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Carboranes as Lewis acids: Tetrel bonding in CB11H11 carbonium ylide

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Abstract: High-level quantum-chemical computations (G4MP2) are carried out in the study of complexes featuring tetrel bonding between the carbon atom in the carbenoid CB₁₁H₁₁ - obtained by hydride removal in the C-H bond of the known *closo*-monocarbadodecaborate anion CB₁₁H₁₂⁽⁻⁾ and acting as Lewis acid (LA) - and Lewis bases (LB) of different type; the electron donor groups in the tetrel bond feature carbon, nitrogen, oxygen, fluorine, silicon, phosphorus, sulphur and chlorine atomic centers in neutral molecules as well as anions H⁽⁻⁾, OH⁽⁻⁾ and F⁽⁻⁾. The empty radial 2*p_r* vacant orbital on the carbon center in CB₁₁H₁₁, which corresponds to the LUMO, acts as a Lewis acid or electron attractor, as shown by the molecular electrostatic potential (MEP) and electron localization funcion (ELF). The thermochemistry and topological analysis of the complexes {CB₁₁H₁₁:LB} are comprehensively analyzed, and classified according to sharing or closed-shell interactions. ELF analysis shows that the tetrel C···X bond ranges from very polarised bonds, as in H₁₁B₁₁C:··O to very weak interactions as in H₁₁B₁₁C···FH and H₁₁B₁₁C···O=C=O.

Keywords: Lewis acid; carborane; carbonium ylide; tetrel bond; quantum chemistry; electron density; ELF

1. Introduction

The very stable B₁₂H₁₂(2-) dianion and its neutral dicarbon counterparts ortho- (1,2-C2B10H12), meta- (1,7-C2B10H12) and para-carborane (1,12-C2B10H12) are icosahedral systems that are closely related to elemental boron. Their isoelectronic analogue, closo-monocarbadodecaborate anion CB11H12(-), first prepared in 1967 [1] and further with other synthesis methods [2,3], is similarly resistant to cage degradation, and many derivatives have been synthesized as described in the literature [4]. The stability and three-dimensional aromaticity of CB11H12(-) has also been explained using quantum-chemical computations [5]. Extraction of hydride H⁽⁻⁾ in the C-H bond from CB₁₁H₁₂(-) leads to a carbocation ylide or carbenoid (1) with a vacant radial $2p_r$ orbital on the cage carbon atom as shown in Figure 1. On the other hand, reaction mechanisms of polyhedral (car)boranes and their derivatives are scarce in the literature and further research is needed in this respect [6-8]. Thus, the permethylated carbenoid analogue CB11Me11 has been postulated as a reaction intermediate during the extraction of the L substituent from L-CB11Me11 carboranes (L = BrCH₂CH₂ or (CF₃)₂CHO) by electrophiles [6,7], further reacting with arenes in the presence of (CF₃)₂CHOH to generate 1-aryl-CB₁₁Me₁₁ products [6-8]. The methyl groups in the permethylated anion CB11Me12(-) have substantial CH3(-) (methide) character according to DFT computations [6-8], and can easily bind to transition metal and main group elements.

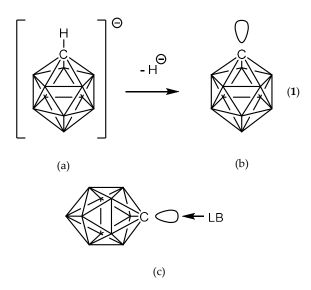


Figure 1. Removal of hydride from the C-H bond in (a) *closo-*monocarbadodecaborate anion $CB_{11}H_{12}^{(\cdot)}$ leads to (b) carbocation ylide or carbenoid $CB_{11}H_{11}$ (1) with a vacant $2p_r$ radial orbital. (c) Complex formation between (1) and a Lewis Base (LB). All vertices correspond to B-H moieties except for the carbon vertex.

On the other hand, in recent years, tetrel bonding - defined as an interaction between any electron donating system (ED) and a group 14 element acting as Lewis acid has called the attention of both experimentalists [9] and theoreticians [10,11]. Here the carbon center in (1) is clearly an acceptor of electrons or Lewis acid, hence we can define a tetrel bonding interaction with an electron donor (ED) or Lewis base, as shown in Figure 1c.

The goal of this work is to study the electronic interaction between the naked carbon vertex in the carbenoid (1) with a series of electron donor molecules and anions, leading to tetrel C-X bonds. The chosen 18 LB systems, including the anions $H^{(\cdot)}$, $F^{(\cdot)}$ and $OH^{(\cdot)}$, are displayed below in Scheme 1 with the corresponding label.

H ⁽⁻⁾ (2)	CH ₂ (3)	CF ₂ (4)	C≡O (5)	N ₂ (6)	NH ₃ (7)
NH=CH ₂ (8)	N≡C-H (9)	OH ⁽⁻⁾ (10)	OH ₂ (11)	O=CH ₂ (12)	O=C=O (13)
F ⁽⁻⁾ (14)	FH (15)	SiH ₂ (16)	PH ₃ (17)	SH ₂ (18)	ClH (19)

Scheme 1. The chosen set of molecules acting as Lewis Base (LB) and forming a tetrel bond with the C atom from carbenoid (1) according to Figure 1c.

