 5**). Among them, fourteen compounds possess two abietane-type units, and eight others are composed of an abietane-type unit and a labdane-type unit. A dimer, 6-(abieta-6',8',11',13'-tetraenyl-12'-oxy)-7-methoxyabieta-8,11,13-trien-12-ol (29), was isolated from the stem barks of Chamaecyparis obtusa [40]. It is composed of two abietanoid units linked via an ether bridge between C-6 and C-12'. In a phytochemical investigation of the black heartwood of Cryptomeria japonica, sugikurojin B (30) was identified as a dimer of 6,7-dihydroxyferruginol and 6,7-dehydroferruginol with a 6-0-12' linkage [41]. In chemical studies of the barks of Calocedrus macrolepis, five abietanoid dimers with the same linkage pattern as sugikurojin B were isolated, including formosadimers A-C (31-33) [42] and calocedimers C and D (34, 35) [43]. Formosaninol (36) and formosanin (37) were isolated from the heartwood of Juniperus formosana [44]. The structure of formosaninol was deduced to be a dimeric ferruginol with 6-0-7' and 7-0-6' linkages on the basis of spectroscopic analysis and chemical evidences. Formosanin was a dimethyl ether of formosaninol. Sugikurojin C (38) was a dimeric ferruginol with the same planar structure as formosa-

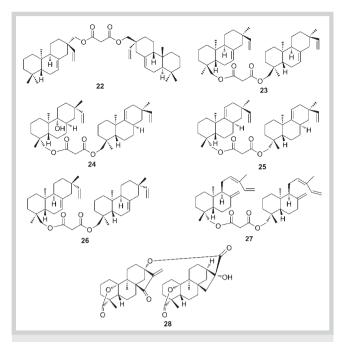


Fig. 4 Diterpenoid dimers from the families Calceolariaceae and Chrysobalanaceae.

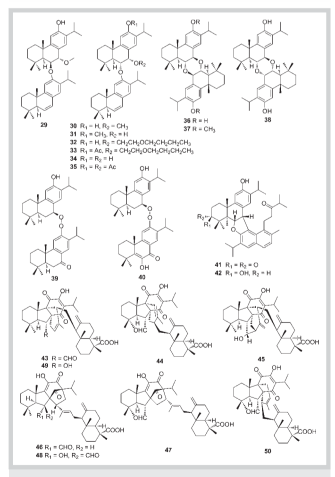


Fig. 5 Diterpenoid dimers from the family Cupressaceae.

ninol, which was isolated from the black heartwood of *Cryptomeria japonica* [41]. The only difference between the two compounds is the configuration at C-7'. Two diterpenoid dimers, namely obtusanols A and B (**39, 40**), were isolated from the heartwood of *Chamaecyparis obtusa*, and characterized by spectroscopic means and chemical degradation [45]. Obtusanols A and B have a rare peroxide bond linking two abietanoid units between C-7 and C-12'. In another study, bicunningines A and B (**41, 42**) were isolated from *Cunninghamia lanceolata* [46]. Their structures were elucidated by spectroscopic measurements. Their absolute configurations were determined by quantum chemical TDDFT (time-dependent density functional theory) ECD calculations, chemical transformations and Mosher's method. They were the first diterpenoid dimers reported to contain a 2,3-dihydrofuran ring fusing an abietanoid and a 4,5-seco-abietanoid.

Taiwaniadducts B–J (**43–50**) were isolated from the leaves of *Taiwania crypomerioides* and identified as dimers composed of an abietanoid unit and a labdanoid unit [47,48]. Taiwaniadducts B, C, H and I were presumably derived from a [4+2] Diels-Alder reaction of the labdane trans-ozic acid and different taiwaniaquinones; while taiwaniadducts E–G appeared to result from the corresponding [5+2] Diels-Alder reaction. Taiwaniadducts B–J are the only naturally occurring heterodimers formed by abietanoid and labdanoid units, and could be considered as taxonomic markers of the species *T. crypomerioides*. They have attracted strong synthetic interest due to their structural complexity.

### Family Euphorbiaceae

Plants from the family Euphorbiaceae are rich in sesquiterpenoids and diterpenoids. At present, eight diterpenoid dimers have been isolated from this family (**○** Fig. 6).

In a chemical study of the twigs and leaves of Croton euryphyllus, a nor-clerodanoid dimer, namely crotoeurin A (51), was isolated, which contains a unique cyclobutane ring formed via [2+2] cycloaddition [49]. The structure was elucidated by spectroscopic analysis and the configuration was confirmed by single crystal X-ray diffraction. Crotoeurin A represents the first nor-clerodanoid dimer with a cyclobutane ring and is of particular significance for the biosynthesis of clerodane-type diterpenoids. Crotoeurin A exhibited neurite outgrowth-promoting activity on nerve growth factor-mediated PC12 cells at a concentration of 10 µM. During a screening program for cytotoxic compounds, two symmetric ent-kauranoid dimers with connectivity at C-17, namely crotonkinensins C and D (52, 53), were isolated from the leaves of the Vietnamese endemic medicinal plant C. tonkinensis [50]. ent-Kauranoid diterpenoids are rarely found from the genus Croton, and these two compounds are the first examples from this genus. Crotonkinensin D showed potent cytotoxic activity against MCF-7, tamoxifen-resistant MCF-7 and adriamycin-resistant MCF-7 breast cancer cell lines, with IC50 values of  $9.4\pm1.7,\,2.6\pm0.9,\,and\,18.9\pm0.6\,\mu M$  , respectively.

In another study, yuexiandajisu D (**54**), an 18-nor-rosane-type diterpenoid dimer, was isolated from the roots of *Euphorbia ebracteolata* [51]. It is the first and to date the only example of 18-norrosane-type diterpenoid dimer isolated from the family Euphorbiaceae. Yuexiandajisu D showed weak cytotoxic activity against HCT-8 and Bel-7402 cancer cell lines, with  $IC_{50}$  values of 2.66 and 3.76 mM, respectively. Langduin C (**55**) was isolated from the roots of *E. fischeriana* and its structure was established by spectroscopic data and single crystal X-ray diffraction analysis [52]. Langduin C is a symmetrical diterpenoid dimer with a five-membered C ring instead of the normal six-membered C ring found in the *ent*-abietane-type diterpenoids. It is the first diterpenoid

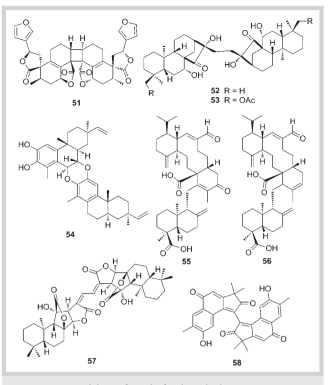


Fig. 6 Diterpenoid dimers from the family Euphorbiaceae.

dimer from the genus *Euphobia*. This dimer is probably derived from jolkinolide B, a major *ent*-abietane-type diterpenoid of this plant, by successive oxidative cleavage of ring C and D, rearrangement, lactonization, and dimerization. Two diterpenoid dimers with a bismagdalenic acid skeleton, namely bisyinshanic acids A and B (**56**, **57**), were isolated from the roots of *E. yinshanica* [53]. Their structures were elucidated on the basis of spectroscopic evidences.

Neoboutomannin (**58**), a degraded diterpenoid dimer, was isolated from the stem barks of *Neoboutonia mannii* [54]. Neoboutomannin was active against *Enterococcus faecalis, Staphylococcus aureus, Proteus mirabilis,* and three *Candida* species, *C. albicans, C. tropicalis* and *C. parapsilosis.* 

#### **Family Fabaceae**

At present, eleven diterpenoid dimers with a cassane-type skeleton have been identified from plants of the family Fabaceae (**• Fig. 7**).

A chemical investigation of the stem barks of *Cylicodiscus gabunensis* resulted in the identification of the cassanoid dimer cyclodione (**59**) [55]. Cyclodione was proposed to be formed through a [4+2] Diels-Alder reaction between two cassanoid units.

Nine diterpenoid dimers were isolated from plants of the genus *Erythrophleum*. They possess an unsymmetrical dimeric structure with two cassanoid units linked through an ester bond at C-16 and C-3' [56–58]. These compounds could be considered as taxonomic markers of this genus. TRAIL is a promising agent for new anticancer therapy as it can induce apoptosis in a variety of cancer cells but not in normal cells [59]. Bioassay-guided fractionation of the extract of *E. succirubrum* for TRAIL resistance-overcoming activity led to the isolation of four cassanoid dimers, namely erythrophlesins A–D (**60–63**) [56]. These four compounds are the first examples of cassanoid dimers linked via an ester

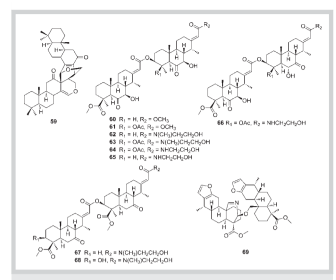


Fig. 7 Diterpenoid dimers from the family Fabaceae.

bond, which are rarely found in nature. Moreover, erythrophlesins C and D possess an amide moiety, which is seldom found in naturally occurring diterpenoids. Erythrophlesins A-C exhibited significant TRAIL resistance-overcoming activity in human gastric adenocarcinoma cells. A detailed phytochemical investigation of the leaves of E. fordii resulted in the isolation of three cassanoid dimers, namely erythrophlesins E-G (64-66), all of which were found to contain an amide group [57]. Their structures were determined by extensive 1D and 2D NMR analyses and ESIMS. Cytotoxic activity of these compounds was evaluated against HCT-8, Bel-7402, BGC-823, A549, and A2780 human cancer cell lines in an MTT assay. All three compounds exhibited significant cytotoxic activity (IC<sub>50</sub> < 10 µM) against these cells. Cytotoxic activity-guided fractionation of E. fordii led to the isolation of two cassanoid amide dimers, namely erythrophlesins H and I (67, 68) [58]. An MTT assay confirmed that erythrophlesin H had significant cytotoxic effect toward the human prostate cancer PC-3 cell line, with an IC<sub>50</sub> value of  $12.5 \,\mu$ M.

