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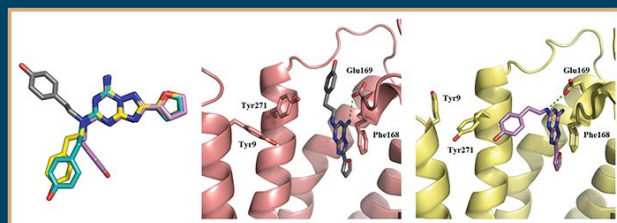
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Lewis acid-base behavior of hypervalent halogen fluorides in gas phase

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Abstract Theoretical studies on Lewis acid-base behavior of hypervalent halogen fluorides, F_3X and F_5X ($X = Cl, Br, I$) have been instrumental in guiding this work. We have also examined whether the hole-lump concept explains the formation of the $F_nX \cdots CO$ complexes. Calculations of proton affinities (PA) and gas-phase basicity (GB) on hypervalent halogen fluorides show that F_3X and F_5X molecules can act as Lewis bases in gas phase. Moreover, theoretical calculations indicate that F_3X and F_5X molecules can act as Lewis acids forming stable complexes with a Lewis base as CO. The quantum theory of atoms in molecules (QTAIM) shows that the electrostatic interaction between the lone pair of the Lewis base (CO) and nucleus of the hypervalent halogen atom (X) plays a key role in stabilizing and determining the optimal geometry

of the $F_nX \cdots CO$ complexes, as in conventional XBs. The localized molecular orbital energy decomposition analysis (LMOEDA) reveals that electrostatic component plays an important role in the stability of the $F_nX \cdots CO$ complexes.

Keywords Hole-lump concept · Laplacian · QTAIM · Halogen bond

Introduction

Various hypervalent halogen fluorides are known. These have been conveniently employed to introduce a few fluorine atoms into the desired place in the substrate [1]. Hypervalent derivatives of the iodine are now routinely used in organic synthesis, probably due to its oxidizing properties, their benign environmental character and commercial availability [2]. Akiba has summarized some general aspects about structure and bonding of hypervalent halogen compounds [3].

Recently, the geometric features of some interactions given by di-, tri-, and tetravalent iodine atoms have been discussed [4]. According to Catalano et al. the fingerprint of XBs is the directionality of the attractive interaction with Lewis base because of anisotropic distribution of the electron charge density around the halogen-atom bridge (Lewis acid).

Moreover, it is well known that monovalent halogen atoms can function as Lewis bases [5–8] (equatorial region). As an example, a search in the Cambridge Structural Database (CSD) for $C-X \cdots M^+$ interactions (with $M^+ = Li^+, Na^+, K^+, Rb^+, Cs^+$ and $X = Cl, Br, I$) reveals that the median value of the $C-X \cdots M^+$ angle is 103.08° . That is to say, cations enter the most negative region of the halogen atom (equatorial region to the $C-X$ bond).⁴ In addition, a monovalent halogen atom can also act as Lewis acid and form attractive interactions with Lewis bases [8–11], these are called halogen bond (XB). All

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these interactions are understood when the anisotropic distribution of the electron density of the monovalent halogen atoms is taken into account.

Landrun et al. have reported an analysis of intermolecular interactions between hypervalent molecules Ph_2IX and F_3X ($\text{X} = \text{Cl}, \text{Br}, \text{I}$) dimers, using a combination of Density Functional Theory (DFT) calculations and qualitative arguments [12]. These authors concluded that the secondary bonding between these species can be understood using the language of donor-acceptor interactions. There is also a strong electrostatic contribution to the secondary bonding. The calculated strengths of these halogen-halogen secondary interactions are all less than 10 kcal mol^{-1} .

Wang has found in the CSD that hypervalent halogen centers behave as acceptors of electron density (Lewis acids) [13]. In addition, Wang showed, through accurate computational results, that the halogen bond involving hypervalent halogen may be weaker than the corresponding halogen bond involving monovalent halogen even in the case that the hypervalent halogen is more positively charged than the monovalent halogen. According to this author the AIM analysis shows that there is no difference between the halogen bond involving hypervalent halogen and the halogen bond involving monovalent halogen [13]. We must make it clear that in any of these research papers, any halogenated compounds having a pentavalent halogen centers were considered.

Grabowski [14] has studied the acidic characteristics of some hypervalent halogen fluorides. In this work, he explains the geometry of the complexes $\text{F}_n\text{Br}\cdots\text{B}$ ($n = 3, 5$ and B : Lewis base) by means of the location of the positive maxima of the electrostatic potential on the bromine atom surface. Through this analysis, this author concluded that a hypervalent bromine atom should always act as the Lewis acid center, while a monovalent halogen atom may act as the Lewis acid and a Lewis base simultaneously. Grabowski also showed that the location of the positive maxima of the molecular electrostatic potential on the surface of both monovalent and polyvalent halogen compounds, predicts the geometry of the studied complexes [14].

