

Intramolecular hypervalent O→Cl interaction in the chloronium cations: an *ab initio* study

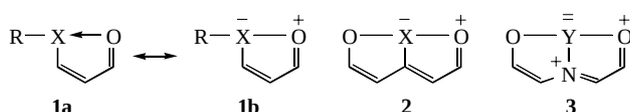
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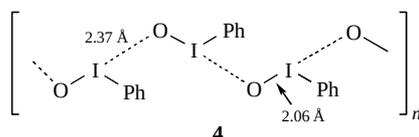
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The *ab initio* [MP2(fu)/6-31G**] and DFT (B3LYP/6-31+G**) calculations predict that the strong hypervalent O→Cl interaction stabilises the cyclic and bicyclic heteropentalene structures of chloronium cations.

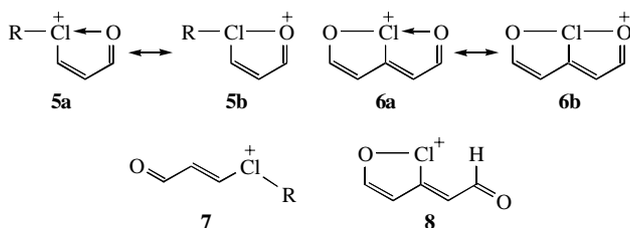
Attractive inter- and intramolecular interactions of the hypervalent type play an important role in the stabilization of sterically hindered conformations of organoelement compounds¹ and are responsible for secondary and tertiary structures of proteins, which are crucial for recognition processes.^{2,3} In the last decade, both experimental and theoretical works have been devoted to the elucidation of the nature of the intramolecular hypervalent O→X(Y) interactions in compounds **1–3**, where X and Y are chalcogens (X = S, Se, Te; R = H, Me, Cl)^{1,4} or pnictogens (Y = N, P, As, Sb, Bi).^{5,6}



The energy of these hypervalent interactions was found to strongly depend on the electronegativity of chalcogen atoms X and substituents R.^{4,6} Similar hypervalent interactions were also observed in organoelement compounds.^{7–9} At the same time, weak attractive intermolecular interactions between chalcogens (O, S) and chlorine were experimentally observed in the bimolecular complexes H₂O...ClF,¹⁰ SO₂...ClF¹¹ and H₂S...ClF.¹² The noncovalent O...I contact length observed in a crystal of PhIO **4** with the T-shaped geometry around the iodine centres is considerably shorter than the sum of the van der Waals radii of O and I (3.32 Å).¹³



Therefore, it may be expected that similar intramolecular attractive O→Hal hypervalent interactions also exist in halogen-containing organic compounds **5** and **6** (X = Hal⁺ and R = H, Me, Ph, F, Cl), iso-electronic to **1** or **2**. Here, we report on the *ab initio* [MP2(fu)/6-31G**]¹⁴ and DFT (B3LYP/6-31+G**) calculations of chloronium cations **5** (R = H, F) and **6**, which evidenced for rather strong attractive O→Cl interactions in these cations.



To estimate the energy of the O→Cl interaction, cations **5** (R = H, F) and **6** were compared with their *trans-trans*-isomers **7** (R = H, F) and **8**, which are free of the O→Cl interaction.

According to the calculations, all structures of **5–8** correspond to genuine minima ($\lambda = 0$, hereafter λ designates the number of hessian negative eigenvalues at a given stationary point) on the

Table 1 *Ab initio* [MP2(fu)/6-31G**] and DFT (B3LYP/6-31+G**) data for cations **5–8**.^a

Structure	Method	E_{tot}	ΔE	ZPE	ΔE_{ZPE}	ΔH	ω_1
5 , C _s R = H	MP2	-65 0.638323	0	0.063294	0	0	142
	DFT	-65 1.763734	0	0.061428	0	0	148
5 , C _s R = F	MP2	-74 9.583135	0	0.057537	0	0	163
	DFT	-75 0.939596	0	0.056113	0	0	162
6 , C _{2v}	MP2	-80 1.638849	0	0.082396	0	0	179
	DFT	-80 3.171914	0	0.080209	0	0	194
7 , C ₁ R = H	MP2	-65 0.627010	7.1	0.061814	6.2	6.6	100
	DFT	-65 1.754787	5.6	0.059694	4.5	5.0	91
7 , C ₁ R = F	MP2	-74 9.550728	20.3	0.055466	19.0	19.4	62
	DFT	-75 0.908761	19.3	0.053964	18.0	18.8	56
8 , C _s	MP2	-80 1.584553	34.1	0.078684	31.7	31.9	72
	DFT ^b	-80 3.098488	46.1				

^a E_{tot} (a.u.) and ΔE (kcal mol⁻¹) are the total and relative energies (1 a.u. = 627.5095 kcal mol⁻¹); ZPE (a.u.) is the harmonic zero-point correction; ΔE_{ZPE} (kcal mol⁻¹) is the relative energy including the harmonic zero-point correction; ΔH (kcal mol⁻¹) is the relative enthalpy under standard conditions $P = 1$ atm and $T = 298.1$ K; ω_1 (cm⁻¹) is the smallest harmonic vibration frequency. ^bSingle-point calculations at the MP2 geometry.

corresponding potential-energy surfaces (PES). Figures 1 and 2 and Table 1 demonstrate the calculated molecular structures, geometries and energy parameters of cations **5–8**.

