The Nature of Triel Bonds, a Case of B and Al Centres Bonded with Electron Rich Sites

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Abstract: The second-order Møller–Plesset perturbation theory calculations with the aug-cc-pVTZ basis set were performed on complexes of triel species: BCl$_3$, BH$_3$, AlCl$_3$, and AlH$_3$ acting as Lewis acids through the B or Al centre with Lewis base units: NCH, N$_2$, NH$_3$, and Cl$^-$ anion. These complexes are linked by triel bonds: B/Al···N or B/Al···Cl. The Quantum Theory of ´Atoms in Molecules´ approach, Natural Bond Orbital method, and the decomposition of energy of interaction were applied to characterise the latter links. The majority of complexes are connected through strong interactions possessing features of covalent bonds and characterised by short intermolecular distances, often below 2 Å. The BCl$_3$···N$_2$ complex is linked by a weak interaction corresponding to the B···N distance of ~3 Å. For the BCl$_3$···NCH complex, two configurations corresponding to local energetic minima are observed, one characterised by a short B···N distance and a strong interaction and another one characterised by a longer B···N distance and a weak triel bond. The tetrahedral triel structure is observed for complexes linked by strong triel bonds, while, for complexes connected by weak interactions, the structure is close to the trigonal pyramid, particularly observed for the BCl$_3$···N$_2$ complex.

Keywords: triel bond; boron centre; aluminium centre; tetrahedral structure; quantum theory of atoms in molecules; natural bond orbital method; π-hole bond

1. Introduction

The number of experimental and theoretical studies concerning inter- and intramolecular interactions increased rapidly in recent years [1–7]. The concept of σ-holes and π-holes is one of the most important trials to explain nature of the broad spectrum of interactions [2,3]. The aerogen (group 18), halogen (group 17), chalcogen (group 16), pnicogen (group 15), and tetrel (group 14) atoms [1–7] may act as the Lewis acid centres interacting with electron-rich sites; these interactions taking their names from names of corresponding centres are often classified as σ-hole bonds [2,3]. In the above-mentioned centres, the depletion of the electron charge may be observed on their edges in directions of their bonds with other atoms. These sites are just named as σ-holes; the above-described electron charge outflow is often sufficient to lead to the positive electrostatic potential (EP) at the σ-holes and, thus, to their Lewis acid properties.

It may be confusing that the centres described here play a role of Lewis acids since the majority of them are electronegative atoms. Some of these centres may be characterised by π-holes that are also regions of the depletion of the electron charge and, consequently, often of the positive electrostatic potential. The π-holes occur for planar species or planar molecular fragments in directions perpendicular to the corresponding planes [2,3]. The elements of the 13th group in their numerous compounds are characterised by the occurrence of π-holes. The corresponding interactions named as triel bonds are also classified as π-hole bonds [8–10]. The triel bonds usually designate interactions
between the triel centre characterised by the positive electrostatic potential, i.e., by the Lewis acid properties, and the electron-rich site, lone electron pair, π-electrons, etc. This term covers the broad spectrum of interactions, from very weak to those of a covalent nature, but most of them are quite strong [8–10]. The boron trihydrides and trihalides are examples of simple planar species with the boron centre possessing positive electrostatic potential that may interact with nucleophiles. It was found that the acidity of the boron centre increases in the following order: BF$_3$ < BCl$_3$ < BBr$_3$ < BI$_3$ [11], which seems to be surprising since the electronegativity of the halogen substituent decreases in this order. Such increase of acidic properties was explained by the back bonding effect [12], where it is the greatest for the BF$_3$ species leading to the significant decrease of the positive charge of boron, i.e., to the decrease of its acidic properties. However, other studies showed that this effect is negligible [13,14]. This led to numerous discussions and polemics concerning the nature of the triel centre interactions [15,16]. It seems that the Lewis base units interacting with these boron trihalides also play a crucial role since it was found that the above-mentioned order depends on the nature of the electron-donating centres [17,18].

Another topic related to boron trihalides interacting with Lewis bases concerns the double potential energy minimum for some of such complexes [19–22]. In the case of the BF$_3$···CH$_3$CN complex, two configurations corresponding to the energetic minima were found characterised by N···B distances amounting to 1.818 Å and ~2.3 Å. The corresponding binding energies for these configurations are equal to −7.7 kcal/mol and −8.7 kcal/mol, respectively [20,21]. For the BCl$_3$···CH$_3$CN complex, two local energetic minima were found that are related to configurations with the N···B distances equal to 1.601 Å and 2.687 Å; the binding energies amount to −12.0 and −4.9 kcal/mol, respectively [22]. It seems that it may be a more general problem for boron centres since, in numerous molecular systems, they are obscured by various types of substituents. This may concern other centres since the similar situation of the double potential minimum was observed for complexes of acetonitrile with tetrahalides of elements of the fourth and 14th groups [23]. It was analysed in this study if such double minima occur for the aluminium trichlorides and trihydrides.

The triel bonds discussed in this study were compared with hydrogen bonds recently [24]. It was also discussed that different types of triel bonds are observed, similarly as for the other interactions. This means that the triel centres may interact with single electron-rich sites, as well as with π-electron systems, like acetylene and ethylene and their derivatives. The complexes of simple triel species with benzene were also analysed recently [25]. The intramolecular triel bonds occur in crystal structures. For example, the interactions in hydrides and halides of 1,2-bis(dichloroboryl)benzene and 1,8-bis(dichloroboryl)naphthalene were analysed theoretically, where these systems and similar ones were found in crystal structures; the corresponding interactions may be classified as intramolecular triel bonds or bifurcated triel bonds [26].

The next matter concerns different properties of different triel elements. It was discussed that characteristics of triel bonds depend on the kind of triel centre in contact with the electron-rich site, i.e., boron, aluminium, gallium, or other 13th group element [8]. These interactions are often very strong, and they possess characteristics of covalent bonds. Cambridge Structural Database [27,28] searches were performed recently, and it was found that the boron centre possesses coordination four most often, in ~64% of all cases, while this coordination for the aluminium centre occurs for ~70% of structures [24]. Coordination three is not so common in both cases in crystal structures, with occurrences of 13.9% and 3.3%, respectively [24]. This is because the trivalent triel centres do not obey the octet rule and, consequently, they are very reactive, strongly interacting with Lewis bases to form tetrahedral structures. The aim of this study is to characterise properties of triel bonds where boron and aluminium centres are in contact with electron-rich sites. The possibility of the existence of the double potential energy minima is also analysed here.
2. Results and Discussion

2.1. The Strength of Triel Bonds

The calculations were performed for complexes of the BCl₃, BH₃, AlCl₃, and AlH₃ species acting as the Lewis acid units with the following Lewis bases: NCH, N₂, NH₃, and Cl⁻ anion. Figure 1 and Scheme 1 present a few examples of systems analysed here.

