A novel stibacarbaborane cluster with adjacent antimony atoms exhibiting unique pnictogen bond formation that dominates its crystal packing†

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We have prepared nido-7,8,9,11-Sb2C2B7H9, the first cluster with simultaneous Sb−B, Sb−C and Sb−Sb atom pairs with interatomic separations with magnitudes that approach the respective sums of covalent radii. However, the length of the Sb−Sb separation in this cluster is slightly less than the sum of the covalent radii. Quantum chemical analysis has revealed that the crystal packing of nido-7,8,9,11-Sb2C2B7H9 is predominantly dictated by pnictogen (Pn) bonding, an unconventional σ-hole interaction. Indeed, the interaction energy of a very strong Sb2−H⋯B Pn-bond in the nido-7,8,9,11-Sb2C2B7H9 dimer exceeds −6.0 kcal mol−1. This is a very large value and is comparable to the strengths of known Pn-bonds in Cl2Pn−⋯− complexes (Pn = As, Sb).

Introduction

Boron atoms have a remarkable ability to form 3-center-2-electron (3c2e) bonds, which results in an astonishing variety of stable three-dimensional molecular structures composed of triangular B−B−B and B−H−B faces. An important class of boranes comprises closo-BnHn2− and nido-BnHn4−. BH vertices can be formally replaced by CH+, S2+, Se2+, P+, As+ and Sb+ units which result in a variety of heteroboranes. While carboranes and thiaboranes are common, heteroboranes with a pnictogen (Pn) atom. Consequently, σ-hole interactions are often called X-, Y- or Pn-bonds. These interactions are well known in organic chemistry. A σ-hole is most often characterized by its magnitude (Vx,max); the higher the Vx,max value, the stronger the X-bond. The Vx,max value can be increased by increasing the atomic number of the X atom or by changing...
the chemical environment. Recently, σ-hole interactions within heteroboranes with positive σ-holes on heteroatoms incorporated have received attention due to the remarkably strong σ-hole interactions. This is counterintuitive when one considers the very low electronegativity of B atoms. It was a phenyl-exo-substituted thiaborane molecule where the σ-hole interactions were first noticed with the resulting formation of a Y-bond that was responsible for the overall packing motif in the crystal structure. A strong X-bond has been observed in the crystal packing of Br-exo-substituted carbaborane. Although a series of heteroboranes containing Pn atoms is already known experimentally such as 1 and 2, no experimental evidence for Pn-bonding in heteroboranes has been reported. Therefore, we used 3 to validate the ability of heteroboranes to form noncovalent Pn-bonds.

Results and discussion

Structural characterization by NMR

The properties of 3 can be inferred from NMR experiments and QM calculations. Table 1 compares experimental and computed 11B NMR chemical shifts, showing excellent agreement between the experimental and computed NMR values with the exception of B(3,6), which are antipodally coupled to the Sb atoms. The discrepancy, which is not unusual for the third row and other heavier elements, indicates that B(5,6) atoms are strongly influenced by Sb(7,8).

Aromatic ring currents are observed in 2D and 3D aromatic molecules such as benzene and closo-B12H12. The nucleus-independent chemical shift (NICS) is a computational method that expresses the absolute magnetic shielding at the center of a 2D or 3D molecule. The NICS values for various parts of molecule 3 are computed to be −41.7, −44.5, −41.7, −44.9, −22.5 and −17.4 ppm for the Sb–B–B, B–B–B, Sb–Sb–B, and Sb–B–C triangles, center of the molecule, and open pentagonal belt, respectively. These results indicate that the cluster as a whole is of aromatic nature. The fact that the triangles that are closest to antimony are highly aromatic indicates that the Sb atoms donate valence electrons to the corresponding atomic triangles. Even the NICS values that are less negative than −40 ppm are comparable to the aromaticity displayed by the 1,3-dehydro-5,7-adamantenediy1 dication and closo-B12H12 (NICS values of −49.6 and −34.4 ppm, respectively). The NICS value for the Sb–Sb–C–B–C pentagon of −17.4 ppm is comparable to the values associated with pyrrole and thiophene (NICS values of −17.3 and −14.7 ppm, respectively; more details in ref. 13). The 7,8,9,11-Pn2C2B7H9 (Pn = P, As, and Sb) compounds display similar trends of NICS values.

Crystal structure of 3

In the crystal structure of 3 (see Fig. 2), four molecules were found in the chiral unit cell. A striking feature of this structure is the well-determined Sb–Sb separation (2.7346(6) Å), which seems to be in line with analogous distances in reduced or over-reduced Sb compounds containing two or more metal atoms. There are several different classes of compounds that could be used for comparison where the Sb–Sb distances are a bit longer than the sum of the covalent radii of 2.8 Å.

