

# The S<sub>N</sub>2 Reaction: A Theoretical-Computational Analysis of a Simple and Very Interesting Mechanism <sup>†</sup>

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**Abstract:** Bimolecular nucleophilic substitution (S<sub>N</sub>2) reaction is one of the most frequently processes chosen as model mechanism to introduce undergraduate chemistry students to computational chemistry methodology. In this work, we performed a computational analysis for the ionic S<sub>N</sub>2 reaction, where the nucleophile charged (X<sup>−</sup>; X=F, Cl, Br, I) attacks the carbon atom of the substrate (CH<sub>3</sub>Cl) through a backside pathway, and simultaneously, the leaving group is displaced (Cl<sup>−</sup>). The calculations were performed applying DFT methods with the Gaussian09 program, the B3LYP functional, the 6-31+G\* basis set for all atoms except iodine (6-311G\*), and the solvents effects (acetonitrile and cyclohexane) were evaluated with the PCM model. We evaluated the potential energy surface (PES) for the mentioned reaction considering the reactants, the formation of an initial complex between the nucleophile and the substrate, the transition state, a final complex where the leaving group is still bound to the substrate and the products. We analyzed the atomic charge (ESP) and the bond distance throughout the process. Gas phase and solvent studies were performed in order to analyze the solvation effects on the reactivity of the different nucleophiles. We observed that increasing solvent polarity, decreases reaction rates. On the other hand, we thought it would be enriching, to carry out a reactivity analysis from the point of view of molecular orbitals. Therefore, we analyzed the MOs HOMO and the MOs LUMO of the different stationary states on PES, both in a vacuum (gas phase) and in acetonitrile as the solvent.

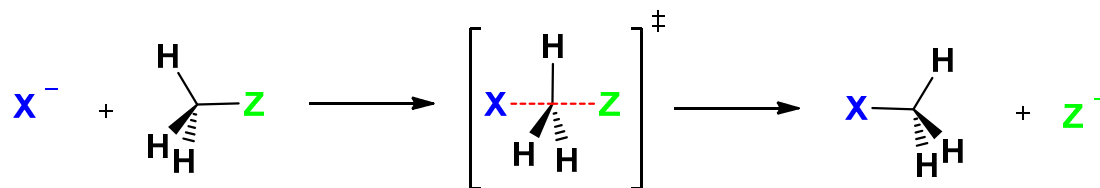
**Keywords:** S<sub>N</sub>2 reaction; DFT; computational chemistry

## 1. Introduction

Computational chemistry has become a very useful technique in organic chemistry. Since it is not part of the contents of the degree subjects in our institution, students approach the research groups to acquire basic knowledge about computational chemistry. In this way, they learn to look for reaction intermediates and transition states, evaluate bond distances and atomic charges, graph and analyze molecular orbitals; the energy calculations help them to conceptualize thermodynamic and kinetic factors in a reaction coordinate, integrating computational theoretical chemistry with the concepts acquired during the undergraduate degree.

Bimolecular nucleophilic substitution (S<sub>N</sub>2) reaction is one of the most frequently processes chosen as model mechanism to introduce undergraduate chemistry students in the computational chemistry methodology [1]. Initially, simple computational calculations are proposed, to then

increase their complexity. In Scheme 1 can be seen the general mechanism of the  $S_N2$  reaction, where X is the nucleophile and Z, is the leaving group.