![Figure 1](image1.png)

**Figure 1.** Examples of two complexes analysed in this study: (a) BCl₃···NCH₃; (b) AlH₃···NCH.

![Scheme 1](image2.png)

**Scheme 1.** The definition of the α-angle; examples of three species with the corresponding values of α are presented: (a) BCl₃···NCH (configuration with the longer B···N distance), (b) BCl₃···NCH (configuration with the shorter B···N distance), and (c) BCl₄⁻ species.

Figure 2. presents maps of the electrostatic potential (EP) for the molecular surfaces of Lewis acid units characterised by the electron density of 0.001 au. One can see regions of positive EP at the triel centres that correspond to the π-holes; values of the maximum EP are also presented. These positive EPs at π-holes determine the Lewis acid properties of triel centres that may be connected with electron-rich sites. The EP value is greater for trihydride than for the corresponding trichloride; moreover, EP is greater at the aluminium centre than at the boron centre of the corresponding species. The lowest maximum value of EP for the BCl₃ species may be explained by the influence of chlorine substituents on the boron centre. However, there is no clear dependence of the EP value on the strength of the corresponding triel bond, as results discussed further here show. Thus, this means that not only electrostatic forces determine the strength of these interactions.
where the trivalent centre obeys the octet rule. The interaction energy, \( E_{\text{int}} \), and the binding energy, \( E_{\text{bin}} \), may be understood as "the energetic cost" of the complex formation. There are systems where \( E_{\text{bin}} \) is positive, showing that the formation of the complex is not energetically preferred [32]; this may correspond to an endothermic reaction. This is not a case for complexes analysed here, where both interaction and binding energies are negative for all cases (Table 1).

The energetic characteristics of complexes linked by triel bonds and analysed here are presented in Table 1. The interaction and binding energies, \( E_{\text{int}} \) and \( E_{\text{bin}} \) [29], respectively, are included, as well as the deformation energy, \( E_{\text{def}} \) [30], and the basis set superposition error correction, BSSE [31]. The interaction energy is the difference between the energy of the complex and the sum of energies of interacting units with their geometries taken from the geometry of the complex. One may say that it describes the strength of interaction of the link in complex. In the binding energy, the energies of the interacting units correspond to their geometries optimised separately. In such a way, \( E_{\text{bin}} \) takes into account the deformation of these units resulting from the complexation, \( E_{\text{bin}} = E_{\text{int}} + E_{\text{def}} \). The latter term, the deformation energy, may be understood as "the energetic cost" of the complex formation. There are systems where \( E_{\text{int}} \) is negative, indicating the local stabilising interaction, while \( E_{\text{bin}} \) is positive, showing that the formation of the complex is not energetically preferred [32]; this may correspond to an endothermic reaction. This is not a case for complexes analysed here, where both interaction and binding energies are negative for all cases (Table 1).

There is a sub-group of complexes analysed here, with anions being products of interactions between the trivalent triel species and the chloride anion. The large -\( E_{\text{int}} \) and -\( E_{\text{bin}} \) values indicate that they are stable systems with interactions corresponding to exothermic reactions. Large deformation energies are observed for these anionic systems. In the \( \text{BCl}_4^- \) anion, this energy is close to 40 kcal/mol. One can see that the formation of these anions leads to great deformations of geometries of interacting units where the planar trichlorides and trihydrides are transformed into the tetrahedral structures. For example, the \( \text{BCl}_4^- \) (see Scheme 1) and \( \text{AlCl}_4^- \) anions are characterised by the \( \text{Cl}^-/\text{B}/\text{Al}^-/\text{Cl}^- \) angle of 109.5° as for methane and other ideal tetrahedral species. The corresponding \( \text{H}/\text{B}/\text{Al}^-/\text{Cl}^- \) angle for the \( \text{BH}_3\text{Cl}^- \) and \( \text{AlH}_3\text{Cl}^- \) anions is equal to 106.5° and 107.1°, respectively (Table 2). These anions, being the result of interaction of trivalent B and Al species with chloride, are stable tetravalent systems where the triel centre obeys the octet rule.

**Figure 2.** The electrostatic potential maps for molecular surfaces with the electron density of 0.001 a.u. The following species are presented: \( \text{BCl}_3 \) and \( \text{BH}_3 \) (up), as well as \( \text{ACl}_3 \) and \( \text{AlH}_3 \) (below); the colour scale is fixed separately for each of the species presented, from maximum positive values (blue) to minimum negative ones (red).
The deformation energy for neutral complexes is greater for stronger interactions. For the BCl complex, the E from Molecules 2020 is negligible, equal to 0.02 kcal/mol.

Table 1 shows that other, neutral complexes are characterised by a broad spectrum of interactions, from ~E_{int} amounting to 1.5 kcal/mol for the BCl···N2 complex, to 50.4 kcal/mol for the BCl···NH3 one. The deformation energy for neutral complexes is greater for stronger interactions. For the BCl···NH3 complex, the E_{def} value amounts 23.6 kcal/mol. For the very weak interaction in the BCl···N2 complex, E_{def} is negligible, equal to 0.02 kcal/mol, while, for another weak interaction in the AlH3···N2 complex,

| Complex          | E_{bin} | E_{int} | BSSE  | E_{def} |
|------------------|---------|---------|-------|---------|
| BCl3···NCH        | -3.2    | -3.5    | 0.8   | 0.3     |
| BCl3···NCH       | -3.8    | -23.6   | 2.5   | 19.8    |
| BCl3···N2         | -1.5    | -1.5    | 0.6   | 0.0     |
| BCl3···NH3       | -26.7   | -50.4   | 2.7   | 23.6    |
| BCl4⁻          | -45.6   | -84.9   | 2.8   | 39.2    |
| BH3···NCH        | -18.6   | -30.4   | 1.2   | 11.8    |
| BH3···N2          | -6.1    | -14.2   | 1.3   | 8.1     |
| BH3···NH3        | -30.8   | -44.0   | 1.1   | 13.2    |
| BH3Cl⁻        | -34.1   | -50.9   | 1.7   | 16.8    |
| AlCl3···NCH      | -23.8   | -29.5   | 1.5   | 5.7     |
| AlCl3···N2        | -7.0    | -9.0    | 1.3   | 2.0     |
| AlCl3···NH3      | -38.8   | -45.7   | 1.6   | 6.9     |
| AlCl4⁻           | -76.0   | -98.5   | 1.7   | 22.5    |
| AlH3···NCH        | -17.7   | -20.6   | 0.7   | 2.9     |
| AlH3···N2         | -5.8    | -6.5    | 0.7   | 0.7     |
| AlH3···NH3       | -28.7   | -32.8   | 0.6   | 4.1     |
| AlH3Cl⁻         | -52.2   | -66.6   | 1.1   | 14.4    |

