Dalton Transactions



PAPER

View Article Online
View Journal | View Issue



Cite this: *Dalton Trans.*, 2015, **44**, 11819

Received 17th April 2015, Accepted 2nd June 2015 DOI: 10.1039/c5dt01460c

www.rsc.org/dalton

A computational analysis of the apparent *nido vs. hypho* conflict: are we dealing with six- or eight-vertex open-face diheteroboranes?†:

João Pedro F. Nunes, a Josef Holub, David W. H. Rankin, Derek A. Wann*a and Drahomír Hnyk*b

A series of computational studies have been undertaken to investigate the electronic structures and bonding schemes for six hetero-substituted borane cages, all of which have been presented in the literature as potential hypho structures. The six species are $hypho-7,8-[C_2B_6H_{13}]^-$ (1a), $hypho-7,8-[CSB_6H_{11}]^-$ (1b), $hypho-7,8-[S_2B_6H_9]^-$ (1c), $hypho-7,8-[NSB_6H_{11}]$ (1d), $exo-7-Me-hypho-7,8-[NCB_6H_{12}]$ (1e), and $endo-7-Me-hypho-7,8-[NCB_6H_{12}]$ (1f) and the so-called $ext{mno}$ rule has been applied to each of them. As no structural data are known for the carbathia-, azathia-, and dithiahexaboranes, we have also applied the $ext{ab initio}/GIAO/NMR$ structural tool for 1b-1d, with 1c having been prepared for this purpose. We conclude that an $ext{mno}$ count of 10 means that 1a, 1b, 1d, 1e, and 1f should be termed $ext{pseudo-nido}$ or $ext{pseudo-hypho}$. Only 1c can be considered to be correctly termed $ext{hypho-7,8-[S_2B_6H_9]}^-$.

Introduction

Polyhedral boranes and heteroboranes have long provided interest covering a range of chemical research, where their unexpected structures, innovative bonding schemes, and unusual chemical properties have resulted in them being considered promising candidates for use in medicinal and materials applications. For decades now an understanding of the molecular architectures of these species has led directly and indirectly to practical uses for boranes and future structural studies will undoubtedly lead to potential further uses being identified.

The variety of bonding schemes exhibited by polyhedral boranes and heteroboranes was initially noted by Lipscomb, ²

with Williams³ and Wade⁴ also making significant contributions. Williams reported a major breakthrough by recognising that the experimentally isolated *nido*, *arachno*, and *hypho* boranes could be derived from the nearest *closo* structures by the removal of one, two, and three $\{BH\}^{2-}$ vertices, respectively. This led to the derivation of Wade's n+1 electron-pair rule for a *closo*-type cluster, where n is the number of vertices.^{4,5} Subsequently, the number of skeletal electron pairs required for stable *nido*, *arachno*, and *hypho* skeletons was determined to be n+1+p, where p is the number of missing vertices; it follows that *closo*- $[B_nH_n]^{2-}$ requires such a formal charge to comply with Wade's rule. It should be noted that *closo* systems are known experimentally for n=5-12, where the icosahedral cluster with I_h point-group symmetry (n=12) is the most stable of the *closo* series.

The unusually high stability of closo- $[B_{12}H_{12}]^{2-}$, as well as the existence of condensed B_{12} units [for example, in the so-called macropolyhedral boron clusters $B_{20}H_{16}$ and $B_{21}H_{18}^{-}$ (ref. 7)], meant that a generalisation of the electron-counting rules was required since such structures are beyond the scope of the Williams–Wade formalism. A generally applicable electron-counting rule – the so-called mno rule – was derived by Jemmis; based on Hückel's rule it allows the structures of macropolyhedral boranes and metallaboranes to be characterised alongside simple boranes. According to the mno rule, m+n+o electron pairs are necessary for a macropolyhedral system to be stable, where m is the number of individual polyhedral subclusters from which a macropolyhedral cluster is composed, n is the number of vertices, and n is the number of

^aDepartment of Chemistry, University of York, Heslington, York YO10 5DD, UK. E-mail: derek.wann@york.ac.uk

^bInstitute of Inorganic Chemistry of the Academy of Sciences of the Czech Republic, v.v.i., 250 68 Husinec-Řež, Czech Republic. E-mail: hnyk@iic.cas.cz

^cSchool of Chemistry, University of Edinburgh, Joseph Black Building, David Brewster Road, Edinburgh, EH9 3FJ, UK

[†] Dedicated to the memory of Professor Ken Wade.

