Studies on the Alkaloids of the Calycanthaceae and Their Syntheses

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Abstract: Plants of the Calycanthaceae family, which possesses four genera and about 15 species, are mainly distributed in China, North America and Australia. Chemical studies on the Calycanthaceae have led to the discovery of about 14 alkaloids of different skeletons, including dimeric piperidinoquinoline, dimeric pyrrolidinoindoline and/or trimeric pyrrolidinoindolines, which exhibit significant anti-convulsant, anti-fungal, anti-viral analgesic, anti-tumor, and anti-melanogenesis activities. As some of complex tryptamine-derived alkaloids exhibit promising biological activities, the syntheses of these alkaloids have also been a topic of interest in synthetic chemistry during the last decades. This review will focus on the structures and total syntheses of these alkaloids.

Keywords: biological activity; biosynthesis; Calycanthaceae alkaloids; structure; synthesis

1. Introduction

The small family of the Calycanthaceae comprises four genera, namely Chimonanthus Lindley, Sinocalycanthus Cheng & S. Y. Chang, Calycanthus L., and Idiospermum, which globally include ca. 15 species [1–3]. The plants of the Chimonanthus and Sinocalycanthus genera are ornamental shrubs endemically distributed in China, and those of the Calycanthus and Idiospermum originate from North America and Australia, respectively. The classification of the species of Chimonanthus genus is still a tough task and has been a subject of debate for a long time [4]. The early literature categorized this
genus into three species, *i.e.*, *Ch. nitens* Oliv., *Ch. praecox* (Linn.) Link, and *Ch. salicifolius* S.Y. Hu [5]. Recently it was proposed that this genus be classified in 10 species based on morphological evidence, including *Ch. nitens* Oliv., *Ch. praecox* (Linn.) Link, *Ch. salicifolius* S.Y. Hu, *Ch. nitens*, *Ch. zhejiangensis* M.C. Liu, *Ch. campanulatus*, *Ch. baokangensis*, *Ch. anhuiensis*, *Ch. caespitosa*, *Ch. campanulatus* var. guizhouensis [6]. In addition, only one species, *C. chinensis* Cheng et S.Y. Chang, is attributed to the *Sinocalycanthus* genus. Three plants named *C. floridus* var. floridus, *C. floridus* var. laevigatus, and *C. occidentalis* Hook. et Arn. pertain to the genus *Calycanthus*. The plant *Idiospermum australiense* (Diels) S. T. Blake a rare tree that occurs only in the North Queensland region of Australia is the sole member of the *Idiospermum* genus. These Calycanthaceae plants are primitive angiosperms and popular ornamental flowers with a pleasant aroma. The Calycanthaceae plants have long been used in Traditional Chinese Medicines (TCMs), to treat rheumatic arthritis, coughs, throat wounds, dizziness, nausea, fever, detoxification, and enteral disease [7–9].

The chemical investigation of the Calycanthaceae plants started more than one hundred years ago in 1888, which led to the isolation of a large amount of alkaloids, flavonoids [10], lignans [6,11], coumarins [12–14], terpenoids [15–17], and essential oils [7,18–20]. It is important to note that the discovery of the Calycanthaceae alkaloids was reminiscent of the history of development of science. Looking back to this history, only 14 alkaloids (Figure 1) whose discoveries were full of hardship and arduousness were characterized from this family. During past decades, there has been a trend towards synthetic approaches of these structurally interesting and bioactive alkaloids. This review attempts to provide timely and comprehensive coverage of the chemical and biological studies related to the Calycanthaceae alkaloids, with a specific focus on summarizing the great amount of synthetic work performed in this area.

