Production of a 15-membered Ring Orifice in Open-cage Fullerenes by Double Photooxygenation of Azafulleroid

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Photooxygenation of silyl-modified azafulleroids **1b** and **1c** gave open-cage fullerenes having keto lactam **2b** and **2c** or diketo imide **3b** and **3c** functional groups on 11- or 15-membered ring orifices. The high reactivity of the bridgehead double bonds was estimated by π -orbital axis vector (POAV) analysis comparing the bisfulleroid derivatives **5d**, **5e**, **6d**, and **6e**.

Among surface modification reactions of C₆₀, singlet oxygen $({}^{1}O_{2})$ oxygenation of azafulleroid and bisfulleroid is the key reaction in the production of open-cage-,¹ hetero-,² and endohedral-fullerenes.³ The [2 + 2] addition of ${}^{1}O_{2}$ to the electron-rich bridgehead double bonds of these fulleroids, followed by symmetrical ring opening, results in the opening of a hole in the fullerenes surface.^{1a,1d-1i} When the first cage-opening reaction of azafulleroid 1a was reported by Wudl and coworkers,^{1a} diketo imide **3a** was suggested as an alternative possible structure in addition to the keto lactam 2a, as shown in Scheme 1. However, the diketo imide 3a has remained unobserved to date. Recently, we reported the selective synthesis of silvl benzene azafulleroids.⁴ⁱ Herein, we report the photooxygenation of these silvl-modified azafulleroids 1b and 1c undergoing stepwise oxidation to afford the first diketo imide 3b and 3c via the keto lactam 2b and 2c as shown in Scheme 1. It was demonstrated that whether the double photooxygenation of azafulleroid and the related bisfulleroid takes place or not can be predicted by π -orbital axis vector (POAV) analysis⁵ of the bridgehead double bonds.

A solution of either **1b** or **1c** in *o*-dichlorobenzene in a pyrex tube was irradiated with a halogen lamp under bubbling oxygen for 5 h. The ¹H NMR signals of methylene protons for **1b** and **1c** at 4.23 and 3.44 ppm disappeared completely (**1b** is shown in Figure 1). The new signals for **2b** and **2c** at 4.52/5.68 and 3.83/4.93 ppm with doublet couplings were observed together with aziridinofullerenes **4b** and **4c** at 4.03 and 3.25 ppm (**2b** and **4b** are shown in Figure 1). After column chromatography through silica gel (eluent: CS₂), **2b** and **2c** were obtained in 85 and 90% yields, as dark brown solid together with 11 and 10% yield of the aziridinofullerene **4b** and **4c**, formed by the photo-



Scheme 1.



Figure 1. ¹H NMR spectra in CDCl₃ solution; (a) 1b, (b) 2b, (c) 3b, and (d) 4b.



Figure 2. MALDI-TOF mass spectra of 2b (left) and 3b (right) in negative-ion reflectron mode.

chemical isomerization of **1b** and **1c**.⁴ The MALDI-TOF mass spectra of **2b** and **2c** show molecular ion peaks at m/z 1039 (M⁻) and 853 (M⁻) (**2b** is shown in Figure 2), indicating that the product was formed by the addition of O₂ to **1b** and **1c**. The ¹³C NMR spectra display two carbonyl carbons at 161/194 and 161/194 ppm for keto lactams **2b** and **2c**. In the fullerene skeleton regions, 59 and 56 signals were observed, respectively, indicating C_1 symmetry for **2b** and **2c**. The IR spectra show two pairs of carbonyl groups at 1682/1726 and 1681/1726 cm⁻¹, respectively.

Next, a solution of either **1b** or **1c** in *o*-dichlorobenzene in a pyrex tube was irradiated for 30 h. The reaction mixtures were monitored by ¹H NMR spectroscopy. During the prolonged irradiation, the signals for **2b** and **2c** decreased in intensity with a concomitant increase in the singlet peaks at 5.24 and 4.49 ppm (**3b** is shown in Figure 1). After chromatographic separation (eluent: CHCl₃), **3b** and **3c** were isolated in 9 and 11% yields, with 58 and 44% yields of **2b** and **2c** recovered.⁶ The final products are suggested to be the diketo imide **3b** and **3c** on the basis of the observed ¹H and ¹³C NMR and MALDI-TOF mass spectra. The molecular ion peaks at m/z 1071 (M⁻) and 885 (M⁻)



Figure 3. Partial atomic numbering of 1a–1c, 2a–2c, 5d, 5e, 6d, and 6e.

