Total Synthesis of Dictymal, a B-seco-Dictymenoid Aldehyde from an Alga, *Dictyota dichotoma* 

Nobuo KATO,\* Shinya TANAKA,<sup>†</sup> Hideo KATAOKA,<sup>†</sup> and Hitoshi TAKESHITA\* Institute of Advanced Material Study, 86, Kyushu University, Kasuga-koen, Kasuga, Fukuoka 816 <sup>†</sup>Graduate School of Engineering Sciences, 39, Kyushu University, Kasuga-koen, Kasuga, Fukuoka 816

A B-seco-dictymenoid aldehyde, dictymal, was synthesized from two optically active iridoid synthons; the required ring system was constructed by lactol-regulated Cope rearrangement and stereoselective Li-<u>tert</u>-BuOH-in-HMPA reduction of the tetrasubstituted C=C.

Next to the recent achievement<sup>1)</sup> of optically active cycloaraneosene (A), a 5-8-5-membered tricyclic diterpene,<sup>2)</sup> we wish to show the total synthesis of dictymal (1),<sup>3)</sup> a unique B-secoderivative of stereochemically different tricyclic diterpenoid from Dictyota dichotoma Lamouroux. 4) 16 1 Dictymal For the synthesis of 1,<sup>5)</sup> the  $CrCl_2$ -mediated condensation strategy of two iridoid synthons<sup>6)</sup> seems to be applicable on the following retro-synthetic pathway: The relation of Me on C-11 ( f in  ${f B}$  ) and the C3-substituent on C-14 ( g ) is allways cis, and the configuration of C-6 ( c ) and C-14 is inherited from the optically active starting iridoids. This means two (3S)-iridoids must be selected as chair the starting materials. CrC1,

Further, the stereochemistry of C-ll in the Cope rearrangement products can be determined by the geometry of the transition state; i.e., the retention via the boat form and the inversion via the chair form.<sup>7)</sup>

However, the condensates of two  $(3\underline{S}^*)$ - or  $(3\underline{R}^*)$ -iridoids are known to cause the rearrangement via both boat and chair transition states,<sup>8)</sup> and since the correct configuration of C-ll for 1 can only be generated via a chair transition Cope rearrangement of the condensate, it is desirable to control the transition state of the rearrangement as the chair form. "The lactol-regulated Cope rearrangement" is satisfactory for the purpose; an incorporation of the 1,5-diene to a part of cyclic systems should give only the thermolysate having <u>E</u>-formed enol ether. Another point in this strategy, chemical reduction of the isolated tetrasubstituted C=C in A ring, could be operated with a slightly modified procedure employed previously.<sup>3)</sup>



a: CrCl<sub>3</sub>-LAH/THF-DMF(**4**, 62%, **5**, 17%); b: TMSC1/Py(85%); c: i) Disiamylborane, H<sub>2</sub>O<sub>2</sub>/OH<sup>-</sup>(96%), i1) PCC(84%); d: KF-Florisi1/MeOH(84%); e: PPTS/aqTHF, TMSC1/Py(**7--8**, 45%; **9-10**a+1**0**b, 76% (2:1)); f: 180 °C/C<sub>7</sub>H<sub>8</sub>(**10**a-**-**11a+11b, 81%(2:1), **10**b-**-**11a+11b, 73%(1:4)); g: i) PPTS/aqTHF(95%), i1) NaBH<sub>4</sub>/aqNaHCO<sub>3</sub>-MeOH(93%); h: i) Ac<sub>2</sub>O/Py(99%), i1) H<sub>2</sub>/Pd-C(98%), i1i) DHP, PPTS/CH<sub>2</sub>Cl<sub>2</sub> (99%), iV) LAH/THF(97%); i: L1-Me<sub>3</sub>COH/HMPA(83%); j: BuLi, Me<sub>3</sub>CCOC1/THF(**15**; 47%, **16**; 11%; **17**, 40%); k: LAH/THF(98%); 1: TsOH/MeOH(**18**a, 52%; **18**b, 37%); m: i) <u>o</u>-nitrophenylselenocyanate, Bu<sub>3</sub>P/THF (97%), i1) H<sub>2</sub>O<sub>2</sub>/THF(86%); n: LAH/THF(95%); o:(COC1)<sub>2</sub>-DMSO, Et<sub>3</sub>N/CH<sub>2</sub>Cl<sub>2</sub>(95%).