2. Computational Methods

Electronic structure quantum-chemical computations were carried out using the G4MP2 model [12,13], which is a fourth-generation method available in the Gaussian16(c) scientific software [14]. This method combines density-functional theory [15,16] and second-order perturbation theory [17] and provides an accurate and economical method for thermochemical predictions. The G4(MP2) model works as follows: The geometries of the molecules are optimized at the B3LYP/6-31G(2df,p) level of theory and then a series of single point energy calculations at higher levels of theory are computed. The zero-point energy, E(ZPE), is based on B3LYP/6-31G(2df,p) frequencies scaled by 0.9854, the same as in G4 theory. The first energy calculation is at the triples-augmented coupled cluster level of theory, CCSD(T), with the 6-31G(d) basis set, i.e., CCSD(T)/6-31G(d). This energy is then modified by a series of energy corrections to obtain a total energy E_0 . For more details on the G4(MP2) method the reader es referred to References [12,13]. In the particular case of the 1:LB complexes, we computed the enthalpy and free energy differences between the complex and separated systems 1 and LB at room temperature as indication of stability of the complex. All complexes included in this work correspond to energy minimum structures, checked with frequency computations. The quantum theory of atoms-in-molecules (QTAIM) [18,19] was used in the topological analysis of the electron density of the 1:LB complexes, with the scientific software AIMAll [20]. This method is based on the analysis of the electron density ρ_r its gradient $\nabla \rho$ and the corresponding Laplacian $\nabla^2 \rho$. For further aspects of this methodology, the reader is referred to the above References [18,19] and Section 3.3.1 below. The electron localisation function (ELF) [21,22] was also used in the topological analysis of the complexes. The ELF is a distribution function which measures the probability of finding an electron near to another electron with an opposite spin, as further described in Section 3.3.2 below.

3. Results

3.1. Geometries of complexes (1:n), n = 2-19

In Figures 2a-b we display the G4MP2 optimized geometries of CB11H12(-) with C5v symmetry - the (1:2) complex - and the carbonium ylide CB11H11 (1), with the different tetrel complexes (1:n), n = 2-19, and structural parameters when necessary, in order to highlight the atomic rearrangements undergone due to the complexation process. The loss of $H^{(\cdot)}$ in the C-H leads also to a structure with C_{5v} symmetry – with a geometrical change which involves a considerable flattenning of the CB₅ pentagonal pyramid with expansion of the corresponding B₅ pentagon, since there is an increase of the B-B bond distance of Δ = +0.022 A. As we proceed down the cage from the top (C atom), the B-B bond differences are +0.022 Å, +0.022 Å and -0.015 Å. Quite noticeable is the shortening of the apical intracage C···B distance ($\Delta \sim -0.3$ Å). The B-H bond distances hardly change at all, with a very slight shortening upon loss of H⁽⁻⁾ with $\Delta = -0.007$ A, -0.008 A and -0.001 A, from top to bottom respectively. If we take the C and B cage nuclei as point charges and define a distorted C_{5v} icosahedron, the corresponding volumes are V(CB₁₁H₁₂(·)) = 12.03 A³ and V(CB₁₁H₁₁) = 11.90 Å³ and therefore there is a shrinkage of the cage by $\Delta V = -0.13$ Å³ (1%). In summary the extraction of hydride in the C-H bond from CB11H12(-) implies a minor change in the cage volume with a flattening of the top CB₅ pentagonal pyramid and minor changes as we proceed down the cage from the top C atom. In Figures 2c-s we display the optimised geometries for the remaining complexes, with the coordinates gathered in the Sup. Info. (SI, Tables S1-S10). The shortest and longest C···X tetrel interactions correspond to the original anion CB11H12(-) or complex (1:2) and the CB11H11···O=C=O tetrel complex (1:13), respectively. In the latter case the attractive interaction is clearly non-covalent in origin with $d(C\cdots O) = 2.693$ Å, as will be discussed later on. We should also emphasize the C···X interaction of the CH₂ and SiH₂ complexes (1:3) and (1:16) respectively, with the LB groups tilted from the C5 axis of rotation. In all other systems, including the CF2 complex (1:4), the C···X bond is aligned with the C_5 axis of rotation.

