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The Pentaoxa[5] peristylanes. A Novel Oxa-Cage System

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Abstract: The synthesis of pentaoxa[5] peristylanes, a novel oxa-cage system, has been accomplished via ozonolysis of 7-anti-2,3-bis-endotriacylbicyclo[2.2.1]-5-heptenes and via a direct chemical transformation of the tetraacetal tetraoxa-cages 5a-c and 6a-c. © 1997 Elsevier Science Ltd.

The synthesis of peristylanes, such as [5]peristylane¹ and [4]peristylane,² has been accomplished and attempts to roof [5]peristylane has been made.³ On the other hand, the synthesis of heterocyclic analogs of peristylanes has received much less attention.⁴ Recently, we conceived that some heterocyclic cage systems might be viewed as novel classes of cagebackboned coronands (crown ethers) and might exhibit interesting cation-binding properties. We also visualized that the "creation" of oxa-cage compounds from carbocyclic cages might be achieved by replacing the skeletal carbon atoms with oxygen atoms at the proper positions and by extending the skeletal backbone.⁵ Thus, we have accomplished the synthesis of tetraacetal tetraoxa-cages,⁵ tetraacetal pentaoxa-cages,⁵ diacetal trioxa-cages,⁵ and triacetal trioxa-cages. We report in this communication the synthesis of pentaoxa[5]peristylanes, a novel oxa-cage system, via ozonolysis of 2,3-bis-endo-7-anti-triacylnorbornenes. We also wish to demonstrate for the first time the direct transformation of tetraacetal tetraoxa-cages to pentaoxa[5]-peristylanes.

Diels-Alder reaction of compound 1^{10} with *cis*-enediones 2a- $f^{5,6}$ in dichloromethane at 0 °C for 72 h gave the *anti-endo* adducts 3a-f in 70-75% yields. Treatment of 3a-f with $Cu(BF_4)_2$ or methanesulfonic acid in dichloromethane at 25 °C gave the hydrolysis products 4a-f in 75-80% yields (Scheme 1). Compounds 4g-i were prepared from (Z)- γ -oxo- α , β -unsaturated thioesters

at -78 °C followed by reduction with dimethyl sulfide gave the tetraacetal tetraoxa-cages 5a-c (30-34%) and 6a-c (34-38%) and the pentaacetal pentaoxa-cages 7a-c (18-22%), the pentaoxa[5]-peristylanes. Ozonolysis of 4d-f under the same reaction conditions gave the pentaacetal pentaoxa-cages 7d-f in 75-80% yields. The by-products 5d-f and 6d-f were too small amount to be isolated. Ozonolysis of 4g-i under the same reaction conditions gave the tetraacetal tetraoxa-cages 8a-c in 85-90% yields. No detectable amount of the pentaoxa[5]peristylanes 9a-c or the tetraacetal tetraoxa-cages 10a-c or 11a-c was obtained. The thioester group may exhibit much less reactive than the acyl groups for the cyclization reaction.

Scheme 1

Compounds 7a-f are white solid.¹¹ The IR spectra of 7a-f lacked carbonyl absorptions and showed strong absorptions near 1050 cm⁻¹ for the ether C-O bonds. The ¹H NMR spectrum of 7a revealed one doublet at δ 5.91 for the acetal proton on C-5 and one doublet at δ 5.85 for the two acetal protons on C-3 and C-7. The absorption at δ 2.09 (a singlet) for the methyl ketone protons of 4a shifted to δ 1.50 for the angular methyl protons of 7a. The ¹³C NMR spectrum of 7a lacked any carbonyl absorption and displayed one peak at δ 113.33 for the acetal carbon C-5, one peak at δ 112.46 for the acetal carbons C-3 and C-7, one singlet at δ 120.30 for the quaternary carbons, and one peak at δ 26.94 for the angular methyl carbons. The IR spectra and ¹H and ¹³C NMR spectra of 7b-f revealed that these compounds possess the same skeleton as 7a.

Treatment of the tetraacetal tetraoxa-cages **5a-c** with catalytic amount of TiCl₄ in dichloromethane at 25 °C for 4 h gave the pentaoxa[5]peristylanes **7a-c** in 70-75% yields and the hydride rearrangement products **12a-c** in 20-15% yields (Scheme 2). Reaction of **6a-c** under the same reaction conditions gave **7a-c** in 85-90% yields. The amount of **12a-c** was too small to be isolated.

Scheme 2

$$5a-c \xrightarrow{\text{TiCl}_4} 7a-c + R \xrightarrow{\text{CH}_2\text{Cl}_2} 7a-c$$

$$12a-c \xrightarrow{\text{CH}_2\text{Cl}_2} 7a-c$$

Thus, we have accomplished for the first time the synthesis of pentaoxa[5]peristylanes, a novel and interesting oxa-cage system.

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- (11) Selected spectral data for 7. 7a: white solid; mp 214-215 °C; ¹H NMR (300 MHz, CDCl₃) δ 5.91 (d, J = 5.4 Hz, 1H), 5.85 (d, J = 5.1 Hz, 2H), 3.70-3.65 (m, 3H), 3.39-3.35 (m, 2H), 1.50 (s, 6H); ¹³C NMR (75 MHz, CDCl₃) δ 120.30 (2C), 113.33 (CH), 112.46 (2CH), 62.72 (2CH), 58.91 (2CH), 58.53 (CH), 26.94 (2CH₃); MS m/z (rel int.) 238 (M⁺, 12), 208 (100). 7e: white solid; mp 178-179 °C; ¹H NMR (300 MHz, CDCl₃) δ 5.90 (d, J = 6.0 Hz, 2H), 5.86 (d, J = 5.1 Hz, 2H), 3.68-3.62 (m, 4H), 3.38-3.34 (m, 1H), 1.76-1.70 (m, 2H), 1.40-1.26 (m, 4H), 0.92 (t, J = 6.9 Hz, 3H); ¹³C NMR (75 MHz, CDCl₃) δ 123.59 (C), 113.51 (2CH), 112.81 (2CH), 59.96 (CH), 58.76 (2CH), 58.27 (2CH), 39.30(CH₂), 26.04(CH₂), 22.66 (CH₂), 13.98 (CH₃); MS m/z (rel int.) 266 (M⁺, 14), 209 (100).