

1,1,3,3-Tetramethylindantellone, the First Telluroketone Stable in Solution

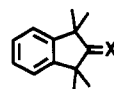
Mao Minoura, Takayuki Kawashima, and Renji Okazaki*

Department of Chemistry, Faculty of Science
The University of Tokyo, Hongo 7-3-1
Bunkyo-ku, Tokyo 113, Japan

Received April 23, 1993

In recent years the chemistry of multiple-bond compounds of heavier typical elements has been a subject of continuing interest.¹ Among these are compounds with a carbon–chalcogen double bond like thio- and selenoketones² (or aldehydes³), which have been intensively studied and include some stable species. For a telluroketone, however, neither the synthesis of a stable species nor the spectroscopic observation of such has been described,⁴ although some scattered examples of stable telluroesters^{5a–c} and telluroamides^{5d–f} as well as the transition-metal complexes of telluroketones (or aldehydes)⁶ have been reported.

We now report the first example of a telluroketone, 1,1,3,3-tetramethylindantellone (**1a**), stable in solution at ambient temperature, which enables the first observation of interesting



1a: X=Te 1c: X=S
1b: X=Se 1d: X=O

spectroscopic properties (¹³C, ¹²⁵Te NMR, and UV/vis) and novel reactivities (1,3-dipolar cycloaddition and ene reaction) of a genuine telluroketone.

We took advantage of the thermal cycloreversion of a novel heterocycle, 1,3,4-telluradiazoline (**2**),⁷ for the synthesis of a telluroketone (Scheme I). Although **2** was stable at 80 °C for 6 h in the solid state without any appreciable decomposition, it gradually decomposed in solution even at room temperature. A carefully degassed CDCl₃ solution of **2** (ca. 0.02 M) in a sealed tube was subjected to thermolysis in the NMR probe. The thermolysis proceeded best around 80 °C although it was carried out in the temperature range of 60–100 °C. The reaction could similarly be performed in benzene. Monitoring by ¹H NMR revealed the completion of the thermolysis for about 4 h and the quantitative formation of telluroketone **1a** and olefins **4** and **5** produced from diazo compound **3**, indicating clean cycloreversion of **2** into **1a** and **3**.⁸

Surprisingly, **1a** is very stable in a dilute solution prepared in the above-mentioned way, surviving after heating at 80 °C for as long as 6 h. However, **1a** is extremely sensitive to oxygen and light, decomposing to give the corresponding ketone **1d** and ditelluride **6**, respectively. The formation of **1a** was confirmed by the following spectral data and chemical reactivities: ¹H NMR δ (CDCl₃) 1.61 (s, 12 H), 7.30 (s, 4 H); ¹³C NMR δ (CDCl₃) 30.3 (q), 79.4 (s), 124.0 (d), 127.9 (d), 147.2 (s), 301.0 (s); ¹²⁵Te NMR δ (CDCl₃) 2858.3; UV/vis λ_{max} (CHCl₃) 825 (ε 100) nm.⁹

The tellurocarbonyl carbon resonates at 301 ppm, which is, to our knowledge, the most deshielded carbon ever observed in a neutral molecule.^{2a,10} This signal is shifted significantly downfield relative to those for C = X (X = Se, S, O) of the other analogs **1b** (δ 294.0),^{11a} **1c** (δ 282.1),^{11b} and **1d** (δ 226.0)^{11a} and for C = Te of tellurocarbonyl compounds perturbed by the mesomeric effect of α-heteroatoms [e.g., *t*-Bu(C=Te)OCH₂(*t*-Bu) δ 229.4,^{5a,b} *t*-Bu(C=Te)OSiMe₃ δ 251.2^{5c}].

