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Boron beats nitrogen: strained boron–boron bonds as (molecular) proton sponges†‡

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Although boron usually behaves as a Lewis acid, some molecular structures containing B–B bonds can act as electron donors. Inspired in reported crystalline structures, the basicity and the hydrogen bond (HB) acceptor capability of diborane derivatives of dipyrazole **1** have been studied theoretically using M06-2x and CCSD(T) computational methods. The topology of the electron density and molecular electrostatic potential of compound **1** reveal that the richest electron region is located above the B–B bond, making it suitable to be a strong donor and a very effective proton catcher. A key finding is the remarkably high proton affinity of the parent derivative, which exceeds that of very strong nitrogen-containing organic bases such as guanidines. In line with this finding, the hydrogen-bonded complexes exhibit binding energies up to 37 kJ mol^{−1}, which is a significant interaction considering the electronegativity of boron in comparison with elements typically involved in hydrogen bonds. We have also designed substituted structures where the inductive effect improves the proton affinity and HB acceptor capabilities. The proton affinities of **1b** reach 1096 kJ mol^{−1}, among the highest reported for molecules in the gas-phase.

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1. Introduction

The small boron atom is one of the most versatile of the periodic table, showing a rich variety of bonding patterns beyond the common trivalent monomeric compounds. For instance, the diborane (B₂H₆) structure with bridging hydrogens is well known for being the paradigm of three centre-two electron (3c–2e) bonds.¹ Another interesting example is the highly stable species B₁₂H₁₂^{2−}, closoborane, which exhibits an icosahedral I_h symmetry.^{2,3} In this structure, each boron atom is surrounded by five atoms: four other borons and one hydrogen.

Single B–B bonds, are found, among other systems, in diboron(4) compounds. The structure of its parent compound, B₂H₄, has been in debate for years. Several experimental and high-level theoretical studies have proved it belongs to the C_{2v} symmetry group.^{4,5} The ability of the B–B bond of this molecule to act as a HB acceptor has been explored by some of us,⁶ and

the reactivity and electronic characteristics of diboron(4) and some of its simple derivatives have been reviewed recently.⁷ The growing interest on boron chemistry and its applications to synthesis made us explore the physicochemical properties of strained boron–boron compounds, with the aim of finding out what were the limits they could reach in particular for the acid–base behaviour.

In the search of promising donors of this kind, we found a set of reported structures achieved by dehydrogenation of dimers of borane derivatives, and characterized by X-ray crystallography, where polycyclic structures **I–III** with B–B bonds are stabilized by dative bonds (Scheme 1).^{8–11} The central core of these structures present two five-membered rings with a common B–B bond. The theoretical analysis of **I** shows that the HOMO orbital is located on this bond, and the protonation of its derivatives proceeds with its breakdown, placing the new hydrogen atom between the two boron atoms.¹² Following these evidences, the dehydrogenation of pyrazobole **IV** could yield compound **1**, which contains a B–B bond leading to two four membered rings (Scheme 2). Because of this, we hypothesized that this highly strained bond could exhibit better donor qualities than the previously described ones.

In the present article, we will study the donor properties of the abovementioned tetracyclic structure **1** with three different substituents on the boron atom (see Scheme 3), to consider a range of inductive effects. For each of them, the protonation and the HB complexes with five typical HB donors (HF, HCl, HBr, HCN and

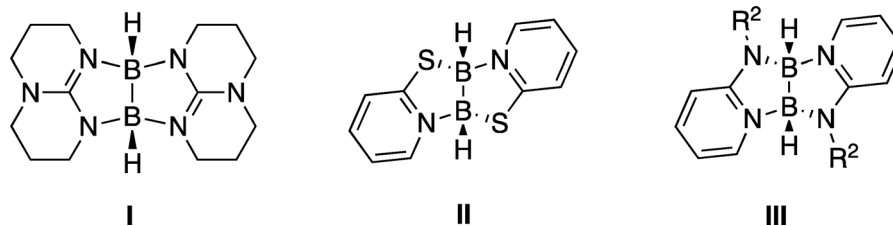
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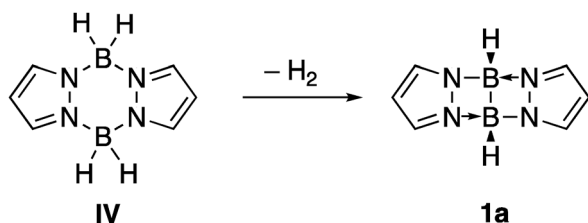
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‡ Dedicated to Professor Resnati, celebrating a career in fluorine and noncovalent chemistry on the occasion of his 70th birthday.