Moreover, the Valence Shell Electron Pair Repulsion (VSEPR) theory [15–17] predicts the presence of one and two lone pairs in hypervalent halogen fluorides F_5X and F_3X ($\text{X} = \text{Cl}, \text{Br}, \text{I}$) respectively. It seems, on the one hand, as previously stated, that F_3X compounds can act as Lewis base [12], but: Can the F_5X compounds act as Lewis base?

On the other hand, the positive σ -hole concept explains many of the characteristics of the conventional XBs [9–11]. According to this concept established by Politzer et al., when a monovalent halogen atom forms a covalent bond, some of its electronic charge is transferred towards the bond region, causing the electronic charge to be diminished in its outer region (along the extension of the covalent bond) but increased in its equatorial region [9–11]. That is to say, the positive region on

X corresponds to the electronically-depleted outer lobe of the half-filled p-type orbital of X that is involved in forming the R-X covalent bond [10, 11, 18]. In the hypervalent halogen fluorides F_5X , as it is mentioned above, there is a lone pair on its outermost portions, centered along the extension of the $\text{F}_e\text{-X}$ bond. Therefore, we can ask ourselves: rigorously, does the positive σ -hole concept explain the formation of the complexes $\text{F}_5\text{X}\cdots\text{B}$ ($\text{X} = \text{Cl}, \text{Br}, \text{I}$ and B : Lewis base)? or is it only valid for XBs that involve monovalent halogens? Is the nature of these $\text{X}\cdots\text{B}$ interactions the same as in the conventional XBs?

Recently, the characteristics of the conventional XBs through Quantum Theory of Atoms in Molecules (QTAIM) have been explained by means of the hole-lump concept [7, 19–23]. It seems that this concept can explain those interactions halogen bonding in which the positive σ -hole concept cannot do it [19, 20]. According to our interpretation of the hole-lump concept the formation of a XB is mainly due to the electrostatic interaction between the charge density provided by the Lewis base and the nucleus of the halogen-atom bridge (Lewis acid) [7, 20, 22, 23].

In this work we firstly investigate the Lewis acid-base behavior of F_3X and F_5X ($\text{X} = \text{Cl}, \text{Br}, \text{I}$) molecules. Secondly, we investigate whether the hole-lump concept explains the formation of the $\text{F}_5\text{X}\cdots\text{CO}$ complexes.

Computational details

Monomers and complexes were optimized without any constraint at the MP2/6–611++G(2d,2p) level of theory using the Gaussian 09 suite of programs [24]. All stationary points were confirmed to be true minima by the absence of any imaginary frequencies. For the study of Lewis base behavior of F_3X and F_5X molecules, we have calculated the proton affinities (PA) and gas-phase basicity (GB). These calculations were performed with G4 [25] method, and we also performed calculations at the MP2/6–311++G(2d,2p) level of theory. The study of Lewis acid behavior of F_3X and F_5X molecules was performed in $\text{F}_n\text{X}\cdots\text{CO}$ ($n = 1, 3, 5$ and $\text{X} = \text{Cl}, \text{Br}, \text{I}$) complexes. The binding energies of these complexes were obtained using the supermolecular approach, and the basis set superposition error (BSSE) was corrected by the counterpoise procedure of Boys and Bernardi [26]. The QTAIM analysis was performed with the AIMAll [27] software.

The energies of interaction have been decomposed following the localized molecular orbital energy decomposition analysis (LMOEDA) [28] formalism, according to the equation below:

$$E_{\text{int}} = E_{\text{elect}} + E_{\text{ex-rep}} + E_{\text{pol}} + E_{\text{disp}}$$

where E_{elect} is the electrostatic component, $E_{\text{ex-rep}}$ is the exchange-repulsion component resulting from the Pauli

exclusion principle, and E_{pol} and E_{disp} correspond to polarization and dispersion terms, respectively. These calculations have been carried out with the GAMESS program (version 2013-R1) [29] at the same calculation level as the optimization.

Results and discussion

Monomers

The $L(\mathbf{r}) = -\nabla^2\rho(\mathbf{r})$ function is of particular interest to the present study due to the fact that its topology shows the regions of the space where the electron density is locally concentrated [basic region $-L(\mathbf{r}) > 0$] or depleted [acidic region $-L(\mathbf{r}) < 0$] [30, 31]. The valence shell of an atom is divided into an inner region where $L(\mathbf{r}) > 0$ and an outer one where $L(\mathbf{r}) < 0$. We will use this function to locate the preferential acidic and basic sites of the interhalogen compounds studied here. According to Bader et al. the topology of $L(\mathbf{r})$ function has been shown to provide information about the spatial localization of electronic charge. The topology of this function provides a physical basis for the VSEPR theory [15–17]. The $L(\mathbf{r})$ function exhibits maxima which indicate the presence of localized concentrations of electronic charge (lump) and shows the regions of the charge depletion (hole), both in the valence shell of an atom. Figure 1 shows at the envelope graph at $L(\mathbf{r}) = 1.0$ au of the F_3Br and F_5Br molecules. The regions of the valence shell of the bromine atom in which there is a higher probability of finding an opposite spin electron pair (lone pairs of the Lewis model) are indicated (lump). In Fig. 1, regions where there is depletion of the electron charge density (hole) are observed. In other words, the $L(\mathbf{r})$ function reveals that hypervalent halogen fluorides can act as Lewis acids or bases. In fig. S1 of the supporting information molecular graph, the critical points (3,-3) (nonbonded maxima) of the $L(\mathbf{r})$ function and the bond angles, $F_e\text{-X-CP}$ [CP: critical point (3,-3) of the $L(\mathbf{r})$ function] are reported.