Table 2. The geometrical and the Natural Bond Orbital (NBO) parameters; B/Al···N/Cl is the distance between the triel centre and the centre of the Lewis base unit (in Å), α-angle (in degrees) is defined in Scheme 1, Ch-shift is the electron charge shift from the Lewis base unit to the Lewis acid that results from complexation (au), while B/Al is the charge (in au) of the triel centre in the complex, \(^1\) and Pol-B/Al is the percentage of the electron density of the B/Al···N/Cl bond orbital at the triel centre; a and b superscripts stand for complexes linked by weak and strong triel bonds, respectively.

| Complex          | B/Al···N/Cl | α-angle | Ch-shift | B/Al | Pol-B/Al |
|------------------|-------------|---------|----------|------|----------|
| BCl3···NCH        | 2.817       | 91.7    | -0.012   | 0.344| -        |
| BCl3···NCH       | 1.628       | 104.0   | -0.292   | 0.265| 22.1     |
| BCl3···N2         | 3.083       | 90.4    | -0.005   | 0.320| -        |
| BCl3···NH3       | 1.614       | 105.0   | -0.411   | 0.312| 22.8     |
| BCl4⁻          | 1.862       | 109.5   | -0.690   | 0.238| 30.4     |
| BH3···NCH        | 1.582       | 103.9   | -0.260   | -0.275| 77.5    |
| BH3···N2          | 1.622       | 101.6   | -0.226   | -0.287| 21.1    |
| BH3···NH3        | 1.652       | 104.7   | -0.396   | -0.234| 79.4    |
| BH3Cl⁻         | 1.968       | 106.5   | -0.549   | -0.289| 74.5    |
| AlCl3···NCH      | 2.024       | 100.0   | -0.126   | 1.327| 90.9     |
| AlCl3···N2       | 2.213       | 96.1    | -0.099   | 1.297| 92.5     |
| AlCl3···NH3      | 1.999       | 100.8   | -0.186   | 1.325| 90.4     |
| AlCl4⁻           | 2.164       | 109.5   | -0.440   | 1.240| 83.5     |
| AlH3···NCH       | 2.099       | 97.7    | -0.107   | 0.903| 91.5     |
| AlH3···N2         | 2.290       | 93.8    | -0.114   | 0.916| 93.5     |
| AlH3···NH3       | 2.071       | 99.2    | -0.179   | 0.899| 8.9      |
| AlH3Cl⁻         | 2.255       | 107.1   | -0.376   | 0.822| 15.1     |

\(^1\) Charges of triel centres in isolated Lewis acid units: BCl$_3$ + 0.312 au, BH$_3$ + 0.299 au, AlCl$_3$ + 1.400 au, AlH$_3$ + 1.125 au.
it amounts to 0.7 kcal/mol. The complexes linked by weak interactions are marked in all tables by superscript a, while those classified as connected by strong interactions are marked by superscript b. However, there is no sharp border between strong and weak interactions, and the division proposed here is rather contractual. These relationships between the deformation energy and the parameters describing strength of interaction were observed earlier for complexes linked by triel bonds [10, 24]. The BSSE correction is located in the range 1–2 kcal/mol for the majority of systems considered here; for a few boron complexes, it is greater than 2 kcal/mol, while, for a few aluminium species, it is lower than 1 kcal/mol.

There are two configurations of the BCl$_3$···NCH complex, characterised by the energies of interaction amounting to $-23.6$ kcal/mol and $-3.5$ kcal/mol with the corresponding B···N distances of 1.628 Å and 2.817 Å, respectively (Table 2). However, the binding energies of these two configurations are almost equal since, for that characterised by the shorter B···N distance, the complexation is connected with great changes of geometries of interacting units. The Cl···B···N angle amounts here to 104°, and the deformation energy is equal to 19.8 kcal/mol. For the configuration with the longer B···N distance, these values are equal to 91.7° and 0.3 kcal/mol, respectively.

The angle mentioned above is designated as $\alpha$ hereinafter; it is between the Cl/H···B/Al bond of the Lewis acid unit and the B/Al···N/Cl intermolecular distance (Scheme 1). In the case of strong interactions, complexes are close to the tetrahedron structure, while, in the case of weak interactions, the Lewis acid unit remains planar, or nearly so, and the $\alpha$ angle defined above is equal or close to 90°. For two configurations of the BCl$_3$···NCH complex, one of them is close to the tetrahedral structure and the other one is close to the trigonal pyramid with the Lewis acid unit close to the planarity.

The $\alpha$ angles are collected in Table 2, while the intermolecular distances are also included there, as well as the selected the Natural Bond Orbital (NBO) [33, 34] parameters. If one refers to the consistent van der Waals radii proposed by Truhlar and co-workers (N—1.55 Å, Cl—1.75 Å, B—1.92 Å, Al—1.84 Å) [35], all intermolecular distances are shorter than the corresponding sum of van der Waals radii. However, the term “intermolecular” should be used with caution since, for the tetrahedral anionic species, all links to the triel centre possess characteristics of covalent bonds. The stability of the similar anion, BF$_4^-$, was discussed recently [36]. The number of structures containing the BF$_4^-$ anion amounts to 16,088 according to search performed through the Cambridge Structural Database (CSD updates up to March 2020). The search performed here for the same version of CSD to find BCl$_4^-$ ions shows only 36 crystal structures. However, there are no structures containing BCl$_3$ neutral species. It was pointed out that trigonal boron trihalides occur only in the gas phase [37]; it seems they are very reactive possessing strong Lewis acid properties, and they react with nucleophiles in more condensed phases.

The electron charge shift from the Lewis base to the Lewis acid unit that results from complexation calculated within the NBO approach is included in Table 2. One can see that greater electron charge shifts are observed for stronger interactions. The greatest shifts between $-0.38$ au and $-0.69$ au occur for anionic complexes, as well as for the BCl$_3$···NH$_3$ and BH$_3$···NH$_3$ complexes, while the smallest ones, lower than $-0.01$ au, occur for complexes linked by the weakest interactions, such as the BCl$_3$···N$_2$ complex and the BCl$_3$···NCH configuration characterised by the longer B···N distance. Table 2 presents the atomic charge of the Lewis acid centre, boron, and aluminium. For all complexes analysed, their formation leads to this charge decrease, up to negative values for the boron centre in the BH$_3$ complexes. There are only two exceptions to the above rule; for the weakest interactions in the BCl$_3$···N$_2$ complex and in one of the BCl$_3$···NCH configurations, the complexation leads to the increase of the positive charge of the boron centre. The Lewis acid properties of the triel centres are supported by the positive electrostatic potentials (EPs) at their surfaces that are related to the above-mentioned $\pi$-holes; the positive EPs for simple species of boron and other triel elements were analysed in former studies in detail [8–10, 38].