The ¹²⁵Te NMR shift of **1a** (2858 ppm) is also unique, being at much lower field than that of a telluroester [e.g., *t*-Bu(C=Te)OSiMe₃ δ 1418].^{5c} This is the most deshielded ¹²⁵Te signal so far observed for organotellurium compounds.¹²

The visible spectrum of **1a** (CHCl₃, 80 °C) showed λ_{max} 825 nm,¹³ which disappeared gradually during several hours even by intermittent irradiation with monitoring light of the spectrometer resulting in the appearance of a new absorption of ditelluride **6** at 400 nm. The absorption at 825 nm, most likely assignable to

(1) For reviews on multiple bond compounds containing group 14 and 15 elements, see the following. Group 14: Wiberg, N. *J. Organomet. Chem.* **1984**, 273, 141–177. Raabe, G.; Michl, J. *Chem. Rev.* **1985**, 85, 419–509. Brook, A. G.; Baines, K. M. *Adv. Organomet. Chem.* **1986**, 25, 1–44. West, R. *Angew. Chem., Int. Ed. Engl.* **1987**, 26, 1201–1211. Raabe, G.; Michl, J. In *The Chemistry of Organosilicon Compounds*; Patai, S., Rappoport, Z., Eds.; John Wiley & Sons: New York, 1989; pp 1015–1142. Barrau, J.; Escudé, J.; Satgé, J. *Chem. Rev.* **1990**, 90, 283–319. Tsumuraya, T.; Batcheller, S. A.; Masamune, S. *Angew. Chem., Int. Ed. Engl.* **1991**, 30, 902–930. Group 15: Cowley, A. H.; Norman, N. C. *Prog. Inorg. Chem.* **1986**, 34, 1–63. Lochschmidt, S.; Schmidpeter, A. *Phosphorus Sulfur* **1987**, 29, 73–109. Regitz, M. *Chem. Rev.* **1990**, 90, 191–213. *Multiple Bonds and Low Coordination in Phosphorus Chemistry*; Regitz, M., Scherer, O. J., Eds.; George Thieme: New York, 1990.

(2) (a) Duss, F. In *Comprehensive Organic Chemistry*; Barton, D. H. R., Ollis, W. D., Eds.; Pergamon: Oxford, 1979; Vol. 3, pp 373–487. (b) Voss, J. *Houben-Weyl Methoden der Organischen Chemie*; Klamann, D., Ed.; George Thieme: New York, 1985; Bd. E11, pp 188–231. (c) Magnas, P. D. In *Comprehensive Organic Chemistry*; Barton, D. H. R., Ollis, W. D., Eds.; Pergamon: Oxford, 1979; Vol. 3, pp 491–538. (d) Paulmier, C. *Selenium Reagents and Intermediates in Organic Synthesis*; Pergamon: Oxford, 1986; pp 58–83. (e) Guziec, F. S., Jr. *The Chemistry of Organic Selenium and Tellurium Compounds*; Patai, S., Ed.; John Wiley & Sons: New York, 1987; Vol. 2, pp 215–273.

(3) Okazaki, R.; Ishii, A.; Fukuda, N.; Oyama, H.; Inamoto, N. *J. Chem. Soc., Chem. Commun.* **1982**, 1187–1188. Vedejs, E.; Perry, D. A.; Wilde, R. G. *J. Am. Chem. Soc.* **1986**, 108, 2985–2989. Okazaki, R.; Ishii, A.; Inamoto, N. *J. Am. Chem. Soc.* **1987**, 109, 279–280. Okazaki, R.; Kumon, N.; Inamoto, N. *J. Am. Chem. Soc.* **1989**, 111, 5949–5951. Ando, W.; Ohtaki, T.; Suzuki, T.; Kabe, Y. *J. Am. Chem. Soc.* **1991**, 113, 7782–7784. For reviews, see: Okazaki, R. *Yuki Gosei Kagaku Kyokai Shi* **1988**, 46, 1149–1163. Usov, V. A.; Timokhina, L. V.; Voronkov, M. G. *Sulfur Rep.* **1992**, 12, 95–158.