These results are in partial agreement with those reported by Grabowski [14] in which it is stated that “For BrF_3 and

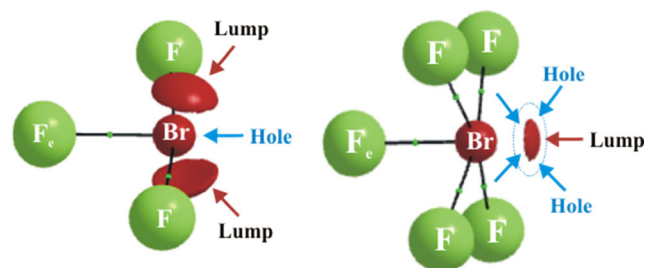


Fig. 1 Envelopes at $L(\mathbf{r}) = 1.0$ au., of F_3Br and F_5Br compounds. Blue arrows indicate electronic depletion regions (Lewis acidic sites) and purple arrows indicate electronic concentration (lone pairs of the Lewis model)

BrF_5 , as well as for their analogs analyzed here, the positive electrostatic potential is observed for the whole hemispheres of bromine centers. It may mean that multivalent bromine atom should always act as the Lewis acidic center. A distinct situation is usually observed for the monovalent halogen atom which possesses a dual character since it may act as the Lewis acid and a Lewis base simultaneously”. However, as discussed below, hypervalent halogen fluorides can act as Lewis bases.

Lewis base behavior

Regularly, the gas-phase basicity (GB) and the proton affinity (PA) are used to characterize the ability of a molecule to accept a proton when in the gas phase. The GB phase basicity is the negative of the free energy change associated with the reaction. The most frequently used index, the PA, is the negative of the enthalpy change at standard conditions. Computational approaches can provide reliable values for GB and PA, which is important since they are hard to determine experimentally. In this work we have investigated these magnitudes and results have been gathered in Table 1.

G4 estimates of CIF molecule are in very good agreement with the existing data of this molecule [32], while MP2 calculation is slightly lower.

According to the values of PA and GB calculated at MP2 level, the ability of these molecules to accept a proton when in the gas phase decreases in the order $\text{FX} > \text{F}_3\text{X} > \text{F}_5\text{X}$ (for the same X) and increases in the order $\text{F}_n\text{Cl} < \text{F}_n\text{Br} < \text{F}_n\text{I}$ (for $n = 1, 3$). These results are according with the strong electron-withdrawing effect of the fluorine atom and with the polarizability of the X halogen atoms respectively. That is to say, when the electron-withdrawing effect on X atom increases its ability to accept a proton decreases, and the higher the polarizability of X atom, the greater its ability to

Table 1 Calculated proton affinity (PA) and gas-phase basicity (GB) at $T = 298.15$ K on X atom (with X = Cl, Br, I)^a

Molecule	PA (kJ mol ⁻¹)		GB (kJ mol ⁻¹)	
	G4	MP2	G4	MP2
FCI	505.1	491.4	479.1	465.4
F ₃ Cl	403.7	368.3	370.9	335.2
F ₅ Cl	406.7	348.4	371.0	311.7
FBr	532.2	525.0	506.5	499.2
F ₃ Br	430.7	411.4	398.1	378.4
F ₅ Br	349.3	312.5	314.2	277.2
FI		576.3		550.9
F ₃ I		499.7		467.1
F ₅ I		345.4		310.6

^a G4 is not implemented for molecule containing iodine atoms

accept a proton. However, the same trend is not observed at G4 level.

In general, values PA and GB are relatively low compared with ammonium (PA = 853.6 kJ mol⁻¹ and GB = 819.0 kJ mol⁻¹). However, they are on the same order of magnitude of that CO (PA = 426.3 kJ mol⁻¹ and GB = 402.2 kJ mol⁻¹) and HF (PA = 484.0 kJ mol⁻¹ and GB = 456.7 kJ mol⁻¹), and they are higher than F₂ molecule (PA = 332.0 kJ mol⁻¹ and GB = 305.5 kJ mol⁻¹) [33].