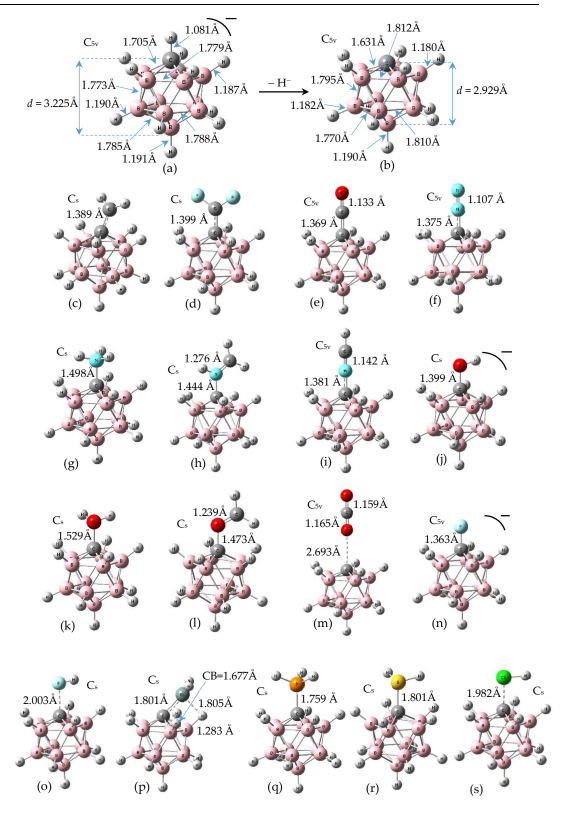


Figure 2. Structures of the G4MP2 optimized greometries for the 18 tetrel complexes (1:n) - with n=2-19 - considered in this work, following Scheme 1: (a) The known anion $CB_{11}H_{12}^{(\cdot)}$ corresponding to complex (1:2), (b) carbenoid $CB_{11}H_{11}$ (1), (c) (1:3), (d) (1:4), (e) (1:5), (f) (1:6), (g) (1:7), (h) (1:8), (i) (1:9), (j) (1:10), (k) (1:11), (l) (1:12), (m) (1:13), (n) (1:14), (o) (1:15), (p) (1:16), (q) (1:17), (r) (1:18) and (s) (1:19).

In Figure 3 we display the $d(C\cdots X)$ distances in the tetrel bonding complexes, ordered from shortest to longest. Clearly, we can classify 5 groups according to the $C\cdots X$ distances, in increasing order: (i) The original anion $CB_{11}H_{12}^{(-)}$ or (1:2) complex, (ii) complexes with $d \sim 1.4$ -1.5 Å including complexes (1: k_2), with $k_2 = (3-12, 14)$, (iii) complexes with $d \sim 1.8$ Å, including complexes (1: k_3), with $k_3 = 16$ -18, (iv) complexes with $d \sim 2.0$ Å, including complexes (1:15) and (1:19), and finally (v) the (1:13) complex with $d \sim 2.7$ Å.

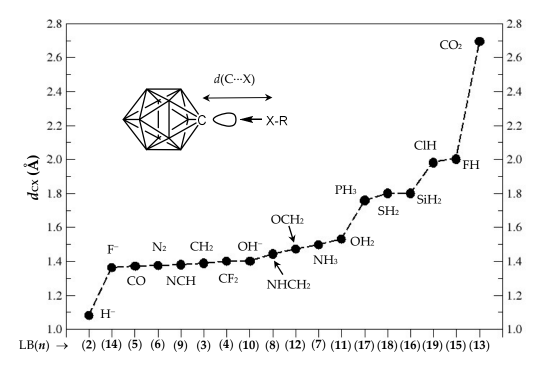


Figure 3. C···X distances d_{CX} (Å) in the tetrel complexes (1:n), n = 2-19, from Figure 2, in increasing order; x axis corresponds to Lewis Base (LB) n and y axis to d_{CX} distances (Å) respectively.

3.2 Thermochemistry of complexes (1:n), n = 2-19

In this subsection we predict the ΔH and ΔG of the tetrel bonding complexes (1:n), n = 2-19. In Table 1 we gather the computed enthalpies and free energies at the G4MP2 level of theory.

Table 1. Enthalpy (Δ H) and free energy (Δ G) of formation for complexes (1:n), n = 2-19, in kJ·mol⁻¹. LB: Lewis base.

	n = 2	3	4	5	6	7	8	9	10
LB	H ⁽⁻⁾	CH_2	CF ₂	C≡O	N_2	NНз	NH=CH2	N≡CH	OH(-)
ΔН	-858.6	-504.9	-326.2	-211.1	-50.5	-268.6	-294.6	-170.6	-659.3
ΔG	-822.3	-457.7	-279.7	-162.9	-2.8	-226.8	-242.7	-120.8	-618.0
	11	12	13	14	15	16	17	18	19
LB	OH ₂	O=CH ₂	O=C=O	F(-)	FH	SiH ₂	PH ₃	SH_2	ClH
ΔΗ	-116.2	-139.8	-14.3	-571.5	-0.7	-418.0	-280.9	-186.1	-47.6
ΔG	-72.1	-87.9	25.7	-529.7	31.3	-366.8	-238.1	-140.9	-8.9

The free energy of formation for complexes (1:*n*) is always negative, except for (1:13) and (1:15) complexes, namely the O=C=O and FH complexes respectively. Small negative values ($|\Delta G| < 10 \text{ kJ} \cdot \text{mol}^{-1}$) are obtained for complexes (1:6) and (1:19) with Lewis bases N₂ and ClH respectively. Therefore, all complexes with negative ΔG should be formed at

room temperature spontaneoulsy, provided an isolated Lewis acid (1) approaches an isolated Lewis base. On the other hand, the enthalpies of formation are negative for all complexes, an indication that the bond energies of a given complex (1:n) have a lower value than the bond energies of separated systems (1) and (n). In order to better visualize the similarities and differences of the thermochemical aspects of complexes (1:n) we plot ΔG and ΔH vs. n in increasing order of each state function.

