Hokkaido Institute of Public Health N-19, W-12, Kita-ku, Sapporo, 060, Japan

Faculty of Pharmaceutical Sciences, Kyushu University 3-1-1 Maidashi, Higashi-ku Fukuoka, 812, Japan

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Makoto Nishizawa Takashi Yamagishi

Gen-ichiro Nonaka Itsuo Nishioka

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Synthesis of (23R)-Calcidiol Lactone (25-Hydroxyvitamin D₃ 26,23-Lactone)¹⁾

(23R)-Calcidiol lactone(25-hydroxyvitamin D_3 26,23-lactone) was synthesized from bisnorcholenic acid. The configuration at C-23 was determined by transformation to 22- and 23-hydroxycholesterols.

Keywords—calcidiol lactone; 25-hydroxyvitamin D_3 26,23-lactone; metabolite of vitamin D_3 ; 22-hydroxycholesterol; 23-hydroxycholesterol

Calcidiol lactone²⁾ is a new metabolite of Vitamen D_3 . Although the preliminary tests of this metabolite showed interesting biological activity,^{2,3)} the configuration at C-23 and C-25 positions as well as the biological role of this metabolite are unknown yet. In order to confirm the reported structure and to elucidate the configuration at C-23 and C-25, it is necessary to synthesize four possible isomers. It would be also urged to prepare the natural calcidiol lactone for biological investigation. We describe herein a synthesis of (23R)-calcidiol lactone.

The 22-aldehyde 3-THP ether 1^4) derived from 22,23-bisnorcholenic acid was coupled with vinylmagnesium bromide to give a mixture of the 22-alcohols (2) in a 6: 1 ratio. The less polar major alcohol, mp 155—156°, possesses the 22*R*-configuration according to the precedents for this mode of reaction.⁵⁾ The 22-alcohol 2 was reacted with ethyl orthopropionate and propionic acid as catalyst in refluxing xylene to give 22,23-trans 26-ethyl ester 3a [mp 105—107.5°; NMR δ 1.11 ppm (3H, d, J=6.6 Hz, 27-H₃), 5.20—5.50 (3H, m, 6-H, 22-H, 23-H); IR 1720 cm⁻¹] in 96% yield. The 26-ester 3a was hydrolysed to the 3-hydroxy-26-acid 3b, mp 175—178°, by HCl-methanol and then KOH-methanol treatment. Iodolactonization⁶) of the acid with iodine in acetonitrile at -0° gave regio- and stereoselectively a single product 4a [mp 220—224°; 94% yield; NMR δ 1.28 (3H, d, J=6 Hz, 27-H₃), 2.50—3.10 (1H, m, 25-H), 4.10 (1H, dd, J=5.1, 5.7 Hz, 22-H), 4.60 (1H, m, 1/2w, 30 Hz, 23-H); IR 1768 cm⁻¹]. The iodolactone 4a was then reduced by freshly distilled tributyltinhydride in dry THF at room temperature to the lactone 5a [mp 223—224°; 85% yield; NMR δ 1.27 (3H, d, J=7.5 Hz, 27-H₃), 4.52 (1H, m, 1/2w 26 Hz, 23-H); IR 1760 cm⁻¹; CD (dioxane) [θ]²⁰ (nm): -19 (249) (negative maximum).

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The C-23 configuration of the lactone 5a was determined as R by transformation of the compound 4a into 22- and 23-hydroxycholesterols as follows. The iodolactone 3-THP ether 4b was converted to 22,23-epoxy-26-methyl ester 6 (mp 184— 185°) with sodium carbonate in methanol. Reduction of the ester with LiAlH₄ in THF at room temperature gave 22,23-epoxy-26-ol 7a. After iodination of the 26-ol by treatment with TsCl-pyridine and then NaI-acetone (reflux), the iodide 7b was reduced with tributyltinhydride to 26-methyl compound 7c. Reduction of the epoxide with LiAlH₄ and subsequent removal of the 3-THP group afforded a 1:1 mixture of 22- and 23-hydroxycholesterols. These hydroxycholesterols were identical with the authentic samples of (22R)-22-hydroxycholesterol $(8)^{7}$ and (23S)-23-hydroxycholesterol $(9),^{8}$ respectively, with respect to the retention times on GLC⁹ and HPLC.¹⁰