2. Structures, Biological Activities, and Biosynthetic Origins of Calycanthaceae Alkaloids

2.1. Structures of Calycanthaceae Alkaloids and Their Discovery

The phytochemical investigation of the plants Calycanthaceae was first described in 1888, which led to isolation of the first Calycanthaceae alkaloid, (+)-calycanthine (1, Figure 1) with a dimeric piperidinoquinoline skeleton, from the seeds of *C. glaucus* Willd. by Eccles. One year later, Wiley proved the high content of this alkaloid in the seeds of the same plant [21]. In 1905, further progress was made by Gordin whereby this principal alkaloid was crystalized in different forms at ordinary temperature and deduced to possess a molecular formula C_11H_14N_2 containing no oxygen atom [21,22]. A few years later, Späth and Stroh expressed their disagreement on Gordin’s work and stated that the empirical formula of (+)-calycanthine should be doubled C_11H_14N_2 [23]. Soon Manske put forward the possibility that this molecular formula might be C_{22}H_{26}N_4 [24], which was finally proved by Barger’s group in 1939 [25]. The structure of this alkaloid (+)-calycanthine, C_{22}H_{26}N_4, was finally established unequivocally by means of X-ray crystal structural analysis of its dihydrobromide dehydrate by Hamor’s group in 1960 [26].
Figure 1. Structures of the Calycanthaceae alkaloids.
In 1905, Gordin also reported a second alkaloid, isocalycanthine, from the seeds of *Chimonanthus* genus, which possessed a different melting point and showed different behavior with respect to the removal of water of crystallization from the hydrated base with those of calycanthine [27,28]. However, Manske expressed doubts about the existence of isocalycanthine. In his study on the species of *C. floridus* L., a great quantity (1.2%) of calycanthine was. Gordin’s seed extract consisted in reality of *C. fertilis*. In the same report, Gordin also described the isolation process of (+)-calycanthine (2.6%) from the seeds of *Meratia praecox* (*C. praecox*) [24]. In 1992, the occurrence of isocalycanthine in a closely related species *Psychotria forsteriana* (Rubiaceae) was reported by Kuballa’s group [29].

In 1938, Barger’s group isolated a minor alkaloid, (−)-calycanthidine (2), from the seeds of *C. floridus*, which molecular formula was deduced to be C_{13}H_{16}N_{2} [30]. This conclusion was corrected by Saxton in 1962, who established both the formula C_{23}H_{28}N_{4} and the structure of (−)-calycanthidine [31]. The alkaloid (−)-chimonanthine (3) was firstly obtained from *Ch. fragrans* Lindle (*Ch. praecox*) by Hodson’s group [32], and its structure was determined on the basis of detailed X-ray analysis of chimonanthine dihydrobromide by Grant’s group [33,34]. As for (−)-folicanthine (4), it was first isolated by Hodson’s group in 1957 and was identified to be a dimeric pyrrolidinoindoline alkaloid, close to (−)-calycanthidine (2), and (−)-chimonanthine (3) [32]. From then on, this seemed to be an end for the structural elucidation of (−)-calycanthidine, (−)-chimonanthine, (−)-folicanthine, until in 2000, the total syntheses of these alkaloids were completed by Overman’s group, proving that (−)-calycanthidine, (−)-chimonanthine, (−)-folicanthine should be drawn as compounds 2, 3, and 4 (Figure 1), respectively; therefore the structures of compounds *ent*-2, *ent*-3, and *ent*-4 reported in the previous papers should be (+)-calycanthidine, (+)-chimonanthine, (+)-folicanthine, respectively [35]. Calycanine (5) with a molecular formula C_{16}H_{10}N_{2} was a product of calycanthine and chimonanthine obtained by Zn dehydrogenation. Its structure was first incorrectly proposed by Barger, and then revised by Woodward’s group via synthesis [32,34,36].

(−)-Idiospermuline (6), a trimeric pyrrolidinoindoline, together with two known dimeric alkaloids, (+)-calycanthine and (−)-chimonanthine were isolated by the bioassay-guided method from the seeds of *Idiospermum australiense* (Diels) S.T. Blake, a native species from North Queensland, Australia. The structure of (−)-idiospermuline was determined by NMR and MS data and the absolute stereochemistry was established by X-ray crystallographic study of idiospermuline trimethiodide (6a) [37].