**Scheme 1.** General  $S_N2$  mechanism.

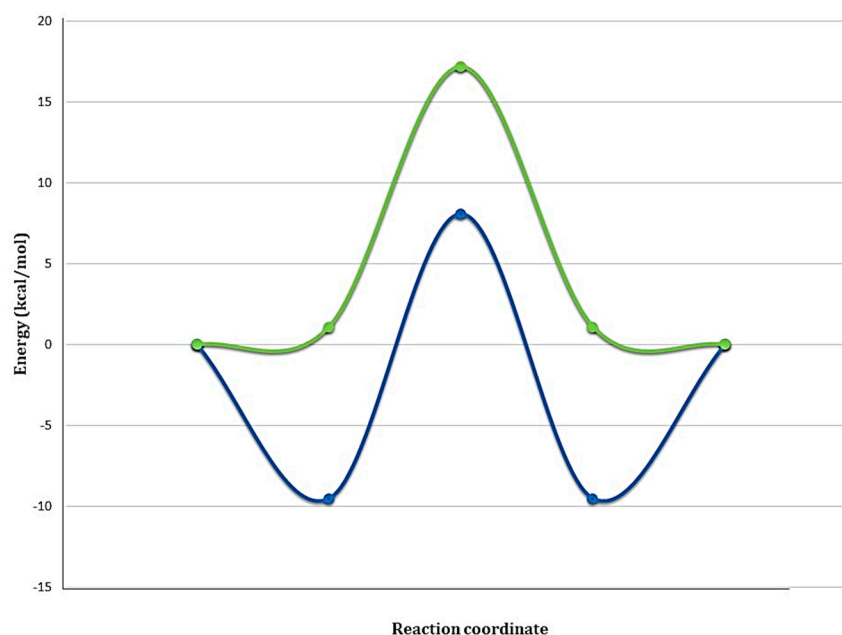
This reaction is continuously being reviewed and analysed by different research groups through computational theoretical studies [2]. The nucleophiles, the leaving groups, the electrophilic centre, the effect of the solvent on the reactivity and the shape of the potential energy surfaces have been evaluated. In this work, we performed a computational analysis for the ionic  $S_N2$  reaction, applying DFT [3] methods with the Gaussian09 program [4]. In this mechanism, the nucleophile charged ( $X^-$ ;  $X=F, Cl, Br, I$ ) attacks the carbon atom of the substrate ( $CH_3Cl$ ) through a backside pathway and simultaneously the leaving group is displaced ( $Cl^-$ ). The calculations were performed using the B3LYP functional [5], applying the 6-31+G\* basis set for all atoms except iodine (6-311G\*) and the solvents effects (acetonitrile and cyclohexane) were evaluated with the polarizable continuum model (PCM) [6] as implemented in Gaussian09. We evaluated the potential energy surface (PES) for the mentioned reaction starting from the reactants, the formation of an initial complex between the nucleophile and the  $CH_3Cl$ , a transition state, a final complex where the leaving group is still bound to the substrate and the products. We analyzed the atomic charge (ESP) [7] and the bond distance throughout the process. Vacuum and solvent study were performed in order to analyze the solvation effects on the reactivity of the different nucleophiles. On the other hand, we thought it would be enriching to carry out a reactivity analysis from the point of view of molecular orbitals. Therefore, we analyzed the MOs HOMO and the MOs LUMO of the different stationary states of the PES, for the  $S_N2$  reaction between methyl chloride and fluoride and bromide as nucleophiles, both in a gas phase and in acetonitrile as the solvent.

## 2. Methods

**Computational Procedure:** The calculations were performed applying DFT methods with the Gaussian09 program. We employed the B3LYP functional, the 6-31+G\* basis set for all atom except iodine (6-311G\*). Zero-point energy was computed at that level for all atoms. The characterization of all stationary points was done by Hessian matrix calculations of geometries obtained with full optimization for the minimums and for the transition state. The energies in solution were obtained within the Tomasi's polarized continuum model (PCM) as implemented in Gaussian09. The atomic charges were derived from the electrostatic potential (ESP-charges). The orbitals were built with the VMD program.

## 3. Results and Discussions

It is known that the  $S_N2$  reaction on a carbon center is a process that occurs through a double-well potential energy surface (PES) in the gas phase (Figure 1, blue line). However, in the aqueous solution, this PES transforms into a unimodal one (Figure 1, green line).



**Figure 1.** Solvent effect on the potential energy surfaces (PES) in a  $S_N2$  mechanism.

Based on this, we decided to study the effect of solvation on the shape of the potential energy surface (PES) for the reaction of  $\text{CH}_3\text{Cl}$  (substrate) and different nucleophiles ( $\text{X}^-$ ;  $\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$ ). As can be seen from Table 1, the studies were carried out in the gas phase in a non-polar solvent such as cyclohexane (Cy;  $\epsilon = 2.02$ ) and in an aprotic polar solvent such as acetonitrile (ACN;  $\epsilon = 38.0$ ).

**Table 1.** Energies (kcal/mol) in the gas phase, Cy and ACN to the stationary points at PES of the ionic  $S_N2$  reaction (B3LYP/6-31+G\* and 6-311G\* for I, PCM = solvent).