It is worth recalling that, for the latter two cases of weakest interactions, the smallest $\alpha$-angles occur; thus, the BCl$_3$ part of the complex is close to the planarity. For all remaining complexes, the
NBO approach detects the σ-bond orbital between the boron or aluminium centre and the nitrogen or chlorine Lewis base site. The polarisation of the latter link is shown in Table 2. It is understood as the percentage of the electron density at the triel (B or Al) centre. One can see that there is no clear tendency here. The greatest concentrations of the electron density at the triel centre occur for the AlCl$_3$ complexes, as well as for the AlH$_3$···N$_2$ and AlH$_3$···NCH complexes, above 90% (except of the AlCl$_4^-$ complex where it amounts to 83.5%). In the case of AlH$_3$···NH$_3$ and AlH$_3$Cl$, the polarisation is equal to 8.5% and 15.1%, respectively, which means that the electron density for the Al···N and Al···Cl links, respectively, is accumulated mainly at nitrogen and chlorine centres. The analysis of these intermolecular links that may be classified as triel bonds, based on results collected in Tables 1 and 2, shows that their properties for aluminium and boron centres differ.

### 2.2. Double Minima for Triel–Lewis Base Potential Energy Curves

It was mentioned earlier here that there are examples of complexes of simple boron species that are characterised by two configurations corresponding to energetic minima, i.e., by the double minimum of potential energy [19–22,38]. This also concerns simple moieties of elements of the fourth and 14th groups [23]. In the case of triel species, two configurations were discussed for the BCl$_3$···CH$_3$CN and BF$_3$···CH$_3$CN complexes; however, in the latter case, one of the potential energy minima is not well separated [20–22]. Two configurations were shortly discussed recently for the BCl$_3$···NCH complex, and two local energy minima were also found for the BBr$_3$···NCH and Bi$_3$···NCH complexes [38]. It was concluded that the existence of two configurations in simple triel species is the result of the balance of interaction energy terms, and that Pauli repulsion plays a crucial role [19]. In a more recent study, it was discussed that the double minimum results from the balance of electrostatic interactions between the BX$_3$ (X = Cl, Br, I) and HCN units [38].

Potential energy curves were constructed for all complexes analysed here, and the double minimum was found only in one case, i.e., for the BCl$_3$···NCH complex. This curve and other potential energy curves for the three additional complexes are presented in Figure 3. Because of differences in the scale for each complex considered, the curves for BH$_3$···NCH, AlCl$_3$···NCH, and AlH$_3$···NCH complexes are restricted only to the proximity of energetic minima. Each of the curve is “normalised” separately. This means that, for each complex, the differences between their energies and the energetic minimum are plotted versus the corresponding Al/B···N distances.

![Figure 3](image-url)  
**Figure 3.** The potential energy curves for the following complexes: BCl$_3$···NCH (full circles), BH$_3$···NCH (open circles), AlCl$_3$···NCH (full squares), and AlH$_3$···NCH (open squares).

It seems that the existence of two potential energy minima for the BCl$_3$···NCH complex results from the influence of chlorine substituents that obscure the boron centre. One may consider that the interaction between the BCl$_3$ and NCH units contains the electrostatic interactions between the nitrogen centre with the boron centre and with the chlorine substituents. The first N···B interaction is attractive, while the N···Cl interactions are repulsive. This is the electrostatic balance mentioned above
here. In contrast, the chlorine substituents cannot obscure the aluminium centre that is characterised by a greater volume than the boron one. However, it seems that the transfer from one configuration characterised by the shallow minimum into the configuration corresponding to the deeper one in the BCl$_3$···NCH complex may be easily forced, for example, by crystal structure forces, since the potential barrier height for such a transfer amounts to 1.6 kcal/mol only.

Figure 4 presents the potential energy curve only for the BCl$_3$···NCH complex. However, the potential energy curve in Figure 4 is more flattened than the corresponding one in Figure 3 because of scale reasons. Another curve presented in Figure 4 also presents energies of this complex for various B···N distances; however, the rigid HCN and BCl$_3$ moieties are considered as possessing geometries corresponding to the isolated species in their energetic minima. This means that these geometries do not change for different B···N distances. This curve is normalised according to the global energy minimum of the complex. One can recall that, in the case of the curve with two minima for each B···N distance, the other geometrical parameters are relaxed (optimised). One can see (Figure 4) that, for the rigid interacting units, a monotonic increase of energy is observed with the shortening of the B···N distance. This means that the increase of repulsion between chlorine centres and nitrogen is not compensated for by the boron···nitrogen attraction. In the case of “relaxed” geometries, the $\alpha$-angle increases with the shortening of the B···N distance. The latter results in weaker Cl···N repulsions than for “the rigid case”. These repulsions for relaxed geometries are compensated for by the B···N attraction, resulting in the occurrence of the second deeper potential energy well for the B···N distance equal to 1.628 Å (see Table 2). However, other interaction energy terms should also be taken into account. The Pauli repulsion is weaker for relaxed geometries than for the rigid interacting species, whereas the orbital energy related to the electron charge shifts is also more important for shorter B···N distances.

![Figure 4](image)

**Figure 4.** The potential energy curves for the relaxed BCl$_3$···NCH complex (full circles) and for the BCl$_3$···NCH complex with rigid Lewis acid and Lewis base units (open circles).

### 2.3. Covalency of Triel Bonds

The former results discussed here show that the majority of triel bonds possess characteristics of covalent bonds. This is in agreement with the experimental data since the X-ray and neutron diffraction results concerning crystal structures indicate that coordination four occurs most often for the triel centres [24]; in other words, the trivalent triel centres are very reactive and they interact strongly with nucleophiles.

Quantum Theory of Atoms in Molecules (QTAIM) [39,40] calculations were performed here at the level corresponding to the optimised geometries, MP2/aug-cc-pVTZ - the second-order Möller–Plesset perturbation theory (MP2) calculations with the aug-cc-pVTZ basis set. Table 3 shows the characteristics of Al/B···N/Cl bond critical points (BCPs) for links between the trivalent triel units and the Lewis base...
species. The electron density at the corresponding BCP, $\rho_{BCP}$, is situated between 0.006 au for the BCl$_3$···N$_2$ complex and 0.131 au for the BCl$_3$···NH$_3$ complex. The first value is typical for weak van der Waals interactions, while the latter one is typical for covalent bonds that are characterised by values of $\sim$0.1 au [39,40]. Few complexes analysed in this study possess $\rho_{BCP}$ values outweighing 0.1 au.