In 2004, Takayama’s group investigated the alkaloidal constituents of the seeds of *Ch. praecox* L., leading to the isolation of two new tryptamine-related alkaloids, chimonamidine (7) and chimonanthidine (8), together with the known (+)-calycanthine, (−)-chimonanthine, (−)-folicanthine, and (−)-calycanthidine. They conducted the total synthesis of (±)-chimonamidine to establish its absolute structure. As a result, natural chimonamidine showed optical rotation $[\alpha]_{D}^{29} = -12.6$, which was significantly different with those of (R)-(−)-chimonamidine ($[\alpha]_{D}^{29} = -178$) and (S)-(−)-chimonamidine ($[\alpha]_{D}^{21} = +171$), suggesting that natural chimonamidine is a mixture slightly enriched with the (R)-(−)-enantiomer. Chimonanthidine (8) was determined to be N_{6}-monodemethylfolicanthine by total synthesis, with the absolute configuration eventually confirmed by a combined strategy of comparing the CD spectrum with that of known (−)-folicanthine [38]. In his paper of 2006, a further phytochemical study on the seeds and rinds of *Ch. praecox* (L.) f. concolor resulted in the isolation of a new pyrrolidinoindoline-type alkaloid, CPC-1 (9), and one new tetrahydroquinoline dimeric alkaloid, CPC-2 (10), together with eight known alkaloids, (+)-calycanthine, (−)-chimonanthine,
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(−)-folicanthine, (−)-calycanthidine, (−)-chimonanthidine, meso-chimonanthine (11), tryptamine (12), and Nα,Nβ-dimethyltryptamine (13) [39].

In 2009, an antifungal activity-guided phytochemical investigation on the defatted seeds of *Ch. praecox*, a species grown in Shaanxi Province of China, afforded two dimeric alkaloids, (+)-calycanthine and (−)-folicanthine [40]. Additionally, an additional indole-derived glycoside, 3-methylcarboxymethyl-indole-1-N-β-D-glucopyranoside (14), and the four known compounds (+)-calycanthine, (−)-chimonanthine, (−)-folicanthine, (−)-calycanthidine were obtained from the fruits and leaves of *C. praecox* in 2011 [15]. A recent phytochemical study on the flower buds of *Ch. praecox* also led to the isolation of five known dimeric alkaloids and several other compounds [41].

2.2. Biological Activities

The Calycanthaceae plants have been used as Traditional Chinese Medicines (TCMs) for the treatment of colds, and as sedative, antitussive, anti-hypertension, antioxidation, anti-inflammatory, and antitumor medicines [14,15]. As the important components of these plants, the Calycanthaceae alkaloids showed biological activities such as anti-convulsant, anti-fungal, anti-viral, analgesic, anti-tumor, and melanogenesis inhibitory properties.

The main representative alkaloid, calycanthine (1), has been recognized as a powerful centrally acting anti-convulsant for a long time [42,43]. It was reported that calycanthine may mediate its convulsant action predominantly by inhibiting the inhibitory neurotransmitter GABA as a result of interactions with L-type Ca2+ channels and by inhibiting GABA-mediated chloride currents at GABAA receptors [44].

(+) Calycanthine (1) and (−)-folicanthine (4) were evaluated for their antifungal activities against five plant pathogenic fungi, *Exserohilum turcicum*, *Bipolaris maydis*, *Alternaria solani*, *Sclerotinia sderotiorum*, and *Fusarium oxysporium*. It turned out to be that *B. maydis* was the most susceptible to 1 with an EC50 value of 29.3 µg/mL, and then *S. sderotiorum* to 4 with an EC50 of 61.2 µg/mL [40].

(−)-Chimonanthine (3) and (−)-folicanthine (4) also showed weak antiviral activities against porcine respiratory and reproductive syndrome virus (PRRSV) with IC50 values of 68.9 ± 3.1 µM and 58.9 ± 10.2 µM, respectively [45].

Chimonanthines were tested on μ- and κ-opioid binding assay, and on the tail-flick and the capsaicin-induced pain models. As a result, (−)-chimonanthine (3), (+)-chimonanthine (ent-3), and meso-chimonanthine (11) showed strong binding affinities towards μ-opioid receptors with Ki values of 271 ± 85 nM, 652 ± 159 nM, and 341 ± 29 nM, respectively, indicating their significant analgesic activities [46,47].