[‡]Electronic supplementary information (ESI) available: Cartesian coordinates for the structures of **cB11**, **nB10**, and **aB9** (Tables S1–S3), **1a–1f** (Tables S4–S9), and nido-[B₆H₁₁]⁺ (Table S10), the s character, p character, and resultant hybridisation of **1a–1f** for all levels of theory (Tables S11–S13), and charges on the heteroatoms of **1a–1f** (Table S14). The molecular structures of **1a–1f** with full atom numbering (Fig. S1), the equivalent for the closo-nido-arachno-ara

single-vertex-sharing condensations. For *nido*, *arachno*, and *hypho* arrangements, one, two, and three additional pairs of electrons are required. Wade's n+1 rule can be considered as a special case of the *mno* rule, where m=1 and o=0.

Some *hypho* and *nido* complexes have very similar structures and electronic structure investigation would seem sensible to ensure clusters are not wrongly classified. In this work we have performed *ab initio* and DFT analyses of the bonding schemes for six potential *hypho* structures $\{hypho-7,8-[C_2B_6H_{13}]^- (1a), hypho-7,8-[CSB_6H_{11}]^- (1b), hypho-7,8-[S_2B_6H_9]^- (1c), hypho-7,8-[NSB_6H_{11}] (1d), exo-7-Me-hypho-7,8-[NCB_6H_{12}] (1e), and endo-7-Me-hypho-7,8-[NCB_6H_{12}] (1f)}, also employing the$ *mno*rule to

see what results it yields. All six structures are shown in Fig. 1, while Fig. S1‡ also gives the hydrogen-atom numbering. As no structural data are currently available for the carbathia-, azathia-, and dithiahexaboranes, we have also applied the *ab initio*/GIAO/NMR structural method to **1b–1d**.

Experimental section

NMR

Experimental ¹¹B NMR chemical shifts for **1b** and **1d** were taken from ref. 9 and 10, respectively; **1c** was prepared according to ref. 11.

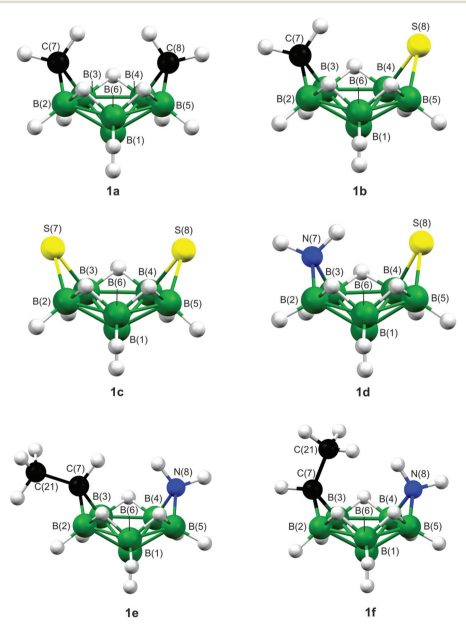


Fig. 1 The molecular structures of eight-vertex diheteroboranes that have been proposed as *hypho*-type clusters. For clarity hydrogen-atom numbering has been omitted. The molecules are $hypho-7.8-[C_2B_6H_{13}]^-$ (1a), $hypho-7.8-[CSB_6H_{11}]^-$ (1b), $hypho-7.8-[S_2B_6H_9]^-$ (1c), $hypho-7.8-[NSB_6H_{11}]$ (1d), $exo-7-Me-hypho-7.8-[NCB_6H_{12}]$ (1e), and $endo-7-Me-hypho-7.8-[NCB_6H_{12}]$ (1f).

Dalton Transactions Paper

Quantum chemical calculations

The geometries of $closo-[B_{11}H_{11}]^{2-}$ (cB11), $nido-[B_{10}H_{10}]^{4-}$ (nB10) and arachno-[B₉H₉]⁶⁻(aB9) were fully optimised using Gaussian09.12 Each of these structures, as well as that of the related hypho species (hB8), is depicted in Scheme 1. (The scheme is reproduced in Fig. S2,‡ where the atom numbering is given.) The geometry optimisations for each of cB11, nB10, and aB9 were performed at the B2PLYP, 13 MP2, 14-18 B98, 19 B97d,²⁰ PBE,^{21,22} PW91,²³⁻²⁷ and HFS²⁸⁻³⁰ levels of theory using the 6-311+ $G(d,p)^{31,32}$ basis set on all atoms; the nature of any stationary points on the potential-energy surfaces were investigated using frequency calculations. The initial geometries for nB10 and aB9 were optimised from geometries obtained by removing one and two vertices from cB11, respectively, using GaussView 5.0.33 The bonding orbitals for cB11, nB10, and aB9 were investigated using natural bond orbital (NBO) analyses to look for three-centre bonds and resonance structures.