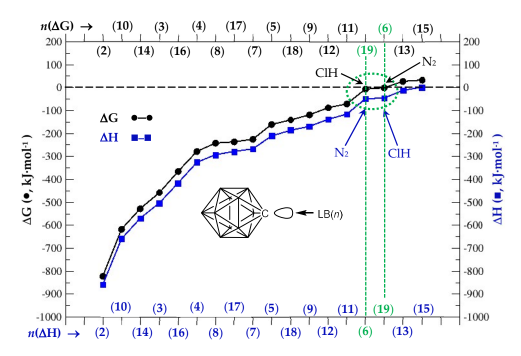


Figure 4. ΔG and ΔH *vs.* Lewis base *n* in respective increasing order, all in kJ·mol⁻¹.

As shown in Figure 4 from left to right in the abscissa, the ΔG (black) and ΔH (blue) in complex formation follow the same order as function of Lewis base number, except for ClH (1:19) and N₂ (1:6) where the order is inverted in the Δ H tendency as compared to Δ G; this is due to the noncovalent and covalent character of each interaction. Hence, the formation of anionic complexes are the most energetic and favourable ones in the order (1:2), (1:10), (1:14) corresponding to Lewis bases $H^{(-)}$, $OH^{(-)}$ and $F^{(-)}$ respectively; then follow complexes (1:3), (1:16) and (1:4) corresponding to Lewis bases CH2, SiH2 and CF2 respectively, namely the carbene series. A plateau with complexes (1:8), (1:17) and (1:7) follows with Lewis bases NH=CH2, PH3 and NH3 respectively. A smaller (positive) slope of $\Delta G/\Delta H$ vs. n appears with complexes (1:5), (1:18), (1:9), (1:12) and (1:11) always in increasing order, which correspond to Lewis bases C≡O, SH2, N≡CH, O=CH2, and OH2 respectively. The weakest bound complexes, with $\Delta G < 0$, correspond to (1:19) and (1:6) with Lewis bases ClH and N₂ respectively. Finally complexes (1:13) and (1:15) with Lewis bases O=C=O and FH respectively show a predicted quantum-chemical value of $\Delta G > 0$ and therefore one should not expect a spontaneous formation of these complexes at room temperature. It is noteworthy to mention the tiny value $\Delta H(1:15) = -0.7 \text{ kJ·mol}^{-1}$ for FH attachment to (1); this number is within the accuracy of the method and therefore a heat of formation for complex (1:15), or the bond energy on both sides of the equation remains unaltered.

In Figure 5 we show for (1), from left to right, the lowest unoccupied molecular orbital (LUMO), the molecular electrostatic potential (MEP) and the electron localization function (ELF). These electronic structure features are computed using the optimized geometry of the system with the G4MP2 method – B3LYP/6-31G(2df,p) model chemistry for structure optimization. As noticed in Figure 5a, the LUMO of the carbonium ylide has considerable wave function amplitude around the C atom – as compared to other regions of the molecule, an indication of the electron acceptor (Lewis acid) nature of the C ylide center; this property is confirmed by the shape of the MEP and the corresponding π -hole just on the top of C ylide center, where negative (surplus) electron density is attracted to. The ELF from Figure 5c shows disynaptic V(B,H) yellow basins corresponding to the the B-H bonds, and the ELF distribution around the CB11 icosahedral cage can be partitioned into green disynaptic and trisynaptic basins as we will describe below in Section 3.3.2.

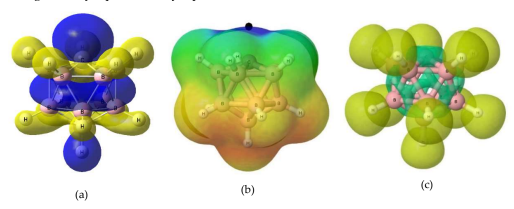


Figure 5. Electronic structure of carbonium ylide (1) CB₁₁H₁₁: (a) LUMO with an isosurface value of the wave-function amplitude $\varphi(LUMO) = \pm 0.02$ au. (b) Molecular electrostatic potential V(r). Red color for V(r) < -0.015 au, Blue color for V(r) > 0.03 au. The black dot indicates the localization of the π -hole (0.061 au), and (c) Electron Localization Function (ELF) with an isosurface of ELF = 0.75. Computations with the G4MP2 level of theory.

The molecular electrostatic potential (MEP) is the potential energy of a proton at a particular location near a molecule. Negative electrostatic potential corresponds to an attraction of the proton by the concentrated electron density in the molecules. The MEP of (1) – Figure 5b – shows that the potential energy of a proton is most positive above the C atom, with a π -hole of +0.061 au, hence a repulsive region for a proton approaching (1), or electron acceptor region. The MEP is smoothly changing from positive to negative values of the potential energy as the proton moves from the C atom down to the B skeleton cage region. A proton would then be attached more favourably to the lower region of the carbonium ylide (1). In other words, Lewis bases, electron donors and nucleophiles should then tend to bind through the C atom of the ylide, hence the study of the tetrel bonding in the complexes (1:n).