It was also recently reported that the methanol extract of the flower buds of *Ch. praecox* showed an inhibitory effect against melanogenesis in ophylline-stimulated B16 melanoma 4A5 cells. A further investigation revealed that the principal alkaloids of (+)-calycanthine (1), (−)-chimonanthine (3), and (−)-folicanthine (4) showed the most potent melanogenesis inhibitory activity, with IC50 values of 0.93, 1.4, and 1.8 µM, respectively. Abutin (174 µM), a commercially tyrosinase inhibitor, was used as a positive control. In this paper, (+)-chimonanthine (ent-3), and meso-chimonanthine (11) showed cytotoxicity at 10 µM [41]. In another assay, compounds 2–4 were screened for the cytotoxicity
against a small panel of human cancer lines, showing cytotoxic effects against gastric carcinoma NUGC3 and hepatocarcinoma SNU739 cancer cells with IC50 values ranging from 10.3 to 19.7 μM [15].

2.3. Biosynthetic Origins

Calycanthine, calycanthidine, chimonanthine, folicanthine, chimonanthidine, and CPC-2 are a series of tryptamine-derived dimeric alkaloids, which were proposed to be originated from \( N_b \)-methyltryptamine (15). The oxidative dimerization of two molecules of \( N_b \)-methyltryptamine forms the key tetraaminodialdehyde intermediate 16, which undergoes several enzyme-catalyzed reactions and modifications yielding calycanthine and CPC-2 (Scheme 1) [39]. A possible biosynthesis of chimonanmidine (7) is shown in Scheme 2. The intermediate 17 was derived from tryptamine (12) by oxidation, and subsequently converted into 18 by introduction a hydroxy function at the benzylic position. Chimonanmidine (7) was finally produced by the transannulation of the lactam ring of 18 [38,48].

![Scheme 1. Potential biogenetic pathway of tryptamine-derived dimeric Calycanthaceae alkaloids.](image1)

![Scheme 2. Potential biogenetic pathway of chimonanmidine (7).](image2)
3. Total Synthesis of Calycanthaceae Alkaloids

3.1. Calycanthines and Chimonanthines

The dimeric piperidinoquinoline and pyrrolidinoindoline alkaloids have long been synthetic topics, and efforts have been made on the total synthesis of these two skeletons. Owing to their characteristic C2-symmetrical bridged bicycles and four chiral centers, three possible calycanthine diastereomers including (+)-calycanthine from Calycanthaceae plants, (−)- and meso-calycanthine from Psychotria forsteriana [29], were identified (Figure 2). Likewise, the C2-symmetrical chimonanthines also comprised (±)- and meso-chimonanthine. The total syntheses of these alkaloids have been intensively studied for decades. Hino speculated that the structure of 1,1′-dimethyl-3,3′-bis(2-aminoethyl)-3,3′-bioxindole was the key intermediate for the syntheses of calycanthine and (±)-folicanthine [49,50]. Some other groups made synthetic efforts to these bis(pyrroloindoline) scaffold by oxidation dimerization of indole [51] and/or oxindoles derivatives [52,53]. The synthetic challenges were attributed to the two structural features of the dimeric pyrroloindole core, including C3a-C3′α σ bond and its vicinal quaternary stereogenic carbons.

In 1964, Scott’s group established an approach using N5-methyltryptamine (15) as starting material to form the magnesium salt of methyl tryptamine in presence of methyl magnesium iodide (CH₃MgI). The intermediate was treated with iron(III) chloride to produce the indolenine dimer (19), which was subsequently transformed to meso- and (±)-chimonanthine in one step. Moreover, (±)-calycanthine was accessible by treating (±)-chimonanthine with aqueous acid, demonstrating that structure of 1 was the thermodynamically preferred scaffold (Scheme 3) [54].

![Figure 2. Structures of calycanthines.](image)
Scheme 3. Biomimetic syntheses of chimonanthines and calycanthines proposed by Scott.

Scheme 4. Stereocontrolled syntheses of meso-chimonanthine and meso-calycanthine by Overman.

