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Unsymmetrical, Cyclic Diborenes and Thermal Rearrangement to a Borylborylene

Tom E. Stennett,[a] James D. Mattock,[b] Ivonne Vollert,[a] Alfredo Vargas*[b] and Holger Braunschweig*[a]

Abstract: Cyclic diboranes(4) based on a chelating monoanionic benzylphosphine linker were prepared by boron-silicon exchange between arylsilanes and B₂Br₄. Coordination of Lewis bases to the remaining sp³ boron atom yielded unsymmetrical sp³-sp³ diboranes, which were reduced with KC₈ to their corresponding trans-diboranes. These compounds were studied by a combination of spectroscopic methods, X-ray diffraction and DFT calculations. PMe₃-stabilized diborane 6 was found to undergo thermal rearrangement to gem-diborone 8. DFT calculations on 8 reveal a polar boron-boron bond, and indicate that the compound is best described as a borylborylene.

Since the report of the first neutral diborone by Robinson and co-workers, which was prepared in low yields via a radical hydrogen scavenging process,[1] the preparation of compounds bearing boron-boron double bonds[5] has been extended by three rational and selective synthetic routes – the reductive coupling of two N-heterocyclic carbene (NHC)-stabilized aryldihaloboranes,[2b, 3] the reduction of B₂X₅R₂ species in the presence of stabilizing phosphine ligands,[5] and the stoichiometric reduction of carbene adducts of B₂Br₄.[5] A very recent development in this field has been the realization of the first unsymmetrical diboranes,[5a, 6] which promise to reveal more information about the fundamental nature of the boron-boron bond, and also allow investigation of the effect of a polarised B-B bond on the species’ behaviour.

We are currently especially interested in the reactivity of phosphine-stabilized diborones. Whereas carbones typically remain bound to boron during the course of the reactions of diborones with small molecules,[5b, 5a, 7] phosphine donors can dissociate under UV light and their diborones display markedly different reactivity.[5b] A drawback of the currently known examples of phosphine-stabilized diboranes is that they are mostly prepared from 1,2-dihoal-1,2-diarylb-oranes, B₂X₅R₂, the syntheses of which are limited in terms of substituent scope.[5b] With convenient routes now available to all of the tetrahalodiboranes(4),[5b, 11] we are therefore keen to develop milder and better-yielding routes for the direct functionalization of B₂X₅ species.

Substituted monoboranes (RBX₃) can often be conveniently prepared by silicon-boron exchange of the corresponding RSiMe₃ species with trihaloboranes,[5b, 11] but equivalent reactions with diboranes(4) are limited to just a handful of examples. In the early 1980s, Haubold et al. prepared bridged sp²-sp³ diboranes by reaction of B₂Cl₄ with 2-silylalkyl- and 2-silyloxypyrindines, forming B-N and B-O bonds, respectively, by Me₃SiCl elimination (Scheme 1).[12] The group of Siebert later succeeded in transferring a Cp⁺ group to B₂Cl₄ via chlorosilane elimination, with a rearrangement giving a cyclopentadienylborylene-boron adduct as the final product.[13]

Inspired by these studies, we selected a readily prepared arylsilane containing a phosphine donor group, with the goal of generating new diboranes capable of being reduced to B-B multiply-bonded compounds. From this starting point, we present a rational synthesis of unsymmetrical diborones and rearrangement of one of these compounds to its gem isomer.[14]