The geometries of the heteroboranes 1a-1f were also fully optimised using Gaussian09 and the method and basis set combinations described above, with the character of each stationary point verified by frequency calculations. Magnetic shieldings were calculated for 1b-1d by running GIAO³⁴ jobs with TZP basis set II by Huzinaga,35 which is well suited for this purpose. In order to investigate the relationships between 1a-1f and their true borane analogues, the heteroatoms for each borane cage were replaced by hydrogen atoms using GaussView 5.0. The geometries of the species formed by the inclusion of hydrogen atoms in place of heteroatoms were optimised using the same levels of theory and basis sets previously employed, and frequency calculations were performed to verify the nature of any stationary points; the distances between the substituted hydrogen atoms and the boron-cage atoms were fixed at values optimised for the B-C/N/S distances. NBO analyses were also performed for all of the H-substituted structures.

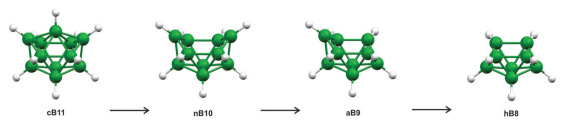
Calculating mno values

The mno rule is used to determine the number of pairs of electrons required for a species to be considered stable by calculating m + n + o (for the meanings of m, n, and o, see above). As an illustration the mno rule is applied to an icosahedral borane by taking the values m = 1, n = 12, and o = 0, yielding m + n + o = 13 electron pairs, the same result as the n + 1 electron-pair rule. Each vertex with one terminal atom contributes all-but-one of its electrons to cluster bonding, with the remaining electron involved in the exo covalent bond. However, in the three isomeric icosahedral carbaboranes closo-C2B10H12, the BH groups each donate one pair of electrons and each CH moiety contributes three electrons (1.5 electron pairs) to the polyhedral bonding, thus satisfying the mno rule [i.e. 10 + $(2 \times 1.5) = 13$ for these neutral carbaboranes]. ³⁶ Similarly, in the icosahedral azaborane *closo*-NB₁₁H₁₂ ³⁷ there are eleven BH groups that contribute eleven electron pairs as well as an NH group that donates two electron pairs to the skeletal bonding. Considering molecules containing third-row elements, the neutral icosahedral closo-SB₁₁H₁₁ thiaborane is known,³⁸ in which BH groups provide eleven electron pairs; sulfur has a tendency to retain one of its lone pairs, contributing four electrons (two electron pairs) to the cluster.

The following closo dianions, presented in order of stability, $B_{12}H_{12}^{2-}$, $B_{11}H_{11}^{2-}$, and $B_5H_5^{2-}$, can be used to derive *nido*, arachno, and hypho heteroboranes via the removal and substitution of vertices.³⁹ However, according to the Williams-Wade concept, heteroboranes cannot exist with two, three, or four BH moieties. Closo- $[B_5H_5]^{2-}$ is, therefore, not a suitable "parent" species for any of the nido, arachno, or hypho heteroborane structures.

There are heteroboranes with molecular shapes based on $nido-[B_{10}H_{10}]^{4-}$ (**nB10**)⁴⁰ and $arachno-[B_{9}H_{9}]^{6-}$ (**aB9**),⁴¹ which are derived from the C_{2v} -symmetric closo- $[B_{11}H_{11}]^{2-}$ (cB11), as shown in Scheme 1. As well as closo, nido, and arachno clusters, Scheme 1 depicts the hypothetical hypho- $[B_8H_8]^{8-}$ (hB8). In contrast, our so-called *hypho* eight-vertex-type heteroboranes resemble the six-vertex arrangement adopted by nido-[B₆H₁₁] (nB6), 42 shown in Fig. 2. (Fig. S3‡ depicts the same structure but also includes hydrogen-atom numbering.)

The molecule $hypho-7,8-[C_2B_6H_{13}]^-$ (1a) has previously been structurally characterised by applying the ab initio/GIAO/NMR structural tool, 43 which confirmed its C_s -symmetric "helmetlike" structure (Fig. 1). Three other proposed hypho species (also shown in Fig. 1) have been calculated to have the same structural motif as 1a; these are hypho-7,8- $[CSB_6H_{11}]^-$ (1b), $hypho-7,8-[S_2B_6H_9]^-$ (1c), 11 and $hypho-7,8-[NSB_6H_{11}]$ (1d). 10 The last of these examples (1d) illustrates how nitrogen can be accommodated in an eight-vertex hypho arrangement, with the same arrangement found in exo- and endo-7-Me-hypho-7,8-



Scheme 1 Schematic representation of the closo-nido-arachno-hypho relationship for n = 11.

Paper

B(3) B(4) B(5) B(1)

Fig. 2 The molecular structure of nido- $[B_6H_{11}]^+$, whose structure resembles a hypho eight-vertex heteroborane. For clarity hydrogenatom numbering has been omitted.

