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Rapid Construction of the ABC Ring System in the *Daphniphyllum* Alkaloid Daphniyunnine C

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ABSTRACT

An efficient and scalable synthesis of the ABC ring system common to the calyciphylline A-type alkaloids has been developed. The tricyclic core of the alkaloids features a bowl-shaped [6-6-5] skeleton with five stereogenic centers including an all-carbon quaternary center. It was constructed rapidly from a readily available carvone derivative through a seven-step sequence involving an aza-Michael addition and Pd-catalyzed enolate α -vinylation as key steps.

The *Daphniphyllum* alkaloids are a group of highly complex polycyclic alkaloids with remarkable structural diversity (Figure 1). More than 200 family members have been isolated over the past 50 years, and they exhibit a vast collection of novel skeletons with unusual ring systems. Historically, the unique structural features and intriguing biosynthesis of these alkaloids have inspired a variety of innovative and elegant studies in total synthesis. One milestone that is well-recognized among organic chemists is Heathcock's biomimetic synthesis of dihydro-*proto*-daphniphylline (1, Figure 1) and other structurally related

members based on his biogenetic proposal.^{2b-d} In recent years, discoveries of unprecedented structures in this alkaloid family have continuously prompted synthetic investigations in this field.³

Among the > 20 subgroups of the *Daphniphyllum* alkaloids, we are particularly interested in calyciphylline A-type alkaloids (Figure 2).⁴ Intrigued by the synthetic challenge posed by these structures, we started a program to study the total synthesis of calyciphylline A-type alkaloids and to explore the possible formation of the core structures in other subgroup members through elaboration of their tricyclic ABC ring system. To achieve these goals, it was necessary to develop an efficient and scalable synthesis for the rapid construction of the tricyclic bowl-shaped [6-6-5] skeleton.

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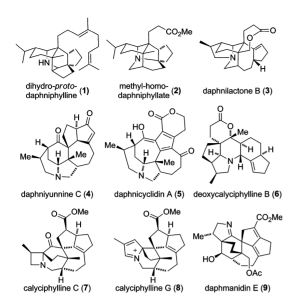


Figure 1. Structural diversities of the *Daphniphyllum* alkaloids.

Our synthetic plan is illustrated in Scheme 1, with daphniyunnine C (4, Figure 1) as an exemplary target. We envisioned that bond disconnections involving an intramolecular Pauson–Khand reaction could greatly simplify the hexacyclic molecule into the less complicated tetracyclic structure 16. This key spiro intermediate could be formed from 17 using a ring-closing metathesis (RCM) strategy, which in turn could be procured through a kinetic alkylation of the unsaturated ketone in 18. We reasoned that the stereochemistry of this step could be tightly controlled by the bowl-shaped architecture of the substrate 18. A short sequence from (S)-carvone (23) was

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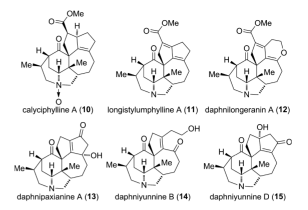


Figure 2. Representative calyciphylline A-type alkaloids.

Scheme 1. Retrosynthetic Analysis of Daphniyunnine C

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expected to readily establish all the stereogenic centers in 18. An intramolecular aza-Michael addition of carvone derivative 22 followed by alkylation and subsequent Pd-catalyzed enolate α -vinylation reaction⁸ could generate 20. Hydrogenation from the convex side of 20 would transform this substrate to the tricyclic product 19 with five stereogenic centers in place.

Herein, we report our success in the facile construction of the common bowl-shaped [6-6-5] skeleton 32 in calyciphylline A-type alkaloids with five stereogenic centers (Scheme 2) from the cost-effective (R)-carvone.

Our synthesis took advantage of a known transformation from (S)-carvone (23) to ent-24 reported by Overman and co-workers. This four-step sequence is very robust and can be readily scaled up to produce over 10 g of 24 in one batch. This transformation neatly controlled the required stereochemistry of the methyl substituent on the B ring. With a sufficient supply of 24 in hand, we started to

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Scheme 2. Synthesis of [6-6-5] Tricyclic Skeleton 32

explore our synthetic strategy by substitution of the hydroxyl group with an amino group. The primary hydroxyl group in 24 was selectively activated as a tosylate, which underwent nucleophilic substitution with sodium azide to afford 26 in excellent yield. Oxidation of the allylic alcohol with PCC¹⁰ and a subsequent Staudinger reaction¹¹ produced a mixture of ent-22 and the aza-Michael addition product 28. With no need for purification, the mixture was directly treated with allyl bromide 298f under typical alkylation conditions to generate a pair of diastereomers 30. In this step, owing to the intrinsic structural feature of the substrate, the aza-Michael addition could only occur from one face of the Michael acceptor and neatly install the C-N bond in a stereoselective manner on the A ring. Because the diastereomeric center in 30 would be lost in the next step, the mixture was used without separation. When 30 was treated under typical conditions for Pd-catalyzed enolate α-vinylation, it was smoothly converted to the tricyclic product 31 with a newly formed all-carbon quaternary center in 84% yield. Again, the intrinsic concavity of the substrate facilitated the stereoselective formation of the challenging all-carbon quaternary center. As we anticipated, hydrogenation of the double bond in 31 with Pd/C afforded 32 in almost quantitative yield with absolute stereochemical control. The stereochemistry of 32 was unambiguously confirmed through single crystal X-ray crystallographic analysis of the oxalate salt of **32** (Figure 3).

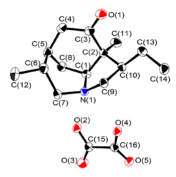


Figure 3. ORTEP depiction of the oxalate salt of **32** (thermal ellipsoids drawn at the 50% probability level).

Given that this synthetic sequence employed straightforward transformations and cost-effective chemicals and reagents in each step, it could be readily scaled up to supply sufficient material for further studies. To date, nearly 15 g of 32 have been prepared in our laboratory.

In summary, we have developed an efficient and scalable synthetic approach for the rapid construction of the ABC ring system commonly found in the *Daphniphyllum* subclass calyciphylline A-type alkaloids. By harnessing the intrinsic structural features of individual intermediates, a high degree of stereochemical control was achieved throughout the sequence. A seven-step sequence allowed us to build the rather complex tricyclic core skeleton from a readily available carvone derivative (24) in an overall yield of 48%. Notably, this core skeleton contains five stereogenic centers including an all-carbon quaternary center, making it a nontrivial synthetic target. Further studies toward the total synthesis of daphniyunnine C and other members of the *Daphniphyllum* alkaloids are currently underway in our laboratory.

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Supporting Information Available. Experimental details and procedures, compound characterization data, copies of ¹H and ¹³C NMR spectra for new compounds, and X-ray crystallographic data. This material is available free of charge via the Internet at http://pubs.acs.org.

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The authors declare no competing financial interest.