(4) For trapping of transient telluroketones and telluroaldehydes, see: Erker, G.; Hock, R. *Angew. Chem., Int. Ed. Engl.* **1989**, 28, 179–180. Segi, M.; Koyama, T.; Takata, Y.; Nakajima, T.; Suga, S. *J. Am. Chem. Soc.* **1989**, 111, 8749–8751. Boese, R.; Haas, A.; Limberg, C. *J. Chem. Soc., Chem. Commun.* **1991**, 1378–1379. Haas, A.; Limberg, C. *Chimia* **1992**, 46, 78.

(5) (a) Barrett, A. G. M.; Barton, D. H. R.; Read, R. W. *J. Chem. Soc., Chem. Commun.* **1979**, 645–647. (b) Barrett, A. G. M.; Read, R. W.; Barton, D. H. R. *J. Chem. Soc., Perkin Trans. 1* **1980**, 2191–2195. (c) Severing, T.; du Mont, W. W. *J. Chem. Soc., Chem. Commun.* **1987**, 820–821. (d) Lerstrup, K. A.; Henriksen, L. *J. Chem. Soc., Chem. Commun.* **1979**, 1102–1103. (e) Lappert, M. F.; Martin, T. R.; McLaughlin, G. M. *J. Chem. Soc., Chem. Commun.* **1980**, 635–637. (f) Segi, M.; Kojima, A.; Nakajima, T.; Suga, S. *Synlett* **1991**, 2, 105–106.

(6) Fischer, H.; Zeuner, S. *J. Organomet. Chem.* **1983**, 252, C63–C65. Fischer, H.; Gerbing, U. *J. Organomet. Chem.* **1986**, 299, C7–C10. Wolf, J.; Zolk, R.; Schubert, U.; Werner, H. *J. Organomet. Chem.* **1988**, 340, 161–178. Headford, C. E. L.; Roper, W. R. *J. Organomet. Chem.* **1983**, 244, C53–C56. Herrmann, W. A.; Weichmann, J.; Serrano, R.; Blechschmitt, K.; Pfisterer, H.; Ziegler, M. L. *Angew. Chem., Int. Ed. Engl.* **1983**, 22, 314–315. Paul, W.; Werner, H. *Angew. Chem., Int. Ed. Engl.* **1983**, 22, 316–317. Hill, A. F.; Roper, W. R.; Waters, J. M.; Wright, A. H. *J. Am. Chem. Soc.* **1983**, 105, 5939–5940. Herrmann, W. A.; Hecht, C.; Ziegler, M. L.; Balbach, B. *J. Chem. Soc., Chem. Commun.* **1984**, 686–687. Werner, H.; Paul, W.; Knaup, W.; Wolf, J.; Müller, G.; Riede, J. *J. Organomet. Chem.* **1988**, 358, 95–121. Fischer, H.; Früh, A.; Troll, C. *J. Organomet. Chem.* **1991**, 415, 211–221.

(7) Okazaki, R.; Minoura, M.; Kawashima, T. *Chem. Lett.* **1993**, 1047–1048.

(8) 1,3,4-Selenadiazolines, synthesized via cycloaddition of sterically hindered selenoketones with diazo compounds, have been reported to undergo thermal cycloreversion to form selenoketones: (a) Back, T. G.; Barton, D. H. R.; Britten-Kelly, M. R.; Guziec, F. S., Jr. *J. Chem. Soc., Perkin Trans. 1* **1976**, 2079–2089. (b) Guziec, F. S., Jr.; Murphy, C. J.; Cullen, E. R. *J. Chem. Soc., Perkin Trans. 1* **1985**, 107–113. (c) Guziec, F. S., Jr.; SanFilippo, L. J.; Murphy, C. J.; Moustakis, C. A.; Cullen, E. R. *Tetrahedron* **1985**, 41, 4843–4852. (d) Tokitoh, N.; Choi, N.; Ando, W. *Tetrahedron Lett.* **1990**, 25, 3571–3574. (e) Guziec, F. S., Jr.; SanFilippo, L. J. *J. Org. Chem.* **1991**, 56, 3178–3181.

(9) The molar extinction coefficient (ε) was obtained by assuming the quantitative formation of **1a** from **2**.