Overman’s group also proposed systemic syntheses for these Calycanthaceae alkaloids. In 1996, he attempted to stereocontrollly synthesize meso-chimonanthine (11) and meso-calycanthine (meso-1) via
a samarium-mediated reductive dialkylation (Scheme 4). Isoindigo (21) was converted to \( N \)-benzyl derivative 22, which then generated 24 by the treatment of 2 equiv. of SmI\(_2\) in the presence of 10 equiv. of LiCl with cis-1,4-dichloro-2-butene. Compound 24 was subsequently transformed into hexacycle 25 with sodium bis(2-methoxyethoxy)aluminum hydride (Red-Al). The cyclohexene of 25 was cleaved to yield 26, which was immediately reduced to diamine 28 via diazide product 27. Intermediate 28 was then treated with excess Me\(_3\)Al at room temperature to provide bis(pyrroloindoline) 29. The desired \( meso \)-chimonanthine (\( meso \)-3) was produced from 29 by a cascade of methylation and deprotection reactions. The final product \( meso \)-calycanthine (\( meso \)-1) was obtained by exposure of \( meso \)-3 to hot dilute acetic acid [55]. In 1999, a flexible approach to \( meso \)-chimonanthine (Scheme 5), \((-)\)-chimonanthine, and \((+)\)-calycanthine (Scheme 6) using an intramolecular Heck reaction cascade was present by Overman \textit{et al.} [56]. One year later, they put forward to a highly efficient synthesis of 3a,3a'-bispyrrolidino[2,3-\( b \)]indolines, a precursor for the \( meso \)-chimonanthine (Scheme 7a) and \((+)\)-chimonanthine (Scheme 7b). This dialkylation route utilized the reactivities of dienolate (chelated or nonchelated) and the chirality of a tartrate-derived dielectrophile to control the relative configuration and absolute stereochemistry, respectively [35].

In 2006, Dalko’s group provided an elegant synthesis of a homologous compound of \( meso \)-chimonanthine, \( N_0 \)-desmethyl-\( meso \)-chimonanthine. They investigated a tandem [4 + 2]-cycloadition-cyclisation of a conveniently functionalized bromoxidole (55) and tryptamine derivative 56 to construct the \( meso \)-chimonanthine core in a highly diastereoselective manner (Scheme 8) [57].
Scheme 6. Double Heck cyclizations for (−)-chimonanthine and (+)-calycanthine by Overman.

In 2007, an alternative strategy for the total synthesis of (−)-calycanthine (ent-1) and (+)-chimonanthine (ent-3) using a reductive CoI-promoted dimerization of endo bromide (+)-60 was proposed by Movassaghi. The vital homodimerization requiring a stoichiometric amount of metal catalyst was the key step to secure the vicinal quaternary stereocenters, which was directed by the stereochemistry at the C8a-position of (+)-60 (Scheme 9a). The further treatment of ent-3 with [D4]acetic acid and deuterium oxide provided (−)-calycanthine (ent-1) (Scheme 9b) [58]. In 2014, Movassaghi’s group also described an enhanced diazene-based method for heterodimerization to enantioselectively synthesize the (−)-calycanthidine (2), meso-chimonanthine (11) and (±)-desmethyl-meso-chimonanthine (58) [59,60].
Scheme 7. Enantioselective dialkylation approach for meso-chimonanthine (a) and (+)-chimonanthine (b) proposed by Overman.

Scheme 8. Synthesis of (+)-N<sub>b</sub>-desmethyl-meso-chimonanthine (58) via a tandem [4 + 2]-cycloaddition-cyclisation by Dalko.
Scheme 9. Synthesis of (+)-chimonanthine (ent-3) (a), (+)-folicanthine (ent-4) and (−)-calycanthine (ent-1) (b) via a reductive CoI-promoted homodimerization by Movassaghi.

In 2012, Kanai and Matsunaga’s group illustrated a straightforward catalytic asymmetric total synthesis to achieve enantioselective (−)-calycanthine (ent-1) and (+)-chimonanthine (ent-3) in seven steps (Scheme 10). A one-pot double Michael reaction from 3,3′-bioxindole with the base catalyst Mn(4-F-BzO)2/Schiff (69) produced the key dialkylated adduct 71 in 69% yield and 95% ee. Then the intramolecular dicyclization of 72 was available by the treatment with LiEtBH in toluene [62]. Recently, a new strategy using a double intramolecular carbamoylketene-alkene [2 + 2] cycloaddition for the synthesis of the racemic chimonanthine was accomplished by Shishido’s group (see Chapter 3.2) [63].