Complexes with Sb–B and Sb–C bonds in a single molecule are very rare. The reported Sb–C separation of 3 is elongated by ca. 0.07 Å when compared to the typical C(aromatic)-SbIII distance. Similarly, the Sb–B distances in 3 (range from 2.312 to 2.458 Å) approach the sum of the covalent radii of the constituent atoms, i.e. 2.25 Å.

Electrostatic potential (ESP) analysis

In order to understand Pn-bonding more generally, we analyzed the charge distributions for a series of PnCl3 and nido-Pn2C2B7H9 (Pn = P, As, and Sb) molecules. The computed Vmax and dipole moment (μ) values are summarized in Table 2. Sb has very low electronegativity comparable to that of B but considerably smaller than those values for C and Cl. In accordance with the concept of electronegativity, the Sb atom has highly positive Vmax values in SbCl3 of 48.6 kcal mol−1. However, the Sb atoms in 3 also have a very high Vmax value of 42.7 kcal mol−1 (the σ-hole in between the adjacent Sb atoms; see Fig. 3) despite being bonded to B and C atoms, which are considerably less electropositive than Cl atoms. Interestingly, the Sb vertices have more positive ESP surfaces than the CH vertices.

### Table 1 NMR results for 3

<table>
<thead>
<tr>
<th></th>
<th>B(1)</th>
<th>B(3)</th>
<th>B(2,4)</th>
<th>B(10)</th>
<th>B(5,6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GIAO</td>
<td>−29.3</td>
<td>−15.5</td>
<td>−6.5</td>
<td>−0.4</td>
<td>1.6</td>
</tr>
<tr>
<td>Exp.</td>
<td>−30.2</td>
<td>−16.8</td>
<td>−9.2</td>
<td>−1.5</td>
<td>−6.9</td>
</tr>
</tbody>
</table>

*All in ppm with respect to BF3·OEt2. Atomic numbering shown in Fig. 1. GIAO-MP2/II(def2-TZVP+ECP on Sb)/MP2/cc-pVTZ(def2-TZVP+ECP on Sb)."
Crystal packing and noncovalent interactions

We have studied pairwise interactions in the crystal structure of 3. It was not possible to compare these interactions to the parent nido-7,8,9,11-Pn2C2B7H9 (Pn = P, As) complexes, as it reveals poor quality crystals, which prevents them from performing reliable X-ray structure determination. Hence, we opted for the halogenated analogues of 1 and 2, for which the crystal structures had already been determined.6,7 We performed highly accurate MP2.5/CBS calculations for all of the bonding motifs as well as benchmark CCSD(T)/CBS calculations for selected motifs. All the computed interaction energies (ΔE) values are summarized in Table 3 (see also Fig. 4). Each molecule of 1 makes two Pn-bonds (P2⋯H–B and P2⋯Cl–B). These two Pn-bonds form the A⋯B binding motif which has ΔE of −6.1 kcal mol⁻¹ and was the dominant binding motif for 1. Besides Pn-bonding, molecule 1 also forms a CH⋯Cl–B H-bond and a C–H⋯B–H HH-bond. The H-bonding of 1 is, however, weaker than its Pn-bonding and the ΔE of the A⋯C motif is less negative (−4.4 kcal mol⁻¹).

The crystal packing of 2 is also dominated by Pn-bonding, as the binding motifs with most negative ΔE values are stabilized by Pn-bonding. The A⋯B motif of 2 is stabilized by two symmetrical As⋯H–B Pn-bonds and has ΔE of −6.0 kcal mol⁻¹. An estimated ΔE of this isolated Pn-bond would thus be about −3 kcal mol⁻¹. The A⋯C motif of 2 has a comparable ΔE of −5.7 kcal mol⁻¹ and is stabilized by As⋯H–B and As⋯I–B Pn-bonds. The other motifs have less negative ΔE values. Among them, the most negative is the D⋯E motif with two As⋯H–B Pn-bonds and ΔE of −4.5 kcal mol⁻¹.

The crystal structure of 3 has two binding motifs with ΔE exceeding −6 kcal mol⁻¹. The A⋯B motif has ΔE of −7.1 kcal mol⁻¹ and is stabilized by multiple interactions, specifically by two Sb⋯H–B Pn-bonds and one C–H⋯H–B HH-bond. On the other side, the A⋯C motif is exclusively formed by a single interaction, namely a Sb⋯H–B Pn-bond. ΔE of −6.5 kcal mol⁻¹ computed for this binding motif can thus be considered as ΔE of the isolated Pn-bond. This is a very large value, exceeding ΔE of isolated Pn-bonding of 2 by about 100%. The Pn-bonding of 3 is even slightly more favorable than Pn-bonding in Cl5SB⋯N(Me)3 and Cl5SB⋯benzene complexes (ΔE of −5.8 and −6.1 kcal mol⁻¹, respectively).15