The starting iridoids,  $(3\underline{S}, 8\underline{R})$ -9-benzyloxy-7-chloro-1-iridene (**2**) and  $(3\underline{S})$ -1,8-iridadien-7-al (**3**),<sup>1)</sup> were treated with CrCl<sub>2</sub> in tetrahydrofuran (THF)-dimethylformamide (DMF) to give a condensate (**4**), 62% yield, and its epimer

## ( **5** ), 17% yield.<sup>9)</sup>

Compound 4 was converted to the corresponding trimethylsilyl (TMS) ether (6),<sup>10</sup>) and further to an aldehyde (7). The cyclic TMS acetal (8), obtained from 7 was subjected to the Cope rearrangement; however, 8 afforded only a trace amount of the thermolysate probably due to a steric hindrance from the axiallyoriented Me on C-15 preventing the overlapping of the 1,5-diene terminals in the transition state. Therefore, 7 was converted to epimerized aldehyde (9), and further to a mixture of TMS acetals (10a and 10b).

When 10a and 10b were independently heated under the Cope rearrangement conditions, a same mixture of 11a and 11b was formed with different ratio, indicating an epimerization at the acetal carbon. Since the same glycol ( 12 ) was obtained from 1 1a and 1 1b by consecutive treatments with PPTS and NaBH, no separation of mixtures of 10 or 11 was necessary. Compound 12 possessed the correct configuration as the ring C found in 1. The tetrahydropyranyl ( THP )-ether (13), prepared from 12 via a four-step sequence, was treated with Li and tert-BuOH in hexamethylphosphoric triamide ( HMPA ) to give a mixture of two isomeric dihydro derivatives (14). In order to partially protect the hydroxyl groups, 14 was treated with pivaloyl chloride to obtain a monoester (15) accompanied by its isomer (16) and diester (17), which were reduced to 14. As a stereoisomeric mixture, 15 was treated with p-toluenesulfonic acid ( TsOH ) to afford the diols ( 18a and 18b ), which were separated by silica-gel column chromatography ( CHCl<sub>2</sub>-AcOEt ). The major isomer 18a was indeed the desired compound judging from the <sup>13</sup>C NMR spectroscopy.<sup>11)</sup> Dehydration of 18a with o-nitrophenylselenocyanate<sup>12)</sup> gave a diene (19), which afforded an alcohol (20).

The Swern's oxidation of **20** yielded an aldehyde  $[\alpha]_D^{29} + 24^{\circ}$  (c 0.46, CHCl<sub>3</sub>) (1it.<sup>6)</sup>  $[\alpha]_D^{18} + 16.4^{\circ}$  (c 0.88, CHCl<sub>3</sub>)) ] whose NMR  $[\delta(H)^{C}6^{D}6=0.87(3H, s), 0.94$  (3H, d, J=6.5 Hz), 1.56(3H, m), 1.62(3H, m), 2.35(1H, dd, J=10, 3 Hz), 2.51(1H, m), 3.00(1H, m), 4.68(1H, m), 4.76(2H, m), 4.87(1H, m), and 9.55(1H, d, J=3 Hz).  $\delta(C)^{CDC1}3=20.5, 21.9, 22.5, 22.9, 28.5, 28.9, 31.8, 39.8, 40.4, 41.2, 45.9$  (2C), 47.8, 50.2, 64.8, 110.2, 112.1, 146.7, 147.0, and 205.5] were consistent with the natural product (1).

Thus, the total synthesis of this unique metabolite is now completed.  $^{14)}$ 

## References

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- 7) For easy understandings, a figure showing stereochemical relationship of the

boat/chair transition geometry and the configurations of the Cope products.