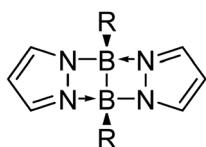




Scheme 1 Examples of polycyclic systems I–III with a B–B single bond.



Scheme 2 Reaction of dehydrogenation of pyrazobole IV to yield the parent compound **1a**.



1a, R = H, **1b**, R = CH₃, **1c**, R = F

Scheme 3 The three derivatives of **1** considered in the present article, to be combined with HF, HCl, HBr, HCN and HCCH hydrogen-bond donors.

HCCH) are studied at the CCSD(T)-F12c and M06-2X computational levels, leading to a survey of 18 complexes fully characterized at a high level of theory.

2. Computational methods

The geometries of the whole set of molecules and complexes have been optimized with the M06-2X method¹³ and the aug-cc-pVTZ basis set,¹⁴ an approach that provides a reliable treatment of electron correlation and noncovalent interactions for this type of complexes.^{15,16} Frequency calculations were carried out at the same computational level to verify that the structures correspond to energetic minima. In addition, the complexes of the parent compound **1a** were reoptimized at the CCSD(T)-F12c/VDZ-F12 level.¹⁷ The CCSD (T) method is known as the gold-standard in quantum chemistry.^{18,19} The DFT calculations were carried out with the Gaussian-16 program²⁰ and the CCSD(T) ones with the Molpro package.²¹ The optimized geometries are collected in Table S1 of the ESI.†

The electronic properties of the systems have been analysed through the molecular electrostatic potential (MEP),²² electron localization function (ELF),^{23,24} natural bond orbital (NBO) theory,²⁵ quantum theory of atoms in molecules (QTAIM),^{26,27} and electron density shift (EDS).²⁸ MEP regions with negative

values (isosurfaces coloured in red) indicate parts of the molecules suitable to react or interact with electron deficient systems, as it is the case in protonation or forming intermolecular hydrogen bonds. The ELF maps reflect localized electron pairs in the space for a given system, offering a Lewis-like picture through the partition of the molecular electron density into basins. They are especially useful to evaluate the electron population of a given bond or a lone pair. The NBO method also provides a Lewis-like description from an orbitalic point of view, including hyperconjugation (charge transfer) between occupied and empty orbitals. The Natural energy decomposition analysis (NEDA),²⁹ which is based on the NBO theory, has been performed to obtain information on the most important components of the interaction energy of the complexes. QTAIM analyses the topology of the electron density, for which the critical points are classified based on the number of positive or negative curvatures as nuclear attractor (3,−3), bond (3,−1), ring (3, +1), and cage critical points (3, +3). In particular, the properties of the bond critical points (BCP) are useful to characterize and quantify the interatomic interactions. Finally, the EDS shows how the electron density is reorganized in the complex with respect to the isolated molecules.

3. Results and discussion

We present our results divided in three parts, where the first section focusses on the properties of the isolated parent compound **1a**, paying attention to electronic characteristics and proton affinity; the second explores the hydrogen bonded complexes of **1a**, and the third and final section discusses the inductive effect of the substituents, specifically **1b** and **1c**, in terms of their proton affinity values and ability to form hydrogen bond complexes. This approach allows a quite systematic exploration of the relationship between structure and donor properties of the prototype strained boron–boron bond.

3.1. Properties of **1** (R = H)

Depicted in Fig. 1, the isolated compound **1a** exhibits *C*_{2v} symmetry with a roof-like shape. According to the MEP and in agreement with what observed for compound **I**, the most negative region in **1** is located above the B–B bond (the ridgepole of the roof), while the positive regions are associated to the hydrogen atoms of the pyrazole rings. The richest electron region between the two borons appears in the ELF analysis represented by a disynaptic basin populated by 2.25 e,