In a recent study, a cassanoid dimer, namely caesanine D (**69**), was isolated from the seeds of *Caesalpinia sappan* [11]. Caesanine D represents the first example of a cassanoid dimer where the subunits are linked via an ether bond. Interestingly, one of the diterpene units of this compound possesses a cassane-type skeleton with an unusual N bridge between C-19/C-20. The structure was determined by various spectroscopic methods and ECD calculation.

# Family Lamiaceae

Plants of the family Lamiaceae contributed a significant number of diterpenoid dimers. At present, 60 diterpenoid dimers have been isolated and structurally characterized. Most are homodimers, composed of two diterpenoids units with the same core skeleton. Twenty-nine kaurane-type diterpenoid dimers were reported from the genus Isodon, and could be considered as taxonomic markers of this genus; sixteen abeitane-type diterpenoid dimers were identified from the genera *Salvia, Clerodendrum, Plectranthus*, and *Teucrium*; four clerodane-type diterpenoid dimers were found from the genera *Salvia* and *Clerodendrum*; three icetexane-type diterpenoid dimers were from the genus *Premna*; and one labdane-type dimer was from the genus *Ballota*.

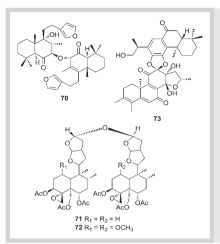


Fig. 8 Diterpenoid dimers from the genera *Ballota* and *Cleroden-drum* (Lamiaceae).

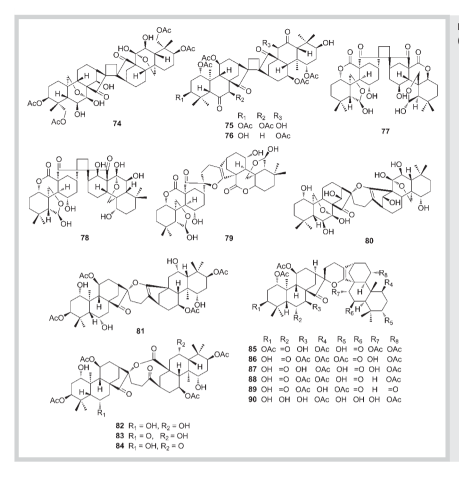
Besides, some are rare heterodimers, consisting of two units belonging to different types of diterpenoids. Four dimers composed of a totarene-type unit and a labdane-type unit, as well as three dimers containing an abietane-type unit and a kaurane-type unit were isolated from the genus Isodon.

**Ballota** genus. A study on the chemical constituents of *B. aucheri* led to the isolation of persianone (**70**) (**• Fig. 8**), a dimer composed of two labdane-type units linked through an ether bond at C-7 [60]. The structure of persianone was elucidated by high field NMR spectroscopy, including NOE difference experiments, and chemical transformations.

Clerodendrum genus. Two compounds, namely inermes A and B (**71, 72**) (**•** Fig. 8), were isolated from *C. inerme* [61]. On the basis of comprehensive spectroscopic analysis, both compounds were elucidated to contain two clerodane units linked through an ether bridge at C-15. Interestingly, a hexahydrofurofuran ring was found in each clerodane unit. The isolation and structural elucidation of trichotomone (73) (OFig. 8) was reported from the roots of the medicinal ornamental plant C. trichotomum [10]. This compound is a rare phenolic ketal derivative consisting of a regular abietanoid and a rearranged abietanoid derivative in a 17  $(15 \rightarrow 16), 18(4 \rightarrow 3)$ -diabeo-abietane framework. The structure was elucidated by extensive spectroscopic methods. The absolute configuration was defined by comparison of experimental and calculated ECD spectra. Trichotomone exhibited significant in vitro cytotoxicity against several human cancer cell lines (A549, Jurkat, BGC-823, and 293 T WT) with IC<sub>50</sub> values ranging from 7.51 to 19.38 µM.

**Isodon genus.** Besides lots of diterpenoid monomers, the genus *Isodon* is also a major source of diterpenoid dimers with a considerable structural diversity. At present, 36 dimers have been isolated from this genus, most of which possess a kaurane-type core skeleton (**• Fig. 9–11**).

Maoecrystal M (**74**) (**•** Fig. 9), the first example of naturally occurring *ent*-kaurane-type dimer, was isolated from *I. eriocalyx* [62]. By means of <sup>1</sup>H-<sup>1</sup>H COSY and ROESY, as well as chemical transformation, the structure of maoecrystal M was determined to be a symmetric dimer of an *ent*-kaurane diterpenoid connected at 16R, 16'R through a cyclobutane ring. The four-membered ring was proposed to be formed by condensation between the olefinic bond in the  $\alpha$ , $\beta$ -unsaturated ketone group of the monomer diterpenoid, probably through a [2+2] cycloaddition [63]. In a chemical study of *I. tenuifolius*, bistenuifolins L and M (**75** and **76**) (**•** Fig. 9) were isolated and found to possess the



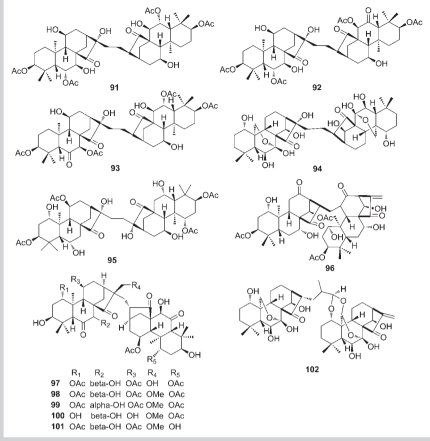


same skeleton as maoecrystal M [64]. In 2008, another two *ent*-kaurane-type dimers connected with a four-membered carbon ring, namely bisjaponins A and B (**77** and **78**) (**• Fig. 9**), were isolated from the aerial parts of *I. japonicus* [65]. These two compounds contain a 6,7-seco-6,20-epoxy-*ent*-kaurane fragment.

An asymmetric ent-kauranoid dimer, namely lushanrubescensin J (79) (O Fig. 9), was isolated from I. rubescens var. lushanensis [66]. Its structure was established by spectroscopic evidences and single crystal X-ray diffraction. It is the first ent-kauranoid dimer found to possess a dihydropyran ring resulting from a [4+2] cycloaddition between the  $\alpha,\beta$ -unsaturated ketone group of one diterpenoid and the olefinic bond of another diterpenoid. Interestingly, this compound contains a 6,7-seco-6,20-epoxyent-kaurane monomer. Lushanrubescensin J exhibited potent inhibitory activity against K562 cells with an IC<sub>50</sub> value of  $0.93 \,\mu g/$ mL. Bisrubescensin C (80) ( Fig. 9), an *ent*-kauranoid dimer with the same linkage pattern as lushanrubescensin J, was isolated from I. rubescens [67]. Four ent-kauranoid dimers, namely biexcisusins B–E (81–84) (O Fig. 9), were reported from *I. excisus* [68]. Their structures which are closely related to bisrubescensin C were established on the basis of detailed spectroscopic analyses. Biexcisusins C-E possess an unprecedented linkage through a nine-membered lactone ring between two *ent*-kaurane-type subunits. The lactone ring was proposed to arise through oxidative cleavage of the double bond of the dihydropyran ring in biexcisusin B. In a chemical study of I. tenuifolius, six ent-kauranoid dimers, namely bistenuifolins A-F (85–90) (O Fig. 9), were identified and found to be linked by a dihydropyran ring [64]. The structures of these compounds were established via spectroscopic analysis. The absolute configurations of bistenuifolins A and D were defined by single crystal X-ray diffraction. Bistenuifolin B exhibited significant cytotoxicity against several human cancer cell lines, including HL-60, SMMC-7721, MCF-7, and SW-480, with  $IC_{50}$  values ranging from 4.0 to 9.9  $\mu$ M.

Three asymmetric dimers, namely xindongnins M-O (91-93) (**• Fig. 10**), have been isolated from *I. rubescens* var. *rubescens* [69]. They represent the first examples of ent-kauranoid dimers with a rare linkage through a single C–C bond between two units. Their structures were characterized by spectroscopic methods including 2DNMR analyses. The relative configuration of xindongnin M was determined by single crystal X-ray diffraction. ent-Kauranoids isolated from the genus Isodon normally have  $\alpha,\beta$ -unsaturated ketone groups. The [4+2] cycloaddition between the  $\alpha,\beta$ -unsaturated ketone of one diterpene unit and the olefinic bond of the second unit might yield a dihydropyran ring. In a further step, hydrolysis and rearrangement at the dihydropyran ring could produce the single C-C bond linkage. Two other ent-kauranoid dimers, namely bisrubescensin B (94) and biexcisusin A (95) ( Fig. 10), connected with a single C–C bond linkage, were isolated from I. rubescens [67] and I. excisus [68], respectively. The co-occurrence of dimers with a single C-C bond linkage (bisrubescensin B and biexcisusin A) and congeners with a dihydropyran ring (bisrubescensin C and biexcisusin B) in the same plant further supports the above proposed biosynthetic pathway. A phytochemical investigation of I. pharicus led to the isolation of an asymmetric dimer, namely bispseurata F (96) (**•** Fig. 10), which is the first and the only example of *ent*-kauranoid dimer connected by direct linkage of C-17 with C-11' [23]. A Michael addition reaction is proposed to be the key step in the biosynthesis of bispseurata F. The dimerization of this type of di-





This document was downloaded for personal use only. Unauthorized distribution is strictly prohibited.

terpenoid dimers is worth further studies. Five *ent*-kauranoid dimers linked by a unique C-16 to C-17' single bond, namely bistenuifolins G–K (**97–101**) (**• Fig. 10**), were identified from *I. tenuifolius* [64]. Bisrubescensin A (**102**) (**Fig. 10**) is an *ent*-kauranoid dimer from *I. rubescens* and contains an unprecedented  $C_{23}$  *ent*-kaurane unit [67].