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- 9) The major product in our previous report was 5.<sup>8)</sup> The condensation under careful exclusion of air and moisture, however, proceeded in favor of desired 4 giving a reversed ratio of products. Details will be discussed elsewhere.
- 10) All the new compounds have been fully characterized. The <sup>1</sup>H NMR data ( in CDCl<sub>3</sub> ) of key compounds were shown as follows:
  - 9: δ=0.03(9H, s), 0.94(3H, s), 0.98(3H, d, J=7 Hz), 1.11(3H, d, J=7 Hz), 1.86(3H, br s), 2.70(1H, m), 3.20(1H, t, J=9 Hz), 3.24(1H, br m), 3.48(1H, dd, J=9, 4 Hz), 4.20(1H, br s), 4.42(1H, d, J=12 Hz), 4.48(1H, d, J=12 Hz), 4.86(2H, d, J=2.5 Hz), 7.27(5H, br s), and 9.65(1H, s).
  - 10a: δ=0.09(9H, s), 0.83(3H, d, J=7 Hz), 1.09(3H, d, J=7 Hz), 1.13(3H, s), 1.73
    (3H, br s), 3.19(1H, t, J=9 Hz), 3.52(1H, dd, J=9, 4 Hz), 3.74(1H, br s), 4.25
    (1H, d, J=8 Hz), 4.43(1H, d, J=12 Hz), 4.50(1H, d, J=12 Hz), 4.81(1H, d, J=2
    Hz), 4.90(1H, d, J=2 Hz), and 7.28(5H, br s).
  - 11a:  $\delta$ =0.18(9H, s), 1.00(3H, s), 1.01(6H, d, J=7 Hz), 4.77(1H, d, J=8 Hz), 6.13 (1H, d, J=2 Hz).
  - 12: δ=0.73(3H, s), 0.89(3H, d, J=7 Hz), 1.02(3H, d, J=7 Hz), 1.59(3H, br s), 2.70 (1H, br m), 3.09(1H, t, J=9 Hz), 3.27(1H, dd, J=9, 4 Hz), 3.29(1H, dd, J=11, 8.5 Hz), 3.42(1H, dd, J=10.5, 5 Hz), 3.61(1H, dd, J=11, 4.5 Hz), 3.71(1H, dd, J= 10.5, 5 Hz), 4.38(1H, d, J=12 Hz), 4.43(1H, d, J=12 Hz), and 7.27(5H, br s).
    18a: δ=0.87(3H, d, J=7 Hz), 0.92(3H, s), 0.95(3H, d, J=7 Hz), 0.96(3H, d, J=6 Hz), 1.20(9H, s), 3.41(1H, dd, J=10.5, 7 Hz), 3.48(2H, d, J=7 Hz), 3.70(1H, dd, J=10.5, 3 Hz), 4.03(1H, dd, J=11.5, 5.5 Hz), and 4.10(1H, dd, J=11.5, 7 Hz).
    19: δ=0.90(3H, s), 1.00(3H, d, J=6.5 Hz), 1.17(9H, s), 1.68(3H, br s), 1.70(3H, br s), 2.46(1H, m), 2.57(1H, m), 3.99(2H, d, J=7 Hz), 4.65(1H, m), 4.66(1H, m), 4.71(1H, br s), and 4.82(1H, br s).
  - **20**: δ=0.85(3H, s), 1.00(3H, d, J=6.5 Hz), 1.70(3H, br s), 1.75(3H, br s), 2.54 (1H, m), 2.59(1H, m), 3.59(1H, dd, J=11, 7.5 Hz), 3.66(1H, br d, J=11 Hz), 4.66 (1H, br s), 4.70(1H, m), 4.82(1H, br s), and 4.85(1H, m).
- 11) C-1 and C-16 of  $18_a$  appeared at  $\delta=39.7$  and 22.8, respectively, while those of  $18_b$  appeared at 50.8 and 21.7. See Ref. 3.
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- 13) We are grateful to Professor Haruhisa Shirahama, Hokkaido University, who kindly performed the identification of both compounds and sent us a copy of manuscript prior to the publication.
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