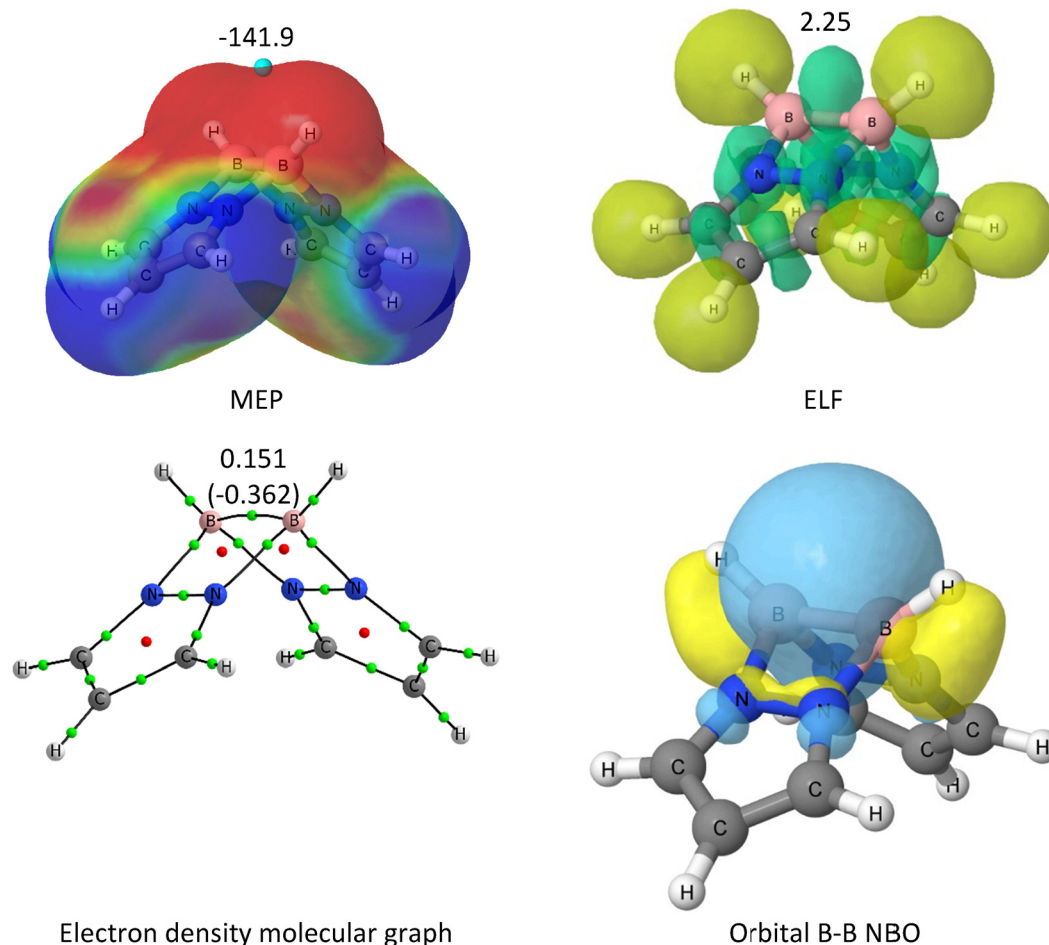


Fig. 1 MEP (0.001 electron density isosurface), ELF (0.80), electron density molecular graph, and natural bond orbital (NBO) of the B–B bond of the parent compound **1a**. In the molecular graph, green and red dots correspond to BCPs and RCPs, respectively. The properties associated to the B–B bonds are indicated in the following units: MEP minimum (kJ mol^{-1}), ELF population of the B–B bond basin (e), ρ_{BCP} and $\nabla^2\rho_{\text{BCP}}$, in parenthesis, (a.u.).

and the same interatomic region viewed through QTAIM analysis presents a B–B BCP with a high electron density ($\rho_{\text{BCP}} = 0.15 \text{ a.u.}$) and negative Laplacian ($\nabla^2\rho_{\text{BCP}} = -0.36 \text{ a.u.}$), which are together the fingerprints of a covalent bond.³⁰ The HOMO corresponds precisely to the same B–B basin region, as shown in the last image of Fig. 1.