Table 3. The Quantum Theory of Atoms in Molecules (QTAIM) parameters (in au), the electron density at the bond critical point (BCP) ($\rho_{BCP}$), the Laplacian of the electron density (V$^2$P$_{BCP}$), and the total electron energy density at BCP (H$_{BCP}$); a and b superscripts stand for complexes linked by weak and strong triel bonds, respectively.

| Complex            | $\rho_{BCP}$ | V$^2$P$_{BCP}$ | H$_{BCP}$ |
|--------------------|--------------|----------------|-----------|
| BCl$_3$···NCH$^a$  | 0.011        | 0.035          | 0.001     |
| BCl$_3$···NCH$^b$  | 0.106        | 0.403          | -0.070    |
| BCl$_3$···N$_2$$^a$ | 0.006        | 0.022          | 0.001     |
| BCl$_3$···NH$_3$$^b$ | 0.131       | 0.256          | -0.111    |
| BCl$_4^-$          | 0.123        | -0.055         | -0.116    |
| BH$_3$···NCH$^b$   | 0.105        | 0.662          | -0.056    |
| BH$_3$···N$_2$$^b$ | 0.088        | 0.652          | -0.038    |
| BH$_3$···NH$_3^b$  | 0.104        | 0.444          | -0.069    |
| BH$_3$Cl$^-$       | 0.084        | 0.132          | -0.065    |
| AlCl$_3$···NCH$^b$ | 0.048        | 0.310          | 0.002     |
| AlCl$_3$···N$_2$$^a$ | 0.028      | 0.164          | 0.003     |
| AlCl$_3$···NH$_3^b$ | 0.060       | 0.340          | -0.005    |
| AlCl$_4^-$$^b$     | 0.063        | 0.292          | -0.011    |
| AlH$_3$···NCH$^b$  | 0.038        | 0.244          | 0.004     |
| AlH$_3$···N$_2$$^a$ | 0.023       | 0.125          | 0.003     |
| AlH$_3$···NH$_3^b$ | 0.048        | 0.274          | 0.000     |
| AlH$_3$Cl$^-$$^b$  | 0.050        | 0.226          | -0.006    |

The typical QTAIM characteristic of the covalent character of the atom–atom link is the negative value of the Laplacian of electron density at the corresponding BCP, V$^2$P$_{BCP}$. This negative value that indicates the concentration of the electron density in the interatomic region is observed only for the BCl$_4^-$ anion. However, it is often assumed that, even for positive V$^2$P$_{BCP}$ values, the negative value of the total electron energy density at the BCP, H$_{BCP}$, confirms the partly covalent character of interaction [41–43]. The majority of the links presented in Table 3 are characterised by the negative H$_{BCP}$ values. The positive H$_{BCP}$ values correspond to the weakest interactions. However, there is no correlation between binding or interaction energy and the QTAIM characteristics: $\rho_{BCP}$, V$^2$P$_{BCP}$, or H$_{BCP}$. For example, the linear correlation coefficient for the dependence between the $\rho_{BCP}$ and interaction energy, E$_{int}$, is equal to 0.517. The $\rho_{BCP}$ parameter is related to the part of the interaction connected with its covalent character and not to the total interaction energy [44]. It seems that, for the part of complexes analysed here, the electrostatic interaction is more important than the term related to covalency—the orbital energy. For example, for the BCl$_3$···NH$_3$ complex mentioned above, the $\rho_{BCP}$ value amounts to 0.131 au, while, for the AlCl$_3$···NH$_3$ complex, it is equal to 0.060 au (Table 3). The interaction and binding energies for these systems are very close to each other (Table 1). However, the electron density at the BCP concerns the covalent character of interaction that is greater for the boron system than for the aluminium one.

The $\alpha$-parameter presented earlier here (Scheme 1) also expresses the covalent character of the interaction since it informs about the transformation from the triel planar trivalent system into the tetrahedral structure. Figure 5 shows the excellent exponential correlation between $\alpha$ and $\rho_{BCP}$. The anionic systems significantly differing from the remaining complexes are excluded from this dependence; they are presented in this figure only for comparison. The results of the decomposition of interaction energies are shown in Table 4. The decomposition was performed for the density functional theory (DFT) calculations. However, there is an excellent agreement between the DFT
and MP2/aug-cc-pVTZ results; for example, a linear correlation between DFT and MP2 interaction energies is observed, $R^2 = 0.981$. Thus, the results of decomposition are well fitted to the MP2 results of calculations discussed here.

![Figure 5](image_url). The exponential relationship between the α-angle (degrees, Scheme 1) and the electron at the B/Al···N BCP (au), $\rho_{BCP}$; the B/Al···Cl contacts of anionic species (open circles) are excluded from this dependence.

**Table 4.** The interaction energy terms (in kcal/mol) for systems analysed here (see Computational Details section for description and designations); a and b superscripts stand for complexes linked by weak and strong triel bonds, respectively.

| Complex       | $\Delta E_{int}$ | $\Delta E_{Pauli}$ | $\Delta E_{elstat}$ | $\Delta E_{orb}$ | $\Delta E_{disp}$ |
|---------------|------------------|--------------------|---------------------|------------------|-------------------|
| BCl$_3$···NCH$^a$ | $-$1.9 | 8.5 | $-$5.8 | $-$2.6 | $-$2.1 |
| BCl$_3$···NCH$^b$ | $-$18.5 | 168.9 | $-$86.9 | $-$98.7 | $-$1.9 |
| BCl$_3$···N$_2$$^a$ | $-$1.0 | 3.7 | $-$1.7 | $-$1.3 | $-$1.7 |
| BCl$_3$···NH$_3$ | $-$43.8 | 202.8 | $-$125.8 | $-$117.7 | $-$3.1 |
| BCl$_3$$^b$ | $-$80.9 | 186.8 | $-$126.6 | $-$140.2 | $-$0.9 |
| BH$_3$···NCH$^b$ | $-$36.7 | 109.9 | $-$59.3 | $-$86.2 | $-$1.1 |
| BH$_3$···N$_2$ | $-$23.6 | 93.6 | $-$42.3 | $-$73.8 | $-$1.1 |
| BH$_3$···NH$_3$ | $-$45.9 | 110.7 | $-$78.1 | $-$76.7 | $-$1.9 |
| BH$_3$Cl$^b$ | $-$53.8 | 88.9 | $-$63.1 | $-$78.8 | $-$0.9 |
| AlCl$_3$···NCH$^b$ | $-$32.7 | 57.4 | $-$47.6 | $-$40.0 | $-$2.6 |
| AlCl$_3$···N$_2$ | $-$10.1 | 35.2 | $-$20.5 | $-$22.4 | $-$2.3 |
| AlCl$_3$···NH$_3$ | $-$43.7 | 82.1 | $-$78.7 | $-$43.6 | $-$3.6 |
| AlCl$_3$$^b$ | $-$95.1 | 93.6 | $-$113.2 | $-$73.8 | $-$1.6 |
| AlH$_3$···NCH$^b$ | $-$21.0 | 41.9 | $-$34.3 | $-$27.2 | $-$1.5 |
| AlH$_3$···N$_2$ | $-$8.5 | 22.3 | $-$13.7 | $-$16.0 | $-$1.2 |
| AlH$_3$···NH$_3$ | $-$32.7 | 59.5 | $-$58.6 | $-$31.2 | $-$2.3 |
| AlH$_3$Cl$^b$ | $-$65.5 | 67.1 | $-$81.2 | $-$50.8 | $-$0.7 |