Interatomic distances and local topological parameters can be interpreted as a measure of the strength of the chemical bonds. Table 2 shows these parameters obtained at the MP2/6-311++G(2d,2p) level of theory. It can be observed that the interatomic distances X–H(+) are relatively short, $d[\text{Cl–H}(+)] \approx 1.31 \text{ \AA}$, $d[\text{Br–H}(+)] \approx 1.45 \text{ \AA}$ and $d[\text{I–H}(+)] \approx 1.62 \text{ \AA}$. But these distances are longer than the respective hydrogen halides [$d(\text{Cl–H}) = 1.275 \text{ \AA}$, $d(\text{Br–H}) = 1.414 \text{ \AA}$ and $d(\text{I–H}) = 1.609 \text{ \AA}$ respectively]. Therefore, we can establish that the X–H(+) bonds in these species are weaker than the respective hydrogen halides.

The equilibrium angles, $F_e\text{–X–H}$ (for the some X), in FX–H(+), F₃X–H(+), F₅X–H(+) species, are near to 95°, 100° and 180° respectively. These values agree with the prediction made by the $L(\mathbf{r})$ topology (see fig. S1 of the supporting information).

It is well known that local topological parameters of the AIM theory allows to characterize the interatomic interactions [30, 31, 34]. That is to say, the values of $\rho(\mathbf{r}_b)$, $\nabla^2\rho(\mathbf{r}_b)$ and $H(\mathbf{r}_b)$ reveal the nature of the interactions. When $\rho(\mathbf{r}_b)$ is relatively large, $\nabla^2\rho(\mathbf{r}_b) < 0$ and $H(\mathbf{r}_b) < 0$ then we have shared interactions (covalent bond), while when $\rho(\mathbf{r}_b)$ is relatively lower, $\nabla^2\rho(\mathbf{r}_b) > 0$ and $H(\mathbf{r}_b) > 0$ then we have closed-shell interactions. Calculated local topological properties at the X–H(+) BCPs, shown in Table 2, present values typical of shared interactions. It is observed that, for the same pair of interacting

Table 2 Selected geometric parameters and selected local topological parameters

Species	$d(\text{X–H})$	$\angle (F_e\text{–X–H})$	Topological parameters		
			$\rho(\mathbf{r}_b)$	$\nabla^2\rho(\mathbf{r}_b)$	$H(\mathbf{r}_b)$
FCl–H(+)	1.310	96.61	0.2395	–0.7883	–0.2289
F ₃ Cl–H(+)	1.317	104.14	0.2471	–0.8932	–0.2487
F ₅ Cl–H(+)	1.311	179.93	0.2635	–1.0018	–0.2756
FBr–H(+)	1.444	94.92	0.1996	–0.4929	–0.1613
F ₃ Br–H(+)	1.452	101.77	0.2078	–0.5630	–0.1713
F ₅ Br–H(+)	1.460	179.97	0.2094	–0.5845	–0.1742
FI–H(+)	1.621	93.90	0.1589	–0.1629	–0.1135
F ₃ I–H(+)	1.628	100.48	0.1664	–0.2250	–0.1227
F ₅ I–H(+)	1.637	179.98	0.1678	–0.2624	–0.1257

Distances in angstrom, angles in degrees and topological parameters in atomic units

atoms, the magnitudes $\rho(\mathbf{r}_b)$, $\nabla^2\rho(\mathbf{r}_b)$ and $H(\mathbf{r}_b)$ increase in the order FX–H(+) < F₃X–H(+) < F₅X–H(+). That is to say, the covalent character of the interactions X–H(+) increases in this order.

It is important to note that geometric analysis gives different results to local topological parameters of charge density at the intermolecular BCP.

Lewis acidic behavior

Table 3 reports the values of the main parameters that describe the geometry of the studied systems. It can be seen that in all the cases, X⋯C intermolecular distances are substantially shorter than the sum of the van der Waals radii [35] of the X and C atoms. This is due to the mutual penetration of X and C atoms electronic densities. In addition, X⋯C intermolecular distances for the same X halogen decrease with the increase of the binding energy. The strength of the interactions F_nX⋯CO ($n = 1, 3, 5$ and X = Cl, Br, I), for the same n, varies with the X halogen, increasing in the order F_nCl⋯CO < F_nBr⋯CO < F_nI⋯CO. These remarks agree with the polarizability of the halogen atoms, which increases in the order Cl < Br < I.

