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Synthesis of Hydrogen-Bridged Tetraborane(6): Substituent Effect of Diaminoboryl Group toward B-B Multiple Bond Character

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A hydrogen-bridging tetraborane(6) was synthesized from boryllithium, a boron nucleophile, in three steps. The structural and spectroscopic analysis of the tetraborane(6) revealed σ -donor and π -acceptor effect of diaminoboryl substituents toward the central B–B moiety having multiple bond character. DFT calculations also supported the experimentally observed substituent effect of boryl group.

Recent development of B-B multiple bond species opened a new area of chemistry due to their characteristic structure and reactivity of π -bond(s).¹ Two-electron reduction of diborane(4) led to a formation of the first B=B double bond species with two anionic charge [A, Figure 1(a)].^{2, 3} The first diborene, a neutral B=B double bond species (B), was synthesized by a reduction of NHC-coordinated BBr₃ derivative.^{4, 5} After the first discovery, propensities of diborene derivatives were extensively studied by an improvement of their synthesis.⁶⁻²⁸ On the other hand, diboryne, a neutral B≡B triple bond species (C),^{7, 29-33} has been synthesized by a reduction of Lewis base-stabilized tetrahalodiborane(4) precursor. Related hydrogen-bridging diborane(4) species possessing B-B multiple bond character were synthesized by using a bulky aryl substituent, Eind group (D).³⁴⁻³⁶ The H-bridged structure of D could be considered as a doubly protonated form of dianionic B≡B triple bond compound as supported with DFT study with a remarkably short B-B bond distance in comparison with those observed in diborene derivatives. Recently, an amino-substituted homocatenated triborane(3) compound (E), which can be considered as a trimer of aminoborylene, was reported to have a B-B multiple bond character.

In the chemistry of multiply-bonded main group element species, characteristic properties of the multiple bond such as

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short bond lengths and narrower HOMO-LUMO gap, have been studied. Connection of these species with boryl substituent(s) has been known to be able to alter the electronic properties of these multiple bonds [Figure 1(b)]. Introduction of dialkylboryl substituent to disilene (F) provided information about π accepting character of dialkylboryl group toward Si=Si double bond.^{37, 38} We reported synthesis and properties of borylsubstituted diphosphene (G), where boryl substituents showed σ -donor and π -acceptor character toward electronic properties of P=P double bond.³⁹⁻⁴¹ Recently, a boryl-substituted diborene (H) was reported to have an effective π -conjugation between B=B double bond and vacant p-orbitals of catecholatoboryl substituents.²³ In this work, we herein report synthesis of borylsubstituted hydrogen-bridged diborane(4), which can be considered as tetraborane(6) species, and the substituent effect of the diaminoboryl substituents toward multiple bond character of the inner B-B moiety.



Figure 1. (a) Examples of multiboron compounds having B-B multiple bond character and (b) multiply bonded p-block element species bearing boryl substituent(s)

Reaction of boryllithyum **1**, which was prepared in accordance with our previous contribution,^{42, 43} with two equivalents of

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trimethylborate to give an unsymmetrical diborane(4) (2) with two amino and two alkoxy substituents in 78% yield (Scheme 1). Treatment of 2 with two equivalents of boron tribromide induced ligand exchange reaction to afford diaminodibromodiborane(4) (3) in 53% yield. Reaction of 3 with five equivalents of potassium graphite in toluene at -35 °C generated a hydrogen-bridged tetraborane(6) 4. Colourless crystals 4 were obtained by recrystallization from its hexane solution in 12% isolated yield. In the ¹H NMR spectrum of 4 (C_6D_6) , the signal of bridging hydrogen atoms was observed as a broad singlet at -2.25 ppm. This chemical shift is higher-field shifted in comparison with that of the aryl-substituted hydrogen-bridged diborane(4) D (2.94 ppm).³⁴ However, the NMR chemical shift calculations by using GIAO method indicated that the higher field shifted signal of bridging hydrogen atoms in 4 would be their peculiar property of B2H2 core structure (Figure S20) rather than the effect of boryl substituents.⁴⁴ In the ¹¹B NMR spectrum, the signals of **4** were observed at 5 and 30 ppm as a broad resonance, in which the former could be assigned as two internal boron atoms in comparison with that in D.34 The IR spectrum of 4 exhibited a series of B–H–B vibrations (1321-1174 cm⁻¹) in the same region to those of **D** (1265-1157 cm⁻¹),³⁴ confirming the existence of the two bridging hydrogen atoms. Reaction of $\mathbf{3}$ with KC₈ in toluene- d_8 resulted in an appearance of small signal (-2.30) ppm) in addition to the signal of the bridging hydrogen atoms in 4 (Figure S14). This new signal could be assigned as a bridging hydrogen atom in partially deuterated 4-d₁. Although ²H NMR spectrum of partially deuterated 4 did not exhibit any detectable signal probably due to significant broadening of the signal caused by coupling between two guadrupolar nuclei of ²H and ¹¹B/¹⁰B, incorporation of deuterium atoms to **4** was also

