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The synthesis and structural aspects of the perbromo-functionalised thiaboranes closo- $SB_nBr_n$  (n = 5, 9, 11): the solid-state structure of the octahedral closo-SB<sub>5</sub>Br<sub>5</sub>, governed by strong dihalogen contacts†

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Co-pyrolysis reactions of B<sub>2</sub>Br<sub>4</sub> with S<sub>2</sub>Br<sub>2</sub> at 350 °C in vacuo yielded the brominated thiaboranes closo-SB<sub>5</sub>Br<sub>5</sub> (1), closo-1-SB<sub>9</sub>Br<sub>9</sub> (2) and closo-SB<sub>11</sub>Br<sub>11</sub> (3), confirmed by high-resolution mass spectrometry, experimental and computational <sup>11</sup>B NMR spectroscopy. The strong  $Br^{\delta+}(\sigma-hole)\cdots Br^{\delta-}(ring)$  attraction has been the decisive energy contribution in the crystal of 1.

Small polyhedral borane and heteroborane clusters of octahedral shape have so far been limited to the borate dianions  $[B_6X_6]^{2-}$ (X = H, Cl or Br), the carboranes<sup>2</sup> 1,2- or 1,6-C<sub>2</sub>B<sub>4</sub>H<sub>6</sub> and the perhalogenated heteroboranes  $P_2B_4X_4$  (X =  $Cl^{3a}$  or  $Br^{3b}$ ),  $As_2B_4X_4$  $(X = Cl^{3c} \text{ or } Br^{3d})$ ,  $SB_5Cl_5$ ,  $^{3e} SeB_5Cl_5$   $^{3f,g}$  and  $TeB_5Cl_5$ .  $^{3h}$  In particular, recent studies of the thermal reactions of B2Cl4 with Se2Cl2 or TeCl<sub>4</sub>, producing the octahedral SeB<sub>5</sub>Cl<sub>5</sub><sup>3f,g</sup> and TeB<sub>5</sub>Cl<sub>5</sub><sup>3h</sup> have shown surprising results: chalcogens heavier than sulphur apparently induce stronger perturbations to the corresponding cluster frameworks, as manifested by the long body diagonal in chlorinated octahedral selena- and telluraboranes, which may prevent their accommodation in octahedral moieties. There are two known pnictogenaboranes, P<sub>2</sub>B<sub>4</sub>Br<sub>4</sub> and As<sub>2</sub>B<sub>4</sub>Br<sub>4</sub>, but no brominated octahedral chalcogenaborane as yet, which has prompted us to carry out the thiaborane reaction<sup>3e</sup> with S<sub>2</sub>Br<sub>2</sub>. On that basis, we wish to report the synthesis and characterisation of closo-SB<sub>5</sub>Br<sub>5</sub> (1) and other new brominated thiaboranes.

We have studied the co-pyrolysis reaction of  $B_2Br_4$  with  $S_2Br_2$ at temperatures above 350 °C. In comparison with chlorinated

The procedure described yielded three new brominated thiaboranes, closo-SB<sub>5</sub>Br<sub>5</sub> (1), closo-1-SB<sub>9</sub>Br<sub>9</sub> (2) and closo-SB<sub>11</sub>Br<sub>11</sub> (3) (Scheme 1), of which we were able to isolate only compound 1 by repeated vacuum-fractionation and crystallisation processes. Compound 1 was obtained in 5% yield based on the idealised eqn (1). The non-separable compounds 2 and 3 (eqn (2) and (3)) were formed in approximately 1% and 3% yields, respectively.

$$11B_2Br_4 + S_2Br_2 \rightarrow 2SB_5Br_5 (1) + 12BBr_3$$
 (1)

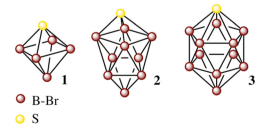
$$19B_2Br_4 + S_2Br_2 \rightarrow 2SB_9Br_9 (2) + 20BBr_3$$
 (2)

$$23B_2Br_4 + S_2Br_2 \rightarrow 2SB_{11}Br_{11} (3) + 24BBr_3$$
 (3)

The new thiaboranes 1-3 are thermally stable under inert atmosphere at least up to their formation in two experiments at 350 °C and 430 °C. In the very high vacuum applied in EI-MS  $(10^{-5} \, \mathrm{mbar})$ , 1 starts to sublime already at 25 °C, 2 at 50 °C and 3 at 85 °C. They are soluble in common aprotic solvents such as benzene, chloroform and methylene chloride.