The strength of the XBs depends on the electron-withdrawing ability of the group to which the halogen is attached. If the substituents of hypervalent halogen are strong electron-withdrawing groups such as fluorine atoms, the hypervalent halogen must be more positively charged than the corresponding monovalent halogen. Consequently, it is reasonable to think that the strength of hypervalent halogen-bonded interactions increases with the number of F atoms added. However, the findings of the current study do not support this idea. In Table 3 it is observed that the strength of the

Table 3 Selected geometric parameters and binding energies

Complexes	Structural parameters				ΔE_{BSSE}
	$d(\text{X}\cdots\text{C})$	$d(\text{F}_e\text{–X})_{\text{Comp}}$	$\Delta d(\text{F}_e\text{–X})$	$\angle F_e\text{–X}\cdots\text{C}$	
FCl⋯CO	2.660	1.680	0.015	179.43	–10.4
F ₃ Cl⋯CO	2.918	1.634	0.003	179.88	–10.2
F ₅ Cl⋯CO	3.089	1.622	–0.004	167.26	–7.6
FBr⋯CO	2.561	1.810	0.024	179.96	–16.8
F ₃ Br⋯CO	2.967	1.750	0.004	179.76	–13.0
F ₅ Br⋯CO	3.211	1.726	–0.001	155.17	–9.1
FI⋯CO	2.456	1.954	0.035	179.96	–24.6
F ₃ I⋯CO	3.101	1.873	0.005	179.87	–13.6
F ₅ I⋯CO	3.345	1.829	0.003	142.55	–9.4

Sum of the van der Waals radii: $r(\text{Cl}) + r(\text{C}) = 3.45$; $r(\text{Br}) + r(\text{C}) = 3.55$; $r(\text{I}) + r(\text{C}) = 3.74$. $d(\text{X}\cdots\text{C})$: intermolecular distance. $d(\text{F}_e\text{–X})_{\text{Comp}}$: F_e–X bond length in complex. $\Delta d(\text{F}_e\text{–X})$: is the change of F_e–X bond length upon complex formation. ΔE_{BSSE} : binding energy corrected by BSSE. Distances in angstrom, angles in degrees and energies in kilojoule per mole

interactions, for the same X, increases in the order $F_5X \cdots CO < F_3X \cdots CO < FX \cdots CO$.

Moreover, it is well established that the bond angle in the conventional XBs is always close to 180° , in the $FX \cdots CO$ and $F_3X \cdots CO$ complexes this is true. But, in the $F_5X \cdots CO$ complexes, the bond angles $F_c-X \cdots C$ are less than 180° . In addition, it is observed that these bond angles decrease in the order $F_5Cl \cdots CO > F_5Br \cdots CO > F_5I \cdots CO$, that is to say, in order inverse to the increase of the strength of the $F_5X \cdots CO$ interactions.

QTAIM analysis is a powerful tool to investigate the electronic properties of the molecular system. In the present study, this theory was systematically applied to obtain a deep insight into different factors defining these interactions. Furthermore, this methodology has been successfully applied in the properties study of a variety of conventional XBs [7, 19–23].

Table 4 reports the selected topological properties of the complexes studied here. The topological analysis of the electron density shows the presence of a BCP and an atomic interaction line of maximum electron density connecting the bridge atom (X) of the Lewis acid F_nX with the carbon atom of the CO molecule (see Fig. 2), confirming that between these two atoms there is a bonded interaction. The topological characteristics at the intermolecular BCP show typical properties of closed-shell interactions. The electron density values range from 0.0100 to 0.0504 au and the Laplacian values range from 0.0299 to 0.1160 au, which correlates fairly well with the values reported for similar interactions [13, 14]. According to the topological analysis of electron density distribution in the QTAIM [30, 31], the electron density at the BCP, $\rho(\mathbf{r}_b)$, is used to describe the strength of a bond. In general, the larger the value of $\rho(\mathbf{r}_b)$, the stronger the bond is. In the complexes studied here it is observed that, for the same X, $\rho(\mathbf{r}_b)$ increases following this order: $F_5X \cdots CO < F_3X \cdots CO < FX \cdots CO$. In other words, $\rho(\mathbf{r}_b)$ increases in the same order that binding energies.

The integration of electron density on the atomic basins, provides a useful tool for analyzing atomic charges, $q(\Omega)$, quadrupole moments, $Q(\Omega)$ and the electrostatic interaction energy between the total charge distribution of A atom and the nucleus of B atom, $V_{e-n}(A,B)$. These values are reported in Table 4.

The integrated atomic charges show that interactions occur between positively charged atoms, $X^{\delta+} \cdots \delta^+C$. The product of the positive charges of these interacting atoms (for the same X) is increased in the order $FX \cdots CO < F_3X \cdots CO < F_5X \cdots CO$. In this order repulsive electrostatic interactions increase and, therefore, the stability of the complexes decreases.

The atomic quadrupole moment in a particular direction (i.e. along the z-axis Q_{zz}) is a measure of the deviation of electron density from spherical symmetry. That is to say, a spherical electron density distribution results in a value of zero for Q_{zz} . This component measures how the electron density is elongated or compressed along the z-axis relative to a direction perpendicular to z. Thus, if Q_{zz} is negative, the electron density is concentrated along this axis and, consequently, the electron distribution is prolate with respect to the z-axis. Similarly, when the component Q_{zz} is positive, the electron distribution is oblate [30, 31]. This property of Q_{zz} allows us to quantify the deformation experienced by the electron density on the atomic basin of interacting atoms. In Table 4, the atomic quadrupole moment values calculated on the basin halogen atom, $Q_{zz}(X)$, measures in the direction of the F_c-X bond are reported. The results show that in the $FX \cdots CO$ and $F_3X \cdots CO$ complexes $Q_{zz}(X) > 0$, which indicates that in this direction an electron charge density depletion is present. However, $Q_{zz}(X) < 0$ for $F_5X \cdots CO$ complexes, which indicates that in this direction an electron charge density accumulation is present.

