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Title: Competition between Halogen, Hydrogen and Dihydrogen Bonding in Brominated Carboranes

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COMMUNICATION

**Competition between Halogen, Hydrogen and Dihydrogen Bonding in Brominated Carboranes**


Abstract: Halogen bonds are a subset of noncovalent interactions with rapidly expanding applications in materials and medicinal chemistry. While halogen bonding is well known in organic compounds, it is new in the field of boron cluster chemistry. We have synthesized and crystallized carboranes containing Br atoms in two different positions, namely bound to C- and B-vertices. The Br atoms bound to the C-vertices have been found to form halogen bonds in the crystal structures. In contrast, Br atoms bound to B-vertices formed hydrogen bonds. Quantum chemical calculations have revealed that halogen bonding in carboranes can be much stronger than in organic architectures. These findings open new possibilities for applications of carboranes both in materials and medicinal chemistry.

Halogen bonds (X-bonds) are nonclassical noncovalent interactions between halogen atoms (X) and electron (e−) donors.[1] X-bonding belongs to the family of σ-hole interactions, and has received a lot of attention during the last decade.[2] Despite the first X-bond being described by Hassel as early as 1954,[3] the nature of the X-bond was only explained in 2007.[4] X-bonding is enabled by a region of partial positive charge (the so-called σ-hole), which is located on the top of the partially negative X atom. A σ-hole can be characterized by its size and magnitude (V_{\text{max}}), which in turn determines the properties of the X-bond.[5] The strength of an X-bond can thus be increased by increasing the V_{\text{max}} value (keeping the same e− donor). This has already been demonstrated in small model systems,[6] as well as in protein-ligand complexes.[6]

Polyhedral boron hydrides are a large group of compounds with unique properties, a wide range of applications,[7] and unusual noncovalent interactions.[8] The group includes compounds displaying 3D aromaticity and an ability to form dihydrogen bonds (HH-bonds)[9] and chalcogen bonds.[10] HH-bonds are unconventional proton-hydride H-bonds characterized by a short distance between H atoms. They generally occur between a positively charged hydrogen of a proton donor (e.g. CH, NH, OH groups) and a σ-bonding electron pair from an MH bond (M = electropositive atom, such as B, Al, Li). Many studies have shown the importance of HH-bonding in crystal packing and supramolecular chemistry.[11] Furthermore, it has been shown that carborane-based inhibitors interact with their biomolecular targets mainly via formation of HH-bonds.[12] Recently, we studied the properties of halogenated carboranes using quantum mechanics (QM) methods. These calculations showed that halogenated carboranes had large and highly positive σ-holes when an X atom was covalently attached to the C-vertex of a neutral carborane molecule.[13][14]

In this current study, we have synthesized the C_{12} symmetrical carboranes 9-Br-closo-1,2-C_{6}B_{10}H_{11} (1) and 1,12-Br_{2}-closo-1,2-C_{6}B_{10}H_{10} (2), which have been characterized by NMR spectroscopy (Table S1), and using X-ray crystallography. 1 and 2 are compared to 1-Ph-2-Br-closo-1,2-C_{6}B_{10}H_{10} (3), which was previously synthesized by Welch et al.[15] who also determined the crystal structure. The pairwise interactions in all three crystal structures (1–3) were studied using high-level QM calculations. We performed highly accurate MP2/S/CBS calculations[16] for all the binding motifs; benchmark calculations using CCSD(T)/CBS were also performed for selected interactions of 1.

Initially, we compared the computed properties of isolated molecules. 1 has a single Br atom bound to a B-vertex, which results in a large computed dipole moment (μ) of 6.4 D. As a consequence, the electrostatic potential (ESP) on the Br atom was found to be highly negative. The negative ESP on the Br atom was due to the fact that it is covalently bound to an electropositive B atom. The V_{\text{max}} of the Br atom on 1 thus had a negative value of −8.8 kcal mol\(^{-1}\), which was, however, less negative than the belt around the center of the atom. The anisotropic charge distribution on the Br atom points to a relative σ-hole. Because the entire surface of the Br atom has negative ESP, the Br atom of 1 can only act as an electron donor when involved in, for example, H-bonding.

Molecule 2 contains two differently bonded Br atoms. One is bound to a C-vertex and the second to a B-vertex. The V_{\text{max}} of the Br atom bound to the B-vertex has a negative value (−5.0 kcal mol\(^{-1}\)), as was the case for 1. However, the Br atom bound to the C-vertex has a positive ESP (V_{\text{max}} = 31.3 kcal mol\(^{-1}\)). Moreover, even the belt around the Br atom was found to have a positive ESP. This is a very large V_{\text{max}} value, particularly if we consider that the V_{\text{max}} values for bromobenzene and pentafluoro-bromobenzene are 11.2 and 29.7 kcal mol\(^{-1}\).
respectively. \textsuperscript{[13]} \textsuperscript{2} could thus form both H- and X-bonds (an H-bond exclusively via the Br bond to B-vertex, and an X-bond exclusively via the Br atom bound to C-vertex). \textsuperscript{2} has a smaller computed \( \mu \) value (5.2 D) than \textsuperscript{1}.