[NCB₆H₁₂] (**1e** and **1f**, respectively). Again, GIAO calculations of the shielding tensors have been used for structural characterisation.⁴⁴

Results and discussion

Calculations were initially performed to investigate the bonding patterns of the parent cB11, nB10, aB9, and hB8 systems, as shown in Scheme 1. Apart from the standard HF calculations perturbed with the MP2 model chemistry, we employed six density functional theory (DFT) approaches to span all the possibilities afforded by DFT, i.e. B98 (stand-alone hybrid functional), B97D (stand-alone pure functional), HFS (exchange-only functional), PW91 (correlation functional), PBE (exchange-combined functional), and B2PLYP (double-hybrid functional), the latter utilising HF exchange and an MP2-like correlation. All seven model chemistries used the 6-31G(d) basis set for the initial geometry optimisations. However, all attempts to identify a minimum on the potential-energy hypersurface of hB8 failed because of the formation of either a series of flattened triangular borane architectures or a random assembly of eight boron atoms. The final basis set used for optimising cB11, nB10, and aB9 was 6-311G(d,p). Cartesian coordinates for these structures are given in Tables S1-S3.‡ Attempts to use extra diffuse functions on these basis sets caused the optimisations to diverge. Second-derivative analyses of these systems showed that they represented minima on the respective potential-energy hypersurfaces. The seven model chemistries were also used to calculate Hartree-Fock and Kohn-Sham natural orbitals and no two-centre two-electron (2c-2e) bonds were predicted for the cage motifs, only for the terminal B-H bonds.

In order to get a deeper insight into the bonding patterns of 1a-1f, we performed entirely the same computational procedures as for cB11, nB10, aB9, and hB8. All of these diheterohexaboranes structures were identified as potential minima at *ab initio* and all DFT levels. Cartesian coordinates relating to the MP2/6-311+G(d,p) calculations are given in Tables S4-S9.‡ The strong resemblance of each of the 1a-1f molecular geometries to that of nido- $[B_6H_{11}]^+$ prompted us to replace each

heteroatom with a hydrogen atom (H_r) , with the B-H_r distances fixed at the values optimised for the B-C/N/S bond lengths. Any substituents originally bonded to C or N were omitted. The new structures generated were termed 1a/H-1e/H (no calculations were required for 1f/H which is identical to 1e/H). The structure of $nido-[B_6H_{11}]^+$ (Fig. 2) was calculated as a comparison as it has the same arrangement of H-bridges; Cartesian coordinates relating to its geometry at the MP2/6-311+G(d,p) level are given in Table S10.‡ As might be expected, frequency calculations performed for each of 1a/H-1e/H showed imaginary frequencies relating to the H_r atoms.

Performing NBO analyses for each of 1a/H-1e/H revealed that three-centre two-electron (3c-2e) bonding is present throughout each of the clusters, i.e. also in B-H_r-B bridges. When these structures were allowed to relax further by optimising the B-H_r distances, 3c-2e bonding persisted for all five species 1a/H-1e/H. The geometry optimisations performed using the 6-311+G** basis sets demonstrate that 3c-2e bonding exists regardless of the lengths of the B-H bridging distances; this was true for all seven methods employed. Similar analyses for 1a-1f (where the heteroatoms are present) showed a different picture, where all B-C/N/S distances were classified as 2c-2e bonds. If the NBO analyses had revealed 2c-2e bonds for B-H_r-B bridges in 1a/H-1e/H, we could have stated that the heteroatoms present in 1a-1f were not part of multicentre bonding. However, as this was not the case, we therefore had to follow another way of finding the nature of bonding in 1a-1f.

Table 1 shows the hybridisation of the heteroatoms in 1a-1f when transforming canonical orbitals to natural ones using NBO analysis. (Such calculations were performed for all levels of theory specified in the Experimental section and then averaged. The range of values is also shown. The characters and hybridisations of the B-C/N/S bonds for each of 1a-1f at all levels of theory are given in Tables S11-S13.‡) Table 1 shows that the heteroatoms in each species are more or less sp³hybridised, as is the case for textbook examples such as CH₄, NH₃, and H₂S. This means that, as well as 2c-2e bonds to boron atoms, all of the heteroatoms are covalently bonded to hydrogen atoms and any carbon atoms that are external to the cage motif. In addition, NBO analyses showed that heteroatoms in 1a-1f are negatively charged. The average calculated charges are presented in Table 2 (with the corresponding values for each level of theory in Table S14th) and are in line with those values for classical covalent species such as CH₄, NH₃, and H₂S. However, such an observation is in contrast to the electron distribution experimentally determined for the icosahedral species, in which the midpoint of CC vector and sulfur atom were found to be positively charged as revealed by vector algebra of experimental dipole moments measured for the exo-substituted icosahedra. 36d,38b

Armed with these results, we applied the *mno* rule to *nido*- $[B_6H_{11}]^+$ as well as to **1a–1f**. For *nido*- $[B_6H_{11}]^+$ m=1, n=6, o=0, and p=1, meaning that eight electron pairs are required to stabilise this system. There are six B–H fragments, each of which contributes one electron pair. The remaining two elec-