Table 4 Selected topological parameters calculated

Complexes	Local			Integrated			
	$\rho(\mathbf{r}_b)$	$\nabla^2\rho(\mathbf{r}_b)$	$H(\mathbf{r}_b)$	$q(X)$	$q(C)$	$Q_{zz}(X)$	$ V_{e-n}(C,X) $
FCI \cdots CO	0.0232	0.0795	0.0023	0.3243	1.1453	6.6377	17.3514
F ₃ Cl \cdots CO	0.0149	0.0529	0.0025	1.1860	1.1349	1.8048	15.8227
F ₅ Cl \cdots CO	0.0113	0.0417	0.0023	1.7950	1.1270	-1.4932	14.9408
FBr \cdots CO	0.0338	0.0965	-0.0001	0.4214	1.1438	8.4402	37.1423
F ₃ Br \cdots CO	0.0163	0.0518	0.0021	1.4339	1.1317	2.3904	32.0342
F ₅ Br \cdots CO	0.0107	0.0363	0.0020	2.3003	1.1232	-1.5207	29.5558
FI \cdots CO	0.0504	0.1160	-0.0072	0.6228	1.0774	12.6167	59.5456
F ₃ I \cdots CO	0.0167	0.0462	0.0014	1.7950	1.1199	3.4431	46.4509
F ₅ I \cdots CO	0.0110	0.0326	0.0014	2.9430	1.1217	-1.1588	42.8536

$\rho(\mathbf{r}_b)$: electron density at the X \cdots C BCP. $\nabla^2\rho(\mathbf{r}_b)$: Laplacian of the electron density at the X \cdots C BCP. $q(X)$: net atomic charge calculated on the basin halogen atom. $q(C)$: net atomic charge calculated on the basin carbon atom. $Q_{zz}(X)$: atomic quadrupole moment values calculated on the basin halogen atom. $V_{e-n}(C,X)$: electrostatic interaction energy between total charge distribution of carbon atom and nucleus of halogen atom. All values in atomic units

When comparing $Q_{zz}(X)$ in the $F_nX\cdots CO$ complexes (for the same X), it is observed that increases in the order $F_5X\cdots CO < F_3X\cdots CO < FX\cdots CO$ and comparing $Q_{zz}(X)$ in the $F_nX\cdots CO$ complexes (for $n = 1, 3$), it is observed that augments in magnitude following the order $F_nCl\cdots CO < F_nBr\cdots CO < F_nI\cdots CO$. In general $Q_{zz}(X)$ augments follow the same order than that of binding energies.

In a previous work it was shown that the formation of the conventional XBs ($D - X\cdots B$) is due to the electrostatic interaction between the charge density provided by the Lewis base and the nucleus of the halogen atom [7, 20, 22, 23, 36]. The halogen atom has an electronic depletion region localized at the outset region in the direction of the $D-X$ bond. This depletion determines the geometry and strength of the conventional XBs. Figure 2 undoubtedly shows that the formation of the $FX\cdots CO$ and $F_3X\cdots CO$ complexes results from the interaction between a region of the space where the electron density is locally concentrated [$\text{lump} - L(\mathbf{r}) > 0$] and the nucleus of the X atom through the region of the space where the electron density is locally depleted [$\text{hole} - L(\mathbf{r}) < 0$]. In $FX\cdots CO$ and $F_3X\cdots CO$ complexes it is clearly observed that these molecules are oriented so that the lump in the valence shell charge concentration (VSCC) of the carbon atom (Lewis base) is aligned with the hole in the VSCC of the halogen atom (Lewis acid). However, on the $F_5X\cdots CO$ complexes the halogen atom X has a cap of electronic charge density concentration on its outermost portions, centered along the extension of the F_c-X bond. This small region of charge concentration causes an interelectronic repulsion with a lone pair of the Lewis base. This repulsion is responsible that the equilibrium angle $F_c-X\cdots C$ is other than 180° . In addition, the $L(\mathbf{r})$ function shows clearly that, in the valence shell of the halogen atom X on the $F_5X\cdots CO$ complexes, there is a region of electronic charge depletion around the cap of charge concentration capable of undergoing a nucleophilic attack. It is this region of electronic charge depletion which produces the formation and determines the geometry of the $F_5X\cdots CO$ complexes. According to the preceding discussion, it seems that the electrostatic interaction between the lone pair of Lewis base (CO) and the nucleus of halogen atom plays a key role in stabilizing and determining the optimal geometry of these hypervalent

halogen bonding interactions, like in the conventional XBs [7, 20, 23, 36] and in the double hole-lump interaction between halogen atoms [22].