and ¹¹B/¹⁰B, incorporation of deuterium atoms to **4** was also supported by APCI-TOF MS and IR spectroscopy (Figures S13, 15, 16). This result indicates the origin of the bridging hydrogen atoms in **4** could be attributed to inter- and intra-molecular hydrogen trapping from toluene solvent and backbone or Dip group in **3**.



Scheme 1. Synthesis of hydrogen-bridged tetraborane(6) 4 (Dip = 2,6-'Pr₂C₆H₃)

The molecular structure of **4** was unambiguously confirmed by X-ray crystallographic analysis (Figure 2). Two bridging hydrogen atoms were placed as a peaks in a difference Fourier map. It was revealed that the inner B–B bond distance of 1.488(4) Å, which is similar to that in **D** [1.4879(7) Å]³⁴ and lies between the reported B=B double and B≡B triple bonds. On the other hand, the outer B–B distances of 1.666(3), 1.661(3) Å are shorter than B–B covalent bond distance (1.75 Å),⁴⁵ and are similar to that of the diboryldiborene **H** (1.645(3) Å-1.676(6)

Å),²³ indicating an effective π -conjugation between internal B-B bond possessing a multiple bond character and vacant porbitals of diaminoboryl substituents in **4**. These results suggest **4** can be described with resonance structures in Scheme 2.



Figure 2. X-ray structure of **4** (left: side view, right: top view); Thermal ellipsoids set at 50% probability. Hydrogen atoms except for the bridging hydrogen atoms are omitted for clarity.

$$\begin{array}{cccc} H & H^{\dagger} & H^{\dagger} & H^{\dagger} \\ B - B - B - B & \leftrightarrow B - B \equiv B - B & \leftrightarrow B = B \equiv B - B & \leftrightarrow B - B \equiv B \equiv B = B \\ H & H^{\dagger} & H^{\dagger} & H^{\dagger} & H^{\dagger} \end{array}$$

Scheme 2. Possible resonance structures of 4

DFT calculations were performed to analyse electronic structure of 4. The experimentally obtained structure of 4 by X-ray analysis was used as an initial structure for optimization at B3LYP/6-31+G(d,p) level of theory. Characteristic molecular orbitals of 4 are illustrated in Figure 3. Two molecular orbitals, HOMO-49 and HOMO-2 would correspond to two $\pi\text{-like}$ orbitals at the central B–B unit. Overlapping of π^* -orbitals of the central B-B unit and two vacant p-orbitals in 4 seems to produce LUMO. Two combinations of two π -type N–B–N orbitals construct energetically degenerated HOMO-1 and HOMO. In contrast, aryl-substituted **D** has slightly higher π^* -like LUMO and π -like HOMO, and two combinations of π -orbitals of Eind groups consist HOMO-1 and HOMO-2 (Figure S19). This result indicates the boryl substituents in ${\bf 4}$ contribute to lower $\pi\text{-}$ and π^* -orbital of the central B-B unit through their π -accepting effect. It should be noted that a bonding combination of π -type orbitals of N–B–N unit and the central B–B unit constructs a linear and delocalized π -conjugation over four boron atoms in HOMO-12. This π -conjugation is consistent with resonance structures in Scheme 2. The NPA charges on the central B–B unit in **4** are negative (B1, B2: -0.239), while those of **D** were positive (0.043, 0.074), supporting the σ -donating effect of boryl substituents to increase the electron density at the central B-B unit.

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Figure 3. Selected molecular orbitals of **4** calculated at the B3LYP/6-31+G(d,p) level of theory. Hydrogen atoms omitted for clarity except for the bridged hydrogens.

Conclusions

By using boryllithium as a starting material, the hydrogenbridging tetraborane(6) derivative **4** was synthesized in three steps. The structural and spectroscopic analysis revealed σ donor and π -acceptor effect of diaminoboryl substituents toward B–B multiple bonding character. DFT calculations further supported the observed substituent effect of boryl groups.

Conflicts of interest

There are no conflicts to declare.

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