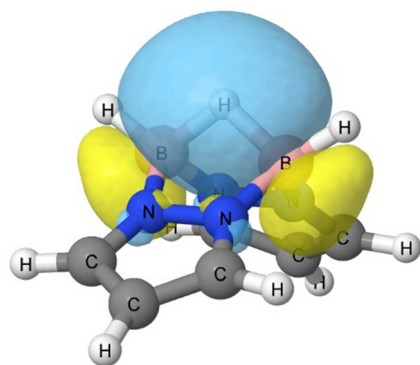


Fig. 2 Orbital associated to the 3c–2e bond of protonated **1a**, labelled as **1a-H⁺**.

It is evident from this set of results that the B–B bond is the most basic region and protonation occurs at this bond, leading to its cleavage. In fact, the calculated proton affinity (PA), 1065 kJ mol^{-1} at the CCSD-F12c computational level (1060 kJ mol^{-1} at M06-2X), is considerably larger than that of guanidine (986 , 979 , and 982 kJ mol^{-1} for the experimental,³¹ M06-2X, and CCSD(t)-F12c values, respectively), and even larger than that of 1,8-bis(dimethylamino)naphthalene ($1028.2 \text{ kJ mol}^{-1}$), which is considered a prototype proton sponge.³¹ The protonated geometry we obtained for **1a** is consistent with the reported experimental protonated structure of **I**, where the additional hydrogen is positioned between the two boron atoms.³² This protonation can be reversed, regenerating the neutral form of molecule **I**.³³ Attempts to protonate **1a** on the B–H bond with dissociation of H_2 , as described for other borane derivatives,³⁴ rearrange to the structure shown in Fig. 2.

The B–B bond cleavage induced by protonation results in an energetic distortion of the structure amounting to 95 kJ mol^{-1} , calculated as the energy difference of the isolated monomer and the protonated structure excluding the additional proton atom. Nevertheless, the substantial stabilization observed upon protonation, as indicated by the value of PA, can be attributed



to the formation of a 3c–2e bond that links the B–H–B group, as depicted in Fig. 2. According to NBO calculations, this 3c–2e bond is richly populated with 1.94 e, with each boron contributing 25% and the bridging hydrogen accounting for 49%, and only 1% from remaining atoms.

At this point, it is interesting to compare neutral **1a** with others mentioned in the Introduction, such as diborane(4). Recalling the disynaptic ELF B–B basin described in Fig. 1, diborane(4) exhibits a monosynaptic basin between boron atoms populated with 2 e,⁶ but instead of a BCP diborane(4) presents a non-nuclear attractor at the same place. The MEP of both structures are pretty similar, but the curvatures of the density are not ((3, –1) vs. (3, +3) critical points). In this sense, borons in diborane(4) are tricoordinated and the monosynaptic basin between them resembles a lone pair, whereas tetracoordinated boron atoms behaviour in **1a** are closer to π donors. Nonetheless, the topology when connecting with HB donors such as HF is completely analogous, as we will see in the following section.

3.2. Hydrogen bonded complexes with **1a**

In view of the basicity of the system, it is not surprising that hydrogen bond donors interact with **1a** through the strained B–B bond. Consequently, complexes with linear proton donors H–X belong as well to the C_{2v} symmetry group, with the H–X molecule pointing towards the middle of the B–B bond. This latter point is a BCP of the B–B bond, as shown by the molecular graph in Fig. 3(a), which is connected through an intermolecular bonding path with the hydrogen atom of HF. This path is similar to that described for HB complexes with π -systems as HCCH and H_2CCH ,³⁵ as in those cases, the description of the electron density topology corresponds to a conflict catastrophe, since a small perturbation of the systems breaks the $H \cdots (B-B)$ BCP bond path. Two examples are included in Fig. S2 (ESI†).

Despite being boron an archetype of electrodeficient atom, the accumulation of electron density due to the formation of the B–B bond results into quite high dissociation energies, as it is important to remind that we are just dealing with neutral complexes in the gas phase. The dissociation values are shown in Table 1, ranging between 34 and 13 kJ mol^{–1} at the CCSD(T)-F12c level, and between 37 and 12 kJ mol^{–1} at the M06-2X level. The similarity between both methods is not surprising, given that the geometries are almost identical (linear regression, “M06-2x distance” = 0.0285 + 0.9874 “CCSD(T)F12c distance”, $R^2 = 0.999$, $n = 5$). The stability order of the complexes follows the sequence HF $\gg$ HCl $>$ HCN $>$ HBr $\gg$ HCCH, in line with the dipole moment and polarizability exhibited by the set of HB donors, what is reflected in a significantly shorter intermolecular distance (2.03 Å) for the strongest complex with respect to the others.