Table 2  Magnitude of e-holes (Vmax) on the surfaces of Group V and VII atoms and dipole moments (μ) computed at the HF/cc-pVDZ level in various nictogen compounds

<table>
<thead>
<tr>
<th>Vmax/kcal mol⁻¹</th>
<th>Group V atoms</th>
<th>Group VII atoms</th>
<th>μ/D</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCl3</td>
<td>3 × 29.2</td>
<td>3 × 6.7</td>
<td>1.1</td>
</tr>
<tr>
<td>AsCl3</td>
<td>3 × 38.6</td>
<td>3 × 2.9</td>
<td>2.1</td>
</tr>
<tr>
<td>SbCl3</td>
<td>3 × 48.6</td>
<td>3 × −3.3</td>
<td>3.3</td>
</tr>
<tr>
<td>nido-7,8,9,11-Pn2C2B7H9 (1)</td>
<td>24.8</td>
<td>—</td>
<td>2.0</td>
</tr>
<tr>
<td>3-Cl-nido-7,8,9,11-Pn2C2B7H9</td>
<td>25.4</td>
<td>−3.1</td>
<td>2.8</td>
</tr>
<tr>
<td>nido-7,8,9,11-As2C2B7H9</td>
<td>31.1 and 2 × 24.4</td>
<td>—</td>
<td>2.8</td>
</tr>
<tr>
<td>3-Cl-nido-7,8,9,11-As2C2B7H9 (2)</td>
<td>32.2 and 2 × 29.0</td>
<td>8.2</td>
<td>3.5</td>
</tr>
<tr>
<td>nido-7,8,9,11-Sb2C2B7H9 (3)</td>
<td>1 × 42.7 and 2 × 25.7</td>
<td>—</td>
<td>4.0</td>
</tr>
</tbody>
</table>

(see Fig. 3) and the Sb atoms lie at the center of a partial positive charge within 3. A very large μ of 3 (μ of 4.0 D) also confirms that the center of the Sb–Sb vector is positively charged.

Methods

Synthesis

A solution of arachno-4,6-C2B7H13 (0.226 g, 2 mmol, see also Fig. 5) in dichloromethane (20 mL) was treated with proton sponge (1.5 g, 7 mmol). SbCl3 (2.0 g, 16 mmol) was added under stirring and cooling to 0 °C. Stirring was continued at room temperature for 4 h and the mixture was then cooled to 0 °C and decomposed by the dropwise addition of water (20 mL). Column chromatography was carried out on a silica gel as the stationary phase (3 g). The solids were mounted onto a silica gel column and the column was eluted with dichloromethane. The chromatography led to the separation of a major yellow fraction of Rf = 0.73 (CH2Cl2), which was evaporated to dryness. 0.255 g of yellowish nido-7,8,9,11-Sb2C2B7H9 was obtained (yield 36%, with respect to the starting material 4,6-C2B7H13).

Computational modeling

Magnetic shielding was calculated using the GIAO-MP2 method incorporated into Gaussian0916 utilizing the IGLO-II basis with
the MP2/cc-pVTZ geometry (with the TZVP basis set on Sb by Weigend\(^{17}\) with ECP by Metz\(^{18}\)) and frozen core electrons. Electrostatic potentials were computed at the HF/cc-pVDZ level for 1 basis set in ref. 19) using Gaussian09 and Molekel4.\(^{20}\)

Interactions energy (\(\Delta E\)) values were calculated for all pairwise interactions of the crystal structures of 1, 2 and 3. All hydrogen atoms were optimized using the DFT-D3/BLYP/DZVP method prior to the energy calculations.\(^{11}\) \(\Delta E\) for the central molecule with each surrounding molecule was evaluated at MP2.5/CBS using the Turbomole 6.6\(^{22}\) and Cuby\(^{21}\) programs. MP2.5/CBS was calculated as the sum of MP2/CBS energy and MP2.5 correction. MP2/CBS was approximated by RI-MP2-F12/cc-pVTZ-F12 for 1.\(^{24}\) In the case of 2 and 3, we used MP2 extrapolation to the CBS from cc-pVTZ to cc-pVQZ (for I and Sb atoms cc-pVTZ-PP and cc-pVQZ-PP pseudopotentials were used).\(^{25}\) The MP2.5 correction term was calculated using the aug-cc-pVDZ basis set. MP2.5/CBS interaction energies were compared to benchmark CCSD(T)/CBS for the interaction motifs of 1 and the most favourable motifs of 3. The CCSD(T) correction term was also determined using the aug-cc-pVDZ basis sets. Counterpoise corrections for basis set superposition error (BSSE) are used for all energy calculations.