Let us return to the BCl$_3$···NH$_3$ and AlCl$_3$···NH$_3$ pair of complexes discussed above here. They are characterised by a similar strength of interaction; however, for the former one, the $\rho_{BCP}$ is over two times greater than for the latter complex. A negative $H_{BCP}$ is observed for the former complex and it is also negative for the latter one, but the latter value is very close to zero. This may suggest that, for the boron moiety, the B···N contact is covalent in nature, at least partly, while the Al···N link for the latter complex is electrostatic in nature. This also supports the suggestion of Gillespie and Popelier [44] that electron density at the bond critical point expresses the covalent nature of interaction. Furthermore, for the aluminium complex, the electrostatic interaction energy term is almost two times greater than the orbital energy, which concerns the absolute values (Table 4). In the case of the BCl$_3$···NH$_3$ complex, another situation is observed, where orbital and electrostatic energies are comparable to each other. The results of Table 4 show that there is no specific difference between boron and aluminium complexes or between the triel centres connected with chlorine or hydrogen substituents. One can also see that
the dispersion energy is much less important with respect to stabilising the systems considered than the electrostatic and orbital terms; this is observed for all complexes analysed.

Figure 6 presents linear correlations between the electron density at the B/Al⋯N/Cl BCP, ρ_{BCP}, and two terms of the interaction energy resulting from its partitioning: Pauli repulsion (ΔE_{Pauli}) and orbital energy (ΔE_{orb}) (see Computational Details section). These are good correlations despite them concerning the sample of complexes containing various kinds of interactions. These two terms, ΔE_{Pauli} and ΔE_{orb}, are often related to the covalent character of interaction [43], especially the orbital energy, which is connected with the electron charge shifts resulting from complexation. It was pointed out in several studies that the dominance of the ΔE_{orb} term testifies to the covalent character of interaction [43]. It is worth mentioning that ρ_{BCP} does not correlate with ΔE_{elstat}, with the ΔE_{disp} term, or with the total interaction energy, as mentioned above.

![Figure 6](image)

**Figure 6.** The linear correlations between the electron density at the B/Al⋯N/Cl BCP (au), ρ_{BCP}, and the interaction energy terms, Pauli repulsion (circles) and orbital energy (squares); the anionic species are designated by open circles and open squares.

### 3. Conclusions

The simple complexes of boron and aluminium trichlorides and trihydrides with Lewis base units such as hydrogen cyanide, molecular nitrogen, ammonia, and chloride anion were analysed. These complexes are linked through interactions that were named in recent studies as triel bonds. The majority of interactions analysed here are classified as strong ones that possess characteristics of covalent bonds. For the corresponding complexes of the latter interactions, the complexation leads to the transformation of the planar trigonal trichlorides and trihydrides into structures containing a tetrahedral triel centre (boron or aluminium). In particular, such a situation occurs for complexes of the chloride anion, while perfect tetrahedron structures are observed for BCl₃⁻ and AlCl₄⁻ species.

In a few cases, weak triel bonds are observed, especially for the BCl₃⋯N₂ complex and one of configurations of the BCl₃⋯NCH complex. The interactions linking the AlCl₃⋯N₂ and AlH₃⋯N₂ complexes may also be classified as weak ones. For these weak interactions, the Lewis acid unit, BCl₃, AlCl₃, or AlH₃, retains almost a flat structure in the complex.

Different parameters may describe the character of triel bonds. Among numerous findings presented in this study, it is worth recalling that the angle parameter, α, describes the transformation from the trigonal planar structure into the tetrahedron; this parameter corresponds to the strength of interaction. It is discussed here that the electron density at the BCP corresponds rather to the part of the energy of interaction that is related to the electron charge shifts, and not to the total interaction energy.
4. Materials and Methods

Computational Details

The calculations were carried out with the use of Gaussian16 set of codes [45]. These are the second-order Møller–Plesset perturbation theory (MP2) [46] calculations with the aug-cc-pVTZ basis set [47]. Frequency calculations were performed at the same MP2/aug-cc-pVTZ level to confirm that the optimised structures of complexes correspond to the energetic minima.

The Quantum Theory of Atoms in Molecules (QTAIM) [39,40] was also used to analyse bond critical points (BCPs) of the intermolecular B/Al···N and B/Al···Cl contacts. The AIMAll program [48] was applied to carry out QTAIM calculations, as well as to calculate the electrostatic potentials (EPs). The analysis of the electron charge density shifts, being the result of complexation, was performed with the use of the Natural Bond Orbital (NBO) method [33,34]. The NBO calculations were performed at the BP86-D3/TZ2P level; i.e., with the use of the BP86 functional [49,50] in conjunction with the Grimme dispersion corrections (BP86-D3) [51] and the uncontracted Slater-type orbitals (STOs) as basis functions with triple-ζ quality for all elements [52]. The same BP86-D3/TZ2P level was applied to perform decomposition energy calculations. The NBO [53] and decomposition energy [54,55] calculations were carried out with the use of the ADF2017 program package [55,56] and using geometries of complexes optimised previously at the MP2/aug-cc-pVTZ level. The total interaction energy for the ADF partitioning is composed according to equation given below.

$$\Delta E_{\text{int}} = \Delta E_{\text{elstat}} + \Delta E_{\text{Pauli}} + \Delta E_{\text{orb}} + \Delta E_{\text{disp}}.$$ (1)

The term $\Delta E_{\text{elstat}}$ corresponds to the quasi-classical electrostatic interaction between the unperturbed charge distributions of atoms; it is usually attractive (negative). The Pauli repulsion, $\Delta E_{\text{Pauli}}$, is the energy change associated with the transformation from the superposition of the unperturbed electron densities of the isolated fragments to the wave function that properly obeys the Pauli principle through antisymmetrisation and renormalisation of the product wave function. This repulsive term (positive) comprises the destabilising interactions between electrons of the same spin on either fragment. The orbital interaction, $\Delta E_{\text{orb}}$, corresponds to the charge transfer and polarisation effects.