The corresponding reaction of B<sub>2</sub>Cl<sub>4</sub> with S<sub>2</sub>Cl<sub>2</sub> has resulted in a much higher number of individual chlorinated thiaboranes closo-SB<sub>n</sub>Cl<sub>n</sub> (with n = 4, 5, 6, 9, 10, 11, 12) than in the

<sup>†</sup> Electronic supplementary information (ESI) available. CCDC 2403282. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.



Scheme 1 Molecular diagrams of closo-SB<sub>5</sub>Br<sub>5</sub> (1), closo-1-SB<sub>9</sub>Br<sub>9</sub> (2) and closo-SB<sub>11</sub>Br<sub>11</sub> (3).

thiaboranes, prepared at 280 °C, this reaction required significantly higher temperatures because the bromo compounds are less volatile than the Cl-synthons.<sup>3e</sup>

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present study. This may be attributed to the fact that the isodesmic heats of formation of the bromo-products are less negative than those of the chloro-products. 3e However, computational studies have confirmed in both cases that octahedral, bicapped square-antiprismatic and icosahedral cages are the most stable in the entire series. The computed isodesmic heats of formation in the origination of closo-SB<sub>n</sub>Br<sub>n</sub> from closo- $SB_{n-1}Br_{n-1}$  (n = 5, 9, 11) at the B3LYP/def2-QZVP//BP86/DZVP-DFT level (in kcal mol<sup>-1</sup>) are as follows:

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$$closo$$
-1-SB<sub>4</sub>Br<sub>4</sub> + B<sub>2</sub>Br<sub>4</sub>  $\rightarrow closo$ -SB<sub>5</sub>Br<sub>5</sub> (1) + BBr<sub>3</sub> -15.6

$$closo$$
-4-SB<sub>8</sub>Br<sub>8</sub> + B<sub>2</sub>Br<sub>4</sub>  $\rightarrow closo$ -1-SB<sub>9</sub>Br<sub>9</sub> (2) + BBr<sub>3</sub> -19.6

$$closo$$
-2-SB<sub>10</sub>Br<sub>10</sub> + B<sub>2</sub>Br<sub>4</sub>  $\rightarrow closo$ -SB<sub>11</sub>Br<sub>11</sub> (3) + BBr<sub>3</sub> -34.3

The new thiaboranes have been characterised by <sup>11</sup>B NMR spectroscopy, high- and low-resolution mass spectrometry, X-ray structure determination for 1 and SO-DFT/ZORA/NMR computations (taking spin-orbit coupling (SO) into account), which have confirmed the experimental <sup>11</sup>B NMR chemical shifts. With SO not included in 11B NMR chemical-shift computations, the values have shifted upfield by about 5 ppm<sup>4</sup> with respect to the SO-DFT/ZORA/NMR results. Based on these results, we have derived the molecular structures of 1, 2 and 3 in their solutions (see also Table 1).

The <sup>11</sup>B chemical shifts of 1, 2 and 3 are within the normal range for halogenated heteroborane clusters<sup>3a-h</sup> and closely follow the trends established for non-halogenated heteroboranes.<sup>5a</sup> They also reflect the symmetries of  $C_{4v}$ ,  $C_{4v}$  and  $C_{5v}$ , respectively.