Table 1. The π -orbital misalignment angles (Φ°) of **1a–1c**, **2a–2c**, **5d**, **5e**, **6d** and **6e** (calculated geometries^a)

Fulleroid/ketolactam	1 (C1–C9, C5–C6)	2 (C1–C9)
a : $R_1 = CH_2OCH_2CH_2OCH_3$	29.7, 30.1	28.8
b : $\mathbf{R}_1 = \mathbf{C}\mathbf{H}_2\mathbf{S}\mathbf{i}\mathbf{P}\mathbf{h}_3$	30.2, 30.7	29.5
\mathbf{c} : $\mathbf{R}_1 = \mathbf{C}\mathbf{H}_2\mathbf{S}\mathbf{i}\mathbf{M}\mathbf{e}_3$	30.1, 30.3	29.6
Bisfulleroid/diketone	5 (C5–C6, C7–C8)	6 (C7–C8)
d : $E = N, R_2 = Py, R_3 = Ph$	29.5, 29.9	14.1
$\mathbf{e}: \mathbf{E} = \mathbf{C}\mathbf{C}\mathbf{O}_2\mathbf{M}\mathbf{e}, \mathbf{R}_2 = \mathbf{R}_3 = \mathbf{C}\mathbf{O}_2\mathbf{M}\mathbf{e}$	29.3, 29.7	13.6

^aAll geometries optimized using the B3LYP/6-31G** basis set.

are due to the addition of O_2 to **2b** and **2c** (**3b** is shown in Figure 2). The ¹³C NMR spectra show two pairs of carbonyl carbons at 161/188 and 162/189 ppm with 32 and 32 signals in the fullerene skeleton regions supporting C_s symmetry for **3b** and **3c**. The IR spectra show two carbonyl groups at 1651/1744 and 1647/1743 cm⁻¹, respectively. The observed stepwise reaction of molecular oxygen with the azafulleroid observed is attributed to the self-sensitive ability of C_{60} and its derivatives.⁷ Comparing **1b**, **1c** to **2b**, **2c**, the fullerene first acts as an O_2 sensitizing agent in ambient light and then reacts with the ¹O₂ generated to give **2b**, **2c** and **3b**, **3c** respectively. Of note is the fact that in order to make the second oxidation more facile, when using C_{60} or dyes (TPP or rose bengal) as sensitizer, nothing need be changed.

The structures of the products having been established, the reactivity of the bridgehead double bond (1a-1c and 2a-2c) toward ${}^{1}O_{2}$ was estimated by the POAV misalignment angles (Φ°) ⁵ as shown in Figure 3 and Table 1. These values are defined as the two-center dihedral angles between the two carbon dihedral angles on bonded pairs of carbons. Komatsu and Murata have reported the single photooxygenation of bisfulleroids 5d and 5e to afford diketones 6d and 6e in spite of the bridgehead double bond remaining. The geometries utilized for the azafulleroids 1a-1c, keto lactams 2a-2c, bisfulleroid 5d and 5e, and diketone 6d and 6e were obtained at the B3LYP/6-31G** level of theory. The misalignment angles of the bridgehead double bonds of **1a–1c**, **5d** and **5e** are far larger than in C_{60} ($\Phi = 0^{\circ}$), as shown in Table 1. However, in the keto lactams 2a-2c, the misalignment angle of the remaining C1-C9 double bond is similar to those of **1a–1c**.⁸ In the diketones **6d** and **6e** the angle of the remaining C7-C8 double bond is far less than those of 5d and 5e. However, it would appear that some elaboration is in order.

In conclusion, we have demonstrated that the double photooxygenation of silyl-modified azafulleroid **1b** and **1c** provides the first diketo imide **3b** and **3c** via the keto lactam intermediate **2b** and **2c**. The sizes of 15-membered ring orifice of **3b** and **3c** were theoretically estimated to be 5.7 Å along the long axis and 3.3 Å along the short axis.⁶ The reactivity of the bridgehead double bond of the fulleroid and related bisfulleroid were estimated by POAV misalignment angles to predict the difference between the keto lactam 2a-2c and the diketones 6d and 6e.

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- 8 Our preliminary experiment of double photooxygenation of 1a was carried out at the same condition of 1b and 1c for 30 h. The resulting crude product is suggested to be a mixture of keto lactam 2a and diketo imide 3a observed by ¹H NMR and MALDI-TOF mass spectra. This preliminary result to be published, disclosed the remaining C1–C9 double bond of 2a having the same reactivity as 2b and 2c predicted by POAV analysis.