3.3.1. Atoms-in-Molecules (AIM) topological analysis of complexes (1:n), n = 2-19

The Quantum Theory of Atoms in Molecules (QTAIM) [18-19] is a useful tool for analyzing the electronic structure of a polyatomic many-electron system, with the electron density $\rho(\mathbf{r})$ as the central function. The topological properties of $\rho(\mathbf{r})$ are analyzed with tha gradient $\vec{\nabla}\rho$ and Laplacian $\nabla^2\rho$, with the eigenvalues of the latter being $(\lambda_1, \lambda_2, \lambda_3)$. The critical points are those with $\vec{\nabla}\rho = \vec{0}$ and a bond critical point (BCP) has $\lambda_3 > 0$ associated with the bond path direction, and $\lambda_1 < 0$, $\lambda_2 < 0$ the two latter associated to two directions where $\nabla^2\rho$ is a maximum; the BCP (-,-,+) appears at the intersection of the bond

path with the interatomic surface S. Other critical points are classified according to the signs of λ_i : Nuclei positions with (-, -, -); ring critical points with (-, +, +); cage critical points with (+, +, +). We should also introduce the local electron kinetic (G > 0), potential (V < 0) and total (H) energy densities, with H = G + V, also useful parameters at the BCP for the description of the type of bonding interaction between atoms in a many-electron system [23]. In the SI (Table S11) we provide the computed values of $\rho(\mathbf{r})$, $\nabla \rho(\mathbf{r})$, $\nabla^2 \rho(\mathbf{r})$, G, V and H for the BCP found between the X atom of the Lewis base in contact with the C ylide center in (1), for all complexes (1:n), n = 2-19.

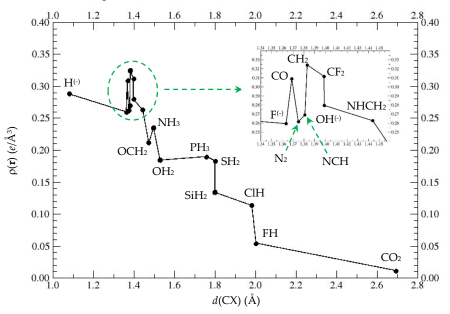


Figure 6a. ρ (**r**) at the BCP of the C···X interaction in complexes (1:*n*) *vs.* d(CX), with n = 2-19. We label each point with the corresponding LB.

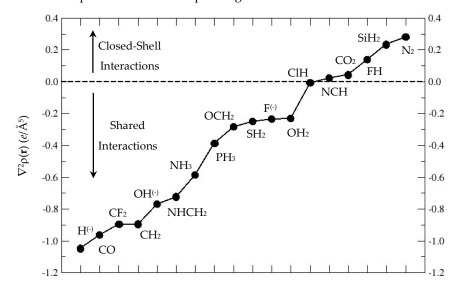


Figure 6b. $\nabla^2 \rho(\mathbf{r})$ at the C···X BCP for complexes (1:*n*), n = 2-19.

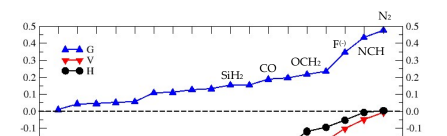


Figure 6c. Plot of G (▲), V (▼) and H (•) vs. Lewis base LB(n) in increasing order of G, V and H respectively, at the Bond Critical Point (BCP) in the C···X interaction of complexes (1:n), n = 2-19, as described in Scheme 1.

In Figure 6a we plot the electron density at the BCP for the C····X interaction $vs.\ d(CX)$ distance. The largest values of ρ_{BCP} correspond to CH₂ ($\rho_{BCP}(CH_2) = 0.32\ e/a0^3$), CF₂, and CO, followed by H⁽⁻⁾, OH⁽⁻⁾, NHCH₂, N₂ and F⁽⁻⁾. Another group follows with lower values, NH₃, OCH₂, PH₃, OH₂, SH₂ and further down SiH₂ and ClH with similar values. Finally, the lowest ρ_{BCP} correspond to FH and CO₂, the latter with $\rho_{BCP}(CO_2) = 0.01\ e/a0^3$. We should emphasize that the ratio $\rho_{BCP,max}(CH_2)/\rho_{BCP,min}(CO_2) = 32$ gives an idea of the topological differences in these BCPs. Given the different type of C···X interactions in the complexes, the $\rho_{BCP}\ vs.\ d(CX)$ can be fit to an approximate negative exponential curve with $\rho_{BCP}\ (dcx) = a + b \cdot \exp(-c \cdot dcx)$, with a = -0.022, b = +3.222 and c = -1.751, and a correlation factor of R² = 0.99 for closed-shell interacting complexes: CO₂, FH, SiH₂, N₂ and NCH. This curve is displayed in the SI file (Figure S1). In general, very good correlations appear if we fix the two interacting atoms, both belonging to the same row of the Periodic Table [24-26].