All of the calculated bond lengths and angles are consistent with the available experimental data on chloronium cations (see ref. 15). As can be seen in Table 1 and Figure 1, the *cis-cis* forms of compounds **5** (R = H or F) are stabilised by the O→Cl interaction, whose energy substantially depends on the substituent at the chlorine. For compound **5** with an electropositive substituent R = H, the O→Cl interaction energy was predicted to be 7.1 (MP2) and 5.6 (DFT) kcal mol⁻¹, whereas, in the case of an electronegative substituent R = F, this energy increases up to 20.3 (MP2) and 19.3 (DFT) kcal mol⁻¹. The nonbonded O...Cl dis-

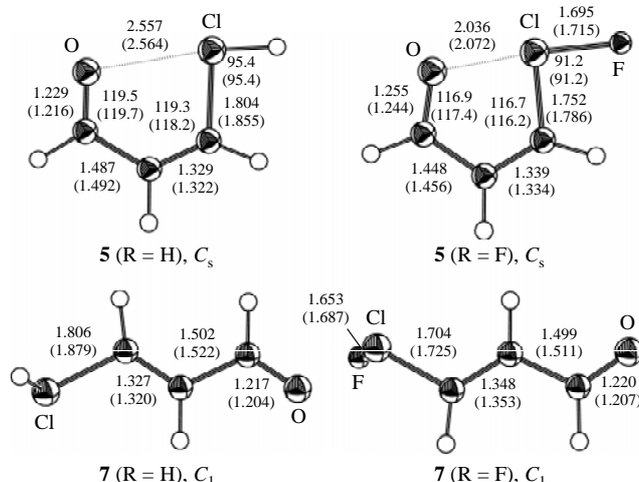


Figure 1 Geometry parameters of cations **5** (R = H, F) and **7** (R = H, F) calculated by the *ab initio* [MP2(fu)/6-31G**] and DFT (B3LYP/6-31+G**) methods (in parentheses). The bond lengths and angles are indicated in angstrom units and degrees, respectively.

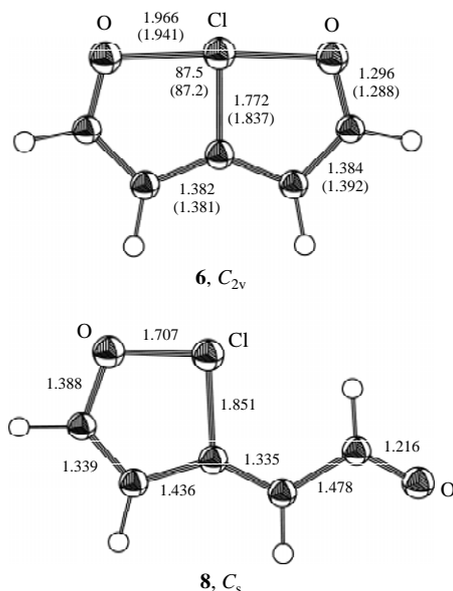


Figure 2 Geometry parameters of cations **6** and **8** calculated by the *ab initio* [MP2(fu)/6-31G**] and DFT (B3LYP/6-31+G**) methods (in parentheses). The bond lengths and angles are given in angström units and degrees, respectively.

tance in **5** (R = F) is ~ 0.5 Å shorter than that in **5** (R = H). The C–Cl lengths calculated for **5** (R = H or F) indicate that the structure of **5** (R = F) is also more delocalised than that of **5** with R = H. Note that the degree of equalization of C–C bonds in **5** (R = H or F) is higher than that in unstrained *trans–trans* conformers **7** (R = H or F) (Figure 1). This fact is indicative of the partially aromatic character of five-membered pseudo-heterocycles **5**. The structures of **7** (R = H or F) have C₁ symmetry since the substituent at the chlorine is out of the molecular plane. The corresponding planar *trans–trans* structures of C_s symmetry are transition states for the internal rotation of the substituent R around the Cl–C bond. The energy barriers for this rotation were found to be about 8 kcal mol^{−1}.

The heteropentalene system of **6**, in which the effects of both the 10π-electronic stabilization and the hypervalent O→Cl interaction act cooperatively, may exhibit a stronger hypervalent bonding than that of compounds **5**. Indeed, according to the calculations, chloronium cation **6** corresponds to a minimum ($\lambda = 0$), the smallest harmonic frequency is $\omega_1 = 179$ cm^{−1}, see Table 1) on the PES of C₃H₄O₂Cl⁺. The structure of **8**, in which no effects of the 10π-electronic stabilization and the hypervalent

O→Cl bonding are present, is predicted to be 34.1 (MP2) or 46.1 (DFT) kcal mol^{−1} thermodynamically less stable than that of **6**. Note that DFT calculations do not reveal a minimum corresponding to a stable structure like **8**. To estimate the energy difference between the structures of **6** and **8**, the latter was calculated by DFT in a single point with the MP2 optimised geometry.

In conclusion, we found that the strong attractive hypervalent O→Cl interaction exists in chloronium cations **5** and **6**. It may be expected that this interaction will increase in similar bromonium and iodonium cations.

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