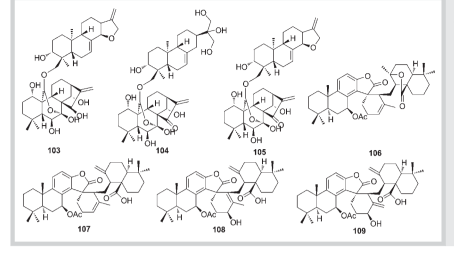
Rubescensin M (**103**) (**• Fig. 11**) was isolated from *I. rubescens* [70]. By detailed spectroscopic analysis, it was deduced to be a dimer linked by an oxygen bridge between C-18 of an abietanoid and C-20' of a kauranoid. Abietane-type diterpenoids are very rare in the genus *Isodon*, and rubescensin M is the first heterodimer from this genus. In a chemical study of *I. rubescens*, hebeiabinins E and F (**104, 105**) (**• Fig. 11**) were identified with the same linkage pattern as rubescensin M [71]. Hebeiabinin F showed significant inhibitory activity against A549 and HT-29 cells with  $IC_{50}$  values of 0.91 and 1.81 µM, respectively.

Hispidanins A–D (**106–109**) (**• Fig. 11**) are four unprecedented heterodimers formed by the bonding of totarane-type and labdane-type diterpenoids. They were obtained from the rhizomes of *I. hispida* [72]. Their structures were elucidated by extensive spectroscopic analyses, and the structure of hispidanin A was further confirmed by single crystal X-ray diffraction. Totarane-type diterpenoids are rarely found in nature. Hispidanins A–D are the first and the only naturally occurring heterodimers composed of a labdane-type and a totarane-type diterpenoid. The biosynthetic pathway of hispidanins A–D was proposed to involve an intermolecular Diels–Alder reaction between totarane-type and labdane-type derivatives. Hispidanin B showed significant cytotoxicity against tumor cell lines SGC7901, SMMC7721, and K562, with IC<sub>50</sub> values of 10.7, 9.8, and 13.7  $\mu$ M, respectively. **Plectranthus genus.** An abietanoid dimer linked by a ketal, namely grandidone A (**110**) (**•** Fig. 12), was isolated from *P. grandidenta-tus* [73]. This compound showed slight antiproliferative activity against five human cancer cell lines MCF-7, NCI-H460, SF-268, TK-10, and UACC-62, with Gl<sub>50</sub> values of  $9.6 \pm 1.8$ ,  $19.2 \pm 3.1$ ,  $25.8 \pm 4.0$ ,  $40.9 \pm 3.7$ , and  $35.7 \pm 1.5 \mu$ M, respectively.

Premna genus. Premnalatifolin A (111) (O Fig. 12), a unique icetexanoid dimer, was isolated from the stem barks of the Indian medicinal plant P. latifolia [74]. Its structure and relative configuration were elucidated on the basis of detailed spectroscopic analyses, including HRESIMS and 2D NMR spectra. This compound is composed of two icetexanoid units linked via an ether bridge. The formation of premnalatifolin A was proposed to follow a radical reaction. A phenoxyl radical of one subunit reacted with a phenyl radical of the other subunit to result in the ether bridge. Premnalatifolin A displayed potent cytotoxicity against HT-29 and MCF-7 cell lines with  $\ensuremath{\text{IC}_{50}}$  values of 12.15 and 1.11 µg/mL, respectively. In 2013, two icetexanoid dimers, namely obtusinones D and E (112 and 113) (O Fig. 12), were isolated from the roots of *P. obtusifolia*, and were suggested to be formed via a hetero-Diels-Alder type dimerization reaction [75]. Obtusinone D represents the first example of a linearly fused icetexanoid dimer, whereas obtusinone E is an angularly fused icetexanoid dimer. The structures of obtusinones D and E were elucidated on the basis of 1D and 2D NMR spectroscopic analyses. Icetexanoid dimers could be considered as taxonomic markers of the genus Premna.

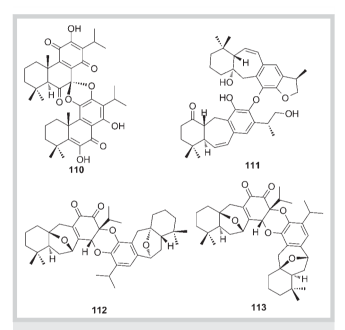
*Salvia* genus. Fourteen diterpenoid dimers have been isolated from the genus *Salvia*, including twelve abietane-type dimers and two clerodane-type dimers (**• Fig. 13**). Abietanoid dimers

**Fig. 11** Diterpenoid dimers from the genus *Isodon* (Lamiaceae) – part III.



from the genus *Salvia* are linked via C–C single bond, ether bridge, dioxane ring, or ketal moiety.

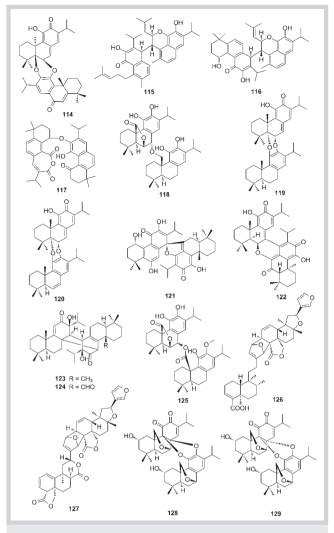
The abietanoid dimer hongencaotone (114) was isolated from the roots of S. prionitis and its structure was determined by spectroscopic data interpretation and X-ray analysis [76]. From the same species, three further abietanoid dimers, namely bisprioterones A-C (115-117), were identified by Zhang and his colleagues [77]. Bisprioterone A possesses two 4,5-seco abietanoid subunits linked via a C-C single bond at C-14 and C-1'. In bisprioterone B the subunits are connected via a C-C single bond between C-14 of an abietanoid subunit and C-1' of a 4,5-seco abietanoid subunit. Bisprioterone C possesses an ether bridge linked between C-12 of an abietanoid subunit and C-1' of an 11,12-seco abietanoid subunit. Their structures were characterized by analysis of 1D and 2D NMR spectroscopic data. The structure of bisprioterone A was further confirmed by single crystal X-ray diffraction. In contrast to their monomers these diterpenoid dimers did not exhibit obvious cytostatic, antiphlogistic, or antibacterial activities. The disappearance of some functional groups during the dimerization process might account for the decrease of the bioactivities. In 1987, rosmanoyl carnosate (118), a dimer composed of two abietanoid units linked via an ether bond between C-7 and C-20', was isolated from the flowers of S. canariensis [78]. It was the first ether-linked diterpenoid dimer identified from the genus Salvia. Two other dimers, namely salviwardins A and B (119 and 120), were isolated from the roots of S. wardii [79]. In both compounds, two abietanoid subunits are connected via a dioxane ring. Salvialeriafone (121), a diterpene-norditerpene conjugate, was isolated from S. leriaefolia and its structure was determined by spectroscopic data analysis [80]. Salvialeriafone which contains a spiro-dihydrofuran moiety attached to ring C of the norditerpenoid unit is the first example of norditerpene-diterpene conjugated abietanoid dimer. The probable origin of the spiro-dihydrofuran group is proposed to be through a nucleophilic addition/substitution between the 1,6,12-trihydroxy derivative of sibiriquinone B [81] and the 6-deoxo analogue of 14-hydroxytaxodion [82]. This compound exhibited antiproliferative activity against HeLa cells with an IC50 value of 10.91 µM. Salvialeriicone (122), isolated from S. leriifolia, is an abietanoid dimer connected via a dihydropyran ring [83]. The structure was determined using mass spectrometry and NMR spectroscopy. In a chemical study of S. broussonetii, two abietanoid dimers, namely broussonetones A and B (123, 124), were isolated [84]. Their structures were deter-



**Fig. 12** Diterpenoid dimers from the genera *Plectranthus* and *Premna* (Lamiaceae).

mined based on spectroscopic data and confirmed by X-ray analysis. These dimers could be formed by a [4+2] cycloaddition of two molecules of 13 $\beta$ -hydroxyabieta-8(14),9(11)-dien-12-one. Broussonetones A and B are the first non-phenolic or quinonic abietanoid dimers to be isolated from natural sources. In a study aimed to the identification of nuclear peroxisome proliferatoractivated receptor (PPAR)- $\gamma$  activators from *S. officinalis*, the epirosmanol ester of 12-O-methyl carnosic acid (**125**), was identified. This compound contains two abietanoid subunits linked by an ester bond [85]. As the only example of abietanoid dimer resulting from the formation of an ester bond, it was considered as an artefact formed during extraction and isolation. This was further supported by the fact that this compound was not detectable in the crude extract by HPLC analysis.