In order to estimate the type of interaction we need to go beyond the electron density at the BCP and analyze the second derivative, the Laplacian $\nabla^2 \rho$, and the kinetic, potential and total energy, G, V and H respectively, of the BCPs in complexes (1:n). In Figure 6b the Laplacian is plotted vs. LB(n) in increasing order. Clearly, we can distinguish the shared interactions for $\nabla^2 \rho > 0$ in the lower left corner and closed-shell interactions for $\nabla^2 \rho > 0$ in the upper right corner of Figure 6b. The two-electron sharing in complexes with H(-), CO, CF2 and CH2 is large, and it is diminished up to OH2. For the complex with HCl, $\nabla^2 \rho = -0.0067~e/\text{Å}^5$, namely in the limit between sharing and closed-shell interactions. For positive Laplacians, in increasing order, the LB in the (1:n) complexes correspond to: NCH, CO2, FH, SiH2 and N2; in these systems the closed-shell interactions are important.

A further analysis of the BCP in the C···X interactions of the (1:n) complexes can be found in the values of G, V and H - with H = G + V being the total energy - as displayed in Figure 6c. The kinetic energy G is associated with repulsion in the bonding region, and the potential energy V shows the effect of the electric field in the internuclear region. Thus, according to Figure 6c, the G and V profiles are inverted for N₂, NCH, F(·) and OH(·), but due to the nature of different nuclei in the C-X interactions, this is not always the case, as seen when we follow the profiles as function of LB(n).

Thus, H can be used for characterising the covalent character of an attractive interaction betweens atoms [21], and a combination of the Laplacian and H for the characterisation of hydrogen bonds [22].

3.3.2 Electron Localisation Function (ELF) analysis of complexes (1:n), n = 2-19

In order to better understand, from the electronic structure point of view, the tetrel bonding in the (1:n) complexes, we further computed the Electron Localization Function (ELF) [27-28], a measure of the likelihood of finding an electron in the neighborhood space of a reference electron located at a given point and with the same spin; therefore ELF is a measure of the Pauli repulsion or exchange interaction [29-30]. The ELF for the carbonium ylidene CB₁₁H₁₁ (1) is shown in Figure 4c. ELF values ranges from 0 to 1 (normalized and without units). In the SI file we provide the ELF for all complexes (1:n) not shown here (Table S12) and below we have selected four cases with short, medium, long and very long C···X distances, according to Figure 3 above: (i) LB = H(·), complex (1:2), (ii) LB = N₂, complex (1:6), (iii) LB = PH₃, complex (1:17), and (iv) LB = CO₂, complex (1:13). In Table 2 we gather the ELF function for these four complexes.

Table 2. ELF (isovalue 0.83 au) for complexes H₁₁B₁₁C:H^(·) → (1:2), H₁₁B₁₁C:N≡N → (1:6), H₁₁B₁₁C:PH₃ → (1:17), and H₁₁B₁₁C:O=C=O → (1:13).

H(-)	N≡N	РНз	CO ₂
11 C(B) = 2.07 1 C(C) = 2.11 11 V(B,H) = 2.05 10 V(B,B,B) = 0.81 15 V(B,B) = 0.76 5 V(C,B) = 1.01 1 V(C,H) = 2.05	11 C(B) = 2.06 1 C(C) = 2.06 2 C(N) = 2.07 11 V(B,H) = 2.05 1 V(N,N) = 4.01 5 V(B,B) = 0.46 5 V(B,B,B) = 2.43 5 V(B,B,B) = 0.97 5 V(C,B) = 1.02 1 V(C,N2) = 2.62 1 V(N1) = 3.51	11 C(B) = 2.07 1 C(C) = 2.07 1 C(P) = 9.89 11 V(B,H) = 2.05 3 V(P,H) = 2.05 5 V(C,B) = 1.01 1 V(C,P) = 2.21 5 V(B,B) = 0.51 5 V(B,B,B) = 2.44 5 V(B,B,B) = 0.91	11 C(B) = 2.07 2 C(C) = 2.07 2 C(O) = 2.13 11 V(B,H) = 2.03 1 V(C2,O2) = 2.26 1 V(C2,O1) = 2.51 5 V(C,B) = 1.20 15 V(B,B) = 0.54 7 V(B,B,B) = 1.08 3 V(C,B,B) = 1.08 1 V(O2) = 5.64 1 V(O1) = 3.44 1 V(O1) = 1.82

Below each ELF function of a given complex (1:n), we also report the function value and average population for the different types of basins. A threshold of 0.2 electrons is considered as to include or not a basin in a group. For example if a V(B,B) basin has a population of 0.8 electrons, the latter is not included in the group of the V(B,B) basins with a population around 1.5. In bold letters we report the value of the disynaptic basin corresponding to the tetrel C···X interaction.