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References

1. Novoa, J.J. Intermolecular Interactions in Crystals: Fundamentals of Crystal Engineering; The Royal Society of Chemistry: London, UK, 2018.
2. Politzer, P.; Murray, J.S.; Clark, T. Halogen bonding: An electrostatically-driven highly directional noncovalent interaction. Phys. Chem. Chem. Phys. 2010, 12, 7748–7758. [CrossRef] [PubMed]
3. Politzer, P.; Murray, J.S.; Clark, T. Halogen bonding and other $\sigma$-hole interactions: A perspective. Phys. Chem. Chem. Phys. 2013, 15, 11178–11189. [CrossRef] [PubMed]
4. Scheiner, S. Detailed Comparison of the Pnicogen Bond with Chalcogen, Halogen, and Hydrogen Bonds. Int. J. Quantum Chem. 2013, 113, 1609–1620. [CrossRef]
5. Scheiner, S. The Pnicogen Bond: Its Relation to Hydrogen, Halogen, and Other Noncovalent Bonds. Acc. Chem. Res. 2013, 46, 280. [CrossRef] [PubMed]
6. Bauzá, A.; Frontera, A. Aerogen Bonding Interaction: A New Supramolecular Force. *Angew. Chem. Int. Ed.* 2015, 54, 7340–7343. [CrossRef]

7. Grabowski, S.J. Hydrogen bonds, σ-hole and π-hole bonds – mechanisms protecting doublet and octet electron structures. *Phys. Chem. Chem. Phys.* 2017, 19, 29742–29759. [CrossRef]

8. Grabowski, S.J. Boron and other Triel Lewis Acid Centers: From Hypovalency to Hypervalency. *ChemPhysChem* 2014, 15, 2985–2993. [CrossRef]

9. Grabowski, S.J. π-Hole Bonds: Boron and Aluminum Lewis Acid Centers. *ChemPhysChem* 2015, 16, 1470–1479. [CrossRef]

10. Grabowski, S.J. Triel Bonds, π-Hole-π-Electrons Interactions in Complexes of Boron and Aluminium Trihalides and Trihydrides with Acetylene and Ethylene. *Molecules* 2015, 20, 11297–11316. [CrossRef]

11. Beckett, M.A.; Strickland, G.C.; Holland, J.R.; Varma, K.S. A convenient n.m.r. method for the measurement of Lewis acidity at boron centres: correlation of reaction rates of Lewis acid initiated epoxide polymerizations with Lewis acidity. *Polymer* 1996, 37, 4629–4631. [CrossRef]

12. Hiberty, P.C.; Ohanessian, G. Comparison of minimal and extended basis sets in terms of resonant formulas. Application to 1,3 dipoles. *J. Am. Chem. Soc.* 1982, 104, 66–70. [CrossRef]

13. Brinck, T.; Murray, J.S.; Politzer, P. A computational analysis of the bonding in boron trifluoride and boron trichloride and their complexes with ammonia. *Inorg. Chem.* 1993, 32, 2622–2625. [CrossRef]

14. Kutzelnigg, W. Chemical Bonding in Higher Main Group Elements. *Angew. Chem. Int. Ed.* 1984, 23, 272–295. [CrossRef]

15. Rowsell, B.D.; Gillespie, R.J.; Heard, G.L. Ligand Close-Packing and the Lewis Acidity of BF₃ and BCl₃. *Inorg. Chem.* 1999, 38, 4659–4662. [CrossRef] [PubMed]

16. Bessac, F.; Frenking, G. Why Is BCl₃ a Stronger Lewis Acid with Respect to String Bases than BF₃? *Inorg. Chem.* 2003, 42, 7990–7994. [CrossRef] [PubMed]

17. Jonas, V.; Frenking, G.; Reetz, M.T. Comparative Theoretical Study of Lewis Acid-Base Complexes of BH₃, BF₃, BCl₃, AlCl₃ and SO₂. *J. Am. Chem. Soc.* 1994, 116, 8741–8753. [CrossRef]

18. Van der Veken, B.J.; Sluyts, E.J. Reversed Lewis Acidity of Mixed Boron Halides: An Infrared Study of the Van der Waals Complexes of BF₃Cl₂ with CH₃F in Cryosolution. *J. Am. Chem. Soc.* 1997, 119, 11516–11522. [CrossRef]

19. Phillips, J.A. Structural and energetic properties of nitrile–BX₃ complexes: substituent effects and their impact on condensed-phase sensitivity. *Theor. Chem. Acc.* 2017, 136, 16. [CrossRef]

20. Giesen, D.J.; Phillips, J.A. Structure, Bonding, and Vibrational Frequencies of CH₃CN-BF₃: New Insight into Medium Effects and the Dispersion between the Experimental and Theoretical Geometries. *J. Phys. Chem. A* 2003, 107, 4009–4018. [CrossRef] [PubMed]

21. Phillips, J.A.; Cramer, C.J. B-N Distance Potential of CH₃CN-BF₃ Revisited: Resolving the Experiment-Theory Structure Discrepancy and Modeling the Effects of Low-Dielectric Environments. *J. Phys. Chem. B* 2007, 111, 1408–1415. [CrossRef] [PubMed]

22. Wrass, J.P.; Sadowsky, D.; Bloomgren, K.M.; Cramer, C.J.; Phillips, J.A. Quantum chemical and matrix-IR characterization of CH₃CN-BCl₃: a complex with two distinct minima along the B-N bond potential. *Phys. Chem. Chem. Phys.* 2014, 16, 16480–16491. [CrossRef]

23. Helminiaik, H.M.; Knauf, R.R.; Danforth, S.J.; Phillips, J.A. Structural and Energetic Properties of Acetonitrile-Group IV (A & B) Halide Complexes. *J. Phys. Chem. A* 2014, 118, 4266–4277.