3.2. Folicanthines

The structures of the folicanthines are also representatives of the dimeric hexahydropyrroloindole alkaloid family. (+)-Folicanthine (ent-4) could be easily obtained in a quantitative yield from (+)-chimonanthine (ent-3) by treatment with formaldehyde and sodium triacetoxyborohydride [NaBH(OAc)3] (Schemes 9b and 10) [58,62]. In striking contrast to the elegant synthetic procedures towards calycanthines and chimonanthines, Gong’s group have developed a highly enantioselective
nucleophilic substitution reaction of 3-hydroxyoxindoles with an enecarbamate catalyzed by chiral phosphoric acids.

Scheme 10. Synthesis of (+)-chimonanthine (ent-3), (+)-folicanthine (ent-4) and (−)-calycanthine (ent-1) via double Michael reaction of bisoxindole by Matsunaga.

In Scheme 11, the vital enantioselective substitution reaction of 3-hydroxy-3,3′-bisindolin-2-one (73) with 75 was catalyzed by a special chiral phosphoric acids 74 to give 76. The dimethylation of 76 provided 77, which was followed with a Beckmann rearrangement to convert the ketones into amide derivatives 78a and 78b. A second Beckmann rearrangement of 78b by the treatment with mercury(II) chloride (Hg2Cl) could furnish amide 79. After introducing a methyl group at the amide nitrogen atom, the product 80 underwent an alkylation to produce key intermediate 81, which was finally converted to the desired product (+)-folicanthine (ent-4) after several step sequences [64]. Contemporaneously, Liang’s group described a concise three-step synthesis of (+)-folicanthine using two molecules of 2-(1-methyl-1H-indole-3-yl)-N-tosylethaneamine (84, Scheme 12). The core structure 85 of (±)-folicanthine could be easily obtained by a one-step cyclization-dimerization of substituted tryptophan in high yield. In general, this synthetic route had the advantages of being highly efficient, atom-economic, and metal-free, but it has no enantioselectivity [65]. Recently, Shishido’s group also developed a total synthesis route to access the folicanthines and chimonanthines in racemic form,
employing a double intramolecular carboamoylketene-alkene \([2 + 2]\) cycloaddition reaction as the key step (Scheme 13). 2,2'-(Buta-1,3-diene-2,3-diyl)bis(nitrobenzene) (90), obtained by a Pd-catalyzed preparation, underwent the key double \([2 + 2]\) cycloaddition to yield bis-carboxylic acid 92. After a three-step sequence, the mixture of (+)-folicanine and meso-folicanine (meso-4) was obtained, which was separated by preparative TLC in 36% and 20% yields, respectively [63].

Scheme 11. Synthesis of (+)-folicanine (ent-4) via asymmetric organocatalytic substitution reaction by Gong.
Scheme 12. Concise synthesis of (±)-ficanthine by Liang.

Scheme 13. Synthesis of ficanthines and (±)-chimanthine via a double intramolecular carbamoylketene-alkene [2 + 2] cycloaddition by Shishido.

3.3. (−)-Idiospermuline (6)

(−)-Idiospermuline (6), a typical trispyrroloidinoindoline alkaloid, contains three all-carbon quaternary centers, consisting of one (−)-chimanthine unit and one pyrroloidinoindoline unit with a C3"a–C7" σ bond. Its retrosynthetic analysis was similar to that of hodgkinsines [66] and is presented in Scheme 14, which guided the total enantioselective synthesis of (−)-idiospermuline (Scheme 15). The first step in the synthesis focused on the dimeric pyrroloidinoindoline derivative 97 similar to that
of (+)-chimonanthine in Scheme 7. The introduction of the readily available stannyl butenilide 102 furnished the intermediate 96. Heck cyclization of 96 treating with chelating diphosphane ligands like bis(1,4-diphenylphosphanyl)butane (dppb) diastereoselectively furnished 3a'R precursor 95. Catalytic hydrogenation of 95 with Pd(OH)2 and H2, followed by reduction of the carbonyl group with Red-Al in toluene, then Na in NH3, provided the desired (−)-idiospermuline (6) [67–70].

Scheme 14. Retrosynthesis of (−)-idiospermuline (6).