\textsuperscript{3} has a single Br atom bound to a C-vertex. The ESP of the Br atom of \textsuperscript{3} is less positive than the ESP of the corresponding Br atom on \textsuperscript{2}. The \( V_{\text{max}} \) value is smaller than that for \textsuperscript{2} but is still highly positive (25.7 kcal mol\(^{-1}\)). The ESP of the belt of the Br atom is, however, negative. The single Br atom of \textsuperscript{3} could thus form both X- and H-bonds simultaneously. The calculated \( \mu \) value for \textsuperscript{3} is about 4.8 D, which is in acceptable agreement with the experimental value of 4.1 D.\textsuperscript{[17]} The experimental \( \mu \) value for \textsuperscript{3} has been explained by its vector analysis.\textsuperscript{[17]} The value of \( \mu \) for \textsuperscript{3} was thus smaller than that calculated for \textsuperscript{1} and \textsuperscript{2}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure1.png}
\caption{ORTEP views and computed electrostatic potential (ESP) on 0.001 a.u. molecular surface at the HF/cc-pVDZ level. Color range of ESP in kcal mol\(^{-1}\).}
\end{figure}

Secondly, we solved the crystal structures of \textsuperscript{1} and \textsuperscript{2}. In the crystal structure of \textsuperscript{1}, two independent molecules were found in the unit cell. The shortest skeletal interatomic distance is attributed to the C–C moiety, which also corresponds to the supramolecular connectivity of one B–Br fragment with two C–H groups building thus the global array (Figure 1). In the previously reported but incomplete (hydrogen atoms not determined) crystal structure of \textsuperscript{2},\textsuperscript{[18]} the positions of the skeletal C atoms were based on simple comparison of interatomic distances between all pairs of skeletal atoms (the shortest separation was attributed to the HC–CB fragment). We have now entirely redetermined the structure of \textsuperscript{2} at lower temperature using state-of-the-art methods; the connectivity in the supramolecular architecture of \textsuperscript{2} is distinguished on the basis of connectivity between the E–Br (E = C or B atoms) and appropriate E–H moieties from the neighboring carborane. This approach takes into the account the difference in Allen electronegativities\textsuperscript{[19]} of elements and groups involved; e.g., \( \Delta \gamma \text{Allen}(\text{B–H}) = -0.249, \) while the differences for C–H, C–Br and B–Br groups are +0.244, –0.141 and –0.634, respectively.

Comparing Allen electronegativities also confirms the widely accepted fact the B–Br···H–C hydrogen bond is highly preferred (\( \Delta \gamma \text{Allen} = 0.878 \)). In general, the supramolecular architecture of 2Br\(_2\)-C\(_2\)B\(_10\) is best described as a helix-like linear chain of two units interconnected by the B–Br···H–C interactions. Moreover, each level of the helix is interconnected with another one of the same helix or neighboring helix by an C–Br···H–B X-bond or HH-bond. Because of substitutional/static disordering of the skeletal atoms within the structure of \textsuperscript{2}, averaging of atomic positions occurs. This leads to the shortest interatomic distance being between BrC–BH fragments. On the other hand, the elongation of the B–Br separation in comparison to the C–Br one by approximately 0.1 Å corresponds to the appropriate bond distance in \textsuperscript{1}.

Thirdly, we analyzed interactions in the crystal structures. The crystal packing and the binding motifs are shown in Figure 2. The interaction energy (\( \Delta E \)) values are summarized in Table 1. Each Br atom of \textsuperscript{1} made two H-bonds with C–H vertices of other molecules. These B–Br···H–C h-bonds were 2.7 Å long and had \( \Delta E \) of –6.0 and –5.7 kcal mol\(^{-1}\). The strength of the H-bond could be caused by the large partial negative charges on the Br atom and large \( \mu \).

The crystal packing of \textsuperscript{2} clearly demonstrates the two different interactions that the Br atoms can make. The Br atom bound to B-vertex forms well-known B–Br···H–C H-bonds, while the Br atom bound to the C-vertex is shown to form novel C–Br···H–B X-bonds. The H-bonding in \textsuperscript{2} is weaker than in that for \textsuperscript{1}, which might be due to the smaller \( \mu \) and sub-optimal arrangement. In this case, C–H interacts with the less negative top part of X atom, and not with the highly negative belt. Even though the \( V_{\text{max}} \) for \textsuperscript{2} was highly positive, \( \Delta E \) of the X-bond was only –2.9 kcal mol\(^{-1}\). These observations could also be due to the suboptimal arrangement of the X-bond.\textsuperscript{[20]} The optimal X-bond arrangement is linear, while the C–Br···H angle in this architecture is 166°. Moreover, the e–‘deficient carbonase cage is likely to be a bad e– donor. Interestingly, there are five binding motifs in the crystal structure with strengths that lie within a narrow range of 0.5 kcal mol\(^{-1}\). This could lead to significant competition between the different interactions during the growth of the supramolecular structure.
HH-bonds can be found in the A-C binding motif (see Figure 2), which is characterized by two symmetrical HH-bonds formed between B–H and the H atoms of the Ph rings. That A-C HH-bond was shorter (2.3 Å) and thus probably stronger. Taking half the ΔE value of the A-C binding motif could thus be considered as an upper limit for the strength of the additional interactions in the A-B motif. The estimated ΔE of the isolated X-bond would thus be about −4 kcal mol$^{-1}$, which is close to the ΔE value calculated for the 1···benzene complex (−3.3 kcal mol$^{-1}$). The reported X-bond can, however, be considered a particularly strong X-bond, when compared with X-bonds for organic compounds. For example, ΔE for the benzene···BrCH$_3$ and benzene···BrCF$_3$ complexes were reported to be about −1.8 and −3.1 kcal mol$^{-1}$, respectively. This comparison confirms the remarkable ability of heteroboron clusters to form very strong σ-hole interactions. Another exceptionally strong σ-hole binding motif has been reported in the crystal structure of 12-Ph-closo-1-SB$_3$H$_{10}$, where the chalcogen binding motif had ΔE of −8.6 kcal mol$^{-1}$.10