Table 1 Hybridisation of the heteroatoms (C/N/S) bonded to the boron atoms. The bond characters were calculated using NBO analyses and are averaged across all levels of theory specified in the experimental section^a

	Bond	s character/%	p character/%	Hybridisation
1a	B(2)-C(7)	28.45	71.53	2.51 ± 0.01
	B(3)-C(7)	26.84	73.14	2.73 ± 0.06
	B(4)-C(8)	26.84	73.14	2.73 ± 0.06
	B(5)-C(8)	28.45	71.53	2.51 ± 0.01
1b	B(2)-C(7)	28.62	71.34	2.49 ± 0.01
	B(3)-C(7)	26.08	73.89	2.83 ± 0.07
	B(4)-S(8)	16.96	82.87	4.91 ± 0.37
	B(5)-S(8)	18.49	81.22	4.40 ± 0.24
1c	B(2)-S(7)	18.50	81.31	4.40 ± 0.23
	B(3)-S(7)	16.72	83.10	4.99 ± 0.36
	B(4)-S(8)	16.72	83.10	4.99 ± 0.36
	B(5)-S(8)	18.50	81.31	4.40 ± 0.23
1d	B(2)-N(7)	29.38	70.61	2.40 ± 0.02
	B(3)-N(7)	27.46	72.53	2.64 ± 0.06
	B(4)-S(8)	16.61	83.20	5.02 ± 0.32
	B(5)-S(8)	17.08	82.71	4.86 ± 0.32
1e	B(2)-C(8)	26.28	73.70	2.80 ± 0.03
	B(3)-C(8)	25.80	74.19	2.88 ± 0.05
	B(4)-N(7)	28.69	71.31	2.49 ± 0.05
	B(5)-N(7)	29.48	70.52	2.39 ± 0.02
1f	B(2)-C(8)	26.12	73.85	2.83 ± 0.03
	B(3)-C(8)	26.00	73.98	2.85 ± 0.05
	B(4)-N(7)	28.49	71.50	2.51 ± 0.05
	B(5)-N(7)	29.41	70.59	$\textbf{2.40} \pm \textbf{0.02}$

^a The characters and hybridisations from each of the individual calculations are given in Tables S11–S13. ^bThe errors quoted are the standard deviation of the values calculated for the different model chemistries.

Table 2 Atomic charges of the heteroatoms (C/N/S) within the borane cage motif. The charges are averaged across all levels of theory specified in the experimental section^a

Atom	Charge ^b	
C(7)	-0.92 ± 0.02	
C(8)	-0.92 ± 0.02	
C(7)	-0.91 ± 0.02	
S(8)	-0.27 ± 0.03	
	-0.22 ± 0.03	
	-0.22 ± 0.03	
	-0.95 ± 0.02	
	-0.18 ± 0.04	
	-0.93 ± 0.02	
	-0.73 ± 0.02	
()	-0.94 ± 0.02	
\$ 2	-0.74 ± 0.02	
	C(7) C(8)	

^a The charges from each of the individual calculations are given in Table S14. ^b The errors quoted are the standard deviation of the values calculated for the different model chemistries.

tron pairs are available from the five bridging hydrogen atoms with one spare electron present; the total number of electron pairs is 6 + 2.5 = 8.5. As there is one excess electron the hypothetical nido- $[B_6H_{11}]^+$ is obviously a cation. (For nido- B_6H_{10} we have eight electron pairs and, consequently, the molecule is neutral.)

Table 3 Computed (GIAO) and experimental NMR chemical shifts for 1b-1d

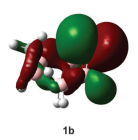
	Vertex					
	B(1)	B(3)	B(6)	B(4)	B(2)	B(5)
1b GIAO ^a Exp. ^{b,c} 1c GIAO ^a Exp. ^{b,d} 1d GIAO ^a Exp. ^{b,e}	-56.5 -54.8 -55.8 -52.2 -56.3 -55.1	-36.3 -33.2 5.3 6.8 -26.4 -25.6	-33.5 -29.4 -26.1 -22.4 -24.9 -22.4	-25.1 -23.9 -31.4 -25.0 -21.7 -20.8	-4.6 -3.1 -26.1 -22.4 -2.0 -1.6	3.1 2.0 5.3 6.8 2.2 1.8

 $[^]a$ GIAO-MP2/II//6-311+G**. b Measured in CDCl $_3$. c Ref. 9. d This work. e Ref. 10.

To apply the *mno* rule to **1a–1f** we need to determine whether they adopt a *hypho* or a *nido* electron count. Let's start by supposing that they are considered to be *hypho*, giving m = 1, n = 8, o = 0, p = 3, meaning that 12 electron pairs are required. In order to comply with a *hypho* electron count the heteroatoms must contribute the following number of electron pairs: sulfur 2, carbon 2, and nitrogen 2.5 for the charges to be "*hypho*" correct.