Scheme 15. Total synthesis of (−)-idiospermuline (6).
3.4. Chimonamidines (7)

Takayama’s group conducted a biomimetic synthesis (Scheme 16) in order to confirm the absolute structure of chimonamide (7). Based on its plausible biogenetic pathway shown in Scheme 2, the precursor \(N_a,N_b\)-dimethyltryptamine (13) was treated with benzyl chloroformate (Cbz-Cl) and \(Na_2CO_3\) to give \(N_a,N_b\)-dimethyl-\(N_b\)-carbobenzyloxytryptamine (103). Oxidation of 103 with dimethyl sulfoxide and hydrochloric acid yielded 104, which was then converted to the racemic mixture of (±)-hydroxyketones 105 by introduction of a hydroxy group at the benzylic position. The target racemic mixture of (±)-chimonamide (7) was obtained by removing the \(N_b\)-Cbz protecting group and forming a new lactam ring under the condition of trimethylsilyl iodide (TMSI). Meanwhile, the chiral synthesis of chimonamide was also performed by Takayama. After several attempts, a strategy that involved the separation of racemic mixtures by (+)-MTPA chloride and SiO2 column chromatography succeeded to yield two diastereomeric esters 106 and 107. Two more steps that involved the hydrolysis with aqueous alkaline solution and cyclization with TMSI of 106 and 107 were conducted to afford two enantiomerically pure compounds (R)-(−)-chimonamide and (S)-(+) chimonamide, respectively. A comparison of the optical rotation between synthesized ([\(\alpha\])\(_{D}^{23}\) = −178 for (R)-(−)-7; [\(\alpha\])\(_{D}^{23}\) = +171 for (S)-(+)7) and natural chimonamide ([\(\alpha\])\(_{D}^{19}\) = −12.6) came to the conclusion that natural chimonamide was a mixture slightly enriched with (R)-(−)-chimonamide [38].

Scheme 16. Total synthesis of (±)-chimonamide (7).
3.5. Chimonanthidine (8)

A synthetic approach that uses hypervalent iodine(III) reagents for the dimerization of indole derivatives was developed for the total synthesis of (+)-chimonanthidine (8) by Takayama’s group (Scheme 17). The methylation of the known compound 108 with methyl iodide (CH₃I) and sodium hydride (NaH) in DMF produced Nα-Methyl-Nβ-trimethylsilyloxy-carbonyl (Teoc) tryptamine (109), which was then treated with 0.5 equiv. phenyliodine(III) bis(trifluoroacetate) (PIFA) in CF₃CH₂OH to yield two dimeric diastereoisomers 110 and 111. (+)-Chimonanthidine (8) was obtained from the monodeprotected amine 112, which was transformed from 111 by treatment with tetrabutylammonium fluoride (TBAF) in THF [38].

\[ \text{Scheme 17. Total synthesis of (+)-chimonanthidine (8).} \]

3.6. Rac-CPC-1 (Rac-9)

The total synthesis of racemic CPC-1 was initially performed to confirm the structure of CPC-1 (Scheme 18). Compound 113 was treated with m-CPBA in the presence of excess trifluoroacetic acid (TFA) in CH₂Cl₂ to afford 3α-hydroxypyrrolidinoindoline (114). Methylation of the hydroxy group of 114 and then removal of the Nβ-Teoc group yielded the intermediate 115, which was finally treated with formalin and then NaBH₃CN to give rac-9. To further establish the absolute configuration of 9, its chiral total synthesis was conducted (Scheme 19). This synthetic approach started from isatin (66), which was treated with (R)-(+)−binol, Ti(OiPr)₄, and tetraallylstannane to yield allylated compound 116. Two recrystallizations of 116 from EtOAc afford enantiomerically pure (S)-(−)-116. Methylation

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**Scheme 17. Total synthesis of (+)-chimonanthidine (8).**

**Scheme 18. Total synthesis of racemic CPC-1 (Rac-9).**

**Scheme 19. Chiral total synthesis of CPC-1 (Rac-9).**
of Na and hydroxy group of \((S)-(\ldots)\text{-}116\) gave the dimethyl compound \(117\). The intermediate \(117\) was treated with OsO\(_4\) and \(N\)-methylmorpholine \(N\)-oxide (NMO) followed by NaIO\(_4\) to provide the aldehyde, which was directly subjected to reductive amination by condensing with CH\(_3\)NH\(_2\) and then reduced with NaBH\(_3\)CN to give \((S)-(\ldots)\text{-}118\). The desired product \((3aS, 8aS)-9\) was obtained by the reductive cyclization of \(118\). The optical rotation value of \((3aS, 8aS)-9\) ([\(\alpha\])\(_D\text{24}\) = +101) was determined to be opposite of that of \(9\) ([\(\alpha\])\(_D\text{24}\) = −88), indicating the absolute configuration of natural CPC-1 to be 3aR,8aR [39].

