Thermodynamic Hydricity of Small Borane Clusters and Polyhedral closo-Boranes †

Igor E. Golub 1,*, Oleg A. Filippov 1, Vasilisa A. Kulikova 1,2, Natalia V. Belkova 1,*, Lina M. Epstein 1 and Elena S. Shubina 1,*

1 A. N. Nesmeyanov Institute of Organoelement Compounds and Russian Academy of Sciences (INEOS RAS), 28 Vavilova St, 119991 Moscow, Russia; h-bond@ineos.ac.ru (O.A.F.); kullisa99@gmail.com (V.A.K.); nataliabelk@ineos.ac.ru (N.V.B.); epst@ineos.ac.ru (L.M.E.)
2 Faculty of Chemistry, M.V. Lomonosov Moscow State University, 1/3 Leninskiye Gory, 119991 Moscow, Russia
* Correspondence: seraph347@gmail.com (I.E.G.); shu@ineos.ac.ru (E.S.S.)
† Dedicated to Professor Bohumil Štibr (1940-2020), who unfortunately passed away before he could reach the age of 80, in the recognition of his outstanding contributions to boron chemistry.

Abstract: Thermodynamic hydricity (HDA MeCN) determined as Gibbs free energy (ΔG[H−]) of the H− detachment reaction in acetonitrile (MeCN) was assessed for 144 small borane clusters (up to 5 boron atoms), polyhedral closo-boranes dianions [BnHn]2−, and their lithium salts Li2[BnHn] (n = 5–17) by DFT method [M06/6-311++G(d,p)] taking into account non-specific solvent effect (SMD model). Thermodynamic hydricity values of diborane B2H6 (HDA MeCN = 82.1 kcal/mol) and its dianion [B2H6]2− (HDA MeCN = 40.9 kcal/mol for Li2[B2H6]) can be selected as border points for the range of borane clusters’ reactivity. Borane clusters with HDA MeCN below 41 kcal/mol are strong hydride donors capable of reducing CO2 (HDA MeCN = 44 kcal/mol for HCO2−), whereas those with HDA MeCN over 82 kcal/mol, predominately neutral boranes, are weak hydride donors and less prone to hydride transfer than to proton transfer (e.g., B2H6, B4H10, B5H11, etc.). The HDA MeCN values of closo-boranes are found to directly depend on the coordination number of the boron atom from which hydride detachment and stabilization of quasi-borinium cation takes place. In general, the larger the coordination number (CN) of a boron atom, the lower the value of HDA MeCN.

Keywords: polyhedral boranes; borane clusters; borohydrides; Lewis acidity; hydride donating ability; DFT calculations

1. Introduction

Boron-based chemistry is vast, diverse, and fascinating due to the ability of boron to form electron-deficient structures of various shapes (such as cages, clusters, etc.) with delocalized electrons and multicenter bonding. Boron hydrides, i.e., boranes (e.g., BH4−, [B3H9]−, [B8H8]2− n = 6–12), are of great interest because of their use as ligands in inorganic chemistry [1–5], as building blocks in material chemistry [3,6–8], and as materials for the energetic purposes—components of batteries [9,10], rocket fuels [11–13] and systems of hydrogen storage [14–17]. Polyhedral boranes are widely used as sources of boron (10B) in boron neutron capture cancer therapy [18–23], in the creation of luminescent materials [24–26], thermally stable polymers [5], liquid crystals and nonlinear optical materials [27,28], as well as precursors of nanostructured materials [29,30].

Thermodynamic hydricity, i.e., hydride donating ability (HDA), determined as Gibbs free energy (ΔG[H−]) for the reaction of hydride ion, H−, detachment, is a very important characteristic of transition metal hydrides [31] and main group hydrides [32–34] that describes their reactivity and
Molecules 2020, 25, 2920 is used for the rational design of catalytic reactions. In our recent work, we have demonstrated the existence of an inversely proportional dependence of the hydride-transfer ability of Li[L3B–H] on Lewis acidity of L3B [34]. High values of HDA indicate high Lewis acidity of parent borane L3B, whereas low HDA values indicate high hydride transfer ability of Li[L3B–H]. For that reason, the hydride donating ability (HDA) of boranes can be used as a measure of the Lewis acidity of parent neutral borane or boron cations.

However, there are significant problems in the experimental determination of the thermodynamic hydricity for hydrides of main group elements (E–H, where E = Si, C, B, Al) because of their instability in polar solvents (MeCN, H2O), which are typically used for that [32,35]. Besides, the E–H bond of main group hydrides is characterized by low polarizability compared to transition metal hydrides and, therefore, in many cases, the detachment of the hydride ion (H−) is challenging even in the presence of a large excess of strong Lewis acid. Due to these problems, there is a huge gap in our knowledge about the reactivity of boranes towards hydride transfer [32,35,36].

In our previous paper [34], we focused on the DFT investigation of thermodynamic hydricity of tetracoordinate borohydrides Li[L3B–H]. However, despite the attempts to evaluate the electron-donating properties in polyhedral boranes by 1H NMR [37], there is a lack of knowledge of the reactivity of B–H bond in small borane clusters and polyhedral boron hydrides [36]. Thus, in this paper, we report on the results of the DFT analysis of thermodynamic hydricity in MeCN (HDA_{MeCN}) for both well-known, structurally characterized boranes and prospective reaction intermediates, some of which were previously known from theoretical works.

2. Results and Discussion

Borane clusters (such as B2H6, [B2H7]−, [B3H8]−, [B12H12]2−, [B10H10]2−, etc.) were discovered as intermediate products of borohydrides thermal decomposition [38–50]. These compounds are widely used in direct synthesis of other boranes [51,52], organoboron compounds [53], and new materials [5]. For example, hydride ion abstraction from borane anions (such as BH4−, [B3H8]−, [B4H9]−, etc.) by Lewis acids BX3 (X = F, Cl, Br) is the most convenient route for the preparation of higher boranes [54,55]. Many small borane clusters are highly reactive and unstable species, so their reactivity is hard to assess experimentally. Others, like polyhedral boron hydrides, require the presence of an excess of Lewis or Brønsted acids for generation of boron-centered quasi-borinium cations [36]. Thus, the reactivity of such compounds can be characterized in terms of their ability to hydride transfer as thermodynamic hydricity (HDA) through DFT calculations [32,34].