### Table 1. Interaction energies in kcal mol$^{-1}$

<table>
<thead>
<tr>
<th>Motif</th>
<th>Interaction</th>
<th>MP2.5/CCSD(T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9-Br-closo-1,2-C$_2$B$<em>3$H$</em>{11}$</td>
<td>H-bond</td>
<td>−5.85/−6.00</td>
</tr>
<tr>
<td>A-B</td>
<td>H-bond</td>
<td>−5.55/−5.69</td>
</tr>
<tr>
<td>A-D</td>
<td>Stacking</td>
<td>−2.32</td>
</tr>
<tr>
<td>1,12-Br$_2$-closo-1,2-C$_2$B$<em>3$H$</em>{10}$</td>
<td>Stacking</td>
<td>−3.37</td>
</tr>
<tr>
<td>A-B</td>
<td>Stacking</td>
<td>−3.32</td>
</tr>
<tr>
<td>A-C</td>
<td>(Parallel displaced)</td>
<td>−3.22</td>
</tr>
<tr>
<td>A-D</td>
<td>Stacking</td>
<td>−3.12</td>
</tr>
<tr>
<td>A-E</td>
<td>H-bond</td>
<td>−2.91</td>
</tr>
<tr>
<td>A-F</td>
<td>X-bond</td>
<td>−6.73</td>
</tr>
<tr>
<td>1-Ph-2-Br-closo-1,2-C$_2$B$<em>3$H$</em>{10}$</td>
<td>X-bond</td>
<td>−5.09</td>
</tr>
<tr>
<td>A-B</td>
<td>X-bond</td>
<td>−3.52</td>
</tr>
<tr>
<td>A-C</td>
<td>H-bond</td>
<td>−2.82</td>
</tr>
</tbody>
</table>

To sum up, we have compared the crystal packing of carboranes containing Br atoms attached to different cage atoms. When Br is bound to C-vertices X-bonds are formed, while Br bound to B-vertices formed H-bonds. QM analyses revealed the strength of the X-bonds in carboranes. The knowledge about noncovalent interactions of halogenated carboranes can be employed in various applications. Besides crystal engineering, it can be used, for example, to increase binding affinity of carborane-based inhibitors towards their biomolecular targets. Furthermore, X-bonds might play an important role in chemical reactions that involve halogenated carboranes. The knowledge obtained from this study might, therefore, also be useful when designing novel synthetic paths.

### Experimental Section

See Supporting Information for full experimental details.
The synthetic procedure for 1 and 2 was based on the gradual bromination of o-carborane. At first, the o-carborane reacted with Br₂ and AlBr₃ and 1 was obtained. Subsequently, 1 underwent magnesiation-based bromination with C₅H₅MgBr and Br₂ to obtain 2. For further details see Reference 23.

As compounds 1 and 2 have not so far been characterized by using NMR spectroscopy, the principal experimental structural tool in boron cluster chemistry, we have performed such spectroscopic studies to determine the architectures of 1 and 2 unambiguously. The corresponding shielding tensors were also computed.

Crystals of 1 and 2 were grown by slow evaporation of hexane solutions. Crystallographic data for structural analysis have been deposited with the Cambridge Crystallographic Data Centre (1459071 and 1459072 for 1 and 2, respectively).

Quantum mechanical (QM) calculations were performed using the X-ray crystal structures. Hydrogen atoms of the crystal structures were optimized using the newly parameterized DFT-D3/BLYP/DZVP method. ΔE values were obtained using MP2.5 and CCSD(T) complete basis set calculations. Electrostatic potentials were computed at HF/cc-pVDZ level. It has recently been shown that larger basis sets are not needed for this purpose.[24]

Acknowledgements

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Keywords: carborane•sigma hole•halogen bond•bromine•X-ray crystal structure


Substituted carboranes can contain Br atoms in two positions, namely bound to C- and B-vertices. When Br atoms are bound to C-vertices C–Br···H–B halogen bonds can be formed in the crystal structure. This is in contrast to Br atoms bound to B-vertices where B–Br···H–C hydrogen bonds are observed.