In Fig. 2 it is also observed that in the $F_3Br\cdots CO$ complex, the two lone pairs of the bromine atom, $LP(Br)$, are confronted with the lone pair of the Lewis base, $LP(C)$. In addition, the two bonding pairs $F-Br$, $BP(F-Br)$, are also faced to the lone pair of the Lewis base. In the $F_5Cl\cdots CO$ complex it is observed one lone pair of the bromine atom and four bonding pairs $F-Cl$ which are confronted with the lone pair of the Lewis base. That is, despite that the hypervalent halogen has a higher positive charge than the monovalent halogen, the halogen bonding interaction involving hypervalent halogen are weaker than the corresponding XBs involving monovalent halogen. This can be explained by the strong interelectronic repulsions of the lone pair of the Lewis base [$LP(C)$] with the lone pair/s [$LP(X)$] and the bonding pairs $F-X$ [$BP(F-X)$] of the hypervalent halogen atom.

These repulsive interactions $LP(Br)\leftrightarrow LP(C)$ and $BP(F-Br)\leftrightarrow LP(C)$ make that $Br\cdots C$ intermolecular distances increases in the order $FBr\cdots CO < F_3Br\cdots CO < F_5Br\cdots CO$.

According to Politzer et al. a halogen bond is a highly directional, electrostatically-driven noncovalent interaction between a region of positive electrostatic potential (positive σ -hole) on the outer side of the halogen X in a molecule $D-X$ and a negative site B , such as a lone pair of a Lewis base or the p -electrons of an unsaturated system [9–11, 18]. There is no doubt that this is true to the monovalent halogens attached to a D group with medium or high electron-withdrawing power. However, as we have seen, in hypervalent halogen bonding interactions $F_5X\cdots CO$ the halogen atom X (Cl, Br or I) has a cap of electronic charge density concentration on its outermost portions, centered along the extension of the F_c-X bond. That is to say, in the region where there would be the positive σ -hole proposed by Politzer. Therefore, we consider that the concept positive σ -hole used to explain the formation and directionality of halogen bonding should continue to be reviewed in other systems.

Moreover, the positive σ -hole concept is not suitable in some special situations, for example in the CH_3Cl molecule [19]. The chlorine atom in this molecule does not have a

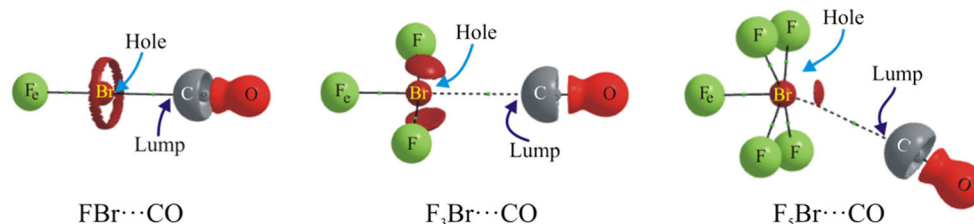


Fig. 2 Envelops at $L(\mathbf{r}) = 0.0125$ au. Note that in $FBr\cdots CO$ and $F_3Br\cdots CO$ complexes, the molecules are oriented so that the lumps in the VSCC of the carbon atom are aligned with the hole in the VSCC of the bromine atom. However, in $F_5Br\cdots CO$ complex, the lump in the

VSCC of the carbon atom interacts with a belt of electronic charge depletion located around a cap of charge concentration. In addition, the bond paths of $\rho(\mathbf{r})$ are shown

positive electrostatic potential on its surface and consequently should not be able to form XBs. However, Riley et al. [37] showed that this molecule is able to form a stable XB with OCH₂ molecule. Recently, a similar situation has been observed in the framework of the pnictogen bond [38]. According to Eskandari et al. “the lump-hole concept is more useful than the σ -hole in which the electrostatic part of potential is only considered. It is shown that the existence of hole in the VSCC of pnictogen atom is responsible for the formation and (in the absence of other interactions) the geometry of pnictogen bonded complexes” [38]. For this reason we consider it more appropriate to analyze, in the context of the QTAIM, the electrostatic interaction energy between the electron cloud of the carbon atom and the nucleus of the halogen atom, $|V_{e-n}(C,X)|$ (these values are shown in Table 4). It is observed that $|V_{e-n}(C,X)|$ (for a same X) increases with the module of the binding energy. These findings allow us to establish that, the electrostatic interaction between the lone pair of C atom and the nucleus of X atom plays a key role in stabilizing these halogen bonding interactions.

The electron-nuclear attractive contribution to virial field measured at the X \cdots C BCP, $|V_{e-n}(\mathbf{r}_b)|$, is a measure of the electrostatic force exerted by the nuclei X and C on the electronic cloud of the intermolecular region. Figure 3 shows a good quadratic correlation between $|V_{e-n}(\mathbf{r}_b)|$ and the binding energies. Therefore, the electrostatic interaction between the electron cloud of the intermolecular region and the nucleus of the halogen and carbon atoms play a key role in stabilizing these complexes.