In this paper, we performed the DFT calculations of the thermodynamic hydricity in MeCN of small borane clusters (containing up to 5 boron atoms) and polyhedral boron hydrides [BnHn]2− (n = 5–17). For anionic species we use their lithium salts to offset the effect of ionic species and to make a correct comparison with the HDA_{MeCN} values for neutral boranes.

2.1. Thermodynamic Hydricity of Small Borane Clusters

Recently, the decomposition pathways of [LiBH4]n, n = 2–12 clusters were investigated by DFT calculations [48]. In the case of dimeric [LiBH4]2 clusters, the release of up to 4 equivalents of H2 was found, as well as [LiBHm]2 (m = 6, 4, 2) reaction intermediates, during the decomposition. Our calculations show that during the decomposition of [LiBH4]2 the first H2 release leads to the decrease of HDA_{MeCN} by 15.9 kcal/mol (Scheme 1). Further H2 release leads to an increase of HDA_{MeCN} due to formation unsaturated diboranes with double and triple bonds.
2.1.1. General Pattern in Thermodynamic Hydricity of Borane Clusters

To gain insight into an effect of borane (BH₃) aggregation in small clusters, we used the monomers LiBH₄, Li₃BH₃, Li₂BH₂ and Li₂BH along with neutral BH₃ and BH species to construct a homologous series of boron clusters (containing up to 5 boron atoms) by consequent addition of BH₃ (Table 1, Schemes S1 and S2). Another two homologous series were constructed based on Li[B₂H₅] and B₂H₂. In each series there are the most stable boranes (namely BH₄⁻, B₂H₆, [B₂H₇]²⁻, [B₃H₈]⁻, B₄H₁₀, [B₅H₁₁]⁻, [B₆H₁₂]⁻, and B₇H₁₄), which are generally observed during thermal decomposition of metal borohydrides [38-50].

Table 1. Calculated hydride donating ability in MeCN (HDA(MeCN) in kcal/mol) for homologous series of neutral and anionic boranes clusters.

| n  | Li[B₃H₆n+1] | B₇H₈n | Li₂[B₄H₆n] | Li[B₆H₆n-1] | B₈H₆n-2 | Li₂[B₈H₈n-2] | Li[B₈H₈n-3] | B₉H₈n-4 |
|----|-------------|--------|------------|-------------|---------|--------------|-------------|--------|
| 1  | LiBH₄       | BH₃    | Li₂BH₃     | LiBH₂       | BH      | Li₃BH        | -           | -      |
| 2  | Li[B₂H₅]   | B₂H₆  | Li₂[B₂H₆]  | Li[B₂H₅]   | B₂H₆   | Li₂[B₂H₆]    | Li[B₂H₆]   | B₂H₆   |
| 3  | Li[B₅H₁₀]  | B₇H₈  | Li₂[B₅H₁₀] | Li[B₅H₁₀]  | B₇H₈   | Li₂[B₅H₁₀]   | Li[B₅H₁₀]  | B₇H₈   |
| 4  | Li[B₇H₁₃]  | B₉H₁₂ | Li₂[B₇H₁₃] | Li[B₇H₁₃]  | B₉H₁₂  | Li₂[B₇H₁₃]   | Li[B₇H₁₃]  | B₉H₁₂  |
| 5  | Li[B₉H₁₆]  | B₁₁H₁₄| Li₂[B₉H₁₆] | Li[B₉H₁₆]  | B₁₁H₁₄ | Li₂[B₉H₁₆]   | Li[B₉H₁₆]  | B₁₁H₁₄ |

The HDA(MeCN) values are given for the most energetically favorable isomer. The most stable boranes in the series according to calculated heats of formation [50,56] are shown by the violet shading of the cells.

According to computed heats of formation [50], neutral boranes (such as B₂H₆, B₄H₁₀, and B₇H₁₄) are less stable than anionic ones (BH₄⁻, [B₂H₆]⁻ and [B₄H₆]⁻); however, the former may be derived from the latter as a result of hydride abstraction by Lewis acids BX₃ (X = F, Cl, Br) [55]. This lower stability of neutral boranes compared to anionic boranes can be explained by higher Lewis acidity. In our previous research, we demonstrated that Lewis acidity of parent borane (R₃B) can be estimated by the analysis of thermodynamic hydricity of their product of hydride addition [R₃BH]⁻. Higher HDA(MeCN) values of [R₃BH]⁻ correspond to higher Lewis acidity of R₃B.

Thermodynamic hydricity values of diborane B₂H₆ (HDA(MeCN) = 82.1 kcal/mol) and its dianion [B₂H₄]²⁻ (HDA(MeCN) = 40.9 kcal/mol for Li₂[B₂H₆]) can be selected as border points for the range of borane clusters’ reactivity. Previously [57], the formal division of the thermodynamic hydricity scale into weak (above 80 kcal/mol), medium (between 80 and 50 kcal/mol) and strong hydride donors (less 50 kcal/mol) was suggested.

Our study and literature analysis show that borane clusters with HDA(MeCN) below 41 kcal/mol are strong hydride donors capable of reducing CO₂ (HDA(MeCN) = 44 kcal/mol for HCO₂⁻ [31]); however, those with HDA(MeCN) over 82 kcal/mol, predominately neutral boranes, are weak hydride donors and less prone to hydride transfer than to proton transfer (e.g., B₂H₆ [58], B₄H₁₀, B₅H₁₁, etc. [59]). Moreover, in higher boranes, the deprotonation of B–H–B bridges leads to B–B bond formation, in which boron is nucleophilic and susceptible to the attack of electrophiles [59]. Thus, higher borane anions could be generated by the insertion of such electrophile molecules as BH₃, B₂H₂ or BH.  