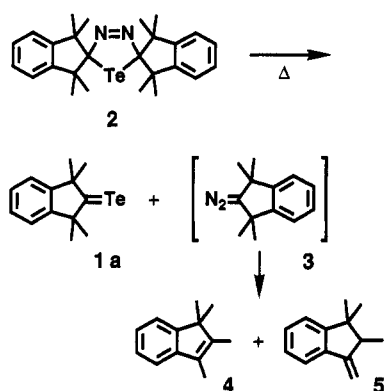
(10) Kalinowski, H. O.; Berger, S.; Braun, S. *Carbon-13 NMR Spectroscopy*; John Wiley & Sons: New York, 1988.

(11) (a) Cullen, E. R.; Guziec, F. S., Jr.; Murphy, C. J.; Wong, T. C.; Andersen, K. K. *J. Chem. Soc., Perkin Trans. 2* **1982**, 473–476. (b) Krebs, A.; Rüger, W.; Ziegenhagen, B.; Hebold, M.; Hardtke, I.; Müller, R.; Schütz, M.; Wietzke, M.; Wilke, M. *Chem. Ber.* **1984**, 117, 277–309.

(12) McFarlane, H. C. E.; McFarlane, W. In *Multinuclear NMR*; Mason, J., Ed.; Plenum: New York, 1987; pp 417–435.

(13) The color of the solution was green soon after it was taken out of the dark but turned yellow very rapidly even under a fluorescent lamp.

Scheme I



the $n\text{--}\pi^*$ transition of the $\text{C}=\text{Te}$ bond, is the longest among the carbon—chalcogen double-bond compounds (1b 667 nm,¹⁴ 1c 527 nm¹⁴).²

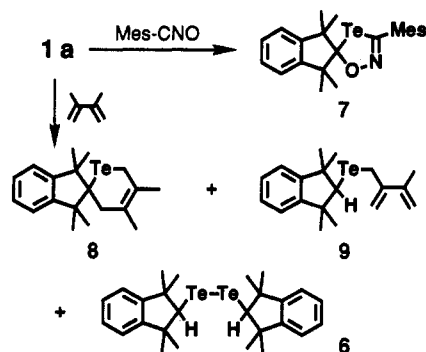
These spectroscopic observations clearly indicate the monomeric nature of 1a, which was also supported by its chemical reactivities (Scheme II).¹⁵ Reaction of 1a with an equimolar amount of mesitronitrile oxide at 80 °C afforded a novel heterocycle (oxatellurazole 7) regioselectively in a good yield (70%).¹⁶ This represents the first example of 1,3-dipolar cycloaddition of a telluroketone. The Diels–Alder reaction of 1a with 2,3-dimethyl-1,3-butadiene proceeded under much milder conditions (100 °C,

(14) Klages, C. P.; Voss, J. *Chem. Ber.* **1980**, *113*, 2255–2277.

(15) All manipulations were performed in the dark with a subdued red lamp.

(16) The regiochemistry of 7 was determined by its mass spectrum, which gave fragmentation of indanone 1d and MesNCTe. The thermolysis of 7 afforded 1d and Mes—N $\equiv$ C, the mechanism of which will be reported elsewhere.

Scheme II



2 h) than the corresponding selenoketone 1b (150 °C, 16 h)¹⁷ to give cycloadduct 8 (24%), ene product 9 (34%), and ditelluride 6 (36%). The formation of 9 is the first demonstration of the ene reactivity of a tellurocarbonyl compound.

Further investigation of the physical and chemical properties of 1a as well as the application of the telluradiazoline methodology to the synthesis of other telluroketones is currently in progress.

Acknowledgment. This work was supported by the Grant-in-Aid for Scientific Research No. 04403005 from the Ministry of Education, Science and Culture, Japan.

Supplementary Material Available: Physical and spectral data of products 6–9 (1 page). Ordering information is given on any current masthead page.

(17) Minoura, M.; Kawashima, T.; Okazaki, R., unpublished results. The products were the corresponding Diels–Alder adduct (57%) and ene product (19%).