As expected, 1 shows two broad signals in a 4:1 ratio at -2.3ppm  $(h_{1/2} \approx 106 \text{ Hz}, B(2-5))$  and 19.4 ppm  $(h_{1/2} \approx 113 \text{ Hz}, B(6))$ , which are in agreement with the signals for closo-SB<sub>5</sub>Cl<sub>5</sub> at −0.3 ppm and 19.4 ppm.<sup>3e</sup> With the application of the Lorentz-Gaussian transformation with a  ${}^{1}J({}^{11}B-{}^{11}B)$  of 25 Hz, the signals for B(2-5) are split into a quartet, indicating the coupling with

Table 1 The computed<sup>a</sup> and experimental <sup>11</sup>B NMR chemical shifts of 1, 2 and 3

1 Octahedral 1	motif		
		B(6)	
Comp. Exp.		20.5 19.4	
2 Bicapped sq	uare-antiprismatio	e motif	
	B(2-5)	B(6-9)	B(10)
Comp. Exp.	8.9 7.5	-4.7 -3.7	53.5 49.6
3 Icosahedral	motif		
	B(2-6)	B(7-11)	B(12)
Comp.	-1.7 -1.4	-6.0 -3.7	14.6 14.2

<sup>&</sup>lt;sup>a</sup> SOC-PBE0/T2ZP//B3LYP/6-311 +  $G^{**}$ .

B(6) and showing the expected cross-peak in the COSY <sup>11</sup>B<sup>11</sup>B NMR with the low-field signal.

The <sup>11</sup>B NMR of 2 consists of three signals in a 4:4:1 ratio at -3.7 ppm (B6-9), 7.5 ppm (B2-6) and 49.6 ppm (B10), which are in good agreement with the signals for closo-SB<sub>9</sub>Cl<sub>9</sub> at -0.3 ppm, 13.1 ppm and 48.8 ppm.  $^{3e}$  The signal at -3.7 ppm shows <sup>11</sup>B<sup>11</sup>B cross-peak correlation with the other two signals and thus can be assigned to B(6-9), the signal at 7.5 ppm to B(2-5), and the poorly-resolved intensity-one signal at 49.6 ppm can be connected with the antipodal downfield shift<sup>5</sup> of B(10).

The <sup>11</sup>B NMR of 3 consists of three signals in a 5:5:1 ratio at -3.7 ppm (B7-11), -1.4 ppm (B2-6) and 14.2 ppm (B12), with expected cross-peaks between B(7-11)/B(2-6) and B(7-11)/B(12). The signals are in agreement with the chemical shifts established for closo-SB<sub>11</sub>Cl<sub>11</sub> at -3.9 ppm, 1.0 ppm and 14.4 ppm.<sup>3e</sup>

The 70 eV EI mass spectra of all compounds show a strong molecular ion. In the case of thiaborane 1, the major cut-off indicates the abstraction of a SBBr edge from SB5Br5 to leave a  $[B_4Br_4]^+$  fragment. The formation of such a  $[B_4Hal_4]^+$  fragment has been analogously found in the mass spectra of closo-SB<sub>5</sub>Cl<sub>5</sub><sup>3e</sup> and closo-1,2-P<sub>2</sub>B<sub>4</sub>Cl<sub>4</sub>.<sup>3a</sup> The MS of thiaborane 2 shows the stepwise abstraction of S, Br, BBr<sub>2</sub> and BBr<sub>3</sub> fragments, whereas 3 exhibits only minimal fragmentation with the abstraction of a single Br atom, which indicates the high stability of the icosahedral cluster framework.

All mass-spectroscopy patterns are consistent with the computed spectra based on natural isotopic abundances. On the basis of simple skeletal electron-counting rules, 6 1, 2 and 3 should adopt octahedral, bicapped square-antiprismatic and icosahedral geometries, respectively, with sulphur contributing four electrons and each BBr unit two electrons to the cluster bonding. The employment of intrinsic-bond orbitals (IBOs)<sup>3e,g,h</sup> has revealed the nature of bonding in 1, 2 and 3. According to the expansion coefficients (ECs: contributions from individual atoms to a particular IBO to reveal the nature of such orbitals), the sulphur involvement in 1 consists of two IBOs with the ECs (the contributing atom in parentheses) of 1.15 (S), 0.51 (B) and 0.25 (B) and one IBO with the ECs of 1.20 (S) and 0.53 (B). Whereas the first pair of IBOs can be assigned to 3c-2e bonding, the second IBO is of 2c-2e type. The rest of the octahedral cage is assembled through 3c-2e IBOs (Fig. 1). The bonding schemes in 2 and 3 are shown in ESI† (Fig. S2).

