

Communications

Synthesis and Characterization of Two Isomers of 14 π -Electron Germaaromatics: Kinetically Stabilized 9-Germaanthracene and 9-Germaphenanthrene

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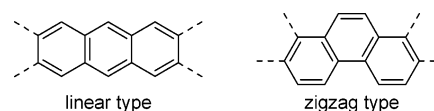
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Received May 1, 2006

Summary: The first stable 9-germaanthracene and 9-germaphenanthrene were successfully synthesized by taking advantage of kinetic stabilization utilizing effective steric protection groups, Tbt and Bbt groups (Tbt = 2,4,6-tris[bis(trimethylsilyl)methyl]phenyl, Bbt = 2,6-bis[bis(trimethylsilyl)methyl]-4-[tris(trimethylsilyl)methyl]phenyl). The similarities and differences of the structures and properties between the two isomers of germaaromatics are discussed.

There has been much interest in the chemistry of “heavy aromatics”, $[4n+2]$ π -electron ring systems containing a heavier group 14 element (Si, Ge, Sn, Pb),¹ from the standpoint of comparison with the parent aromatic hydrocarbon compounds, which play very important roles in organic chemistry. Although heavy aromatics have been known to be highly reactive and undergo ready dimerization or oligomerization under ambient conditions so far, we have succeeded in the synthesis of the first stable neutral sila- and germaaromatics,^{1g} i.e., silabenzene,² 1- and 2-silaphthalenes,^{3,4} 9-silaanthracene,⁵ germabenzene,⁶ and 2-germanaphthalene,⁷ by taking advantage of kinetic stabilization using efficient steric protection groups, Tbt and

Scheme 1



Bbt. The experimental and theoretical studies on the structures, spectroscopic properties, and reactivities of these kinetically stabilized sila- and germaaromatics evidenced the high aromaticity of the monosila- and monogermaaromatic systems.^{1g} Very recently, the stable stannaromatics,⁸ 2-stannaphthalene, was also isolated and fully characterized, suggesting the considerable aromaticity of a 10 π -electron ring system containing a much heavier group 14 atom, an Sn atom.⁹ Thus, the concept of kinetic stabilization is evidenced to be of great use for the construction of stable heavy aromatics. On the other hand, anthracene and phenanthrene are attractive 14 π -electron aromatic systems due to their unique electrochemical and photochemical properties from the viewpoint of material science.¹⁰ Hence, it should be of great importance to elucidate the similarities and differences between two types of extended π -electron aromatic systems containing a heavier group 14 atom, i.e., the linear type (acene system) and the zigzag type (phen system) such as anthracene and phenanthrene (Scheme 1). We report here the synthesis of the two structural isomers of 14 π -electron germaaromatic systems, 9-germaanthracene **1** and 9-germaphenanthrene **2**, utilizing a Tbt or Bbt group as a steric protection group.

In consideration of previous reports,^{1g} the reactions of the corresponding precursors bearing a leaving group (halogeno or

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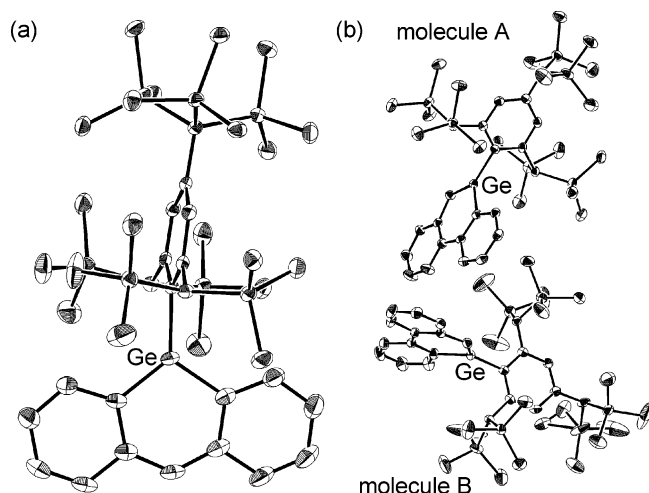


Figure 1. ORTEP drawing of **1b** (a) and **2** (b) with thermal ellipsoid plots (50% probability). Hydrogen atoms are omitted for clarity.

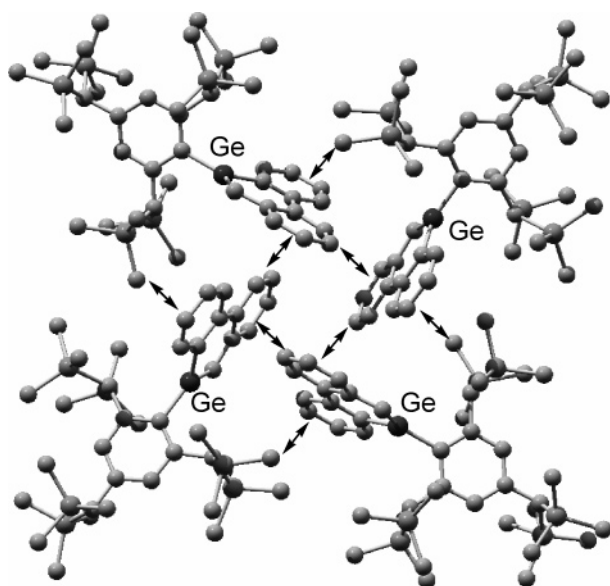


Figure 2. Tetrameric packing structure of 9-germaphenanthrene **2**.

canonical structure of a 9-germaphenanthrene skeleton.¹⁶ Thus, the Ge—C bond of the 9,10-position of 9-germaphenanthrene **2** features higher double-bond character than the other Ge—C bond of **2**.