For a reader unfamiliar with the strength expected for gas-phase neutral complexes of this kind, it might seem that the **1a**:HCCH interaction is quite weak. However, it is important to put this result in an appropriate context. A comparison with results reported in the literature at the CCSD(T)-F12c level³⁶ for nonpolar HCCH and cyclopropane as HB acceptor (paradigm of strained cycle), shows that **1a** is a stronger HB acceptor. For instance, the dissociation energy values for the complexes of HF with HCCH, cyclopropane and **1a** are 17.6, 17.9 and 33.9 kJ mol^{–1}, respectively. Regarding the impact of the interaction, it should be mentioned that the donor molecule essentially keeps its geometrical features and suffers very small distortions, whereas the largest deformation energy is only 2.30 kJ mol^{–1} in the HF complex. The complex formation produces the elongation of the B–B bond and a small increment of the roof angle of **1a** (see Table S2 and Fig. S1, ESI†). Excellent linear correlations ($R^2 > 0.97$) are obtained between the three geometrical parameters analysed ($H \cdots CB$ dist., $\Delta B-B$ bond, and Δ roof angle) (Fig. S1, ESI†). Shorter $H \cdots X$ distances

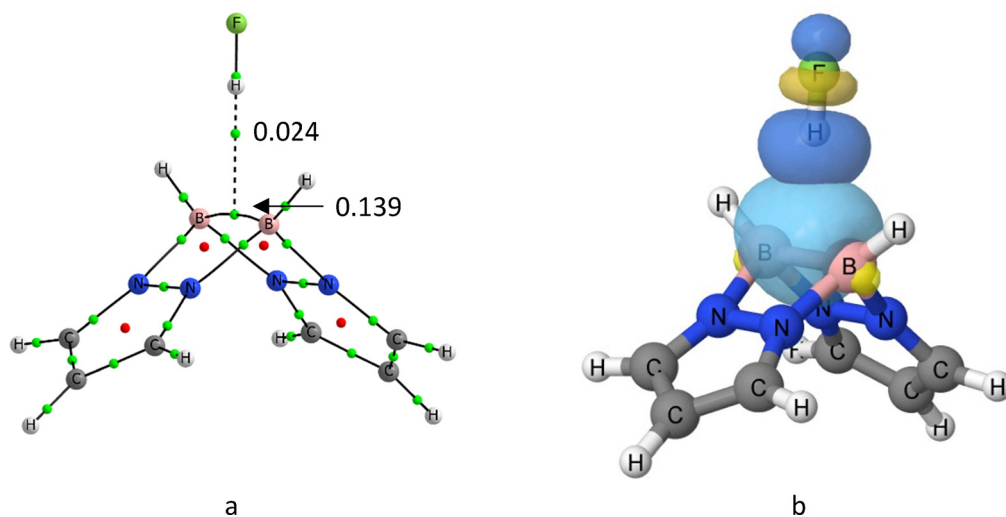


Fig. 3 (a) Molecular graph of **1a**:HF. (b) NBO orbitals involved in the formation of the **1a**:HF complex. Green and red dots of the molecular graph correspond to the location of the BCPs and RCPs, respectively. The molecular graphs of the set of complexes are illustrated in Table S1 (ESI†). The values of the electron density at the intermolecular and the B–B bcp's are indicated in au.



Table 1 Dissociation energy, D_e (kJ mol⁻¹), and intermolecular distance (Å) of the **1a**:HX complexes

Complex	D_e		H...CB* distance ^a	
	CCSD(T)-F12c	M06-2X	CCSD(T)-F12c	M06-2X
1a :HF	33.9	37.2	2.037	2.031
1a :HCl	25.4	24.8	2.192	2.200
1a :HBr	23.7	22.7	2.200	2.206
1a :HCN	24.8	24.6	2.405	2.404
1a :HCCH	13.0	12.0	2.585	2.577

^a CB* is a dummy atom located at the middle of the B-B bond.

correspond to larger increments of the B-B bonds (up to 0.035 Å in the strongest complex, the HF one, along with an increment of 1.6° in the roof angle of **1a**).