From the aerial parts of *S. wagneriana*, two clerodanoid dimers (**126** and **127**) were obtained with their structures established by 1D- and 2D-NMR spectroscopic analyses [86]. They are the on-



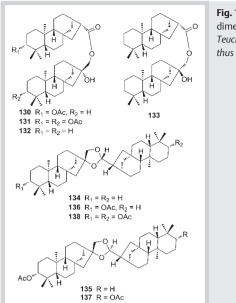
**Fig. 13** Diterpenoid dimers from the genera *Salvia* and *Teucrium* (Lamiaceae).

ly clerodane-type diterpenoid dimers reported from the genus Salvia.

**Teucrium genus.** A pair of dimeric abietanoid stereoisomers connected via a dioxane ring, namely biteuvisones A and B (**128** and **129**) (**• Fig. 13**), were isolated from *T. viscidum* [87]. These two compounds are proposed to be formed through a hetero-Diels-Alder reaction of the o-quinone of teuvisone.

#### **Family Liliaceae**

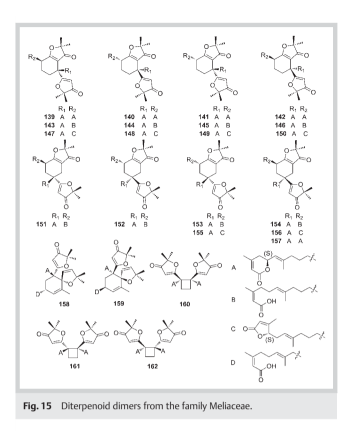
Nine diperpenoid dimers were isolated from the bulbs of *Fritillaria ebeiensis* (Liliaceae). They contain two *ent*-kauranoid units linked through an ester bond or a dioxolane ring (**• Fig. 14**). In 1995, Wu and colleagues [88] found two compounds, namely fritillebins A and B (**130, 131**), which possess an *ent*-kauranoid dimer skeleton linked via an ester bond between C-17 and C-17'. These compounds are the first diterpenoid dimers identified from the family Liliaceae. Later, the same group [89] reported other two dimers, namely fritillebins C and D (**132, 133**), from the same plant. These two dimers share the same core skeleton as fritillebin A. An acetal diterpenoid dimer with *ent*-kauranoid skeleton, namely fritillebinide A (**134**), was isolated from the bulbs of *F. ebeiensis* [22]. The structure of fritillebinide A was elu-



cidated by spectroscopic analysis and chemical synthesis. It represents the first *ent*-kauranoid dimer possessing a dioxolane ring formed by aldol condensation. In subsequent studies, two pairs of further *ent*-kauranoid dimers containing a dioxolane ring, namely fritillebinides B and C (**135, 136**) and fritillebinides D and E (**137, 138**), were isolated from the same plant [90–92]. Fritillebinides B and D have a R configuration at C-17' while fritillebinides C and E have a S configuration at this position.

#### **Family Meliaceae**

Diterpenoids are not widely found in plants of the family Meliaceae. In fact, Aphanamixis grandifolia is the only source of diterpenoid dimers in this family, which has contributed 25 congeners in recent years (**> Fig. 15**). A. grandifolia is an arbor tree mainly distributed in the tropical and subtropical areas of Asia [93]. Its leaves and roots are used as folk medicine in China to treat rheumatism and alleviate pain [94]. As part of a search for new DGAT inhibitors, the ethanolic extract of A. grandifolia was found to exhibit significant inhibition against DGAT-1. Bioassay-guided isolation resulted in identification of four diastereoisomers possessing an unprecedented carbon skeleton, namely aphadilactones A–D (139–142) [7]. Their structures and absolute configurations were determined by a combination of spectroscopic data, chemical degradation, partial synthesis, experimental CD spectra and ECD calculations. Aphadilactones A-D were proposed to be formed from two molecules of nemoralisin-type diterpenoid through an enzyme-catalyzed [4+2] cycloaddition reaction, which leads to a cyclohexene ring with a 2,2-dimethylfuran-3 (2 H)-one ring and the substituents attached at the para-position [7]. According to further biological evaluation, aphadilactone C is a potent DGAT-1 inhibitor (IC<sub>50</sub> =  $0.46 \pm 0.09 \,\mu$ M) with marginal activity against DGAT-2 (IC<sub>50</sub> > 100  $\mu$ M). In addition, these compounds have weak antimalarial activity with IC<sub>50</sub> values ranging from 120 to 190 µM. In a later study, eight diterpenoid dimers with the same skeleton as aphadilactone A, namely aphanamenes C-F and K-M (143-150), were isolated from the root barks of A. grandifolia [8]. In this study, other six diterpenoid dimers with a 2,2-dimethylfuran-3(2H)-one ring and two substituents attached at the meso-position, namely aphanamenes G-J, O and P



(151-156), were identified [8]. The structures of these compounds were elucidated by spectroscopic analysis, and their absolute configurations were determined using the CD exciton chirality method. As shown in a further study, these compounds exhibited significant inhibition of LPS-induced NO production in RAW264.7 macrophages, with IC<sub>50</sub> values ranging from 7.75 to 19.31 µM. The isolation and structural elucidation of aphanamene B (157) was reported as part of an investigation of A. grandifolia [95]. This compound shares the same skeleton with aphanamene G. Aphanamene A (158) was also reported in this study and found to possess a spiro 2,2-dimethyl dihydroxyfuran ring on the cyclohexene ring [95]. It was proposed to be formed through a different [4+2] cycloaddition reaction. Both structures were elucidated by spectroscopic analysis, and the absolute configuration of aphanamene A was determined by ECD calculations. These two compounds inhibited LPS-induced NO production in RAW264.7 cells with IC<sub>50</sub> values of 9.72 and 7.98 µM, respectively. Recently, a chemical investigation into the minor constituents of A. grandifolia yielded one diterpenoid dimer, namely aphadilactone I (159), which was found to be a diastereoisomer of aphanamene A [9]. Besides, three diastereomeric diterpenoid dimers, namely aphadilactones E-G (160-162), were also isolated from this species and found to contain a new carbon skeleton incorporating a 1,1,2,2-tetrasubstituted cyclobutane moiety. Their structures and absolute configurations were fully established by comprehensive spectroscopic data analysis and ECD calculations. It was proposed that aphadilactones E–G were formed through a [2+2] cycloaddition reaction in a head-to-head and tail-to-tail way. Aphadilactones E and F exhibited remarkable antimalarial activity with IC<sub>50</sub> values of  $1.03 \pm 0.13$  and  $2.86 \pm 0.47 \mu$ M, respectively. These dimers could be considered as taxonomic markers of the species A. grandifolia.

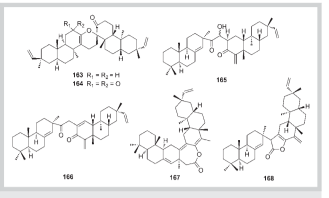


Fig. 16 Diterpenoid dimers from the family Rhizophoraceae.

## Family Rhizophoraceae

The occurrence of diterpenoid dimers in the family Rhizophoraceae was only reported from the mangrove plant Ceriop tagal. At present, six dolabrane-type dimers have been isolated (O Fig. 16). By means of extensive spectroscopic analysis and single crystal X-ray diffraction, two dolabrane-type dimers, namely tagalsins I (163) and J (164), were identified. They represent the first examples of diterpenoid dimers from the family Rhizophoraceae [96]. In later studies, four dimers, namely tagalsins L-N (165-167) and 8(14)-enyl-pimar-2'(3')-en-4'(18')-en-15'(16')endolabr-16,15,2',3'-oxoan-16-one (168), were isolated from the roots of C. tagal [97,98]. 8(14)-enyl-pimar-2'(3')-en-4'(18')-en-15'(16')-endolabr-16,15,2',3'-oxoan-16-one exhibited antifouling activity against cyprid larvae (Balanus albicostatus) of the barnacle without significant toxicity. Dolabrane-type dimers could be considered as taxonomic markers of the species C. tagal. The stem barks of Xylopia acutiflora yielded a dimeric diterpene derived via Diels-Alder condensation of kaurene and labdane monomers. The structure of the dimer, which has been given the trivial name acutifloric acid, was assigned on the basis of detailed spectroscopic analysis.

### **Family Taxaceae**

The isolation and structure elucidation of grandione (**169**) (**• Fig. 17**) was reported in the course of an investigation of Chinese specimens of *Torreya grandis* [99]. Grandione is formed by two icetexanoid units linked via a hetero-Diels-Alder dimerization reaction and shares the same skeleton as obtusinone D (**112**). Grandione represents the first and the only example of a linearly fused icetexanoid dimer from the family Taxaceae. Diabietane ether (**170**) (**• Fig. 17**), an abietanoid dimer connected by an ether linkage, was isolated from the needles of *Taxus cuspidata* [100].

#### Family Velloziaceae

An unusual bis-diterpenoid diacid, bismagdalenic acid (**171**) (**• Fig. 17**), was isolated from the hexane extract of the Brazilian plant *Vellozia magdalenae* [101]. Bismagdalenic acid is a dimer formed via a Diels-Alder condensation of magdalenic acid and a regular labdane diterpenoid, cis-ozic acid. This is the first report of the isolation of diterpenoid dimer from the family Velloziaceae.

# Family Zingiberaceae

The rhizomes of Alpinia pahangensis yielded the labdanoid dimer pahangensin C (**172**) (**• Fig. 17**) [102]. This dimer is formed via an ester bond between C-15 and C-15'. The structure of pahangensin C was elucidated by spectroscopic methods including 1D and 2D NMR and LCMS-IT (ion trap)-TOF analyses. It is the only diterpenoid dimer reported from the family Zingiberaceae.