In the ¹H and ¹³C NMR spectra of **1b** and **2**, all of the signals corresponding to the central 9-germaanthracene and 9-germaphenanthrene skeletons were observed within the aromatic region, 7–8 ppm for the ¹H NMR spectra and 110–150 ppm for the ¹³C NMR spectra. In particular, the protons of the 10-positions of **1b** and **2** appeared as characteristic singlet signals at 7.73 and 8.13 ppm, respectively, which are apparently lower

(16) In the case of anthracene and phenanthrene, the slight bond alternations can be seen, reflecting their most dominant canonical structures as shown below. Similarly, one can see their slight bond alternations in the germaaromatic rings of **1b** and **2**, which should be reasonably explained in terms of their dominant canonical structures as well as their all-carbon analogues.

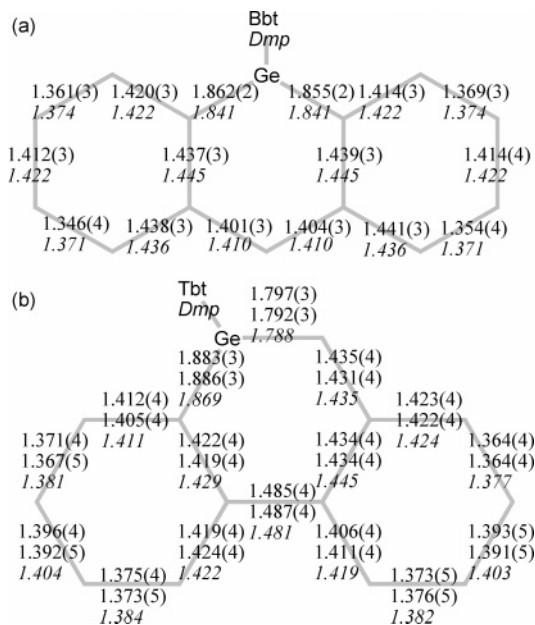
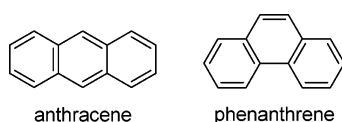


Figure 3. Observed and calculated (*italic*) bond lengths (Å) for (a) 9-germaanthracenes and (b) 9-germaphenanthrenes. The calculated values are obtained for the Dmp (2,6-dimethylphenyl)-substituted model compounds at the B3LYP/6-31G(d) level.

than those of olefins (ca. 4.5–7.0 ppm).¹⁷ In addition, the calculated chemical shifts for the model compounds bearing less hindered substituents on the Ge atoms are in good agreement with the observed values, indicating that the steric effect of Tbt and Bbt on the central germaaromatic rings should be negligible.¹³ Thus, the NMR spectral properties of **1b** and **2** are similar to each other. Electronic spectra should be of great use in the evaluation of the extension of π -electrons on the aromatic rings. In the UV/vis spectra in hexane, the longest absorption maxima (λ_{max}) corresponding to the HOMO–LUMO electron transitions of **1b** (520 nm) and **2** (409 nm) are red-shifted as compared with those of 1-Tbt-germabenzene⁶ (326 nm) and 2-Tbt-2-germanaphthalene⁷ (386 nm), respectively. That is, the π -electron conjugation can be extended as the n value of $[4n+2]$ π -electron aromatic systems increases.¹⁸ Furthermore, the longest λ_{max} of 9-germaanthracene **1b** is apparently longer than that of **2**, reflecting their colors in solution (red for **1b** and pale yellow for **2**; see Figure S6 in the Supporting Information). Hence, it is evidenced that the 14π -electron systems of germaaromatics can be extensively conjugated in the linear type but not so effectively in the zigzag type.

In conclusion, two isomers of 14π -electron germaaromatic systems, 9-germaanthracene **1b** and 9-germaphenanthrene **2**, were synthesized as stable crystalline compounds. Their similarities and differences are revealed on the basis of the structural and spectroscopic analyses. Further investigations on their photochemical and electrochemical properties, reactivities, and applications as arene ligands to transition metals are currently in progress.

(17) For example, the olefinic proton of triphenylethylene is observed at 6.96 ppm [SDBSWeb: <http://www.aist.go.jp/RIODB/SDBS/> (National Institute of Advanced Industrial Science and Technology, 2006)].

(18) The longest absorption maxima of silaaromatic systems are apparently red-shifted as compared with those of the corresponding all-carbon systems. However, those of germaaromatic systems are observed in the region similar to that for the corresponding silaaromatic systems. The systematic comparisons of the longest absorption maxima between the kinetically stabilized sila- and germaaromatic systems with all-carbon systems are shown in the Supporting Information.

Acknowledgment. This work was partially supported by Grants-in-Aid (Nos. 17GS0207, 12CE2005, 14078213, and 16750033) and the 21st Century COE on Kyoto University Alliance for Chemistry from the Ministry of Education, Culture, Sports, Science and Technology, Japan. T.S. thanks the Kinki Invention Center for the research grant.

Supporting Information Available: X-ray crystallographic data of **1** and **15** in CIF format, experimental procedures, and spectral data. These materials are available free of charge via the Internet at <http://pubs.acs.org>.

OM060371+