Therefore, for 1a we have 7.5 + 4 = 11.5 electron pairs, thus requiring one extra electron to stabilise the system. This C_s skeleton is therefore characterised with a single negative charge. The same applies to the other non-nitrogen-containing compounds (1b and 1c), where sulfur and/or carbon contributes 4 electron pairs, resulting in an overall single negative charge for these species. However, such electron pair contributions from the heteroatoms are in conflict with the NBO results shown in Table 1, where C, N, and S are more or less sp³-hybridised and are connected to the boron atoms by conventional 2c–2e bonds. Moreover, the carbon and nitrogen atoms are not naked, but rather are bonded also to two *exo* atoms or groups, *i.e.* the corresponding one electron pair from each of these heteroatoms cannot contribute to the skeletal moieties.

Secondly, we may formally consider 1a-1f to be nido clusters, where values of n = 6, m = 1, o = 0, and p = 1 yields an mno value of eight and, consequently, eight electron pairs are required to stabilise these molecules, as was the case for nido- $[B_6H_{11}]^+$. In the latter, six B-H bonds contribute six electron pairs, and five hydrogen bridges contribute 2.5 electron pairs, i.e. this system must have a single positive charge (6 + 2.5 = 8.5)and so one electron must be removed to get 8). There is also another hypothetical system, nido-[B₆H₉]-, which has three H-bridges and complies with the mno rule (mno = 8; 6 + 1.5 = 7.5, so the formal charge is -1). On that basis, there is no scope for the two additional bridging bonds that would be required for this to be a nido system with a single negative charge. Such a charge is, however, unambiguously observed for 1a, 1b, and 1c, in which there are five bridges (three B-H-B bridges and two B-C/S-B bridges). The same argument can be applied to relate the neutral molecules 1d, 1e, and 1f Paper Dalton Transactions





10

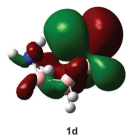


Fig. 3 HOMO for 1b-1d at HF/6-311+G(d,p).

to nido-B₆H₁₀ (6 + 4/2 = 8; there is no scope for further bridge-type bonding to comply simultaneously with the nido electron count requirement and to keep the system neutral).

Only **1c** can truly be considered to be correctly classified as hypho-7,8- $[S_2B_6H_9]^-$, since **1c** can afford to accept four electron pairs from the sulfur atoms into cluster bonding leaving one lone pair of electrons on each sulfur atom.

We must therefore conclude that **1a**, **1b**, **1d**, **1e**, and **1f** are, in reality, neither formally *hypho* skeletons nor *nido* structures. To be formally *hypho*, an *mno* value of 12 should be satisfied for **1a**, **1b**, **1d**, **1e**, and **1f**. Moreover, we would actually need to have an *mno* value of 10 to "accommodate" two extra B-C/N/S-B bridges and to comply with *nido* requirement. Since each of **1a**, **1b**, **1d**, **1e** and **1f** is analysed in terms of having a *mno* value of 12 or 8 (and not 10), we might call these systems *pseudo-nido* or *pseudo-hypho*. As stated above, **1c** can be correctly classified as *hypho-*7,8-[S₂B₆H₉]⁻.

Small clusters are known to be very sensitive to the inclusion of electron dynamic correlation when shielding tensors are being calculated. The GIAO-MP2 calculations predict ¹¹B chemical shifts that compare well with experimental values. Table 3 compares computed and experimental ¹¹B chemical shifts for **1b–1d**.

While the computed and experimental ^{11}B NMR chemical shifts generally compare well, the presence of sulfur in **1b–1d** makes these fits for some atoms slightly worse than for clusters reported in the literature that do not contain a third-row element. Such discrepancies can be attributed to the inadequacy of using a triple-zeta Huzinaga type-II basis set on sulfur. 46 The most striking features of the individual spectra are shifts to low frequencies for the "bottom" boron atom, B(1). When comparing these values we clearly see the difference between the nature of the ^{11}B chemical shifts for B(2) and B(3) in **1b** and **1d** and those of **1c**. Inspecting the highest occupied molecular orbitals (HOMOs; see Fig. 3) for **1b–1d** offers an explanation in terms of the entirely different shieldings of B(2) and B(3) when comparing **1b** and **1d** with C_8 -symmetric **1c**.

The decent agreement between theory and experiment suggests that MP2/6-311+G(d,p) geometries serve as valid representations of the molecular geometries in solution, which are characterised by very long B-B distances (values are between 1.94 and 1.98 Å) bridged by heteroatoms.