Reaction of 2-trimethylsilylbenzyl(dicyclohexyl)phosphine (2a), prepared in quantitative yield by treating the corresponding aryllithium compound, 2-LiC₆H₄CH₂PCY₂,Et₂O (1), with Me₃SiCl,[15] with B₂Cl₄ at 0 °C led to the formation of sp³-sp³ diborone 3a in high yield (98%; Scheme 2). The ¹¹B NMR spectrum of 3a shows the low- and high-field signals expected for the sp² (73.3 ppm) and sp³ (~10.5 ppm) boron atoms, respectively. A broad ³¹P NMR signal at 2.6 ppm is also indicative of coordination to boron. X-ray diffraction provided confirmation of the expected structure (Figure 1), with the B-B bond length of 1.678(9) Å slightly shorter than published sp³-sp³ diborones,[16] possibly as a result of steric constraints imposed by the chelating benzophosphine group. Addition of a further equivalent of 2a to 3a resulted in no reaction at room temperature, while higher temperatures (80 °C) resulted in slow, unselective conversion to several species, none of which could be identified. The corresponding reaction with 2-trimethylsilylbenzyl(di-tert-butyl)phosphine (2b) also yields the equivalent sp³-sp³ diborane, 3b. In this case, an intermediate species is observed when the reaction is performed at room temperature; observation of signals in the ¹¹B NMR spectrum at 68.6 and ~5.7 ppm, a broad ³¹P signal at 21.6 ppm and a signal at 0.31 ppm in the ¹³C NMR spectrum for the SiMe₃ group indicate formation of adduct A, which can be completely converted to 3b by heating to 60 °C for 2 h. This observation explains the regioselectivity of the formation of 3a and 3b. Silicon-boron exchange reactions of haloboranes are, to the best of our knowledge, limited to sp²-hybridized boranes, hence the selective formation of the 1,2-isomer.[17] The structure of 3b was also determined by X-ray crystallography (Figure 1), with the B-B bond length (1.681(9) Å) and other parameters not differing significantly from those in 3a.

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Supporting information for this article is given via a link at the end of the document.
Scheme 2. Synthesis of sp²-sp³ diboranes 3a and 3b via Me₂SiBr elimination.

Treatment of 3a with small Lewis bases, L, led to their coordination to the remaining sp² boron atom and formation of sp²-sp³ diboranes (L = PMe₃ (4), N,N-diisopropylimidazol-2-ylidene (IPr) (5), Scheme 3), as indicated by a shift of the low-field resonance for the sp² boron atom to high field. The ³¹P NMR spectrum of 4 shows two broad, mutually coupled doublets at −7.9 and −12.4 ppm (JₚB = 40 Hz), whereas the sole signal of 5 appears at −10.5 ppm. The structure of 4 was confirmed by X-ray diffraction (Figure 1). The B-B bond length displays the expected slight elongation, to 1.737(3) Å, with the structural parameters being otherwise unremarkable.


Reaction of 4 with KC₈ in benzene gave a stark color change to dark yellow and formation of a new species with a single broad signal in its ¹¹B NMR spectrum at 20.4 ppm, within the range of known diboranes. After two days conversion was complete, and after workup diborene 6 was isolated as a yellow solid in 68% yield. Single crystals suitable for X-ray diffraction were grown from hexane, allowing confirmation of the formation of 6 (Figure 2, Table 1). At 1.563(3) Å, the B-B distance is in the expected range for a boron-boron double bond, while the P-B bonds are both contracted by roughly 0.07 Å compared to 4. Interestingly, the benzylphosphine chelate appears to cause a distortion of the substituents from ideal double-bond geometry, as the P₁-B₁-B₂-C₁ unit has a torsion angle of 17.3°. A similar procedure resulted in the reduction of 5 to diborene 7, isolated as an orange solid in 34% yield, which is the first reported non-allenic diborene stabilized by two different classes of Lewis base – a phosphine and an NHC. In this case, the ¹¹B NMR signals at 30.0 and 12.5 ppm are at the extreme ends of the range for previously reported diboranes, and therefore suggest a reasonable degree of polarity of the B-B bond. The structure of 7 was also confirmed by X-ray diffraction (Figure 2), and the B-B double bond (1.562(6) Å) found to be the same length as that in 6, within error.
In which the monovalent B1 boron atom bitrals of 2 charge distribution is inversed, i.e. s remained coordinated to boron. After 2 d the HOMO and LUMO are found to be at lower energies in phosphine derivative exclusively perpendicular to the B=B bond. Both homoarene substituents in previously reported diborenes sit B=B plane by the chelate. This conjugation is noteworthy as from the π higher C:→B2→B1 charge flow compared to the P:→B2→B1 to the situation in 6. These observations suggest that there is a higher σ C1/B2→B1 charge flow compared to the P:→B2→B1 path. The LUMO of both compounds has a large contribution from the π-system of the arene moiety, which is held close to the B=B plane by the chelate. This conjugation is noteworthy as homoarene substituents in previously reported diborenes sit exclusively perpendicular to the B=B bond. Both the HOMO and LUMO are found to be at lower energies in phosphine derivative 6 than in 7 due to the higher σ-donor strength of the NHCl.