Reactives	Initial Complex	TS	Final Complex	Products	Medium
$\text{CH}_3\text{Cl} + \text{Cl}^-$	$\text{ClCH}_3 \cdots \text{Cl}$	$\text{Cl} \cdots \text{CH}_3 \cdots \text{Cl}$	$\text{Cl} \cdots \text{CH}_3\text{Cl}$	$\text{CH}_3\text{Cl} + \text{Cl}^-$	gas phase Cy ACN
−960,3862476	−960,4014362	−960,3875537	−960,4014362	−960,3862476	
−960,4434855	−960,4493855	−960,4299163	−960,4493855	−960,4434855	
−960,4979295	−960,4981126	−960,4705547	−960,4981126	−960,4979295	
$\text{CH}_3\text{Cl} + \text{F}^-$	$\text{ClCH}_3 \cdots \text{F}$	$\text{F} \cdots \text{CH}_3 \cdots \text{Cl}$	$\text{Cl} \cdots \text{CH}_3\text{F}$	$\text{CH}_3\text{F} + \text{Cl}^-$	gas phase Cy ACN
−599,9712194	−599,9967551	−599,9962676	−600,0393030	−600,0258111	
−600,0416609	−600,0489560	−600,0448631	−600,0886326	−600,0832416	
−600,1078282	−600,1008882	−600,0916783	−600,1365553	−600,1379508	
$\text{CH}_3\text{Cl} + \text{Br}^-$	$\text{ClCH}_3 \cdots \text{Br}$	$\text{Br} \cdots \text{CH}_3 \cdots \text{Cl}$	$\text{Cl} \cdots \text{CH}_3\text{Br}$	$\text{CH}_3\text{Br} + \text{Cl}^-$	gas phase Cy ACN
−3071,914705	−3071,9309271	−3071,916889	−3071,927285	−3071,907127	
−3071,967437	−3071,9753328	−3071,957629	−3071,973850	−3071,964306	
−3072,017946	−3072,0194136	−3071,996614	−3072,020962	−3072,018676	
$\text{CH}_3\text{Cl} + \text{I}^-$	$\text{ClCH}_3 \cdots \text{I}$	$\text{I} \cdots \text{CH}_3 \cdots \text{Cl}$	$\text{Cl} \cdots \text{CH}_3\text{I}$	$\text{CH}_3\text{I} + \text{Cl}^-$	gas phase Cy ACN
−7419,737743	−7419,747647	−7419,728013	−7419,731969	−7419,718586	
−7419,789004	−7419,790584	−7419,767876	−7419,776964	−7419,775654	
−7419,837345	−7419,832233	−7419,806318	−7419,821963	−7419,829877	

We observed that all the structures were shown to be more stable energetically under solvated conditions than the ones in vacuum. Despite the notable difference in the solvation capacity between both solvents studied (acetonitrile and cyclohexane), as indicated by the Hughes–Ingold rules, an increase in polarity solvent produces a small decrease on the rates of the reaction due to the dispersal of charges in the transition state. For example, in the case of  $\text{Cl}^-$  as nucleophile, the activation energy ( $E_a$ ) was 12.2 kcal/mol in Cy as the solvent, and  $E_a = 17.3$  kcal/mol in ACN as the solvent, as can be seen from Figure 2. For the other nucleophiles,  $\text{Br}^-$  and  $\text{I}^-$ , values were very similar: for  $\text{Br}^-$   $E_a = 11.1$

and 14.3 kcal/mol in Cy and ACN, respectively (Figure 4), and for  $I^-$   $E_a$  = 14.2 (Cy) and 16.2 kcal/mol (ACN) (Figure 5). As is known, in the gas phase,  $F^-$  was the best nucleophile, with an  $E_a$  = 0.3 kcal/mol; besides, the process was very exothermic  $\Delta H$  = −26.7 kcal/mol (Figure 3). Between  $Cl^-$  and  $Br^-$ , the last one showed to be the best nucleophile in the aprotic polar solvent. The reaction was exothermic (−1 kcal/mol). We can clarify that we were forced to use a different base set for iodine, so a comparison between this and the other halogens would not be very appropriate from the energy point of view.

As can be seen from Figures 3–5, all the studied nucleophiles showed a double-well PES in the gas phase (blue line) and in cyclohexane (orange line) and a unimodal PES in acetonitrile (green line). We think that this effect is due to the changes in solvation of the different stationary points by the solvents.

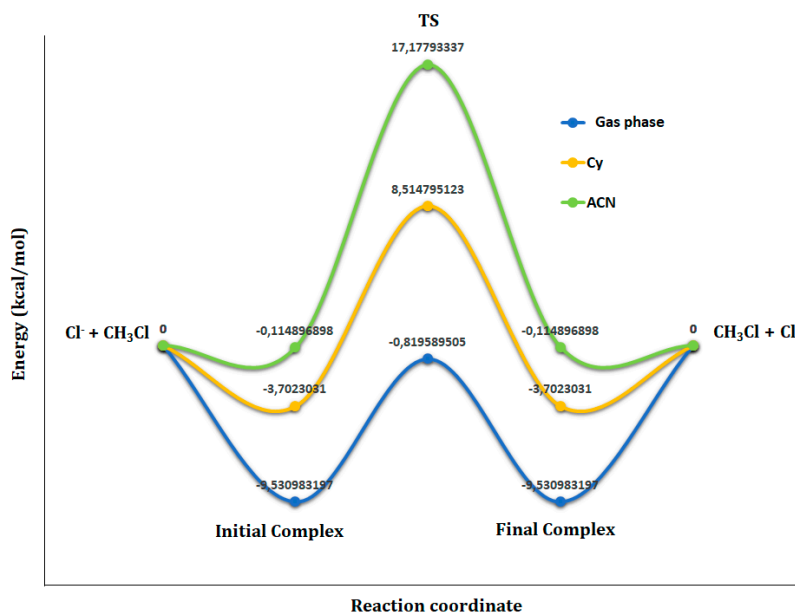


Figure 2. Solvent effect on the PES for the  $S_N2$  reaction of  $Cl^-$  and  $CH_3Cl$ .