Diterpenoid Dimers from Fungi, Liverworts and Gorgonian

V

A few diterpenoid dimers have been reported from sources other than plants, including fungi, liverworts and a gorgonian. These groups of organisms could be potential sources of novel diterpenoid dimers with promising biological activities and are worth further investigation in the future.

# Fungi (family Psathyrellaceae)

Radianspene M (**173**) (**• Fig. 18**), a guanacastane-type diterpenoid dimer, was isolated from a fermentation of the M65 strain of the higher fungus *Coprinus radians* [103]. This is the first report of a diterpenoid dimer from fungi and provides new opportunities to investigate the dimerization mechanisms of diterpenoids as fungi are much easier to be manipulated in the laboratory through cultivation than plants.

# Liverworts (family Scapaniaceae)

Two labdanoid dimers, namely scapaundulins A and B (**174** and **175**) (**• Fig. 18**), were isolated from the diethyl ether extract of the Japanese liverworts *Scapania undulata* [104]. Their structures were characterized by spectroscopic techniques, especially 2D NMR and mass spectrometry. Two identical labdanoid units are connected via ester linkages in scapaundulin A, or hemiacetal linkages in scapaundulin B, from C-8 of one subunit to C-11 of the other subunit. The structures of scapaundulins possess a C<sub>2</sub> axis of symmetry.

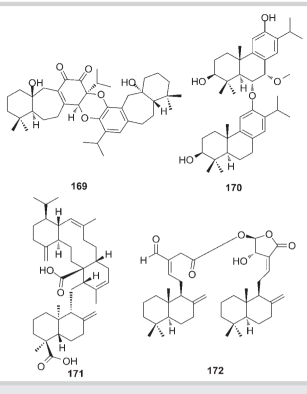
# Gorgonian (family Gorgoniidae)

A chemical study of the hexane extract of the Caribbean gorgonian *Antillogorgia bipinnata* collected in San Andre's Island, Colombia, led to the isolation of an unprecedented heptacyclic diterpenoid dimer, namely bisersolanolide (**176**) (**• Fig. 18**) [105]. The structure of this secondary metabolite was established by spectroscopic studies including 2D NMR, IR, UV, and accurate mass measurements, and was further confirmed by synthesis. Bisersolanolide is the first diterpenoid dimer found to contain two cembranoid units. The generation of the 2,3-dihydro-4H-pyran ring is proposed to occur via a Diels-Alder coupling of two units of gersolane diterpenoids.

# **Synthesis**

#### ▼

Due to their high structural diversity and their biological activities, diterpenoids have attracted remarkable attention from a synthetic perspective. In contrast, only few successful total syntheses of diterpenoid dimers have been reported.



**Fig. 17** Diterpenoid dimers from the families Taxaceae, Velloziaceae and Zingiberaceae.

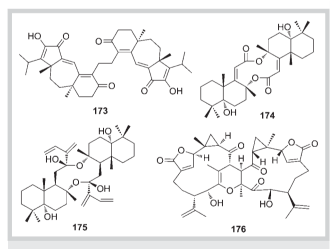


Fig. 18 Diterpenoid dimers from the families Psathyrellaceae, Scapaniaceae and Gorgoniidae.

# Synthesis of grandione

Grandione (**169**) is a unique icetexanoid dimer. In 2005, Kurihara's group [106] first reported the partial synthesis of grandione from demethylsalvicanol via the solid state hetero-Diels-Alder type dimerization reaction. Three years later, Majetich's group [107] reported the total synthesis of (+)-grandione from benzyl bromide (**177**) and 6,6-dimethyl-1,3-cyclohexadione (**178**). The authors developed a two-step process to convert the achiral enone **179** into the 5*S* alkene **180** (**• Fig. 19**). Next, the authors took advantage of Kelecom's approach to convert alkene **180** into alcohol **181** [108]. The epoxidation of the C-1, C-10-trisubstituted double bond occurred from the  $\beta$ -face of **180**, and the subsequent opening of this epoxide with LAH introduced a  $\beta$ -oriented tertiary alcohol at C-10 (**• Fig. 19**). To avoid the solid state hetero-Diels-Alder reaction, the authors carried out the cycloaddition in water at 50 °C overnight, which produced (+)-grandione in good yield (**• Fig. 19**).

#### Synthesis of aphadilactones

Shortly after the isolation of aphadilactones A-D (139-142), a study on the total synthesis of these diterpenoid dimers was reported [13]. A proposed biosynthetic pathway of aphadilactones was put forward by Yue's group, in which the S-dienelactone 184 (**•** Fig. 20) served as a common biosynthetic precursor to these dimers [7]. The diastereomeric aphadilactones A-D were obtained in comparable amounts from the natural source, strongly suggesting that the final [4+2] dimerization was a non-enzymatically catalyzed process. The total syntheses of aphadilactones A-D were accomplished in eleven steps starting from the commercially available 1-methoxy-3-methylbuta-1,2-diene (182) and but-2-y-nal (183) (O Fig. 20). The S-dienelactone 184 reacted with BHT (butylated hydroxytoluene) in toluene at 170° C for 17 hours to form aphadilactones A–D (approx. 1:1:1:1) through the bioinspired [4+2] dimerization/1,3  $\sigma$ -hydrogen migration (**© Fig. 20**).

#### Synthesis of taiwaniadduts

A few members of taiwaniaquinoids, namely taiwaniadducts A-J, possess a characteristic Diels-Alder cycloadduct scaffold. In 2014, Li's group [109] carried out the first total synthesis of taiwaniadducts B-D, which took advantage of an Iridium-catalyzed asymmetric polyene cyclization in the synthesis of the two key fragments, namely taiwaniaquinone F (185) and methyl trans-ozitate (186). Then, the dimerization reaction was carried out with Er (fod)<sub>3</sub> under neat conditions and elevated temperature to produce the cycloadduct 187 (52% yield) and its regioisomer 188 (21% yield) but no other positional or diastereomeric isomers (**© Fig. 21**). The site selectivity toward the C-8 olefin over the C-12 olefin may be attributable to the bulky isopropyl and the electron-donating methoxyl that make the latter olefin a worse dienophile. The facial selectivity may arise from the steric effect of the axial C-20 methyl group. Both cycloadducts were subjected to a three-step sequence of oxidation to furnish taiwaniadducts B and C. Me<sub>2</sub>AlCl-mediated carbonyl-ene reaction formed taiwaniadduct D (91% yield) (**© Fig. 21**).

# Conclusions

▼

As illustrated in this review, naturally occurring diterpenoid dimers have become an important research area in the field of natural products. There have been around 90 publications focusing on chemistry of diterpenoid dimers during the period covered by this review (1981 to January 2016). Up-to-date, 176 diterpenoid dimers have been described, most of which are from higher plants. As shown in **• Table 1**, the family Lamiaceae contributes the greatest number of diterpenoid dimers (60 compounds), and the families Meliaceae (23 compounds) and Cupressaceae (22 compounds) have also afforded numerous compounds. In contrast, only a few examples of diterpenoid dimers have been isolated from fungi, liverworts or marine animals, which might be due to the limited availability of these natural re-

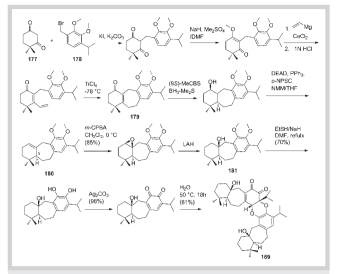


Fig. 19 Majetich's synthesis of (+)-grandione.

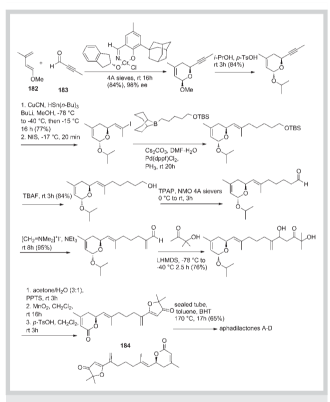
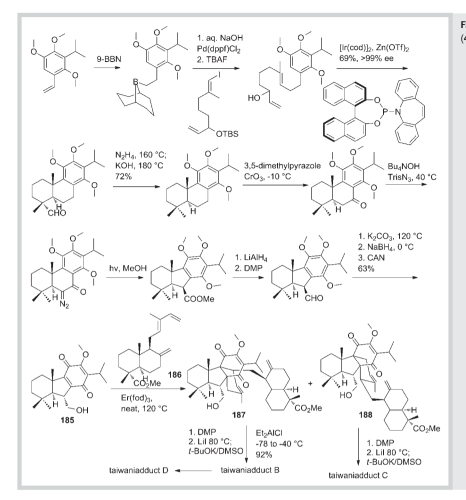


Fig. 20 Nan's synthesis of aphadilactones A–D (139–142).

sources. These organisms are believed to be promising materials for identifying novel diterpenoid dimers and eventually developing lead compounds.

There is a great structural diversity of diterpenoid dimers. As highlighted in this review, diterpenoid dimers can be classified into one of the following skeletons: kaurane-type, abietane-type, nemorallisin-type, labdane-type, clerodane-type, cassane-type, dolabrane-type, pimarane-type, icetexane-type, guanacastanetype, cembrane-type, rosane-type, or a combination of two from the above structural types. With 44 compounds the kaurane-type



# Fig. 21 Li's synthesis of taiwaniadducts B–D (43–45).

dimers contribute the greatest proportion. Most of the diterpenoid dimers are homodimers formed by two units of the same skeleton. Only 23 compounds are heterodimers containing two units of different skeletons. Three dimers composed of abietanetype and kaurane-type units, and four dimers composed of totarane-type and labdane-type units have been identified from plants of the family Lamiaceae. Besides, eight dimers with abietane-type and labdane-type units, and eight compounds with kaurane-type and labdane-type units have been found from plants of the families Cupressaceae and Annonaceae, respectively.