### Table 1. Selected bond lengths (Å) for compounds 6-8.

<table>
<thead>
<tr>
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<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1-B2</td>
<td>1.563(3)</td>
<td>1.562(6)</td>
<td>1.558(7)</td>
</tr>
<tr>
<td>P1-B1</td>
<td>1.879(2)</td>
<td>1.863(5)</td>
<td>1.853(4)</td>
</tr>
<tr>
<td>P2-B</td>
<td>1.909(2)</td>
<td>-</td>
<td>1.869(4)</td>
</tr>
<tr>
<td>C1-B2</td>
<td>1.587(3)</td>
<td>1.573(5)</td>
<td>1.576(6)</td>
</tr>
<tr>
<td>C2-B2</td>
<td>-</td>
<td>1.582(6)</td>
<td>-</td>
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</table>

Calculations based on the Kohn-Sham Density Functional Theory (DFT) at the OLYP/TZ2P level were conducted to further understand the observed results and the underlying rationales. Figure 3 shows the frontier orbitals of 6 and 7 and Table 2 displays the calculated bond lengths, bond orders and partial charges of the two structures in the gas phase. While both species largely display diborene characteristics in terms of electronic structure, as shown by their frontier orbitals and the B-B bond length, the phosphine-stabilized compound 6 exhibits a slightly higher B-B bond order than 7, and a less polarized B-B unit. In addition, this unequal charge distribution is inversed, i.e. in 6 the B1 centre carries the lesser negative charge, in contrast to the situation in 7. These observations suggest that there is a higher C:→B2→B1 charge flow compared to the P:→B2→B1 path. The LUMO of both compounds has a large contribution from the π-system of the arene moiety, which is held close to the B=B plane by the chelate. This conjugation is noteworthy as homoarene substituents in previously reported diborenes sit exclusively perpendicular to the B=B bond. Both the HOMO and LUMO are found to be at lower energies in phosphine derivative 6 than in 7 due to the higher σ-donor strength of the NHCl.

### Table 2. Calculated properties of compounds 6 and 7.

<table>
<thead>
<tr>
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<th>6</th>
<th>7</th>
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<tbody>
<tr>
<td>α(BB)/Å</td>
<td>1.593</td>
<td>1.595</td>
</tr>
<tr>
<td>α(B2-C1)/Å</td>
<td>1.588</td>
<td>1.576</td>
</tr>
<tr>
<td>Bond order(BB)</td>
<td>1.52</td>
<td>1.48</td>
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<tr>
<td>q(B1)</td>
<td>−0.129</td>
<td>−0.149</td>
</tr>
<tr>
<td>q(B2)</td>
<td>−0.155</td>
<td>−0.113</td>
</tr>
<tr>
<td>q(Br)</td>
<td>−0.141</td>
<td>−0.153</td>
</tr>
</tbody>
</table>