Conclusions

According to these computational efforts, we have concluded that hypho-7,8- $[S_2B_6H_9]^-$ (1c) is correctly classified as a formal hypho structure. However, the other compounds studied here should be considered as pseudo-hypho or pseudo-nido, i.e. there is a nido vs. hypho conflict, and also a disagreement with the literature. Conceivably, according to the mno rule all valence electrons of carbon and nitrogen are accounted for in skeletal bonding when assuming hypho electron count. On the contrary, we would need to have 10 electron pairs instead of eight to comply with the formal nido electron count while also having sufficient electrons to allow for the bonding of carbon and nitrogen cage atoms to exo carbon and hydrogens atoms. The fact that hypho- $[B_8H_8]^{8-}$ has been shown not to exist complements such a conclusion.

Acknowledgements

D.A.W. thanks the EPSRC for funding a Career Acceleration Fellowship (EP/I004122), and for partially funding, along with the Department of Chemistry, University of York, a student-ship for J.P.F.N. We acknowledge the use of the EPSRC National Service for Computational Chemistry Software hosted at Imperial College in carrying out this work, which also made use of the York Advanced Research Computing Cluster (YARCC) provided by the University of York. We acknowledge the Czech Science Foundation (project number P207/11/07505) for financial support. All data supporting this study are provided either in the results section of this paper or as supplementary information accompanying this paper.

References

- 1 See, for example, (a) K. Wade, *Polymer*, 1997, 38, 4539; (b) J. M. Oliva, D. J. Klein, P. v. R. Schleyer and L. Serano-Andres, *Pure Appl. Chem.*, 2009, 81, 719; and many other references emerging these days.
- 2 W. N. Lipscomb, Adv. Inorg. Chem. Radiochem., 1950, 1, 117.

- 3 R. E. Williams, *Adv. Inorg. Chem. Radiochem.*, 1976, 18, 67.
- 4 (a) K. Wade, J. Chem. Soc. D, 1971, 792; (b) K. Wade, Adv. Inorg. Chem. Radiochem., 1976, 18, 1.
- 5 K. Wade, Inorg. Nucl. Chem. Lett., 1972, 8, 563.

Dalton Transactions

- 6 D. Hnyk, J. Holub, T. Jelínek, J. Macháček and M. G. S. Londesborough, *Collect. Czech. Chem. Commun.*, 2010, 75, 1115, and references therein.
- 7 E. Bernhardt, D. J. Brauer, M. Finze and H. Willner, *Angew. Chem., Int. Ed.*, 2007, **46**, 2927.
- 8 See, for example, (a) M. M. Balakrishnarajan and E. D. Jemmis, J. Am. Chem. Soc., 2000, 122, 4516; (b) E. D. Jemmis, M. M. Balakrishnarajan and P. D. Pancharatna, J. Am. Chem. Soc., 2001, 123, 4313.
- 9 J. Holub, J. D. Kennedy, T. Jelínek and B. Štíbr, *J. Chem. Soc.*, *Dalton Trans.*, 1994, 1317.
- 10 T. Jelínek, J. D. Kennedy and B. Štíbr, *J. Chem. Soc., Chem. Commun.*, 1993, 1628.
- 11 S. O. Kang and L. G. Sneddon, *J. Am. Chem. Soc.*, 1989, **111**, 3281.
- 12 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr., J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, Gaussian 09, Revision D.01, Gaussian, Inc., Wallingford, CT, 2013.
- 13 S. Grimme, J. Chem. Phys., 2006, 124, 034108.
- 14 M. Head-Gordon, J. A. Pople and M. J. Frisch, *Chem. Phys. Lett.*, 1988, 153, 503.
- 15 S. Saebø and J. Almlöf, Chem. Phys. Lett., 1989, 154, 83.
- 16 M. J. Frisch, M. Head-Gordon and J. A. Pople, *Chem. Phys. Lett.*, 1990, **166**, 275.
- 17 M. J. Frisch, M. Head-Gordon and J. A. Pople, *Chem. Phys. Lett.*, 1990, **166**, 281.
- 18 M. Head-Gordon and T. Head-Gordon, *Chem. Phys. Lett.*, 1994, **220**, 122.
- 19 H. L. Schmider and A. D. Becke, *J. Chem. Phys.*, 1998, **108**,
- 20 S. Grimme, J. Comput. Chem., 2006, 27, 1787.
- 21 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865.
- 22 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, **78**, 1396.