The ESP (electrostatic potential molecular surface) of 1 is similar to that of its chlorine analogue, 3e but the bromine

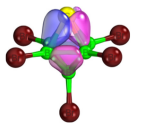


Fig. 1 Visualised IBOs for SB<sub>5</sub>Br<sub>5</sub> (1). The colour coding is as follows: blue classical 2c-2e bonding, pink 3c-2e bonding

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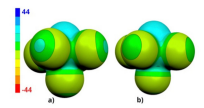


Fig. 2 The ESP molecular surfaces of 1 (a) and SB<sub>5</sub>Cl<sub>5</sub> (b) computed at the HF/cc-pVDZ level.

Table 2 The minimum and maximum values of the ESP molecular surfaces computed at the HF/cc-pVDZ level

	$V_{\rm s,max}$ [kcal mol <sup>-1</sup> ]		$V_{\rm s,min}$ [kcal mol <sup>-1</sup> ]	
Compound	Sulphur atom	Halogen atoms	Halogen atoms	
$SB_5Br_5(1)$ $SB_5Cl_5$	$24.5 \\ 24.8^{3e}$	$4 \times 11.6; 10.2$ $4 \times 6.6; 5.7$	$4 \times -6.0; -5.8$ $4 \times -5.6; -5.1$	

atoms of 1 have more positive  $\sigma$ -holes than the chlorines of SB<sub>5</sub>Cl<sub>5</sub> (see Fig. 2 and Table 2).

To confirm the octahedral structure of 1, single crystals were grown via vacuum sublimation and subjected to X-ray diffraction (see Fig. 3). Compound 1 crystallises in  $P2_1/c$  with three crystallographically independent closo-SB5Br5 octahedrons present in the corresponding asymmetric unit. As a consequence of symmetry, a total of twelve molecules are present in the centrosymmetric monoclinic cell (see ESI,† for details). Crystal packing is dominated by short contacts between the bromine atoms of neighbouring molecules, indicative of dihalogen bonding  $(d(Br \cdot Br) = 339 \text{ to } 368)$ pm, see also Fig. 4). Intermolecular bonding interactions between the sulphur and the bromine atoms, however, seem to be absent because no short contacts (<the sum of vdW radii) have been observed.

All the interaction motifs present in the crystal structure of 1 have been investigated using a cluster model. Two-body interaction energies have been computed for the selected crystallographic molecule using the SAPT method,7 which enables

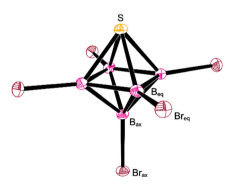


Fig. 3 ORTEP plot (thermal ellipsoids drawn at the 50% probability level) of the molecular structure of closo-SB5Br5 (1). The asymmetric unit contains three independent molecules with very similar metrical parameters; only one of these molecules is shown. Selected averaged bond lengths (pm) and angles (°):  $B_{eq}-Br_{eq}$  189.4(5),  $B_{ax}-Br_{ax}$  190.4(5),  $S-B_{eq}$  $196.8 \textbf{(2)}, \; B_{eq} - B_{eq} \; 177.8 (7), \; B_{eq} - B_{ax} \; 170.1 \textbf{(3)}, \; B_{eq} - S - B_{eq} \; 53.7 \textbf{(2)}, \; B_{eq} - B_{eq} -$  $B_{eq}$  90.0(3),  $B_{eq}-B_{ax}-B_{eq}$  63.0(3),  $S-B_{ax}-Br_{ax}$  179.1(2).