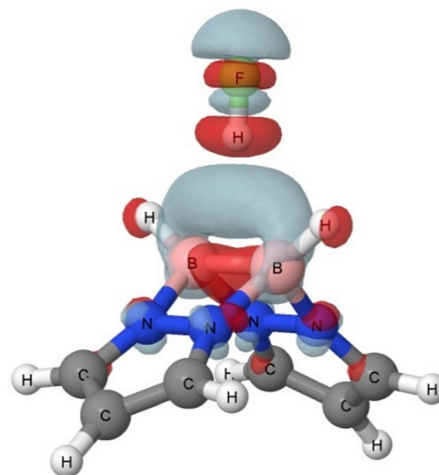
Because of the bond elongation, the electron density of the B-B orbital (Fig. 3(b)) decreases, and accordingly a part of its density is transferred to the antibonding σ^* (HX) molecular orbital, resulting in a net charge transfer. The charge transfer (CT) stabilization is precisely the most important attractive component in the natural energy decomposition analysis (NEDA), shown in Table 2. When looking at the different components of the interaction energy, the CT contribution ranges between the 46–33% of the attractive terms, being larger in the most stable complexes and smaller in the least one [**1a**:HCCH]. The rest of the stabilizing components are mostly electrostatic, but polarization and even more exchange have significant contributions. As expected, the strongest interactions are accompanied by the largest electronic deformation of the monomers, which oppose to binding. The effects of the charge transfer can be quantified as well looking at the increment of the dipole moment of the complex (last column in Table 2), and the molecular regions contributing the most visualized through EDS maps (Fig. 4). For the case illustrated in the latter figure, the dipole moment enhancement $\Delta\mu$ is 1.42 D, which is due to the population of the HB region and the electron gaining in the fluorine lone pairs.

3.3. Enhancing basicity by inductive effect on the boron-boron bond.

Going back to the question of the limits of boron bonds as donors, it is obvious from a chemical point of view that the effects of substitution need to be explored. There is a clear inductive effect evidenced by the PA values of these molecules,

Table 2 Components of the interaction energy (kJ mol⁻¹) obtained with the NEDA method. Column headings: CT = charge transfer, ES = electrostatic, POL = polarization, XC = exchange–correlation, Def = electronic deformation of the interacting monomers **1a** and HX. Last column corresponds to the increment of the dipole moment (D)

Complex	CT	ES	POL	XC	Def 1a	Def HX	$\Delta\mu$ (D)
1a :HF	−65.3	−46.3	−8.1	−22.3	48.0	55.4	1.42
1a :HCl	−57.3	−34.6	−15.1	−24.3	52.6	53.1	1.57
1a :HBr	−63.4	−34.1	−17.3	−26.0	63.4	54.0	1.77
1a :HCN	−34.4	−29.3	−12.9	−16.7	37.8	30.9	1.40
1a :HCCH	−19.1	−14.5	−10.8	−13.6	26.8	19.7	0.84

**Fig. 4** Electron density shift of the **1a**:HF complex. Red and grey regions correspond to poorer and richer electron regions upon complexation, respectively.

which are 1128.4 kJ mol⁻¹ for **1b** and 1064.8 kJ mol⁻¹ for **1c**; in other words, 36 kJ mol⁻¹ larger and 28 smaller, respectively, than the one obtained for the parent compound **1a**, 1092.6 kJ mol⁻¹. It is more than well documented the positive inductive effect that methyl groups have on regular C–C bonds, and the same is observed here on a B–B bond. Regarding the hydrogen-bond acceptor capabilities (see Table 3), and in good agreement with the proton affinities (Table 1), the ranking of dissociation energies is **1b**:HX > **1a**:HX > **1c**:HX. Note that the strongest complex, **1b**:HF, has a dissociation energy of almost 40 kJ mol⁻¹, together with a bonding distance below 2 Å. Even the complex with the weakest HB donor, HCCH, is reinforced, and the corresponding distance decreases by ~0.1 Å.