In Table 2 we use the following notation:

- C : core basin
- V(X): monosynaptic basin, which can be associated to a lone pair
- V(X,X): disynaptic basin
- V(X,X,X): trisynaptic basin
- 5 V(B,B) = 0.42: There are 5 disynaptic basins involving two boron atoms with an average population of 0.42 electrons.

In the ELF representation, the following colors are used:

- Blue : core basin
- Red: monosynaptic basin
- Grey: polysynaptic basin
- Beige: V(X,H) basin
- Atoms: Boron (green), Carbon (black), Hydrogen (silver), Oxygen (red), and Phosphorus (tan).

Thus, according to Table 2, in the complex (1:2), the existing anion CB11H12(-), the 11 core electrons from the boron cage, the 1s2(B) ones, are gathered in the C(B) core basins, which have an average population of 2.07 electrons; the same description applies for the single C atom and the C(C) = 2.11 basin. The 11 B-H bonds on each non-naked B vertex of the icosahedron correspond to the 11 V(B,H) = 2.05, basically a two-electron covalent B-H bond. Then the distribution of the remaining valence cage electrons (corresponding to 2s and 2p electrons from B and C, plus the surplus electron or negative charge of the anion) are distributed in the 10 V(B,B,B) trisynaptic basins with anaverage population of 0.81 electrons, the 15 V(B,B) disynaptic basins with a population of 0.76 electrons, and the 5 V(C,B) disynaptic basins with ~ 1 electron in them. The V(C,H) disynaptic basin, with a population of 2.05, corresponds to the $C \cdot \cdot \cdot X = C \cdot H$ bond in (1:2), a two-center two-electron bond. Addition of the values of all basins leads to the number of electrons in CB11H12(-): 74. We should emphasize that ELF is a function which reports the probability of finding an electron pair with opposite spins in a region of space. Using a certain isovalue, we are able to define regions of space, basins, with a certain probability to find an electron pair. For example, in the plots of the ELF, we used an isovalue of 0.83, in other words, we plot regions of space where we have a high probability to find a pair of electrons. Once the basins are defined, we can integrate the electronic density into those basins, which are the values reported above in Table 2, and correspond to the number of electrons in that basin. In ELF analysis the partition of space is not based on the electron density, as in AIM, but on the ELF probability function.

As reported in Table 2, the C···X interaction in the selected complexes can be described by the presence or absence of V(C,X) valence basins, and its population. For the complexes (1:6) and (1:17), these values are $V_{(1:6)}(C,N) = 2.62$ and $V_{(1:17)}(C,P) = 2.21$ respectively. Therefore, ELF describes the C···X for the N_2 complex as a bond, with a multiplicity close to 1.5, between the C(ylide) and N nuclei, and for the PH₃ complex a C(ylide)P single bond, with additional 0.2 electrons. As regard to the (1:13) complex with CO₂, the ELF does not localize a basin between the C(ylide) and the O=C=O molecule, and thus no bonding is expected neither a complex formation. The lone pairs from N_2 and CO_2 appear as red monosynaptic basins, as displayed in Table 2.

4. Discussion

The presence of a filled or empty lone pair on the C atom in the known anion $CB_{11}H_{12}^{(-)}$, complex (1:2), depens on whether we remove a proton or a hydride from the C-

H bond, leading to a dianion [H₁₁B₁₁C:]⁽²⁻⁾ (**1b**) or a carbonium ylide H₁₁B₁₁C (**1**), respectively, the latter process is shown in Figure 1. The tetrel complexes (**1**:*n*) presented in the previous section show a rich variety of thermochemical and electronic structure features, with tetrel C···X interactions from different nuclei: $X = \{H, C, N, O, F, Si, P, Cl\}$. The C(ylide) center in (**1**) confers to this particular molecule with a Lewis acid (LA) character, hence the tetrel denomination. The strength for electronic attachment in (**1**) is given by the computed free energy of formation (**1**) + (*n*) \rightarrow (**1**:*n*). The strongest complexes correspond to those formed with anions H⁽⁻⁾, OH⁽⁻⁾ and F⁽⁻⁾, and the weakest complexes to those formed with FH, CO₂, N₂ and ClH. The C···X distance varies considerably in all complexes, ranging from 1.081 Å for (**1**:**2**), LB = H⁽⁻⁾, to 2.694 Å for (**1**:**13**), LB = CO₂. The complex strength is not related to the C···X distance, namely, complexes with similar C···X distances may have different free energy of formation; *e.g.* the free energy of formation for complexes (**1**:**14**) LB = F⁽⁻⁾ and (**1**:**6**) LB = N₂ is -530 kJ·mol⁻¹ and -3 kJ·mol⁻¹ respectively, with very similar d(C-X) distances, 1.364 Å and 1.375 Å.