The linkages of diterpenoid dimers include single C–C bonds, ether bonds, ester bonds and ring moieties. Enzyme-catalyzed Diels-Alder cycloaddition reaction is proposed to be a major mechanism involved in the synthesis of diterpenoid dimers. In addition enzyme-mediated Michael addition or aldol condensation is also proposed to form C–C linkages in diterpenoid dimers. At present, there is no direct evidence confirming the proposed biosynthetic pathways of diterpenoid dimers and the putative natural Diels-Alderase still remains unknown. Increased efforts should be made in the future to elucidate the key enzymes and individual steps of the biosynthetic pathway of diterpenoid dimers.

Diterpenoid dimers have been reported with various bioactivities, including cytotoxic, anti-inflammatory, anti-microbial, anti-malarial, and anti-fouling effects. However, all these studies were carried out in in vitro assays. No *in vivo* animal studies or clinical trials have been conducted to evaluate the therapeutic effects of diterpenoid dimers. Moreover, due to low amounts, most diterpenoid dimers have never been biologically tested. Further investigations should be performed in the future.

The studies summarized in this review confirm the potential of diterpenoid dimers for the discovery of novel pharmaceutical agents. It is hoped that chemists, pharmacologists and biologists will intensify research efforts on these complex secondary metabolites as a potential source of novel bioactive lead compounds.

# Acknowledgments

# ▼

Financial support by the Research Fund of University of Macau (MYRG2014-00020-ICMS-QRCM and MYRG2015-00153-ICMS-QRCM) and the Science and Technology Development Fund of Macau (120/2013/A3) is gratefully acknowledged.

# **Conflict of Interest**

The authors declare no competing financial interest.

# References

- 1 Hanson JR. Diterpenoids of terrestrial origin. Nat Prod Rep 2015; 32: 1654-1663
- 2 Vasas A, Hohmann J. Euphorbia diterpenes: isolation, structure, biological activity, and synthesis (2008–2012). Chem Rev 2014; 114: 8579– 8612

- 3 *Gonzalez MA*. Aromatic abietane diterpenoids: their biological activity and synthesis. Nat Prod Rep 2015; 32: 684–704
- 4 Cragg GM. Paclitaxel (Taxol): a success story with valuable lessons for natural product drug discovery and development. Med Res Rev 1998; 18: 315–331
- 5 Siebert DJ. Salvia divinorum and salvinorin A: new pharmacologic findings. J Ethnopharmacol 1994; 43: 53–56
- 6 Roth BL, Baner K, Westkaemper R, Siebert D, Rice KC, Steinberg S, Ernsberger P, Rothman RB. Salvinorin A: a potent naturally occurring nonnitrogenous kappa opioid selective agonist. Proc Natl Acad Sci U S A 2002; 99: 11934–11939
- 7 *Liu J, He XF, Wang GH, Merino EF, Yang SP, Zhu RX, Gan LS, Zhang H, Cassera MB, Wang HY, Kingston DG, Yue JM*. Aphadilactones A–D, four diterpenoid dimers with DGAT inhibitory and antimalarial activities from a Meliaceae plant. J Org Chem 2014; 79: 599–607
- 8 Zhang HJ, Zhang YM, Luo JG, Luo J, Kong LY. Anti-inflammatory diterpene dimers from the root barks of Aphanamixis grandifolia. Org Biomol Chem 2015; 13: 7452–7458
- 9 Zhang H, Liu J, Gan LS, Dalal S, Cassera MB, Yue JM. Antimalarial diterpenoid dimers of a new carbon skeleton from Aphanamixis grandifolia. Org Biomol Chem 2016; 14: 957–962
- 10 Wang WX, Zhu JJ, Zou YK, Hong ZL, Liu ST, Li M, Huang Y, Xiong J, Zhao Y, Yang GX, Xia G, Hu JF. Trichotomone, a new cytotoxic dimeric abietanederived diterpene from Clerodendrum trichotomum. Tetrahedron Lett 2013; 54: 2549–2552
- 11 Zhang JY, Abdel-Mageed WM, Liu MM, Huang P, He WN, Li L, Song FH, Dai HQ, Liu XT, Liang JY, Zhang LX. Caesanines A–D, new cassane diterpenes with unprecedented N bridge from *Caesalpinia sappan*. Org Lett 2013; 15: 4726–4729
- 12 Zi J, Mafu S, Peters RJ. To gibberellins and beyond! Surveying the evolution of (di)terpenoid metabolism. Annu Rev Plant Biol 2014; 65: 259– 286
- 13 Yin JP, Gu M, Li Y, Nan FJ. Total synthesis of aphadilactones A–D. J Org Chem 2014; 79: 6294–6301
- 14 *Oikawa H, Tokiwano T.* Enzymatic catalysis of the Diels-Alder reaction in the biosynthesis of natural products. Nat Prod Rep 2004; 21: 321– 352
- 15 Oikawa H, Katayama K, Suzuki Y, Ichihara A. Enzymatic-activity catalyzing *exo*-selective Diels-Alder reaction in solanapyrone biosynthesis. Chem Commun 1995; 1321–1322
- 16 Katayama K, Kobayashi T, Oikawa H, Honma M, Ichihara A. Enzymatic activity and partial purification of solanapyrone synthase: first enzyme catalyzing Diels-Alder reaction. Biochim Biophys Acta 1998; 1384: 387–395
- 17 Auclair K, Sutherland A, Kennedy J, Witter DJ, Van den Heever JP, Hutchinson CR, Vederas JC. Lovastatin nonaketide synthase catalyzes an intramolecular Diels-Alder reaction of a substrate analogue. J Am Chem Soc 2000; 122: 11519–11520
- 18 Ose T, Watanabe K, Mie T, Honma M, Watanabe H, Yao M, Oikawa H, Tanaka I. Insight into a natural Diels-Alder reaction from the structure of macrophomate synthase. Nature 2003; 422: 185–189
- 19 Xu JA, Deng QL, Chen JG, Houk KN, Bartek J, Hilvert D, Wilson IA. Evolution of shape complementarity and catalytic efficiency from a primordial antibody template. Science 1999; 286: 2345–2348
- 20 Kim SP, Leach AG, Houk KN. The origins of noncovalent catalysis of intermolecular Diels-Alder reactions by cyclodextrins, self-assembling capsules, antibodies, and RNAses. J Org Chem 2002; 67: 4250–4260
- 21 Siegel JB, Zanghellini A, Lovick HM, Kiss G, Lambert AR, Clair JLS, Gallaher JL, Hilvert D, Gelb MH, Stoddard BL, Houk KN, Michael FE, Baker D. Computational design of an enzyme catalyst for a stereoselective bimolecular Diels-Alder reaction. Science 2010; 329: 309–313
- 22 Wu JZ, Ruan HL, Yao NH, Sun HD, Zhao QS, Morizane C, Iida A, Fujita T. Structural elucidation and synthesis of fritillebinide A from bulbs of Fritillaria ebeiensis. Acta Pharm Sin 1999; 34: 600–604
- 23 Zhao Y, Huang SX, Xiao WL, Ding LS, Pu JX, Li X, Yang LB, Sun HD. Diterpenoids from Isodon pharicus. Tetrahedron Lett 2009; 50: 2019–2023
- 24 Matsuda T, Kuroyanagi M, Sugiyama S, Umehara K, Ueno A, Nishi K. Cell differentiation-inducing diterpenes from Andrographis paniculata Nees. Chem Pharm Bull (Tokyo) 1994; 42: 1216–1225
- 25 Hasan CM, Healey TM, Waterman PG. Chemical studies in the Annonaceae .13. acutifloric acid-a diterpene dimer from the stem bark of Xylopia acutiflora. Phytochemistry 1985; 24: 192–194
- 26 Vilegas W, Felicio JD, Roque NF, Gottlieb HE. Diterpenic adducts from Xylopia Species. Phytochemistry 1991; 30: 1869–1872