![Scheme 1](image-url)  

Scheme 1. HDA(MeCN) (in kcal/mol) for (LiBH₄)₂ decomposition intermediates [48] and their monomers.
Based on the data obtained for the homologous series (Table 1, Schemes S1 and S2), an evolution of thermodynamic hydricity can be traced on the example of B₃ clusters transformations (Scheme 2). Thus, neutral boranes B₃H₉, B₃H₇, and B₃H₅ feature the highest Lewis acidity (highest HDAₑMCN values). Monoanionic boranes Li[B₃H₁₀] etc. have lower Lewis acidity than parent neutral boranes. The two-electron reduction of the neutral boranes leads to a significant decrease of HDAₑMCN values (by 41.1 kcal/mol for Li₂[B₃H₉] and by 20.0 kcal/mol for Li₂[B₃H₇]).

Although it is generally acknowledged that, in clusters, borane anions’ hydric hydrogen becomes less reactive due to increasing charge distribution across the cluster [8], our calculations show that HDAₑMCN values are not directly connected with the size of borane cluster (Figure S1). However, if we plot computed HDAₑMCN values for all series of boranes (n = 1–5; Figure 1), it appears that minimum HDAₑMCN values (nucleophilic boron) are typical for the dianionic species and maximum HDAₑMCN (electrophilic boron) for the neutral boranes, whereas borane monoanions are in the intermediate position. Since neutral transition metal hydrides are better hydride donors than cationic ones [31], it is obvious that anionic boranes should be much more better hydride donors than neutral boranes.

An increasing size of borane cluster in neutral BₙH₃ₙ, BₙH₃ₙ₋₂, and BₙH₃ₙ₋₄ and dianionic Li₂[BₙH₃ₙ] and Li₂[BₙH₃ₙ₋₂] series results in a drop of HDAₑMCN values. This leads to a smoothing of the extrema in the graph of HDAₑMCN for larger borane clusters as B₄ and B₅. Moreover, in the B₃ clusters due to triangle form an effective charge distribution occurs, which leads to the systematically lower values of HDAₑMCN.

It is worth noting that a decrease of saturation of the borane cluster results in an increase of HDAₑMCN that is especially pronounced for neutral diboranes.

Scheme 2. Interconversion scheme for Li[B₃H₁₀]. Lithium atoms are omitted for clarity.
2.1.2. Features of Thermodynamic Hydricity in Homologous Series

The Li[B_nH_{3n+1}] series is formed by consecutive BH_3 addition to tetrahydroborate-anion BH_4^- (Scheme 3). The first addition of BH_3 yields [B_2H_7]^— and leads to the HDA MeCN decrease by 16.4 kcal/mol, while the subsequent BH_3 addition leads to a gradual HDA MeCN increase due to the delocalization of electrons between a greater number of atoms.

**Scheme 3.** Formation of Li[B_nH_{3n+1}] series. Black numbers denote HDA MeCN values for B–H groups marked by blue color (in kcal/mol), red numbers in square brackets are the Gibbs free energies (ΔG° MeCN, in kcal/mol) of branched isomers relative to the most stable linear isomer for Li[B_4H_{13}] or Li[B_5H_{16}]. Lithium atoms are omitted for clarity.

Tetrahydroborate anions BH_4^- [4,60–68] and [B_2H_7]^— [69,70] are used as ligands in the synthesis of transition metal complexes. Structures of other borohydride anions were accessed by DFT...
calculations [58,71], but only branched isomers of Li[B_{4}H_{13}] and Li[B_{5}H_{16}] have been reported [71]. However, we found that their linear isomers were energetically more favorable (see $\Delta G_{\text{MeCN}}^0$, Scheme 3, Table S1). It is important to note that regardless of the structure of the isomer, the hydride detachment only occurs from the terminal BH$_3$ groups, due to their increased hydricity.

The B$_n$H$_{3n}$ series represents neutral boranes formed by consecutive BH$_3$ aggregation (Scheme 4), which causes a gradual decrease of HDA$_{\text{MeCN}}$ at each step (from 108.4 to 55.1 kcal/mol). Borane BH$_3$ has high HDA$_{\text{MeCN}}$ value ($\text{HDA}_{\text{MeCN}} = 108.4$ kcal/mol) and cannot be isolated due to its high Lewis acidity; however, it can be stabilized by dimerization into diborane B$_2$H$_6$ molecule ($\text{HDA}_{\text{MeCN}} = 82.1$ kcal/mol) or by interaction with Lewis bases (Scheme 5). The Lewis acid–base complexes of BH$_3$ are characterized by reduced HDA$_{\text{MeCN}}$ and some of them, especially amine and phosphine boranes, are able to form σ-complexes with transition metals [72–77].

![Scheme 4](image)

**Scheme 4.** B$_n$H$_{3n}$ series. Black numbers denote HDA$_{\text{MeCN}}$ values for B–H groups marked by blue color (in kcal/mol), red numbers in square brackets are the Gibbs free energies ($\Delta G_{\text{MeCN}}^0$ in kcal/mol) relative to the corresponding most stable isomer.

$\text{BH}_3 > \text{B}_2\text{H}_6 > \text{H}_3\text{P} \cdot \text{BH}_3 > \text{Me}_2\text{S} \cdot \text{BH}_3 > \text{H}_3\text{N} \cdot \text{BH}_3 > \text{THF} \cdot \text{BH}_3$

108.4  82.1  78.2  76.6  64.8  58.7

**Scheme 5.** The trend of HDA$_{\text{MeCN}}$ (in kcal/mol) for borane Lewis acid–base complexes. The HDA$_{\text{MeCN}}$ values are taken from ref. [34].

The diborane-based structures B$_2$H$_5$(μ-H)($(\text{BH}_2)(μ-H))_m$BH$_3$ (m = 1–3) were found to be the most stable isomers for higher borane clusters (Scheme 4, Table S2). In previous theoretical works, cyclic structures of B$_3$H$_9$ and B$_4$H$_{12}$ have been reported to be the most stable isomers in the gas phase [78–80]. However, according to our optimization, [M06/ and MP2/6-311++G(d,p) theory levels in MeCN using SMD] is the most stable configuration for B$_3$H$_9$ is B$_2$H$_5$(μ-H)BH$_3$, whereas butterfly-like and cyclic structures are slightly less favorable in terms of Gibbs free energy scale ($\Delta G_{\text{MeCN}}^0 = 0.5$ kcal/mol and 2.3 kcal/mol for M06 and 0.7 kcal/mol and 1.3 kcal/mol for MP2, respectively, Table S3).