Unsymmetrical diborenes remain very much a novelty, with the only reported examples coming from the group of Kinjo in the past year and their behavior is as yet largely unexplored. We initially tested the thermal stability of 6 and 7. Heating 6 to 80 °C in benzene led to a slow conversion to a new species displaying a broad signal at 52.2 ppm and a pseudo-triplet at –21.1 ppm in the 11B NMR spectrum. Broad multiplets at 23.3 ppm and −11.9 ppm in the 31P NMR spectrum indicated that both phosphines remained coordinated to boron. After 2 d conversion was complete, and single crystals were grown by concentration of the reaction solution. X-ray diffraction revealed the product to be 8 (Scheme 4, Figure 2), in which an exchange of the diborene substituents had occurred, with both phosphines now bonded to the same boron atom. The B-B bond distance of 1.558(7) Å remains in the range for a boron-boron double bond, unchanged from 6, while the B-P distances are slightly shorter than those in 6. Heating compound 7 in benzene resulted in an even slower, unselective reaction, with no NMR signals comparable to those for 8 observed. As yet we have been unable to identify any of the products.

The nature of the boron-boron bond in 8 is not immediately clear; a classical π-bond would necessitate a formal negative charge on B2 and a positive charge on B1 (Scheme 4). Alternatively, the compound can be viewed as a boryl-substituted borylene, in which the monovalent B1 boron atom carries a lone pair of electrons. Although borylenes were formerly merely postulated as fleeting intermediates, isolable, metal-free borylene compounds stabilized by two (or one) neutral donors are now firmly established. A recent computational study by Ghorai and Jemmis predicted that a simplified diborene, 3B2Br2(PhH)2, would be more stable as its
gem isomer than as its cis and trans isomers (derivatives of both of which have been realized experimentally), while its NHC derivative favors the trans conformation.\(^{[16]}\) The authors proposed a borylborylene type structure, involving a large donor-acceptor contribution to the boron-boron bond, from the P\(_2\)B unit to the B=B boron atom. Having isolated such a compound, we again turned to DFT for more insight into the formation and structure of 8.

The calculated thermal conversion of 6 to 8 corresponds to a stabilization by 3.90 kcal/mol in terms of total electronic energy in the gas phase, and by 5.08 kcal/mol in toluene (within the conductor-like screening model). The more pronounced lowering in energy in the presence of a polarizable medium suggests a highly polarized B-B core in the product 8. Table 3 shows the comparison of 6 and 8 in terms of selected parameters. The B-B bond order is somewhat lower in 8 (1.406) than in 6 (1.544), but nevertheless indicates significant double-bond character. Close inspection of the calculated partial charges (\(q_{BI} = -0.243, q_{B2} = -0.038\)) indicates a highly charge-separated state in 8, in which the polarity is inverted with respect to 6 and the negative charge localized on B1. This analysis favors the borylborylene configuration proposed by Jemmis,\(^{[19]}\) where B1 carries a lone pair. Furthermore, the frontier molecular orbitals shown in Figure 3 indicate a marked difference in the HOMO compared to the trans-diborones. While in both compounds 6 and 7, the HOMO describes a typical (albeit slightly distorted) diborene \(\pi\)-system, in compound 8 the HOMO, though still bearing an overall \(\pi\) symmetry, is more centered on the P1-B1-P2 unit. The LUMO of 8 is of boron-boron \(\pi\)-antibonding character, with significant delocalization into the aren \(\pi\)-system. While the HOMO-LUMO gaps of the compounds are of similar magnitude, both of these orbitals are stabilized by the rearrangement, the HOMO to a slightly greater extent. The HOMO-1 of all three compounds represents the B-B sigma bonding interaction. Compound 8 bears some resemblance to a diboraallene compound recently reported by Kinjo and co-workers;\(^{[6]}\) however, in that case, the HOMO is localized on the B=C double bond, with the lower-lying HOMO-1 describing the boron-boron \(\pi\)-interaction.