### Crystallography

Full sets of diffraction data for 3 were collected at 150(2) K using a Bruker D8-Venture diffractometer equipped with a Mo (Mo/K\(_{\alpha}\) radiation; \(\lambda = 0.71073\) Å) microfocus X-ray (μS) source, a Photon CMOS detector and an Oxford Cryosystems cooling device.

The frames were integrated using the Bruker SAINT software package using a narrow-frame algorithm. Data were corrected for absorption effects using the Multi-Scan method (SADABS). Data were treated by using XT-version 2014/5 and SHELXL-2014/7 software implemented in an APEX3 v2016.5-0 (Bruker AXS) system.\(^{26}\) Hydrogen atoms were mostly localized on a difference Fourier map; however, to ensure uniformity of treatment of the crystal, all hydrogen atoms were recalculated into idealized positions (riding model) and assigned temperature factors \(H_{iso}(H) = 1.2U_{eq}\) (pivot atom). H atoms in were placed with C–H distances of 1.1 Å for B–H and C–H bonds in the carborane cage. Hydrogen atoms in the upper rim of the cage were placed according to the appropriate maxima found on the Fourier difference electron density map.

\[
R_{int} = \frac{\sum |F_o - F_c|}{\sum F_o}; S = \frac{[\sum w(F_o^2 - F_c^2)^{\frac{1}{2}}]/[N_{	ext{diff}} - N_{\text{params}}]}{[\sum w(F_o^2)^{\frac{1}{2}}]} \text{ for all data,} \quad R(F) = \frac{\sum |F_o| - |F_c|/\sum |F_o|} \quad \text{for observed data,} \quad wR(F) = \frac{\sum w(F_o^2 - F_c^2)^{\frac{1}{2}}/\sum w(F_o^2)^{\frac{1}{2}}}]^{\frac{1}{2}}
\]

Crystallographic data for structural analysis have been deposited with the Cambridge Crystallographic Data Centre, CCDC no. 1553208 for 3\(^{\dagger}\).

Crystallographic data and structural refinement parameters for colorless single crystals of 3: C\(_3\)H\(_8\)B\(_{2}\)Sb\(_2\), \(M = 352.26\text{ g mol}^{-1}\),

![Fig. 5](image_url)
orthorhombic, \( Pna2_1 \), \( a = 20.4799(12) \), \( b = 6.6471(4) \), \( c = 6.5803(4) \) \( \AA \), \( \beta = 90^\circ \), \( Z = 4 \), \( V = 895.79(9) \) \( \AA^3 \), \( D_e = 2.612 \) g cm\(^{-3}\), \( \mu = 5.946 \) mm\(^{-1}\), \( T_{\text{min}}/T_{\text{max}} = 0.583/0.745 \); \( -27 \leq h \leq 27 \), \( -8 \leq k \leq 8 \), \( -8 \leq l \leq 8 \); 19,233 reflections measured (\( \Delta \theta_{\text{max}} = 28.35^\circ \)), 2223 independent (\( R_{\text{int}} = 0.1199 \)), 1796 with \( I > 2\sigma(I) \), 106 parameters, \( S = 1.049 \), \( R_1 \) (obs. data) = 0.0271, \( wR_2 \) (all data) = 0.0398; max., min. residual electron density = 1.091, \(-1.107 \) e \( \AA^{-3} \).

Conclusions

To sum up, in compound 3 (\( \text{nido-7,8,9,11-Sb}_2\text{C}_2\text{B}_7\text{H}_8 \)) we have synthesized and crystallized the very first reported cluster containing simultaneously Sb–B, Sb–C and Sb–Sb covalent bonds.

Further quantum chemical analysis revealed that the most stable crystal motifs in the crystal structure of \( \text{Cl}_3\text{Pn} \) occupy the same positions as Sb in \( \text{Cl}_3\text{Pn} \). Phosphorus and arsenic can bond in the same positions as Sb in \( \text{Cl}_3\text{Pn} \). However, in contrast to nido-7,8,9,11-\( \text{Pn}_2\text{C}_2\text{B}_7\text{H}_8 \) (\( \text{Pn} = \text{P}, \text{As} \)), the energy of a very strong Sb–H–B Pn-bond would exceed \( -6.0 \) kcal mol\(^{-1}\). For comparison, the Pn-bonding in \( \text{Cl}_3\text{Pn} \) complexes (\( \text{Pn} = \text{As}, \text{Sb} \)) is less than the sum of Sb covalent radii. Further quantum chemical analysis revealed that the most stable crystal motifs in the crystal structure of \( \text{Cl}_3\text{Pn} \) occupy the same positions as Sb in \( \text{Cl}_3\text{Pn} \). Phosphorus and arsenic can occupy the same positions as Sb in \( \text{Cl}_3\text{Pn} \).

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the research project RVO 61388963 of the Czech Academy of Sciences. We acknowledge the financial support of the Czech Science Foundation (17-08045S).

References