The Li$_2$[B$_n$H$_{3n}$] series is formed by the consecutive addition of BH$_3$ to Li$_2$[BH$_3$] (Scheme S3). Both [BH$_3$]$^2^-$ [81] and [B$_2$H$_4$]$^2^-$ [82–88], which could be obtained by the reduction of B$_2$H$_6$ [89], are used as ligands in transition metal complexes. Boranes of this series formally could be obtained by
two-electron reduction from their neutral analogues $B_nH_{3n}$, leading to a significant decrease in their thermodynamic hydricity by 24–65 kcal/mol. High reactivity of $[B_2H_6]^2$ towards hydride transfer ($HDA_{MeCN} = 40.9$ kcal/mol for $Li_2[B_2H_6]$) results in full substitution of terminal BH hydrides in $(Cp^*M)_2(x^2-B_2H_6)$ ($M = V, Nb, Ta$) in chlorinated solvents ($CH_2Cl_2$ and $CHCl_3$) [86,88]. Higher boranes were not isolated, which is apparently related to their low $HDA_{MeCN}$ (Table 1); however, according to our DFT calculations, the structures of $[B_3H_6]^{2−}$, $[B_4H_{12}]^{2−}$ and $[B_5H_{13}]^{2−}$ are based on $[B_2H_6]^{2−}$ geometry (Scheme S3, Table S4).

The $Li[B_nH_{3n−1}]$ series is formed via the consecutive addition of BH$_3$ to LiBH$_2$ (Scheme S4). Along this series, $HDA_{MeCN}$ decreases from 64.4 kcal/mol for LiBH$_2$ to 44.5–45.1 kcal/mol for Li[B$_3$H$_{11}$] and Li[B$_2$H$_{14}$]. The most stable structure in this series is Li[B$_3$H$_8$], and this anion is used as a ligand in transition metal complexes [70,90–94]. In the case of Li[B$_4$H$_{11}$] and Li[B$_3$H$_{14}$], the isomers, the structures of which are based on $[B_3H_8]^{−}$, are found by 19.2 and 24.0 kcal/mol lower in $\Delta G^{°}_{MeCN}$ scale than the isomers having a cyclic structure (Scheme S4, Table S5).

According to the literature, there are several possibilities for synthesizing $[B_3H_8]^{−}$ (Scheme 6) [52,58,95]. Formation of the octahydrotriborate anion $[B_3H_8]^{−}$ in the reaction of BH$_4^{−}$ with B$_2$H$_6$ or with B$_2$H$_4$ can be explained by the electrophilic nature of neutral boranes (evidenced by higher $HDA_{MeCN}$, Table 1 and Scheme S1) [58,95].

![Scheme 6. Reactions of tetrahydroborate anion with neutral boranes leading to $[B_3H_8]^{−}$. $HDA_{MeCN}$ values (in kcal/mol) are also given.](image)

The $B_nH_{3n−2}$ series is formed via the consecutive addition of BH$_3$ to BH (Scheme S5). In this series, except for the B$_3$H$_{13}$, all compounds were obtained experimentally. The first BH$_3$ addition leads to an increase in $HDA_{MeCN}$ by 14.5 kcal/mol, the second addition causes its reduction by 30.9 kcal/mol, whereas for the next members in the series, B$_3$H$_7$, B$_4$H$_{10}$, and B$_5$H$_{13}$, $HDA_{MeCN}$ varies in a narrow range of 74.5–78.2 kcal/mol.

Boron monohydride radical is known to be generated in the gas phase by photodissociation of BH$_3$CO [96,97]. B$_2$H$_4$ can be obtained by the reaction of B$_2$H$_6$ with F radicals in the gas phase [98]. Due to its high electrophilicity ($HDA_{MeCN} = 105.4$ for B$_2$H$_4$ is comparable to $HDA_{MeCN} = 108.4$ for BH$_3$), there is a lack of transition metal complexes with B$_2$H$_4$ [99]; however, it can be stabilized in the form of a Lewis acid–base complex (Me$_3$P)$_2$B$_2$H$_4$ [100–104]. In turn, (Me$_3$P)$_2$B$_2$H$_4$ can act as a bidentate ligand in transition metal complexes [102–105]. B$_3$H$_7$ can be easily obtained from B$_3$H$_5^{−}$ by reaction with a non-oxidizing acid (Scheme 6) in ether solvents (e.g., THF) or in the presence of other Lewis bases (L = R$_3$N, R$_3$P) [52,106,107] with the formation of L-B$_3$H$_7$ adduct [108]. The most stable borane in this series, B$_4$H$_{10}$, was characterized rather a long time ago, and its structure was determined in the gas phase by gas-phase electron diffraction [109,110]. B$_4$H$_{10}$ geometry is preserved in the structure of B$_5$H$_{13}$ (Scheme S5, Table S6), which is characterized by a slightly lower $HDA_{MeCN}$ value.
In the Li₂[BₙH₃₋₂] series formed by the consecutive addition of BH₃ to Li₂BH (Scheme S6), the HDAMeCN values decrease by 20–52 kcal/mol relative to the corresponding neutral boranes BₙH₃₋₂ (Table 1 and Scheme 2). Both [B₂H₄]²⁻ [99,102,104,111] and [B₃H₇]²⁻ [82,112–114] are used as ligands in transition metal complexes. The most stable structures, [B₄H₁₂]²⁻ and [B₅H₁₅]²⁻, are based on the geometry of [B₃H₇]²⁻, whereas cyclic structures are energetically less favorable by 7.4 and 16.8 kcal/mol (ΔG°MeCN), respectively (Table S7).

The Li₂[BₙH₃₋₃] series is formed by the addition of BH₃ to B⁻ (Scheme S7, Table S8). The change in the thermodynamic hydricity in this series is uneven. The first BH₃ addition leads to a significant drop in HDAMeCN by 41.9 kcal/mol, the second addition causes its reduction by 13.3 kcal/mol, and the third addition leads to an increase of HDAMeCN by 11.3 kcal/mol. The first two members of this series, [B₂H₅]⁻ [115,116] and [B₃H₆]⁻ [80], are observed in the gas phase during collision-induced dissociation [117], and their structures were predicted by DFT calculations. [B₄H₆]⁻ can be synthesized in quantitative yield by deprotonation of B₄H₁₀, whereas the addition of B₂H₆ to [B₄H₆]⁻ yields [B₅H₁₂]⁻ [51,118]. [B₄H₈]⁻ is used as a ligand in transition metal complexes [70,119–122], and it is the most stable structure in this series (HDA MeCN = 60.2 kcal/mol).