Examples of recent related systems is the 3D analogue of phenyllithium, the lithia-carborane $CB_{11}H_{11}:Li^{(-)}$, studied in solution, as a solid and by quantum-chemical computations [31]. Indeed, $Li^{(-)}$ is a very poor Lewis base, but certainly attaches to (1), as recently shown, and defined as the lithiated dianion $[CB_{11}H_{11}]^{(2)}$. On the other hand, this process can also be seen as a carborane dianion (1b) – a very reactive species – attached to $Li^{(+)}$, a described in Reference [31]. Table 8-1 from the book by Grimes, *Carboranes* [4], reports hundreds of compounds derived from $CB_{11}H_{12}^{(-)}$ and therefore this is a rich field not only from a synthetic point of view but also for studying the electronic structure of tetrel $C\cdots X$ bonds in these compounds, and specially if the isolated tetrel complexes (1:n) could ever be synthesized, taking into acount that this work is purely theoretical with predictive quantum-chemical computations.

The electronic structure of the complexes has been analyzed thoroughly with AIM and ELF methods, showing the C···X sharing and closed-shell interactions in the complexes according to the values of the Laplacian of the electron density. In Table 3, we gather the ELF values for disynaptic basins V(C,X) in the C···X region showing values of ELF: we can find very polarised C-F bonds in (1:14) – only one electron in the C···X region – single C-X bonds for H(-) and NH=CH2, and intermediate cases, such as in complex (1:5) with a 1.5 multiplicity C-C bond for the CO complex. No V(C,X) disynaptic basins are found for CO2 and FH, an indication of the poor electron-donating ability of these Lewis bases (LB), with indeed long d(C-X) distances and positive free energies of formation $\Delta G > 0$, hence confirming the unlikely formation of these two complexes.

Table 3. Population of ELF disynaptic basins V(C, X) in tetrel complexes (1:n), n = 2-19, describing the C···X interaction. LB = Lewis base.

n	LB	V(C, X)	п	LB	V(C, X)
2	$H^{(-)}$	2.05	11	OH_2	1.56
3	CH_2	2.53	12	O=CH ₂	1.65
4	CF_2	2.71	13	O=C=O	
5	C≡O	2.78	14	$\mathbf{F}^{(-)}$	0.99
6	N≡N	2.62	15	FH	
7	NH_3	1.76	16	SiH_2	2.40
8	NH=CH2	2.02	17	PH_3	2.21
9	N≡CH	2.34	18	SH_2	1.85
10	OH(-)	1.32	19	ClH	1.11

According to the Cambridge structural database(CSD) [32], several tetrel complexes derived from (1) have been characterised [33-39], where the C(ylide) center interacts with a nitrogen atom from neutral aminoderivatives, including pyridine. These structures are

shown in the SI file as Table S13. The shortest C(ylide)···N distance, $d(C \cdot \cdot \cdot N) = 1.477 \text{ Å}$, corresponds to the pyridine complex 1-(4-methoxypyridinium)-1-carba-*closo*-dodecaborane [33]. The longest C(ylide)···N distance, $d(C \cdot \cdot \cdot N) = 1.554 \text{ Å}$, corresponds to the complex 12-iodo-1-(4-pentylquinuclidine)-1-carba-*closo*-dodecaborane [34]. There is a tetrel complex of (1) with NH₃, 1-amino-2-fluorocarba-*closo*-dodecaborane [35], where one B-H vertex hydrogen atom on position 2 has been substituted by a fluor atom, with $d(C \cdot \cdot \cdot N) = 1.486 \text{ Å}$. Our (1:7) tetrel complex H₁₁B₁₁C \leftarrow :NH₃ has a predicted $d(C \cdot \cdot \cdot N) = 1.498 \text{ Å}$ with the G4MP2 model, hence there is a good agreement with experiment.

5. Conclusions

The results presented in this work show that by means of quantum-chemical computations we can predict the formation of tetrel complexes between the icosahedral carbonium ylide $CB_{11}H_{11}$ - derived from extraction of $H^{(-)}$ in the known anion $CB_{11}H_{12}^{(-)}$ – and a set of simple molecules and anions. The driving force of formation for these complexes can be accounted for from thermochemical quantum-chemical computations, using statistical mechanics implemented in the scientific software Gaussian16 [14] and we predict that all complexes can be formed with the exception of the FH and CO_2 molecules, with N_2 and CIH complexes with indeed very low, though negative, free energies of formation.

The tetrel C···X interactions in all complexes have been thoroughly studied by means of AIM and ELF methods hence defining the type of bond and interaction, ranging from very polarised bonds, with one electron in the C···X moiety, to intermadiate cases as in the carbenes :CH₂ and :CF₂ and silane :SiH₂, with one and a half electrons in the C···X region.

The existence of known tetrel complexes of the carbonium ylide $CB_{11}H_{11}$ with amino derivatives, including pyridine, opens the door toward further experimental and theoretical studies in the electronic structure of unusual bonds and interactions between C(ylide) centers in carboranes and other atoms.

We hope that the results from this work can be used for the isolation of reactive species, such as the recently found dianion derived from proton extraction in the well known carborane anion $CB_{11}H_{12}^{(-)}$, a key molecule in the description of 3D aromaticity within boron chemistry.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Tables S1-S10. G4MP2 optimised goemetries of complexes (1:*n*). Table S11. AIM data for complexes (1:*n*). Table S12. ELF data for complexes (1:*n*) not displayed in the main text.

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