

Construction of Iterative Tetrahydrofuran Ring Units and Total Synthesis of (+)-Goniocin

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Supporting Information

ABSTRACT: Cytotoxic acetogenin (+)-goniocin has been synthesized in 17 steps from (R)-O-tritylglycidol. The core structure of the contiguous C_{22} – C_{10} threo-trans-threo-trans-tris-tetrahydrofuran (THF) ring involving an iterative THF-ring unit was synthesized. An iterative THF ring unit was constructed from an alkenyl-substituted THF ring in four steps including a Pd(II)-catalyzed ring-closing reaction and cross-metathesis. This method is general and allows the preparation of both *trans-threo-trans*- and *trans-threo-cis*-THF ring units flexibly.

Annonaceae acetogenins are a class of natural lipophilic polyketides with C_{32} or C_{34} unbranched fatty acid chains, and they are isolated from the *Annonaceous* plant family. They display a wide array of biological activities including anticancer, anti-inflammatory, pesticidal, antimalarial, immunosuppressive, and neurotoxic. Goniocin (1) was isolated from the bark of *Goniothalamus giganteus* in Thailand by the McLaughlin group in 1994. They discovered the unique structure and potent cytotoxic activity of 1. The structure of 1 consists of a core contiguous tris-*trans*-tetrahydrofuran (THF) ring unit, a left linear hydrocarbon chain unit, and a methyl-substituted γ-butenolactone unit on the right.

The chiral γ -butenolactone moiety is a common pharmacophore of acetogenins. Bioenergetic production has been reported to be hindered by the inhibition of mitochondrial complex I. 3,1b,k In fact, 1 displays remarkable profiles of potent growth inhibition of cancer cells. 1,4 Goniocin is the only acetogenin that has an all*trans* tris-THF ring unit in the $C_{10}-C_{21}$ chain. Although the intriguing structure and potent cytotoxic activity of 1 against cancer cell lines at very low concentrations are attractive, this compound has not been fully investigated. Only one landmark total synthesis has been reported by Sinha and Keinan et al. in 1998. 5,6 However, no additional report for the synthesis of 1 has appeared since then. We are interested in the total synthesis of 1

and its biological activity. Herein, we present a short synthetic route to (+)-goniocin along with a new method for the stereocontrolled construction of the bis- and tris-THF ring units.

The contiguous bis-THF ring unit has often been observed in the structure of many acetogenins, 1g-j and in fact, numerous syntheses have been reported for acetogenins possessing a bis-THF ring unit. 7,8 Nonetheless, flexible and efficient stereocontrolled synthesis of the bis-THF ring unit is desired not only for the synthesis of 1 but also for the synthesis of other members of bis-THF acetogenins. Our synthetic plans for the trans-threotrans-bis-THF (IA) and trans-threo-cis-bis-THF (IB) rings are depicted in Scheme 1. The alkene-substituted bis-THF ring of IA could, in principle, be constructed by Pd(II)-catalyzed stereospecific ring formation from chiral ε -hydroxy allylic alcohol IIA. ⁹ This precursor IIA could be derived by cross-metathesis of chiral 1-tetrahydrofurylpentenol III with chiral allylic alcohol possessing an (S)-chiral center. Meanwhile, the trans-cis-bis-THF ring isomer IB could be derived from III via the same two steps, except that (R)-allylic alcohol is used instead of (S)-allylic alcohol. The common intermediate III is provided from the 1alkenyl THF compound V in two steps (ozonolysis followed by stereoselective butenylation). This method would provide an additional stereodefined THF ring unit from a simple alkene V or an aldehyde IV. Thus, the stereodivergent synthesis of the 2alkenyl-5,2'-bis-THF ring from the 2-alkenyl-mono-THF ring could be achieved in four steps.

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Scheme 1. Synthetic Strategy for *trans-trans-* and *trans-cis-*Bis-THF Ring Units IA and IB

Four step reaction sequence for the stereocontrolled elongation of a THF unit

On the basis of this idea, we began the synthesis with (R)-Otritylglycidol 210 (Scheme 2). Ring opening of 2 with allylmagnesium bromide gave 3 in 96% yield, which was subjected to cross-metathesis with (S)-5-phenylpent-1-en-3-ol (S)-4¹¹ in the presence of Grubbs' II catalyst (0.5 mol %) to obtain 5 in 71% yield. Catalytic cyclization of 5 with PdCl₂(MeCN)₂ (5 mol %)⁹ gave the desired trans-THF product 6 in 90% yield along with cis-THF ring isomer 6' in 5% yield. The second THF ring for trans isomer 10 was constructed via the following four steps: (i) ozonolysis of 6 afforded THF-aldehyde 7 in 86% yield; (ii) addition of butenylmagnesium bromide to 7 in THF gave the chelation-controlled product 8 in 51% yield and its diastereomer 8' in 24% yield; 12 (iii) cross-metathesis of 8 with (S)-4 in the presence of Grubbs' II catalyst (0.1 mol %) gave diol 9 in 81% yield; and (iv) Pd(II)-catalyzed cyclization of 9 furnished the trans-threo-trans-bis-THF ring product 10 in 88% yield and the trans-threo-cis ring isomer 11 in 4% yield. However, cross-metathesis of 8 with (R)-4 under the same conditions gave diol the 12 in 84% yield. Pd(II)-catalyzed cyclization of 12 afforded 11 in 86% yield and 10 in 5% yield.