Additionally, the B-B interaction in 6 and 8 was estimated within the Energy Decomposition Analysis formalism (EDA) under the assumption of equal B1-P1 bond energies in 6 and 8. Table 4 gives the values of the different bonding components. The total interaction energy is of slightly larger magnitude in 6. Both the orbital interactions and repulsive Pauli components of the interaction are larger in 8 — this can be ascribed to borylene lone pair donation to the boron centre, resulting in significant \(\pi\)-\(\pi\) interactions. The electrostatic component is larger in 8, reflecting the more polar nature of the diboron unit. Furthermore, the deformation densities (Figure 4), which depict the charge flow upon interaction of B1 in 6 and 8 with the rest of the corresponding molecule, indicate significant similarities and differences between the two isomers. In both 6 and 8, the main deformation density \(\Delta \rho_1\) shows charge accumulation towards the boron atom stabilized by PMe\(_3\). However, in 8, one can see that charge flows to B1 not only from B1-P1 (dative) but also from B2 (covalent) — confirming the borylene-like nature of B1. The B2\(\rightarrow\)B1 flow in 6 is only a secondary deformation density \(\Delta \rho_2\) and is of \(\pi\)-like symmetry, as expected for diborones; in 8, there is also some charge flow (\(\Delta \rho_2\)) towards the B2-B1 bond of \(\pi\)-like character, further supporting the assignment of B1 as a doubly dative-stabilized borylene.

<table>
<thead>
<tr>
<th>(\Delta \rho_1) properties</th>
<th>6 (gas)</th>
<th>6 (solv)</th>
<th>8 (solv)</th>
<th>8 (solv)</th>
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<tr>
<td>(d(BB))</td>
<td>1.618</td>
<td>1.593</td>
<td>1.612</td>
<td>1.593</td>
</tr>
<tr>
<td>Bond order(BB)</td>
<td>1.349</td>
<td>1.526</td>
<td>1.406</td>
<td>1.544</td>
</tr>
<tr>
<td>(q(B1))</td>
<td>-0.243</td>
<td>-0.129</td>
<td>-0.240</td>
<td>-0.133</td>
</tr>
<tr>
<td>(q(B2))</td>
<td>-0.019</td>
<td>-0.155</td>
<td>-0.038</td>
<td>-0.156</td>
</tr>
<tr>
<td>(q(Br))</td>
<td>-0.131</td>
<td>-0.141</td>
<td>-0.169</td>
<td>-0.167</td>
</tr>
</tbody>
</table>

Table 3. Calculated properties of compounds 6 and 8.

To summarize, we have used boron-silicon metathesis at B\(_2\)Br\(_{4}\) to develop a route to unsymmetrical diborones based upon a chelating benzylphosphine linker, with either a phosphine or an N-heterocyclic carbene as the second stabilizing ligand. DFT calculations confirm the slight polarity of these compounds. We have also reported thermodynamic rearrangement of one of these unsymmetrical trans-diborones to an unprecedented gem-diborone containing a polar boron-boron bond. On the basis of DFT calculations, which reveal large electrostatic and orbital contributions to the bonding, we propose that the compound is best described as a borylborylene.

Acknowledgements

The authors gratefully acknowledge the European Research Council (ERC) under the European Union Horizon 2020 Research and Innovation Program (grant agreement no. 669054) for funding. A.V. thanks the University of Sussex for financial support.

Keywords: boron • borylene • multiple bonds • rearrangement • DFT calculations


After the preparation of this manuscript, Kinjo and co-workers published an example of a gem-diborene as a result of a hydrogen atom transfer from a diborallene: W. Lu, Y. Li, R. Ganguly, R. Kinjo, J. Am. Chem. Soc. 2018, 140(4), 1253-1258. 


Unsymmetrical diborenes were prepared by reduction of cyclic diboranes(4) containing a benzylphosphine bridge. A doubly phosphine-stabilized diborene underwent an unprecedented, thermally induced rearrangement to form a borylborylene.

Tom E. Stennett, James D. Mattock, Ivonne Vollert, Alfredo Vargas* and Holger Braunschweig*

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