Finally, the BₙH₃₋₄ series is formed via the consecutive addition of BH₃ to B₂H₂ (Scheme S8, Table S9). Upon the BH₃ addition, HDAMeCN gradually decreases from an extremely high value of 168.6 kcal/mol for B₂H₂ to 85.5 kcal/mol for B₂H₁₁. B₂H₂ [100,123], B₃H₅ [100] and B₄H₆ [124,125] are obtained as a result of higher borane cleavage (e.g., B₃H₆) and need to be stabilized by the interaction with Lewis bases (Me₃P, Me₃N, Me₂S) due to their high Lewis acidity. Their Lewis acid–base adducts (such as (Me₃P)₂B₂H₂) have used as ligands in transition metal complexes [104,126,127].

### 2.2. Thermodynamic Hydricity of Polyhedral Closo-Boranes

During thermal decomposition of metal tetrahydroborates, the formation of stable metal dodecaborane [B₁₂H₁₂]²⁻ with an admixture of [B₁₀H₁₀]²⁻ via intermediate boron clusters such as [B₂H₇]⁻ and [B₃H₆]⁻ was observed [42,45–47,50]. In octahedral [B₆H₆]²⁻ and icosahedral [B₁₂H₁₂]²⁻ closo-borane diions, all terminal BH groups are equivalent. However, in their lithium salts Li₂[BₙHₙ], cation–anion interaction causes asymmetry in the bond lengths of terminal BH groups interacting with Li atoms. Therefore, thermodynamic hydricity of polyhedral closo-boranes was assessed for both [BₙHₙ]²⁻ diions and their lithium salts Li₂[BₙHₙ] (n = 5–17).

Theoretical calculations show that the thermodynamic stability of polyhedral closo-boranes increases with an increase in the number of boron atoms participating in the cluster formation from [B₂H₂]²⁻ to [B₁₂H₁₂]²⁻ [50]. According to thermodynamic stabilities, closo-borane diions such as [B₂H₁₇]²⁻ and [B₄H₁₅]²⁻ should be even more stable than [B₁₂H₁₂]²⁻ [128]. Indeed, DFT calculations predict that the formation of large clusters (such as [B₄₂H₄₂]²⁻, [B₆₀H₆₀]²⁻, [B₂₀H₂₀]²⁻) is possible [129–132]. However, one should keep in mind that the more B–H bonds present in a polyhedral closo-borane, the higher its stability, since more energy is stored in these chemical bonds [30].

To gain insight in thermodynamic hydricity of polyhedral closo-boranes, we calculated at first stage HDAMeCN of small dianionic boranes [BₙHₙ]²⁻ and their lithium salts Li₂[BₙHₙ] (n = 2–4) (Figure 2), which can be formally viewed as building blocks or prototypes of polyhedral closo-boranes.

![Figure 2. M06-optimized geometries of Li₂[BₙHₙ] (n = 2–4). Lithium atoms are omitted for clarity.](image-url)
Whereas \([B_2H_2]^{2-}\) is linear, and \([B_3H_3]^{2-}\) is planar, \([B_4H_4]^{2-}\) has been described as having an “intermediate configuration between planar and tetragonal geometry” \cite{133}, which is actually disphenoid. In each structure, all B-H\(_{\text{term}}\) bonds are equivalent (\(r_{\text{B-H\(_{\text{term}}}) = 1.191 \text{ Å}\) for \([B_2H_2]^{2-}\); \(r_{\text{B-H\(_{\text{term}}}) = 1.209 \text{ Å}\) for \([B_3H_3]^{2-}\) and \(r_{\text{B-H\(_{\text{term}}}) = 1.206 \text{ Å}\) for \([B_4H_4]^{2-}\). The decrease of HDA\(_{\text{MeCN}}\) values in this series (Scheme 7) is apparently associated with a better delocalization of electron density and with an increase in the number of possible resonance structures as the cluster size grows. In \([B_2H_2]^{2-}\) featuring a triple B-B bond, there are two 2c-2\(\bar{e}\) \(\pi\)-bonds, whereas in \([B_3H_3]^{2-}\) and \([B_4H_4]^{2-}\) featuring \(\pi\)-aromatic systems, there is one 3c-2\(\bar{e}\) and one 4c-2\(\bar{e}\) delocalized \(\pi\)-bond, respectively \cite{133}.

\[
[B_2H_2]^{2-} > [B_3H_3]^{2-} > [B_4H_4]^{2-}
\]

\[100.8 \quad 37.3 \quad 21.1\]

\textbf{Scheme 7.} Order of HDA\(_{\text{MeCN}}\) (in kcal/mol) for borane dianions \([B_nH_n]^{2-}\) (\(n = 2–4\)).

Further increase in borane cluster size results in the formation of polyhedral \textit{closo}-boranes structures \(\text{Li}_2[B_nH_n]\) (\(n = 5–17\)) (Figure 3), which are formed following the Wade’s rules \cite{134,135}. Due to the 3D aromaticity, \textit{closo}-boranes are characterized by higher HDA\(_{\text{MeCN}}\) values (Figure 4, Figure S2) than small \([B_2H_2]^{2-}\) clusters (\(n = 2–4\)). Formal addition of BH to \([B_4H_4]^{2-}\) yields \([B_5H_5]^{2-}\), which is the least stable member of the polyhedral \textit{closo}-boranes \([B_nH_n]^{2-}\) according to formation enthalpy \cite{50}, and has never been synthesized \cite{128,133}. The structure of \([B_5H_5]^{2-}\) is typical for the polyhedral \textit{closo}-boranes—there are two types of boron atoms—two B atoms form caps, and a group of three other B atoms has the geometry of a B\(_5\)H\(_5\) triangle, forming a belt of the polyhedron (for further description of structural features of polyhedral \textit{closo}-boranes, see in SI).