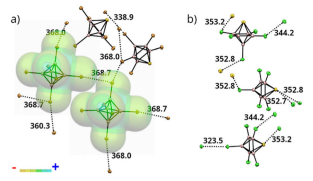


Fig. 4 The short contacts below the sum of vdW radii in the crystal packing of  $\mathbf{1}$  (a) and  $SB_5Cl_5^{3e}$  (b) are indicated by dot lines (in pm). An example of  $\mathrm{Br}^{\delta+}\cdots\mathrm{Br}^{\bar{\delta}-}$  is depicted in term of the corresponding molecular ESP surfaces. The color coding of the ESP is in the range of -26 to 26 kcal mol<sup>-1</sup>.

energy decomposition into individual terms. For comparison, the same analysis has also been performed for SB<sub>5</sub>Cl<sub>5</sub>. Both crystals are dominated by dispersion as expected. Surprisingly, the contribution of dispersion to crystal packing was slightly less pronounced in 1 than in SB<sub>5</sub>Cl<sub>5</sub> (i.e. the dispersion in them formed 63 and 67%, respectively, of the sum of all the attractive terms in the SAPT decomposition). The explanation is that there are considerably more positive  $\sigma$ -holes on Br atoms in 1 than on Cl atoms in SB<sub>5</sub>Cl<sub>5</sub>. Compound 1 thus has a better balance of positive and negative sites for crystal packing - it has six highly positive σ-holes and five negative rings around B-Br bonds, unlike  $SB_5Cl_5$ , which has only one highly positive  $\sigma$ -hole on the S atom to match the five negative rings around B-Cl bonds. Consequently, the electrostatic contribution is more pronounced in the crystal packing of 1, which is dominated by dihalogen bonds, than in SB<sub>5</sub>Cl<sub>5</sub>, dominated by chalcogen bonds (see Fig. 4 and Table 3). This enhanced electrostatic contribution may be ascribed to the fact that the crystallisation process of 1 is more straightforward than that of SB5Cl5.

In a thermodynamically controlled reaction at 350 °C, simple inorganic synthons such as B2Br4 and S2Br2 have provided octahedral, bicapped square-antiprismatic and icosahedral structural motifs referred to as closo-SB5Br5, closo-1-SB9Br9 and closo-SB<sub>11</sub>Br<sub>11</sub>, respectively. Structural features have been elucidated through a combination of experimental and computational approaches including NMR spectroscopy together with MS spectrometry and DFT/ZORA/NMR model chemistry. These methods have confirmed that these three structural arrangements are

Table 3 The interaction energies ( $\Delta E^2$ ) computed at the SAPTO/jun-ccpVDZ level have been decomposed into electrostatic ( $E_{\rm elec}$ ), exchange  $(E_{\text{exch}})$ , induction  $(E_{\text{ind}})$  and dispersion  $(E_{\text{disp}})$  contributions using the SAPT methodology. All the energies are in kcal mol<sup>-1</sup>. The relative values in round brackets indicate the contribution to the sum of all the attractive

Compound	$\Sigma SAPT0$	$\Sigma E_{ m elec}$	$\Sigma E_{\mathrm{exch}}$	$\Sigma E_{\mathrm{ind}}$	$\Sigma E_{ m disp}$
SB <sub>5</sub> Br <sub>5</sub> (1) SB <sub>5</sub> Cl <sub>5</sub>	$-41.5 \\ -32.2$	-38.8 (30%) -24.0 (26%)		( )	-80.4 (63%) -61.5 (67%)

good representations of the molecular geometries in solutions. In addition, crystal-structure determination of the octahedral brominated thiaborane shows that the Br...Br contacts are strong enough to be able to overcome the repulsion between the  $\sigma$ -holes located on the bromines and the sulphur. As a consequence, the crystal-packing forces are dominated by strong  $Br^{\delta^+}(\sigma-hole)\cdots$  $Br^{\delta}$  (ring) attraction, *i.e.* the crystal is formed without chalcogen bonding detected in the chlorine analogue of 1.

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### Data availability

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The data supporting of this article have been included as part of the ESI.† The crystallographic data for compound 1 (closo-SB<sub>5</sub>Br<sub>5</sub>) were deposited with the Cambridge Crystallographic Data Centre with CCDC no. 2403282.†

### Conflicts of interest

There are no conflicts to declare.