![Scheme 18. Total synthesis of rac-9.](image)

**Scheme 18.** Total synthesis of rac-9.

\(9\):

\(\text{Isatin (66)}\)

- Tetraallylatannane, \((R)-(\ldots)\)-binol, Ti\((\text{OPr})_4\), 2-propanol, 0 °C, to rt, 10 h
- Recrystallization from AcOEt

\(\text{i}) OsO}_4\), NMO, CH\(_3\)CN, H\(_2\)O, rt, 65 h
\(\text{ii}) NaIO}_4\), 1,4-dioxane, H\(_2\)O, rt, 30 min
\(\text{iii}) CH\(_3\)NH\(_2\)-HCl, MgSO\(_4\), MeOH, rt 1 h; then NaBH\(_3\)CN, rt, 3h

\(\text{LiAlH}_4\), THF, 0 °C, 1.5 h, rt, 2.5 h

\(\text{CH}_3\text{I, NaH, DMF, 0 °C, 3 h; rt, 4.5 h}\)

\(\text{TBAF, THF, 0 °C, 1.5 h; rt, 16.5 h}\)

\(\text{HCHO aq; NaBH}_3\text{CN, MeOH, rt, 1h}\)

\(\text{CH}_3\text{I, NaH, DMF, 0 °C, 3.5 h}\)

\(\text{NaBH}_3\text{CN, MeOH, rt, 1h}\)

**Scheme 19.** Total synthesis of \((3aS, 8aS)-9\).

4. Conclusions

In conclusion, the Calycanthiaceae plants are rich in promising bioactive dimeric and/or oligomeric piperidinoquinoline and hexahydropyrroloindole alkaloids, which are characterized by unique vicinal quaternary stereocenters. These alkaloids have been a longstanding challenge as total synthetic targets. During the last decades, stereocontrolled total synthetic approaches, such as metal-catalyzed dialkylation, intramolecular double Heck reaction, tandem [4 + 2]-cycloaddition-cyclisation, Co\(^1\)-promoted reductive homodimerization, double Michael reaction, double Beckmann rearrangement, and intramolecular double carbamoylketene-alkene [2 + 2] cycloaddition, have extensively explored. Taken together, these results will keep research on the metabolites of the Calycanthaceae plants as a hot topic for the scientific community in the future.
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Author Contributions

The basic ideas were thought by K.-J. C; the whole paper was written by J.-B. X and polished by K.-J. C.

List of Abbreviations

Ac—acetyl; Bn—benzyl; Boc—tert-butoxycarbonyl; CAN—ceric ammonium nitrate;
CAS—camphorsulfonic acid monohydrate; Cbz—carbobenzoxy; m-CPBA—m-chlorperoxybenzoic acid;
dba—trans,trans-dibenzylideneacetone; dppe—1,2-bis(diphenylphosphino)ethane;
dpb—bis(1,4-diphenylphosphanyl)butane; DMA—N,N-dimethylacetamide; DMF—2,5-dimethylfuran;
DMSO—dimethyl sulfoxide; 2,2-DMP—2,2-dimethoxypropane; DMPU—1,3-dimethylhexahydro-2-
pyridimidone; HMDS—1,1,1,3,3,3-hexamethyldisilazane; HMPA—hexamethylphosphoramide;
NaHMDS—sodium hexamethyldisilazide; NMO—4-methylmorpholine-N-oxide;
PIFA—phenyliodine(III) bis(trifluoroacetate); PMP—1,2,2,6,6-pentamethylpiperidine;
PSTA—p-toulenesulfonic acid; Red-Al—sodium bis(2-methoxyethoxy)aluminum hydride;
TBAF—tertbutylammonium fluoride; Tf—trifluoromethanesulfonyl; TFA—trifluoroacetic acid;
THF—tetrahydrofuran; TMEDA—N,N,N,N-tetramethylethylenediamine; TMS—trimethylsilyl

Conflicts of Interest

The authors declare no conflict of interest.

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