Boron atoms in the belt and caps of the polyhedron have different coordination numbers (CN)—4 and 5, respectively. That in turn leads to a difference in B–H bonds’ lengths (\(r_{\text{B-H\(_{\text{eq}})}} = 1.194 \text{ Å}\) and \(r_{\text{B-H\(_{\text{eq}})}} = 1.202 \text{ Å}\), Figure 3) and anisotropy in the charge distribution across the polyhedron (Figure S2). According to natural bond population analysis (NPA) of \([B_5H_5]^{2-}\) the capping boron atoms are more negatively charged (\(q_{\text{MeCN}} = -0.437\)) than the equatorial ones (\(q_{\text{MeCN}} = -0.331\)). Thus, in the case of \([B_5H_5]^{2-}\), the apical BH groups (82.0 kcal/mol) have higher HDA\(_{\text{MeCN}}\) values than equatorial ones (60.0 kcal/mol) (Figure S2). This HDA\(_{\text{MeCN}}\) (BH\(_{\text{eq}}\))/HDA\(_{\text{MeCN}}\) (BH\(_{\text{ap}}\)) ratio is conserved when going to larger dianions \([B_{10}H_{10}]^{2-}\), \([B_{15}H_{15}]^{2-}\) and \([B_{17}H_{17}]^{2-}\).

To gain insight into the thermodynamic hydricity of polyhedral \textit{closo}-boranes, it is highly important to consider the difference in geometry along with the charge distribution in their dianions (Figure S2, Table S10). These parameters suggest different reactivity of boron atoms forming the polyhedron’s caps and belt (Figure 3). Although different bond lengths of terminal B-H already indicate different properties of these centers, it is not possible to estimate their thermodynamic hydricity, since there is no general relationship between \(r_{\text{B-H\(_{\text{eq}})}}\) and HDA\(_{\text{MeCN}}\) (Figure S4).

Thermodynamic hydricity was assessed for both \([B_nH_n]^{2-}\) dianions (Figure 4, Table S10) and their lithium salts \(\text{Li}_2[B_nH_n]\) (\(n = 5–17\)) (Figure S3, Table S11). In lithium salts, the cation–anion interaction causes asymmetry in BH bond length and leads to different HDA\(_{\text{MeCN}}\) values for the same vertex types, which pushes us to use HDA\(_{\text{MeCN}}\) for dianions to make a correct comparison inside the \([B_nH_n]^{2-}\) series. HDA\(_{\text{MeCN}}\) values for lithium salts (typically higher by 10.2–22.3 kcal/mol than those for dianions) are provided for the comparison with the neutral and monoanion hydrides (Table S11). HDA\(_{\text{MeCN}}\) of \textit{closo}-boranes directly depends on the coordination number (Table S10) of the boron atom, for which the hydride abstraction and stabilization of quasi-borinium cation take place. In general, the larger the coordination number (CN) of a boron atom, the lower the value of HDA\(_{\text{MeCN}}\). Deviation from this rule is observed only for \([B_7H_7]^{2-}\), \([B_{13}H_{13}]^{2-}\) and \([B_{14}H_{14}]^{2-}\), where the boron atoms with the highest CN have the largest HDA\(_{\text{MeCN}}\) values. The probable explanation is that despite that the boron atoms forming the polyhedron belt have a lower CN than the boron atoms of the cape, the interaction between them and the surrounding atoms is stronger. This is suggested by shorter \(r_{\text{B-B}}\) for borons with lower
CN, as in, e.g., \([\text{B}_{11}\text{H}_{11}]^{2-}\) (1.773–1.802 Å), \([\text{B}_{13}\text{H}_{13}]^{2-}\) (1.739–1.826 Å) and \([\text{B}_{14}\text{H}_{14}]^{2-}\) (1.732–1.904 Å), in comparison to longer bonds at boron atoms with higher CN (\(r_{B-B}\)) is 1.738–1.971 Å for \([\text{B}_{11}\text{H}_{11}]^{2-}\), 1.842–1.904 Å for \([\text{B}_{13}\text{H}_{13}]^{2-}\) and 1.904 Å for \([\text{B}_{14}\text{H}_{14}]^{2-}\).

In most cases, except for \([\text{B}_{7}\text{H}_{7}]^{2-}\) and \([\text{B}_{16}\text{H}_{16}]^{2-}\), the equatorial BH groups have the lowest HDA\text{MeCN} values. The trend in the ability to hydride transfer (Scheme 8, for \(	ext{Li}_2[\text{B}_6\text{H}_6]\) see Scheme S10) correlates with the calculated Gibbs free energy per unit (G°/n) for both \(	ext{Li}_2[\text{B}_n\text{H}_n]\) and \([\text{B}_n\text{H}_n]^{2-}\) (\(n = 5–17\)) (Figures S5 and S6) and with previously reported energy per unit by PRDDO calculations [136]. The data obtained indicate that the most stable \(\text{closo}\)-borane clusters. Boron atoms in the belt and caps of the polyhedron have different coordination numbers (CN)—numbers of boron atom. Black numbers are B–H bond length in Å. Dotted lines represent the connection between capes and belts of a polyhedron.

![Figure 3. M06-optimized geometries of \([\text{B}_n\text{H}_n]^{2-}\) (\(n = 5–17\)) dianions of polyhedral \(\text{closo}\)-boranes. Boron atoms given in violet, green or yellow color represent structural fragments with different coordination numbers of boron atom. Black numbers are B–H bond length in Å. Dotted lines represent the connection between capes and belts of a polyhedron.](image-url)
Multi-reference systems are or might be involved since it has been parametrized for both main group elements and transition metals. In this regard, M06 is a more versatile method than M06-2X (generally recommended for calculation thermochemistry of main group elements) and can be used in the cases where multi-reference systems are or might be involved since it has been parametrized for both main group elements and transition metals [138].

Computational Details

In the present manuscript, DFT/M06 was used to allow a comparison to hydride donating ability of tetracoordinated boron hydrides previously calculated by the same method [34]. Additionally, the values obtained can be used as a reference for assessing the effectiveness of the activation of B-H bonds by transition metals. In this regard, M06 is a more versatile method than M06-2X (generally recommended for calculation thermochemistry of main group elements) and can be used in the cases where multi-reference systems are or might be involved since it has been parametrized for both main group elements and transition metals [138].