- 27 Takahashi JA, Boaventura MAD, Bayma JD, Oliveira AB. Frutoic acid, a dimeric kaurane diterpene from Xylopia frutescens. Phytochemistry 1995; 40: 607–609
- 28 Moreira IC, Roque NF, Lago JHG. Diterpene adducts from branches of Xylopia emarginata. Biochem Syst Ecol 2006; 34: 833–837
- 29 Martins D, Hamerski L, Alvarenga SAV, Roque NF. Labdane dimers from Xylopia aromatica. Phytochemistry 1999; 51: 813–817
- 30 Yang NY, Tian LJ, Meng ZM, Han Y. A new diterpenoid dimer from Annona glabra. Chin Chem Lett 2003; 14: 58–61
- 31 Chen CH, Hsieh TJ, Liu TZ, Chern CL, Hsieh PY, Chen CY. Annoglabayin, a novel dimeric kaurane diterpenoid, and apoptosis in Hep G2 cells of annomontacin from the fruits of Annona glabra. J Nat Prod 2004; 67: 1942–1946
- 32 Labbe C, Castillo M, Hernandez M. Diterpenoids from Baccharis lejia. Phytochemistry 1991; 30: 1607–1611
- 33 Chamy MC, Piovano M, Garbarino JA, Gambaro V, Miranda C. Diterpenoids from Calceolaria species 3. foliosate, a bis-diterpene and 9-epient-7,15-isopimaradiene derivatives from Calceolaria foliosa. Phytochemistry 1989; 28: 571–574
- 34 *Piovano M, Chamy MC, Garbarino JA, Gambaro V.* Diterpenoids from Calceolaria species 4.9-epi-*ent*-7,15-isopimaradiene derivatives from *Calceolaria glandulosa*. Phytochemistry 1989; 28: 2844–2845
- 35 Chamy MC, Piovano M, Garbarino JA, Miranda C, Gambaro V. Diterpenoids from Calceolaria species 5. diterpenes from Calceolaria lepida. Phytochemistry 1990; 29: 2943–2946
- 36 Chamy MC, Piovano M, Garbarino JA, Gambaro V. Diterpenoids from Calceolaria species 10. diterpenes from Calceolaria polifolia. Phytochemistry 1991; 30: 3365–3368
- 37 Silva P, Chamy MC, Piovano M, Garbarino JA. Diterpenoids from Calceolaria petioalaris 14. diterpenoids from Calceolaria species. Phytochemistry 1993; 34: 449–451
- 38 Garbarino JA, Molinari A. Diterpenoids from Calceolaria species 9. labdane diterpenes from Calceolaria densifolia. J Nat Prod 1992; 55: 744– 747
- 39 Braca A, Abdel-Razik AF, Mendez J, Morelli I. A new kaurane diterpene dimer from Parinari campestris. Fitoterapia 2005; 76: 614–619
- 40 Hanari N, Yamamoto H, Ooi T, Kusumi T, Kuroda K. A new diterpene dimer from the bark of Chamaecyparis obtusa. J Wood Sci 2001; 47: 36–40
- 41 Arihara S, Umeyama A, Bando S, Imoto S, Ono M, Tani M, Yoshikawa K. A new abietane and two dimeric abietane diterpenes from the black heartwood of *Cryptomeria japonica*. Chem Pharm Bull (Tokyo) 2004; 52: 354–358
- 42 Hsieh CL, Tseng MH, Kuo YH. Formosadimers A, B, and C from the bark of Calocedrus macrolepis var. formosana. Chem Pharm Bull (Tokyo) 2005; 53: 1463–1465
- 43 Hsieh CL, Shiu LL, Tseng MH, Shao YY, Kuo YH. Calocedimers A, B, C, and D from the bark of *Calocedrus macrolepis* var. *formosana*. J Nat Prod 2006; 69: 665–667
- 44 *Kuo YH, Yu MT.* Diterpenes from the heartwood of *Juniperus formosana* Hay var *concolor* Hay. Chem Pharm Bull (Tokyo) 1996; 44: 1431–1435
- 45 *Kuo YH, Chen CH, Wein YS.* New dimeric monoterpenes and dimeric diterpenes from the heartwood of *Chamaecyparis obtusa* var. *formosana*. Helv Chim Acta 2002; 85: 2657–2663
- 46 Hou XF, Yao S, Mandi A, Kurtan T, Tang CP, Ke CQ, Li XQ, Ye Y. Bicunningines A and B, two new dimeric diterpenes from Cunninghamia lanceolata. Org Lett 2012; 14: 460–463
- 47 *Lin WH, Fang JM, Cheng YS.* Diterpenes and related cycloadducts from *Taiwania cryptomerioides.* Phytochemistry 1996; 42: 1657–1663
- 48 Lin WH, Fang JM, Cheng YS. Cycloadducts of terpene quinones from Taiwania cryptomerioides. Phytochemistry 1997; 46: 169–173
- 49 Pan ZH, Ning DS, Wu XD, Huang SS, Li DP, Lv SH. New clerodane diterpenoids from the twigs and leaves of Croton euryphyllus. Bioorg Med Chem Lett 2015; 25: 1329–1332
- 50 Thuong PT, Thi HMP, Thi VTL, Dao TT, Dang TT, Nguyen QT, Oh WK. Symmetric dimers of *ent*-kaurane diterpenoids with cytotoxic activity from *Croton tonkinensis*. Bioorg Med Chem Lett 2012; 22: 1122–1124
- 51 *Fu GM, Qin HL, Yu SS, Yu BY.* Yuexiandajisu D, a novel 18-nor-rosanetype dimeric diterpenoid from *Euphorbia ebracteolata* Hayata. J Asian Nat Prod Res 2006; 8: 29–34
- 52 Zhou TX, Bao GH, Ma QG, Qin GW, Che CT, Lv Y, Wang C, Zheng QT. Langduin C, a novel dimeric diterpenoid from the roots of *Euphorbia fischeriana*. Tetrahedron Lett 2003; 44: 135–137

- 53 Zhang BY, Wang H, Luo XD, Du ZZ, Shen JW, Wu HF, Zhang XF. Bisyinshanic acids A and B, two novel diterpene dimers from the toots of Euphorbia yinshanica. Helv Chim Acta 2012; 95: 1672-1679
- 54 Tene M, Tane P, Tamokou JD, Kuiate JR, Connolly JD. Degraded diterpenoids from the stem bark of Neoboutonia mannii. Phytochemistry Lett 2008: 1: 120-124
- 55 Tane P, Bergquist KE, Tene M, Ngadjui BT, Ayafor JF, Sterner O. Cyclodione, an unsymmetrical dimeric diterpene from Cylicodiscus gabunensis. Tetrahedron 1995; 51: 11595-11600
- 56 Miyagawa T, Ohtsuki T, Koyano T, Kowithayakorn T, Ishibashi M. Cassaine diterpenoid dimers isolated from Erythrophleum succirubrum with TRAIL-resistance overcoming activity. Tetrahedron Lett 2009; 50: 4658-4662
- 57 Du D, Qu J, Wang JM, Yu SS, Chen XG, Xu S, Ma SG, Li Y, Ding GZ, Fang L. Cytotoxic cassaine diterpenoid-diterpenoid amide dimers and diterpenoid amides from the leaves of Erythrophleum fordii. Phytochemistry 2010:71:1749-1755
- 58 Hung TM, Cuong TD, Kim JA, Lee JH, Woo MH, Min BS. In vitro apoptotic effect of cassaine-type diterpene amides from Erythrophleum fordii on PC-3 prostate cancer cells. Bioorg Med Chem Lett 2014; 24: 4989-4994
- 59 Rustaiyan A, Mossleminkupaii MH, Papastergiou F, Jakupovic J. Persianone, a dimeric diterpene from Ballota aucheri. Phytochemistry 1995; 40:875-879
- 60 Pandey R, Verma RK, Gupta MM. Neo-clerodane diterpenoids from Clerodendrum inerme. Phytochemistry 2005; 66: 643-648
- 61 Shen XY, Isogai A, Furihata K, Sun HD, Suzuki A. Maoecrystal M-a naturally-occurring symmetrical ent-kaurane dimer from Rabdosia eriocalyx. Phytochemistry 1994; 35: 725-729
- 62 Yang JH, Wang WG, Du X, He F, Zhang HB, Li XN, Li Y, Pu JX, Sun HD. Heterodimeric ent-kauranoids from Isodon tenuifolius. | Nat Prod 2014; 77: 2444-2453
- 63 Yang LB, Yang J, Li LM, Lei C, Zhao Y, Huang SX, Xiao WL, Han QB, Pu JX, Sun HD. Symmetric and asymmetric ent-kaurane dimers isolated from Isodon japonicus. Tetrahedron Lett 2008; 49: 3574-3577
- 64 Han QB, Lu Y, Wu L, He ZD, Qiao CF, Xu HX, Zheng QT, Sun HD. An asymmetric ent-kauranoid dimer from Isodon rubescens var. lushanensis. Tetrahedron Lett 2005; 46: 5373-5375
- 65 Huang SX, Xiao WL, Li LM, Li SH, Zhou Y, Ding LS, Lou LG, Sun HD. Bisrubescensins A-C: three new dimeric ent-kauranoids isolated from Isodon rubescens. Org Lett 2006; 8: 1157-1160
- 66 Hong SS, Lee SA, Lee C, Han XH, Choe S, Kim N, Lee D, Lee CK, Kim Y, Hong [T, Lee MK, Hwang BY. Dimeric ent-kaurane diterpenoids from Isodon excisus. J Nat Prod 2011; 74: 2382-2387
- 67 Han QB, Lu Y, Zhang LL, Zheng QT, Sun HD. Novel ent-kaurane dimers from Isodon rubescens var. rubescens. Tetrahedron Lett 2004; 45: 2833-2837
- 68 Han QB, Li RT, Zhang JX, Sun HD. New ent-abietanoids from Isodon rubescens. Helv Chim Acta 2004; 87: 1007-1015
- 69 Huang SX, Pu JX, Xiao WL, Li LM, Weng ZY, Zhou Y, Han QB, Peng SL, Ding LS, Lou LG, Sun HD. ent-Abietane diterpenoids from Isodon rubescens var. rubescens. Phytochemistry 2007; 68: 616-622
- 70 Huang B, Xiao CJ, Huang ZY, Tian XY, Cheng X, Dong X, Jiang B. Hispidanins A-D: four new asymmetric dimeric diterpenoids from the rhizomes of Isodon hispida. Org Lett 2014; 16: 3552-3555
- 71 Marques CG, Pedro M, Simoes MF, Nascimento MS, Pinto MM, Rodriguez B. Effect of abietane diterpenes from Plectranthus grandidentatus on the growth of human cancer cell lines. Planta Med 2002; 68: 839-840
- 72 Suresh G, Babu KS, Rao MSA, Rao VRS, Yadav PA, Nayak VL, Ramakrishna S. Premnalatifolin A, a novel dimeric diterpene from Premna latifolia Roxb. Tetrahedron Lett 2011; 52: 5016-5019
- 73 Salae AW, Boonnak N. Obtusinones D and E, linear and angular fused dimeric icetexane diterpenoids from Premna obtusifolia roots. Tetrahedron Lett 2013; 54: 1356-1359
- 74 Li M, Zhang JS, Chen MQ. A novel dimeric diterpene from Salvia prionitis. J Nat Prod 2001; 64: 971-972
- 75 Xu J, Chang J, Zhao M, Zhang JS. Abietane diterpenoid dimers from the roots of Salvia prionitis. Phytochemistry 2006; 67: 795-799
- 76 Gonzalez AG, Rodriguez CM, Luis JG. Diterpenes from the flowers of Salvia canariensis. Phytochemistry 1987; 26: 1471-1474
- Xiao QL, Xia F, Yang XW, Liao Y, Yang LX, Wei YK, Li X, Xu G. New dimeric and seco-abietane diterpenoids from Salvia wardii. Nat Prod Bioprospect, advance online publication 8 April 2015; DOI: 10.1007/s13659-015-0054-6