### Notes and references

1 For selected surveys on polyhedral boranes, see: (a) W. N. Lipscomb, Boron Hydrides, Benjamin, New York, 1963; (b) E. L. Muetterties, Boron Hydride Chemistry, Academic Press, New York, 1975; (c) J. F. Liebman, A. Greenberg and R. E. Williams, Advances in Boron

- and the Boranes, VCH Verlagsgesellschaft, Weinheim New York, 1988; (d) R. B. King, Chem. Rev., 2001, 101, 1119-1152.
- 2 For selected surveys on carboranes, see: (a) Electron Deficient Boron and Carbon Clusters, ed. G. A. Olah, K. Wade and R. E. Williams, John Wiley and Sons, New York, 1991; (b) T. Onak, Polyhedral Carbaboranes, Comprehensive Organometallic Chemistry II, ed. E. W. Abel, F. G. A. Stone and G. Wilkinson, Pergamon, Oxford, UK, 1995, Vol. 1, 217-255; (c) M. A. Fox, Polyhedral Carboranes, Comprehensive Organometallic Chemistry III, ed. R. H. Crabtree and D. M. P. Mingos, Elsevier, 2007, Vol. 3, 49-112; (d) R. N. Grimes, Carboranes, ed., Elsevier, Amsterdam, The Netherlands, 2016.
- 3 (a) W. Haubold, W. Keller and G. Sawitzki, Angew. Chem., Int. Ed. Engl. 1988, 27, 925-926; (b) W. Keller, L. G. Sneddon, W. Einholz and A. Gemmler, Chem. Ber., 1992, 125, 2343-2346; (c) R. Schäfer, W. Einholz, W. Keller, G. Eulenberger and W. Haubold, Chem. Ber., 1995, 128, 735-736; (d) W. Keller, W. Einholz, D. Rudolph and T. Schleid, Z. Anorg. Allg. Chem., 2017, 643, 664-668; (e) W. Keller, F. Lissner, J. Ballmann, J. Fanfrlík, D. Hnyk and T. Schleid, Angew. Chem., Int. Ed., 2024, 63, e202406751; (f) W. Keller and M. Hofmann, Z. Anorg. Allg. Chem., 2017, 643, 729-731; (g) W. Keller, M. Hofmann, H. Wadepohl, M. Enders, J. Fanfrlík and D. Hnyk, Dalton Trans., 2023, 52, 16886-16893; (h) W. Keller, J. Ballmann, M. B. Sárosi, J. Fanfrlík and D. Hnyk, Angew. Chem., Int. Ed., 2023, 62, e202219018.
- 4 See e.g. (a) W. Keller, M. Hofmann, M. B. Sárosi, J. Fanfrlík and D. Hnyk, Inorg. Chem., 2022, 61, 16565-16572; (b) W. Keller, M. B. Sárosi, J. Fanfrlík, M. Straka, J. Holub, M. L. McKee and D. Hnyk, RSC Adv., 2023, 13, 19627-19637.
- 5 (a) S. Heřmánek, Chem. Rev., 1992, 92, 325-362; (b) F. Teixidor, C. Vinas and R. W. Rudolph, Inorg. Chem., 1986, 25, 3339-3346; (c) T. P. Fehlner, P. T. Czech and R. F. Fenske, Inorg. Chem., 1990, 29, 3103-3109; (d) S. Heřmánek, D. Hnyk and Z. Havlas, J. Chem. Soc., Chem. Commun., 1989, 23, 1859-1861; (e) M. Bühl, P. V. R. Schleyer, Z. Havlas, D. Hnyk and S. Heřmánek, Inorg. Chem., 1991, 30, 3107-3111.
- 6 (a) R. E. Williams, Inorg. Chem., 1971, 10, 210-214; (b) K. Wade, Adv. Inorg. Chem. Radiochem., 1976, 18, 1-66; (c) R. E. Williams, Adv. Inorg. Chem. Radiochem., 1976, 18, 67; (d) R. W. Rudolph, Acc. Chem. Res., 1976, 9, 446-452; (e) R. E. Williams in Electron Deficient Boron and Carbon Clusters, ed. G. A. Olah, K. Wade and R. E. Williams, Wiley and Sons, New York, 1991, 11-93; (f) R. E. Williams, Chem. Rev., 1992, 92, 177-207.
- 7 T. M. Parker, L. A. Burns, R. M. Parrish, A. G. Ryno and C. D. Sherrill, J. Chem. Phys., 2014, 140, 094106.