3. Materials and Methods

Figure 4. HDA MeCN of polyhedral closo-borane dianions [BnHn]2− (n = 5–17). Blue columns represent the lowest HDA MeCN values, and red columns represent the highest in the given anion; green columns show HDA MeCN for symmetric [B5H5]2− and [B12H12]2− dianions.

\[
[B_{12}H_{12}]^{2-} > [B_{14}H_{14}]^{2-} > [B_{16}H_{16}]^{2-} > [B_{18}H_{18}]^{2-} > [B_{20}H_{20}]^{2-} > [B_{22}H_{22}]^{2-} > [B_{24}H_{24}]^{2-} > [B_{26}H_{26}]^{2-} > [B_{28}H_{28}]^{2-} > [B_{30}H_{30}]^{2-} > [B_{32}H_{32}]^{2-} > [B_{34}H_{34}]^{2-} > [B_{36}H_{36}]^{2-} > [B_{38}H_{38}]^{2-} > [B_{40}H_{40}]^{2-} > [B_{42}H_{42}]^{2-} > [B_{44}H_{44}]^{2-} \\
79.6 \quad 75.9 \quad 75.6 \quad 74.1 \quad 71.9 \quad 70.8 \quad 70.8 \\
> [B_{11}H_{11}]^{2-} > [B_{17}H_{17}]^{2-} > [B_{25}H_{25}]^{2-} > [B_{33}H_{33}]^{2-} > [B_{41}H_{41}]^{2-} > [B_{49}H_{49}]^{2-} > [B_{57}H_{57}]^{2-} > [B_{65}H_{65}]^{2-} > [B_{73}H_{73}]^{2-} > [B_{81}H_{81}]^{2-} > [B_{89}H_{89}]^{2-} > [B_{97}H_{97}]^{2-} > [B_{105}H_{105}]^{2-} > [B_{113}H_{113}]^{2-} \\
65.3 \quad 62.4 \quad 60.0 \quad 56.8 \quad 56.3 \quad 39.4
\]

Scheme 8. The trend of HDA MeCN for dianions of polyhedral closo-borane [BnHn] (n = 5–17) taking the lowest value in the given anion.

Previously, electron-donating properties of polyhedral boranes were assessed by 1H NMR, which revealed the electron-donating ability decrease in the row [2-B10H9]2− > [B12H12]2− > [1-B12H12]2− [37]. According to our calculations, the HDA MeCN value for [B12H12]2− (79.6 kcal/mol) is higher than HDA MeCN for equatorial BH groups in [B10H10]2− (75.6 kcal/mol), but lower than for apical BH terminal groups in [B10H10]2− (89.0 kcal/mol).

Figure 4.
All calculations were performed without symmetry constraints using the M06 hybrid functional [138] and MP2 implemented in the Gaussian09 (Revision D.01) (Wallingford, CT, USA) [139] software package, using the 6-311++G(d,p) basis set [140].

Vibrational frequencies were calculated for all optimized complexes at the same level of theory to confirm a character of local minima on the potential energy surface. Visualization of the optimized geometries was realized using the Chemcraft 1.8 graphical visualization program [141].

The inclusion of nonspecific solvent effects in the calculations was performed by using the SMD method [142]. Acetonitrile (MeCN, \( \varepsilon = 35.7 \)) was chosen as a solvent for the geometry optimization because a large amount of data on reduction potentials, pKa values, and experimental hydride donating ability (HDA) of transition metal hydride complexes were determined in MeCN [31,143].

The calculations were carried out with an ultrafine integration grid and a very tight SCF option to improve the accuracy of the optimization procedure and thermochemical calculations.

Hydride donating ability in MeCN (HDA\(_{\text{MeCN}}\)) was calculated as Gibbs free energy of hydride transfer [HDA\(_{\text{MeCN}}\) = \( \Delta G^\circ[\text{H}^-_{\text{Solv}} = G^\circ_{\text{Solv}} (E^+\text{)} + G^\circ_{\text{Solv}} (H^-) - G^\circ_{\text{Solv}} (E-H)\)].

From the data obtained during the geometry optimization, for each molecule, the most stable configuration of each of its conformers was chosen. To find the most stable configuration of cationic boranes, the terminal hydrogen atoms (B–H\(_{\text{term}}\)) were torn off from each vertex in optimized molecules. As is widely known, bridge hydrogen atoms (B–H\(_{\text{br}}\)) have an increased acidity [5,58,144–147]; therefore, their participation in the hydride transfer was not considered herein. For most of the small borane clusters, due to the structural rearrangements during the geometry optimization only one stable configuration of cationic boranes was found.

In the case of polyhedral close-boranes due to the rigid frame of the boron cluster [133,148], several quasi-borinium cations were observed, localized on vertices from which hydride was torn off.

4. Conclusions

Our DFT study of thermodynamic hydricity (HDA\(_{\text{MeCN}}\)) revealed that for small borane clusters (up to 5 boron atoms), the hydride detachment occurs only from the ending terminal BH\(_n\) (\( n = 1–3 \)) group having the largest number of hydrogen atoms. The experimental data and the HDA\(_{\text{MeCN}}\) pattern obtained suggest that stable borane clusters have HDA\(_{\text{MeCN}}\) between 48 and 82 kcal/mol. Hydrides with lower HDA\(_{\text{MeCN}}\) would be hydrolytically unstable, and those with higher HDA\(_{\text{MeCN}}\) tend to aggregation in larger clusters. Neutral boranes with high Lewis acidity such as B\(_2\)H\(_2\) (166.6 kcal/mol), B\(_3\)H\(_3\) (109.9 kcal/mol), BH\(_3\) (108.4 kcal/mol), B\(_4\)H\(_4\) (105.4 kcal/mol) and B\(_5\)H\(_5\) (89.2 kcal/mol) could be stabilized in the form a Lewis acid–base complex (L-B,H\(_\nu\)), where L = R\(_3\)N, R\(_3\)P, THF, etc.), the formation of which increases the BH group’s hydricity (lowers HDA\(_{\text{MeCN}}\)) [25,26]. Borane clusters with HDA\(_{\text{MeCN}}\) less than 41 kcal/mol are strong hydride donors capable of reducing CO\(_2\) (HDA\(_{\text{MeCN}}\) = 44 kcal/mol for HCO\(_2^-\)), whereas those with HDA\(_{\text{MeCN}}\) over 82 kcal/mol, predominately neutral boranes, are weak hydride donors and could even serve as proton donors (e.g., B\(_2\)H\(_6\), B\(_3\)H\(_{10}\), B\(_4\)H\(_{11}\) etc.).