- 78 Choudhary MI, Hussain A, Ali Z, Adhikari A, Sattar SA, Ayatollahi SA, Al-Majid AM. Diterpenoids including a novel dimeric conjugate from Salvia leriaefolia. Planta Med 2012; 78: 269-275
- 79 Hussain A, Adhikari A, Choudhary MI, Ayatollahi SA, Atta-ur-Rahman. New adduct of abietane-type diterpene from Salvia leriifolia Benth. Nat Prod Res, advance online publication 6 January 2016; DOI: 10.1080/14786419.2015.1115997
- 80 Fraga BM, Diaz CE, Lopez-Rodriguez M. Two novel abietane dimers from transformed root cultures of Salvia broussonetii. Tetrahedron Lett 2014; 55: 877-879
- 81 Christensen KB, Jorgensen M, Kotowska D, Petersen RK, Kristiansen K, Christensen LP. Activation of the nuclear receptor PPAR gamma by metabolites isolated from sage (Salvia officinalis L.). J Ethnopharmacol 2010; 132: 127-133
- 82 Bisio A, De Tommasi N, Romussi G. Diterpenoids from Salvia wagneriana. Planta Med 2004; 70: 452-457
- 83 Gao C, Han L, Zheng D, Jin HW, Gai CY, Wang JB, Zhang H, Zhang LR, Fu HZ. Dimeric abietane diterpenoids and sesquiterpenoid lactones from Teucrium viscidum. J Nat Prod 2015; 78: 630-638
- 84 Wu JZ, Morizane C, Iida A, Ueda SI, Zhou ZL, Xu M, Zhang M, Li RM, Fujita T. Structures of three new diterpenoids, fritillebic acid and fritillebins A and B, from bulbs of Fritillaria ebeiensis G. D. Yu et G. Q. Ji. Chem Pharm Bull (Tokyo) 1995; 43: 1448-1453
- 85 Wu JZ, Ruan HL, Zeng CL, Cheng HA, Zhang F, Zhao QS, Sun HD, Fujita T. Structures of two new diterpenoid dimers from bulbs of Fritillaria ebeiensis. J Asian Nat Prod Res 1999; 1: 251-257
- 86 Wu JZ, Wen YP, Ruan HL, Yao NH, Zhao QS, Sun HD, Morizane C, Iida A, Fuilta T. Structural elucidations of two ent-kaurane dimers from bulbs of Fritillaria ebeiensis var. purpurea. J Asian Nat Prod Res 2000; 2: 213-218
- 87 Wu JZ, Ruan HL, Yao NH, Zhao QS, Sun HD, Morizane C, Iida A, Fujita T. Structures of two diterpenoid dimers from bulbs of Fritillaria ebeiensis. J Asian Nat Prod Res 2000; 2: 161-167
- 88 Ruan HL, Zhang YH, Wu JZ, Sun HD, Fujita T. Two new diterpenoid dimers, fritillebinide D and E, from bulbs of Fritillaria ebeiensis. J Asian Nat Prod Res 2002; 4: 309-314
- 89 Zhang HJ, Luo J, Shan SM, Wang XB, Luo JG, Yang MH, Kong LY. Aphanamenes A and B, two new acyclic diterpene [4+2]-cycloaddition adducts from Aphanamixis grandifolia. Org Lett 2013; 15: 5512-5515
- 90 Zhang Y, Lu Y, Mao L, Proksch P, Lin WH. Tagalsins I and J, two novel tetraterpenoids from the mangrove plant, Ceriops tagal. Org Lett 2005; 7: 3037-3040
- 91 Chen JD, Qiu Y, Yang ZW, Lin P, Lin YM. Dimeric diterpenes from the roots of the mangrove plant Ceriops tagal. Helv Chim Acta 2008; 91: 2292-2298
- 92 Chen JD, Yi RZ, Lin YM, Feng DQ, Zhou HC, Wang ZC. Characterization of terpenoids from the root of Ceriops tagal with antifouling activity. Int | Mol Sci 2011; 12: 6517-6528
- 93 Galli B, Gasparrini F, Lanzotti V, Misiti D, Riccio R, Villani C, He GF, Ma ZW, Yin WF. Grandione, a new heptacyclic dimeric diterpene from Torreya grandis Fort. Tetrahedron 1999; 55: 11385-11394
- 94 Ni ZY, Wu YB, Dong M, Zhang ML, Wang YF, Sauriol F, Huo CH, Shi QW, Gu YC, Kiyota H, Cong B. Diabietane ether, a new dimeric abietane with an ether linkage from Taxus cuspidata Needles. Z Naturforsch B 2011; 66: 1083-1086
- 95 Pinto AC, Pizzolatti MG, Epifanio RDA, Frankmolle W, Fenical W. The isolation of novel diterpenoids, including a C-40 bis-diterpenoid, from the Brazilian plant Vellozia magdalenae (Velloziaceae). Tetrahedron 1997; 53: 2005-2012
- 96 Sivasothy Y, Ibrahim H, Paliany AS, Alias SA, Md Nor NR, Awang K. A new bis-labdanic diterpene from the rhizomes of Alpinia pahangensis. Planta Med 2013; 79: 1775-1780
- 97 Ou YX, Li YY, Qian XM, Shen YM. Guanacastane-type diterpenoids from Coprinus radians. Phytochemistry 2012; 78: 190–196
- 98 Yoshida T, Toyota M, Asakawa Y. Scapaundulins A and B, two novel dimeric labdane diterpenoids, and related compounds from the Japanese liverwort Scapania undulata (L) Dum. Tetrahedron Lett 1997; 38: 1975-1978
- 99 Rodriguez AD, Shi JG. Isolation, structure elucidation, and synthesis of bisgersolanolide, a novel heptacyclic bis-diterpenoid from the gorgonian octocoral Pseudopterogorgia bipinnata. Org Lett 1999; 1: 337-340
- 100 Johnstone RW, Frew AJ, Smyth MJ. The TRAIL apoptotic pathway in cancer onset, progression and therapy. Nat Rev Cancer 2008; 8: 782-798
- 101 Obenhuber AH, Gianetti TL, Bergman RG, Arnold J. Regioselective [2+2] and [4+2] cycloaddition reactivity in an asymmetric niobium(bisimi-

do) moiety towards unsaturated organic molecules. Chem Commun 2015; 51: 1278–1281

- 102 Gao WY, Zhang R, Jia W, Zhang J, Takaishi Y, Duan HQ. Immunosuppressive diterpenes from *Veronicastrum sibiricum*. Chem Pharm Bull (Tokyo) 2004; 52: 136–137
- 103 Uchida M, Miyase T, Yoshizaki F, Bieri JH, Ruedi P, Eugster CH. Isolation of 14-hydroxytaxodione from *Plectranthus grandidentatus* Gurke and of 7 new dimeric diterpenoids from *Plectranthus grandidentatus*, *Plectranthus myrianthus* and *Coleus carnosus* – structures of grandidone a, 7-epi-A, B, 7-epi-B, C, D and 7-epi-D. Helv Chim Acta 1981; 64: 2227–2250
- 104 *Chen SK, Chen BY, Li H.* Flora Reipublicae Popularis Sinicae (Zhongguo Zhiwu Zhi). Beijing: Science Press; 1997
- 105 Hu XM, Zhang WK, Song LR, Hu L, Zhang GZ, Xie ZW, Wang XT, Xu GJ, Xiao PG, Ling YK, Ding XL, Cao CL, Li YK, Yu WX, Hong X, Wang JH.

Chinese Materia Medica (Zhonghua Bencao). Shanghai: Shanghai Science and Technology Press; 1999

- 106 Aoyagi Y, Takahashi Y, Satake Y, Fukaya H, Takeya K, Aiyama R, Matsuzaki T, Hashimoto S, Shiina T, Kurihara T. Biomimetic synthesis of grandione from demethylsalvicanol via hetero-Diels-Alder type dimerization and structure revision of grandione. Tetrahedron Lett 2005; 46: 7885–7887
- 107 Majetich G, Zou G. Total synthesis of (-)-barbatusol, (+)-demethylsalvicanol, (-)-brussonol, and (+)-grandione. Org Lett 2008; 10: 81–83
- 108 *Kelecom A.* Isolation, structure determination, and absolute configuration of barbatusol, a new bioactive diterpene with a rearranged abietane skeleton from the Labiate *Coleus barbatus*. Tetrahedron 1983; 39: 3603–3608
- 109 Deng J, Zhou SP, Zhang WH, Li J, Li RF, Li A. Total synthesis of taiwaniadducts B, C, and D. J Am Chem Soc 2014; 136: 8185–8188