Finally, we have studied the interaction energy components derived from the LMOEDA method. This partition scheme is a useful tool for a quantitative interpretation of the strength of the intermolecular interactions. In Table 5, the interaction energy components are reported.

It is observed that in all the complexes the most important stabilization term is electrostatic (between 41 and 47% of the stabilizing terms) followed by polarization/dispersion (in smaller proportions). In the complexes with greater interaction

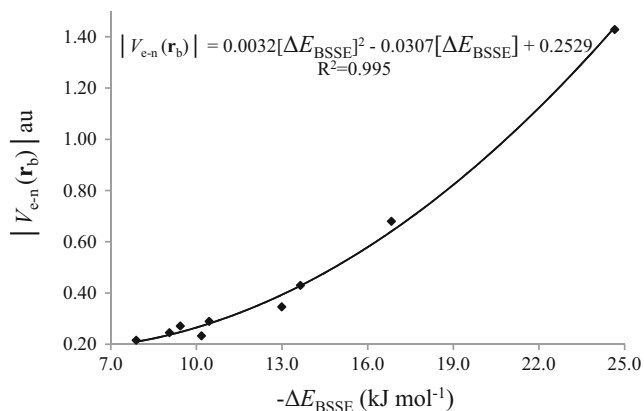


Fig. 3 Correlation between $|V_{e-n}(\mathbf{r}_b)|$ and the binding energies

Table 5 Energy decomposition analysis of the interaction energy obtained with the LMOEDA methodology

Complexes	E_{elect}	$E_{\text{ex-rep}}$	E_{pol}	E_{disp}	E_{int}
FCl \cdots CO	-32.8	55.7	-21.0	-16.4	-14.5
F ₃ Cl \cdots CO	-20.0	29.9	-10.7	-13.4	-14.3
F ₅ Cl \cdots CO	-13.9	21.5	-7.5	-12.4	-12.4
FBr \cdots CO	-62.0	108.2	-45.0	-22.5	-21.4
F ₃ Br \cdots CO	-24.2	36.0	-13.6	-14.4	-16.2
F ₅ Br \cdots CO	-15.0	21.0	-7.6	-13.0	-14.7
FI \cdots CO	-110.1	215.4	-101.8	-35.5	-31.9
F ₃ I \cdots CO	-24.1	37.7	-14.4	-15.3	-16.1
F ₅ I \cdots CO	-15.7	22.5	-7.5	-11.8	-12.5

All values in kilojoule per mole

energies (FX \cdots CO) it is observed that $E_{\text{pol}} > E_{\text{disp}}$, while in the hypervalent halogen-bonded interactions there is larger contribution of the dispersion. This indicates that in F_nX \cdots CO ($n = 3, 5$) complexes the orbital interactions, related to polarization term [28], are less important than in the conventional XBs.

Conclusions

In this work, a systematic theoretical study about Lewis acid-base behavior of hypervalent halogen fluorides in gas phase was carried out on a series of binary complexes. The Lewis base behavior was studied in systems F_nX-H(+) ($n = 1, 3, 5$ and X = Cl, Br, I), while the Lewis acidic behavior was studied in complexes F_nX \cdots CO ($n = 1, 3, 5$ and X = Cl, Br, I).

The topological analysis of the $L(\mathbf{r})$ function reveals that the hypervalent halogen fluorides, F_nX ($n = 3, 5$ and X = Cl, Br, I) can act as either Lewis bases or Lewis acids. The ability of these molecules, according to proton affinities (PA) and gas-phase basicity (GB), to accept a proton decreases in the order FX > F₃X > F₅X (for the same X) and increases in the order F_nCl < F_nBr < F_nI (for $n = 1, 3$).

The analysis of the geometric parameters of the F_nX \cdots CO ($n = 1, 3, 5$ and X = Cl, Br, I) interactions shows that, the X \cdots C intermolecular distances are substantially shorter than the sum of the van der Waals radii of the X and C atoms. The bond angles F_e-X \cdots C are close to 180° in the F_nX \cdots CO ($n = 1, 3$) complexes, while in the F₅X \cdots CO complexes, the bond angles F_e-X \cdots C are considerably less than 180°.

QTAIM reveals that the F_nX \cdots CO interactions are the result of the electrostatic interaction between the charge density provided by the lone pair of the Lewis base [LP(C)] and the nucleus of hypervalent halogen atom, like in the conventional XBs.

LMOEDA reveals that electrostatic component plays an important role in the stability of the F_nX \cdots CO complexes. In

addition, in the hypervalent halogen-bonded interactions $F_nX \cdots CO$ ($n = 3, 5$), the orbital interactions are less important than in the conventional XBs ($FX \cdots CO$).

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