In close-boranes, HDA\(_{\text{MeCN}}\) depends on the coordination number (CN) of the boron atom from which hydride detachment and stabilization of quasi-borinium cation takes place. Thus, HDA\(_{\text{MeCN}}\) for equatorial boron vertices (75.6 kcal/mol, CN = 6) in [B\(_{10}\)H\(_{10}\)]\(^{2-}\) is lower than those for apical boron vertices (89.0 kcal/mol, CN = 5). That could explain the observed experimental reactivity row [2-B\(_{10}\)H\(_{9}\)]\(^{2-}\) > [B\(_{12}\)H\(_{12}\)]\(^{2-}\) > [1-B\(_{10}\)H\(_{9}\)]\(^{2-}\) [37].

Supplementary Materials: The following are available online. Scheme S1. Scheme of homologous series of neutral and monoanionic boranes clusters. Scheme S2. Homologous series of neutral and dianionic boranes clusters, Figure S1. Plots of HDA\(_{\text{MeCN}}\) against the number of boron atoms in borane clusters derived from different starting species. Table S1. Computed [M\(_{06}/6-311++G(d,p)\)] B–H terminal bond length (in Å), hydride donating ability (HDA\(_{\text{MeCN}}\)) add\( \Delta G^\circ[\text{H}^-_{\text{MeCN}}\) in kcal/mol] and enthalpy of hydride detachment reaction (\( \Delta H^\circ[\text{H}^-_{\text{MeCN}}\) in kcal/mol) for [Li[B\(_n\)H\(_{3n+1}\)] series. Table S2. Computed [M\(_{06}/6-311++G(d,p)\)] B–H terminal bond length (in
Å), hydride donating ability (HDA\textsubscript{MeCN} alias \(\Delta G^\circ\textsubscript{MECN}\)) in kcal/mol and enthalpy of hydride detaching reaction (\(\Delta H^\circ\textsubscript{MECN}\)) relative to the most stable isomer of \(\text{B}_2\text{H}_6\) in MeCN (\(\Delta G^\circ\textsubscript{MeCN}\) in kcal/mol) and B-H terminal bond length (in Å), hydride donating ability (HDA\textsubscript{MeCN} alias \(\Delta G^\circ\textsubscript{MECN}\)) in kcal/mol and enthalpy of hydride detaching reaction (\(\Delta H^\circ\textsubscript{MECN}\)) and hydride donating ability (HDA\textsubscript{MeCN} alias \(\Delta G^\circ\textsubscript{MECN}\)) in kcal/mol and B-H terminal bond length (in Å), hydride donating ability (HDA\textsubscript{MeCN} alias \(\Delta G^\circ\textsubscript{MECN}\)) in kcal/mol and enthalpy of hydride detaching reaction (\(\Delta H^\circ\textsubscript{MECN}\)) for \(\text{B}_2\text{H}_{3n-2}\) series. Scheme S4. \(\text{Li}[\text{B}_2\text{H}_{3n-1}]\) series. Table S5. Computed \(\Delta H\) and \(\Delta G\) for \(\text{Li}[\text{B}_2\text{H}_{3n-2}]\) and \(\text{Li}[\text{B}_2\text{H}_{3n-1}]\) series. Table S6. \(\text{Li}[\text{B}_2\text{H}_{3n-3}]\) series. Scheme S7. \(\text{Li}[\text{B}_2\text{H}_{3n-3}]\) series. Table S8. \(\Delta G\) for \(\text{Li}[\text{B}_2\text{H}_{3n-3}]\) series. Scheme S9. \(\Delta G\) for \(\text{Li}[\text{B}_2\text{H}_{3n-3}]\) series. Scheme S9. \(\Delta G\) for \(\text{Li}[\text{B}_2\text{H}_{3n-4}]\) series. Table S9. \(\Delta G\) for \(\text{Li}[\text{B}_2\text{H}_{3n-4}]\) series. Figure S2. NPA charge distribution (showed in blue-green-red scale from -0.40 to 0.05), calculated for M06-optimized geometries of diions \(\text{B}_2\text{H}_4\)\textsuperscript{2−} (n = 5–17) of polyhedral \textit{closo}-boranes in MeCN. Table S10. Coordination numbers (CN) of boron atom in polyhedral \textit{closo}-boranes. Figure S2. NPA charge distribution (showed in blue-green-red scale from -0.40 to 0.05), calculated for M06-optimized geometries of diions \(\text{B}_2\text{H}_4\)\textsuperscript{2−} (n = 5–17) of polyhedral \textit{closo}-boranes in MeCN. Table S10. Coordination numbers (CN) of boron atom in polyhedral \textit{closo}-boranes. Figure S2. NPA charge distribution (showed in blue-green-red scale from -0.40 to 0.05), calculated for M06-optimized geometries of diions \(\text{B}_2\text{H}_4\)\textsuperscript{2−} (n = 5–17) of polyhedral \textit{closo}-boranes in MeCN. Table S10. Coordination numbers (CN) of boron atom in polyhedral \textit{closo}-boranes. Figure S2. NPA charge distribution (showed in blue-green-red scale from -0.40 to 0.05), calculated for M06-optimized geometries of diions \(\text{B}_2\text{H}_4\)\textsuperscript{2−} (n = 5–17) of polyhedral \textit{closo}-boranes in MeCN. Table S10. Coordination numbers (CN) of boron atom in polyhedral \textit{closo}-boranes. Figure S2. NPA charge distribution (showed in blue-green-red scale from -0.40 to 0.05), calculated for M06-optimized geometries of diions \(\text{B}_2\text{H}_4\)\textsuperscript{2−} (n = 5–17) of polyhedral \textit{closo}-boranes in MeCN. Table S10. Coordination numbers (CN) of boron atom in polyhedral \textit{closo}-boranes.
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