Some further chemical insight from the D_e values can be obtained using a combination of a Free-Wilson matrix^{37,38} (Table S3, ESI†) and the nucleophilicity and electrophilicity indexes, whose relationship is based on the eqn (1) proposed by Legon:^{39,40}

$$D_e = c \cdot N_b E_{HX} \quad (1)$$

where N_b is the nucleophilicity of the three molecules acting as HB acceptors (**1a**, **1b** and **1c**) and E_{HX} is the electrophilicity of

Table 3 Dissociation energy (kJ mol⁻¹) and intermolecular distances (Å) of the **1b**:HX and **1c**:HX set of complexes, where HX = HF, HCl, HBr, HCN, HCCH

Complex	1b		1c	
	D_e	H...CB	D_e	H...CB
1b :HF	39.8	1.991	20.8	2.168
1b :HCl	28.4	2.151	15.2 ^a	2.332
1b :HBr	27.0	2.108	14.3	2.348
1b :HCN	27.2	2.324	18.7	2.456
1b :HCCH	15.5	2.481	8.0 ^a	2.634

^a These complexes show almost negligible imaginary frequencies (8i cm⁻¹ in **1b**:HCl and 5i and 3i cm⁻¹ in **1b**:HCCH). The C_{2v} geometries of all the systems will be considered for consistency.



Table 4 Fitted values of nucleophilicity N_b and electrophilicity E_{HX} according to eqn (1) by Legon

Molecule	N_b	Molecule	E_{HX}
1a	5.9	HF	6.0
1b	6.7	HCl	4.2
1c	3.7	HBr	3.9
		HCN	4.2
		HCCH	2.2

HX. Parameter c is chosen to have a value of 1.0 kJ mol^{-1} and thus N_b and E_{HX} are dimensionless. Fitting these values, we obtain the results collected in Table 4.

As long as the N_b values of the ligands follow the sequence **1b** > **1a** > **1c**, and E_{HX} reflects the electrophilicity of H in the HB acceptor according to the order HF > HCl ~ HCN > HBr > HCCH, we can say that the results are fully in line with the dissociation energies previously observed, to the point that the fitted values using the derived from the N_b and E_{HX} vs. the calculated D_e energies are linearly correlated with a R^2 value larger 0.98 (see Fig. S3, ESI[†]). We can hypothesize that the inductive effects we generally predict for sigma carbon bonds are expected to be followed here as well, opening a door for tuning donor abilities in a predictable manner. Similar conclusions could be followed by analysing the topology of the electron density of the complexes, explained in detail in Fig. S4 (ESI[†]).

Again, it is useful to remember that derivatives of diborane(4) compounds, such as those synthesized and characterized by Horn and collaborators,¹² can also be tuned by the choice of the substituents, improving the nucleophilicity of the system.

4. Conclusions

Novel structures including a strained B–B bond are proposed in this article, for which we have explored the protonation and hydrogen bond acceptor properties using the CCSD(T)-F12c and DFT (M06-2X) computational methods. These systems exhibit proton affinities greater than those of well-known organic bases, such as guanidines or 1,8-bis(dimethylamino) naphthalene ($1028.2 \text{ kJ mol}^{-1}$), which is considered a prototype proton sponge. Furthermore, their complexes with proton donors are highly stable, as evidenced by the dissociation energy of -34 kJ mol^{-1} for the **1a**:HF complex. We find these results relevant, as we are dealing with electrodeficient atoms behaving as electron donors and proton sponge qualities better than nitrogen donors, which is a result that was not easily predictable. Moreover, the modulation of their properties seems to be affordable. Replacing the hydrogen atoms bonded to the boron atoms in **1a** with methyl groups or fluorine atoms reveals significant inductive effects, reaching PA values higher than 1096 kJ mol^{-1} . We have also observed that the protonation of these systems leads to the cleavage of the B–B bond and the subsequent formation of a 3c–2e B–H–B bond.

As occurs with classical lone-pair donor atoms, the strained B–B bond can form hydrogen bonds, as observed when studying the hydrogen-bonded complexes with a set of representative compounds. The interaction takes place at the centre of the B–B bond, consistent with the position of the MEP minima and the HOMO orbital of the isolated **1a** molecule. The electron density characteristics of these complexes indicate a catastrophic instability, like that observed in π systems. The stability of the complexes has been correlated with the nucleophilicity and electrophilicity indexes using eqn (1) proposed by Legon. The nucleophilicity results for the three derivatives of **1** align with the characteristics of their substituents in typical carbon chemistry.

Conflicts of interest

There are no conflicts of interest to declare.

Data availability

The data supporting this article have been included as part of the ESI.[†]

Acknowledgements

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