- 23 J. P. Perdew, in *Electronic Structure of Solids '91*, ed. P. Ziesche and H. Eschrig, Akademie Verlag, Berlin, 1991, p. 11.
- 24 J. P. Perdew, J. A. Chevary, S. H. Vosko, K. A. Jackson, M. R. Pederson, D. J. Singh and C. Fiolhais, *Phys. Rev. B: Condens. Matter*, 1992, 46, 6671.
- 25 J. P. Perdew, J. A. Chevary, S. H. Vosko, K. A. Jackson, M. R. Pederson, D. J. Singh and C. Fiolhais, *Phys. Rev. B: Condens. Matter*, 1993, 48, 4978.
- 26 J. P. Perdew, K. Burke and Y. Wang, *Phys. Rev. B: Condens. Matter*, 1996, 54, 16533.
- 27 K. Burke, J. P. Perdew and Y. Wang, in *Electronic Density Functional Theory: Recent Progress and New Directions*, ed. J. F. Dobson, G. Vignale and M. P. Das, Plenum, 1998.
- 28 P. Hohenberg and W. Kohn, Phys. Rev. B, 1964, 136, 864.
- 29 W. Kohn and L. J. Sham, Phys. Rev. A, 1965, 140, 1133.
- 30 J. C. Slater, *The Self-Consistent Field for Molecular and Solids, Quantum Theory of Molecular and Solids*, McGraw-Hill, New York, 1974, vol. 4.
- 31 R. Krishnan, J. S. Binkley, R. Seeger and J. A. Pople, J. Chem. Phys., 1980, 72, 650.
- 32 T. Clark, J. Chandrasekhar and P. v. R. Schleyer, *J. Comput. Chem.*, 1983, 4, 294.
- 33 http://www.gaussian.com/g_tech/gv5ref/gv5ref_toc.htm.
- 34 (a) R. Ditchfield, Mol. Phys., 1974, 27, 789; (b) K. Wolinski,
 J. F. Hinton and P. Pulay, J. Am. Chem. Soc., 1990, 112,
 8251; (c) J. Gauss, J. Chem. Phys., 1993, 99, 3629.
- 35 W. Kutzelnigg, U. Fleischer and M. Schindler, *NMR Principles and Progress*, Springer, Berlin, 1990.
- 36 Structural data accumulated on the three dicarbadodecaboranes 1,2-, 1,7-, and 1,12-C₂B₁₀H₁₂ are exemplified in the following publications: (a) A. R. Turner, H. E. Robertson, K. B. Borisenko, D. W. H. Rankin and M. A. Fox, *Dalton Trans.*, 2005, 1310; (b) S. Samdal, H. Møllendal, D. Hnyk and J. Holub, *J. Phys. Chem. A*, 2011, 115, 3380; (c) A. R. Campanelli, A. Domenicano and D. Hnyk, *J. Phys. Chem. A*, 2015, 119, 205; (d) For the charge distribution in the 1,2-isomer see: D. Hnyk, V. Všetečka, L. Drož and O. Exner, *Collect. Czech. Chem. Commun.*, 2001, 66, 1375.
- 37 D. Hnyk, M. Bühl, P. v. R. Schleyer, H. V. Volden, S. Gundersen, J. Müller and P. Paetzold, *Inorg. Chem.*, 1993, 32, 2442.
- 38 (a) D. Hnyk, E. Vajda, M. Bühl and P. v. R. Schleyer, *Inorg. Chem.*, 1992, 31, 2464; (b) J. Macháček, J. Plešek, J. Holub, D. Hnyk, V. Všetečka, I. Císařová, M. Kaupp and B. Štíbr, *Dalton Trans.*, 2006, 1024.
- 39 P. v. R. Schleyer, K. Najafian and A. M. Mebel, *Inorg. Chem.*, 1998, 37, 6765.
- 40 Decaborane(14), $B_{10}H_{14}$, is one of the building blocks of boron cluster chemistry.
- 41 D. A. Wann, P. D. Lane, H. E. Robertson, J. Holub and D. Hnyk, *Inorg. Chem.*, 2013, **52**, 4502, and references therein.
- 42 In essence, there exists neutral *nido*-B₆H₁₀ structurally characterised in the solid state, see: F. L. Hirshfeld,

Paper

- K. Eriks, R. E. Dickerson, E. L. Lippert Jr. and W. L. Lipscomb, *J. Chem. Phys.*, 1958, 28, 56.
- 43 M. G. S. Londesborough, Z. Janoušek, B. Štíbr, D. Hnyk, J. Plešek and I. Císařová, *Dalton Trans.*, 2007, 1221.
- 44 T. Jelínek, B. Štíbr, J. D. Kennedy, D. Hnyk, M. Bühl and M. Hofmann, *Dalton Trans.*, 2003, 1326.
- 45 D. Hnyk and E. G. Jayasree, *J. Comput. Chem.*, 2013, 34, 656, and references therein.
- 46 In closo-1-SB₁₁H₁₁ (see ref. 38b) and closo-1-SB₉H₉, see D. Hnyk, D. A. Wann, J. Holub, S. Samdal and D. W. H. Rankin, $Dalton\ Trans.$, 2011, 40, 5734, the difference between the computed and experimental $\delta(^{11}B)$ values of B(12) and B(10) atoms, respectively, atoms that are antipodally coupled with S, using almost the same model chemistries and solvent as for 1d is also ca. 5 ppm; when using C_6D_6 for the second system such a difference even increases.