The facial selectivity in the butenylation of 7 was improved through the addition of a chelate complex of 7 with MgBr₂—etherate complex¹³ into an excess of the butenylmagnesium bromide in ether. The ratio of isomers became 6.5:1 favoring of the chelation product 8 (65%) over its isomer 8′ (10%). Additionally, the diastereomeric isomer 8′ was converted into the (R)-isomer 8 in 68% yield by Mitsunobu inversion followed by hydrolysis of the resulting p-nitrobenzoate ester. In the four-step reaction sequence starting from 7 and resulting in either 10 or 11, we efficiently constructed a bis-THF ring system in a stereocontrolled manner. This methodology was then used for the total syntheses of goniocin.

Scheme 3 summarizes our retrosynthetic plan for goniocin. The C_9 – C_{34} unit 13 could be assembled from a right γ -butenolide unit (R)-14¹⁴ by alkene cross-metathesis at the C_8 –

Scheme 2. Stereocontrolled Synthesis of *trans-threo-trans-* and *trans-threo-cis-*Bis-THF Rings

 C_9 bond. The C_{22} -hydroxy group could be created by the addition of a dodecyl unit to the C_{22} -formyl group derived from 15. The consecutive tris-*trans*-THF ring unit in 15 could be installed following the same strategy described for the formation of the bis-THF intermediate 10. The preparation of 10 via the mono-THF intermediate 6 is shown in Scheme 2.

Scheme 4 shows the synthesis of 1 from 10. First, oxidative cleavage of alkenyl bond of 10 by ozonolysis gave aldehyde 16 in 86% yield. Addition of butenylmagnesium bromide to 16 in THF

10 (trans-THF) 5%

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Scheme 3. Retrosynthetic Plan for (+)-Goniocin

gave the desired product 17 and the undesired isomer 17' in a 1:1.2 ratio and in 33% and 37% yields, respectively. However, the formation of a MgBr₂ complex of bis-THF aldehyde 16 and the addition by butenylmagnesium bromide resulted in a worse ratio of 1:1.7, with 17 and 17' in 28% and 47% yields, respectively. This result suggests that chelation control in the addition to the aldehyde was not exerted because of the formation of the MgBr₂ complex with the bis-THF ring of 16 and, instead, control opposite to that expected occurred through the Felkin-Anh model. The influence of the unexpected stereoselectivity by the existence of a neighboring THF group was reported. 15 The undesired (S)-isomer 17' was transformed into the (R)-isomer 17 in 68% overall yield via the Mitsunobu method. Crossmetathesis of 17 with (S)-4 gave 18 in 82% yield. Compound 18 underwent Pd(II)-catalyzed tris-THF ring formation in THF for 20 min at rt, producing 15 in 75% yield along with the cis product 15' in 6% yield. Deprotection of the O-Tr group of 15 followed by Swern oxidation provided the aldehyde 20 via alcohol 19 in 91% overall yield. Addition of the MgBr₂ complex of aldehyde 20 to a large excess of dodecylmagnesium bromide in ether gave the desired isomer 13 in 43% yield and its isomer 13' in 28% yield. 16 Coupling of 13 with an excess of 14 via cross-metathesis gave 21 in 90% yield. HF-promoted deprotection of the O-silyl group and reduction of the alkene with diimide furnished the total synthesis of 1 in 52% yield in two steps. The spectroscopic data for synthetic 1 (i.e., ¹H and ¹³C NMR and mass spectra) are identical to those previously reported in the literature.² In particular, the specific degree of optical rotation was $\left[\alpha\right]^{24}_{D}$ +8.2 (c 0.42, CHCl₃).1

Subsequently, the effect of the synthetic goniocin 1 was measured on the proliferation of four cancer cell lines. The IC50 values of cytotoxicity against HL-60, Paca 2, MCF-7, and A 549 were 2.3, 9.4, 21.8, and 6.6 μ M/mL, respectively.

In conclusion, we have accomplished the total synthesis of (+)-goniocin. The synthesis comprises 17 linear steps from (R)-O-tritylglycidol. The four-step reaction sequence from an alkenyl unit provides short access to the installation of an additional THF unit. The synthetic strategy and technologies developed in this

Scheme 4. Synthesis of (+)-Goniocin

paper can be applied for the synthesis of acetogenins and their analogues possessing a multi-THF ring unit.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.orglett.6b00877.

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Synthesis protocols and characterization of all new compounds including ¹H and ¹³C NMR spectra (PDF)

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Notes

The authors declare no competing financial interest.

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