Introduction of a hydroxy group at C-25 position was achieved by oxidation with MoOPH (MoO₅·Py·HMPA) of the enolate of lactone generated by lithium diisopropylamide¹¹⁾ in THF at -78° to give 25-hydroxy-lactone 10a [mp 184—193°, 65% yield; NMR δ 1.47 (3H, s, 27-H₃);

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⁹⁾ The retention times on GLC using glass capillary column, $30 \text{ m} \times 0.25 \text{ mm}$ i.d.; (22R)-22-OH (8) bis-trimethylsilyl ether (TMSi), 37.11 min; (22S)-22-OH bis-TMSi, 38.11 min; (23R)- and (23S)-23-OH (9) bis-TMSi, 41.22 min.

¹⁰⁾ The retention times on HPLC using Zorbax SIL, 15 cm \times 4.6 mm i.d., hexane–CH₂Cl₂ (6: 1), 20 kg/cm²; (22R)-22-OH (8) bis-(+)- α -methoxy- α -trifluoromethyl-phenylacetate (MTPA), 7.4 min; (22S)-OH bis-MTPA, 9.0 min; (23R)-23-OH bis-MTPA, 9.4 min; (23S)-23-OH (9) bis-MTPA, 7.8 min.

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IR 1762 cm⁻¹; CD (dioxane) $[\theta]^{20}$ (nm) -49510 (230) negative maximum¹²⁾ as a single product. After conversion of the 3-THP ether to 3-acetate, transformation of 10b (mp 224—228°) into calcidiol lactone 12 was carried out by the standard procedure. 13) Bromination with N-bromosuccinimide in CCl₄, followed by debromination (collidine in refluxing xylene) gave a mixture of the 5,7-diene and 4,6-diene. Treatment with p-toluenesulfonic acid to convert the 4,6-diene into the much less polar 2,4,6-triene, purification by preparative TLC, alkaline hydrolysis of the acetyl group and then acidification gave the 5,7-diene 11 (30% yield, λ_{max} 262, 271, 281.5 and 293 nm). Irradiation of the diene 11 with a medium pressure mercury lamp in benzene-EtOH solution at 0° for 2.5 min, refluxing for 1 hr in the same solvent and purification by preparative TLC afforded the (23R)-calcidiol lactone 12 [23% yield, λ_{\min} 226, λ_{\max} 265 nm; MS m/e: 428 (M⁺), 410, 395, 369, 271, 253, 211, 199, 197, 183, 171, 159, 158, 143, 136, 118]. The UV and MS spectra of the synthetic calcidiol lactone were superimposable with the data reported by DeLuca.²⁾ The signals in NMR spectrum [CDCl₃, 200 MHz; δ 0.58 (3H, s, 18-H₃), 1.05 (3H, d, J = 5 Hz, 21-H₃), 3.99 (1H, m, 3α -H), 4.86 and 5.10 (2H, s pair of broad s, 19-H), 6.07 (1H, d, I=12 Hz, 7-H), 6.28 (1H, d, I=12 Hz, 6-H) are almost identical with the data reported for the natural metabolite, except one proton signal at δ 4.78(m) corresponding to C-23 proton, instead of the signal at δ 4.46.2 This might suggest that the natural calcidiol lactone has 23S-configuration. However, it cannot be ruled out that this shift may be due to the difference of the C-25 stereochemistry.

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Laboratory of Chemistry for Natural Products, Tokyo Institute of Technology, Nagatsuta, Midori-ku, Yokohama 227, Japan

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Nobuo Ikekawa Yutaka Hirano Masaji Ishiguro¹⁴⁾ Jun-ichi Oshida Tadashi Eguchi Satoru Miyasaka

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14) Present address: Suntory Institute for Biomedical Research, Suntory Ltd., Wahayamadai, Shimamoto-cho, Mishima-gun, Osaha.