24. Grabowski, S.J. Triel bond and coordination of triel centres – Comparison with hydrogen bond interaction. *Coord. Chem. Rev.* 2020, 407, 213171. [CrossRef]

25. Grabowski, S.J. Triel bonds-complexes of boron and aluminum trihalides and trihydrides with benzene. *Struct. Chem.* 2017, 28, 1163–1171. [CrossRef]

26. Grabowski, S.J. Bifurcated Triel Bonds—Hydrides and Halides of 1,2-Bis(Dichloroboryl)Benzene and 1,8-Bis(Dichloroboryl)Naphthalene. *Crystals* 2019, 9, 503. [CrossRef]

27. Groom, C.R.; Bruno, I.J.; Lightfoot, M.P.; Ward, S.C. The Cambridge structural database. *Acta Cryst.* 2016, B72, 171–179.

28. Wong, R.; Allen, F.H.; Willett, P. The scientific impact of the Cambridge Structural Database: A citation-based study. *J. Appl. Cryst.* 2010, 43, 811–824. [CrossRef]

29. Piela, L. *Ideas of Quantum Chemistry*; Elsevier Science Publishers: Amsterdam, The Netherlands, 2007; pp. 684–691.
30. Grabowski, S.J.; Sokalski, W.A. Different types of hydrogen bonds: correlation analysis of interaction energy components. J. Phys. Org. Chem. 2005, 18, 779–784. [CrossRef]
31. Boys, S.F.; Bernardi, F. The calculation of small molecular interactions by the differences of separate total energies. Some procedures with reduced errors. Mol. Phys. 1970, 19, 553–561. [CrossRef]
32. Grabowski, S.J. Lewis Acid Properties of Tetrel Tetrafluorides—The Coincidence of the σ-Hole Concept with the QTAIM Approach. Crystals 2017, 7, 43. [CrossRef]
33. Reed, E.; Curtiss, L.A.; Weinhold, F. Intermolecular Interactions from a Natural Bond Orbital, Donor-Acceptor Viewpoint. Chem. Rev. 1988, 88, 899–926. [CrossRef]
34. Grabowski, S.J. Hydrogen bonds with BF₃ anion as a proton acceptor. Crystals 2020, 10, 460. [CrossRef]
35. Mantina, M.; Chamberlin, A.C.; Valero, R.; Cramer, C.J.; Truhlar, D.G. Consistent van der Waals Radii for the Whole Main Group. J. Phys. Chem. A 2009, 113, 5806–5812. [CrossRef]
36. Grabowski, S.J. Two Faces of Triel Bonds in Boron Trihalide Complexes. J. Comput. Chem. 2018, 39, 472–480. [CrossRef] [PubMed]
37. Gillespie, R.J.; Hargittai, I. The VSEPR Model of Molecular Geometry, Dover Publications, Inc.: Mineola, NY, USA, 2012.
38. Grabowski, S.J. What is the Covalency of Hydrogen Bonding? Chem. Rev. 2011, 11, 2597–2625. [CrossRef] [PubMed]
39. Bader, R.F.W. Atoms in Molecules, A Quantum Theory; Oxford University Press: Oxford, UK, 1990.
40. Matta, C.; Boyd, R.J. (Eds.) Quantum Theory of Atoms in Molecules: Recent Progress in Theory and Application; Wiley-VCH: Weinheim, Germany, 2007.
41. Jenkins, S.; Morrison, I. The chemical character of the intermolecular bonds of seven phases of ice as revealed by AB initio calculation of electron densities. Chem. Phys. Lett. 2000, 317, 97–102. [CrossRef]
42. Arnold, W.D.; Oldfield, E. The Chemical Nature of Hydrogen Bonding in Proteins via NMR: J-Couplings, Chemical Shifts, and AIM Theory. J. Am. Chem. Soc. 2000, 122, 12835–12841. [CrossRef]
43. Grabowski, S.J. Hydrogen bonds with BF₃ as a proton acceptor. Crystals 2020, 10, 460. [CrossRef]
44. Frisch, M.J.; Trucks, G.W.; Schlegel, H.B.; Scuseria, G.E.; Robb, M.A.; Cheeseman, J.R.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G.A.; et al. Gaussian 16, Revision A.03; Gaussian, Inc.: Wallingford, CT, USA, 2016.
45. Möller, C.; Plesset, M.S. Note on an Approximation Treatment for Many-Electron Systems. Phys. Rev. 1934, 46, 618–622. [CrossRef]
46. Woon, D.E.; Dunning, T.H., Jr. Gaussian Basis Sets for Use in Correlated Molecular Calculations. III. The second row atoms, Al-Ar. J. Chem. Phys. 1993, 98, 1358–1371. [CrossRef]
47. Keith, T.A. AIMAll, Version 13.05.06; TK Gristmill Software: Overland Park, KS, USA, 2013.
48. Becke, A.D. Density-functional exchange-energy approximation with correct asymptotic behavior. Phys. Rev. A 1988, 38, 3098–3100. [CrossRef]
49. Perdew, J.P. Density-functional approximation for the correlation energy of the inhomogeneous electron gas. Phys. Rev. B 1986, 33, 8822–8824. [CrossRef] [PubMed]
50. Grimme, S.; Antony, J.; Ehrlich, S.; Krieg, H. A consistent and accurate ab initio parametrization of density functional dispersion correction (DFT-D) for the 94 elements H-Pu. J. Chem. Phys. 2010, 132, 154104. [CrossRef] [PubMed]
51. van Lenthe, E.; Baerends, E.J. Optimized Slater-type basis sets for the elements 1–118. J. Comput. Chem. 2003, 24, 1142–1156. [CrossRef] [PubMed]
52. Glendening, E.D.; Badenhoop, J.K.; Reed, A.E.; Carpenter, J.E.; Bohmann, J.A.; Morales, C.M.; Landis, C.R.; Weinhold, F. NBO, 6.0; Theoretical Chemistry Institute, University of Wisconsin: Madison, WI, USA, 2013; Available online: http://nbo6.chem.wisc.edu/ (accessed on 18 May 2020)(this web-side refers nowadays to NBO 7.0 version, all pre-NBO7 versions are cease).
53. Ziegler, T.; Rauk, A. CO, CS, N₂, PF₃, and CNCH₃ as σ Donors and π Acceptors. A Theoretical Study by the Hartree-Fock-Slater Transition-State Method. Inorg. Chem. 1979, 18, 1755–1759. [CrossRef]
54. Velde, G.T.E.; Bickelhaupt, F.M.; Baerends, E.J.; Guerra, C.F.; van Gisbergen, S.J.A.; Snijders, J.G.; Ziegler, T. Chemistry with ADF. J. Comput. Chem. 2001, 22, 931–967. [CrossRef]
56. ADF2017, SCM, Theoretical Chemistry, Vrije Universiteit, Amsterdam, The Netherlands. Available online: http://www.scm.com (accessed on 18 May 2020).

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