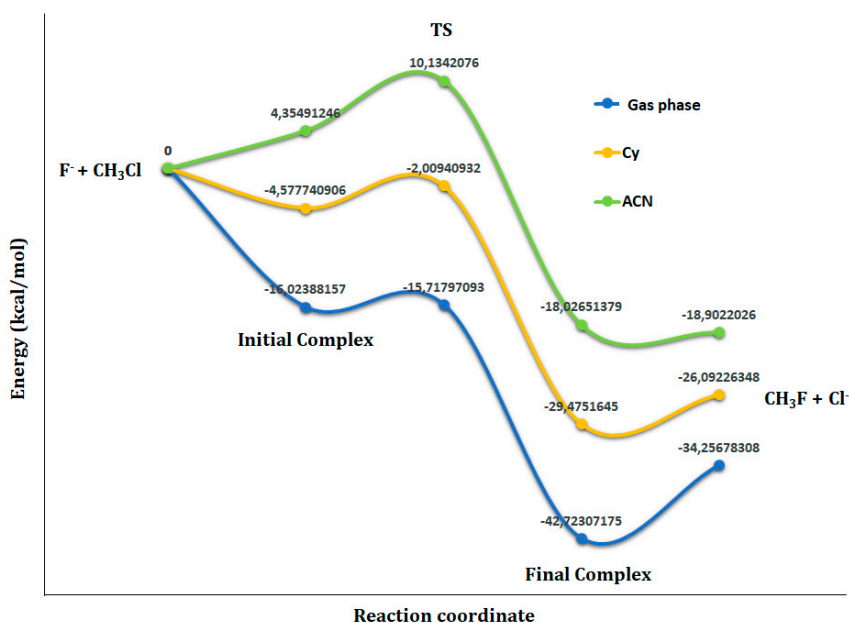


Figure 3. Solvent effect on the PES for the  $S_N2$  reaction of  $F^-$  and  $CH_3Cl$ .

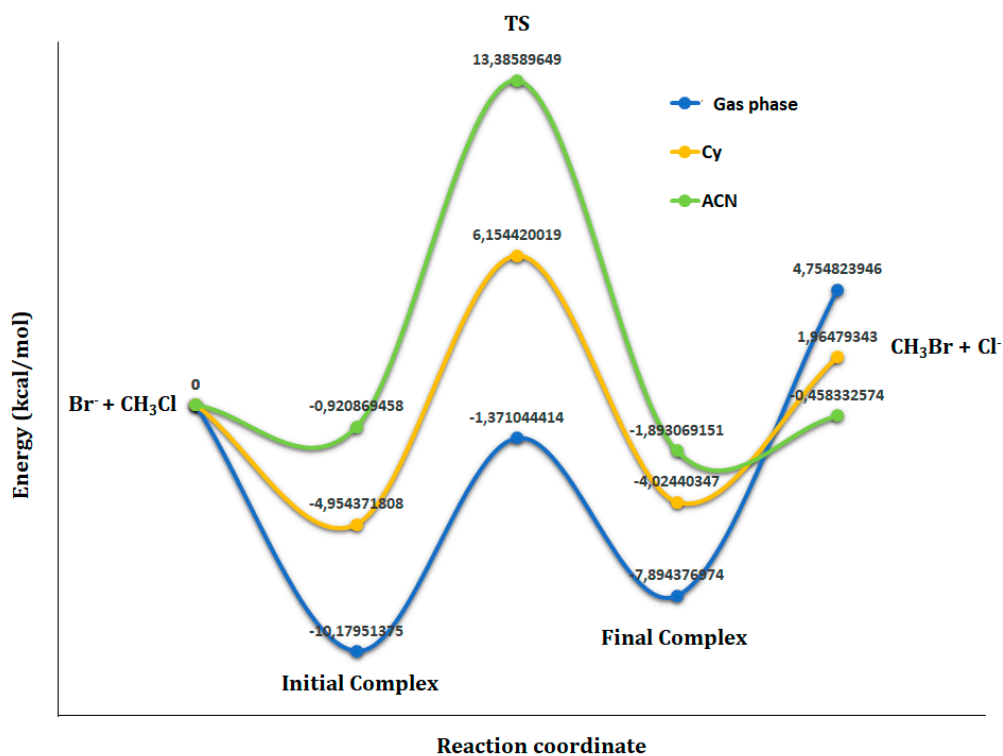


Figure 4. Solvent effect on the PES for the  $S_N2$  reaction of  $Br^-$  and  $CH_3Cl$ .

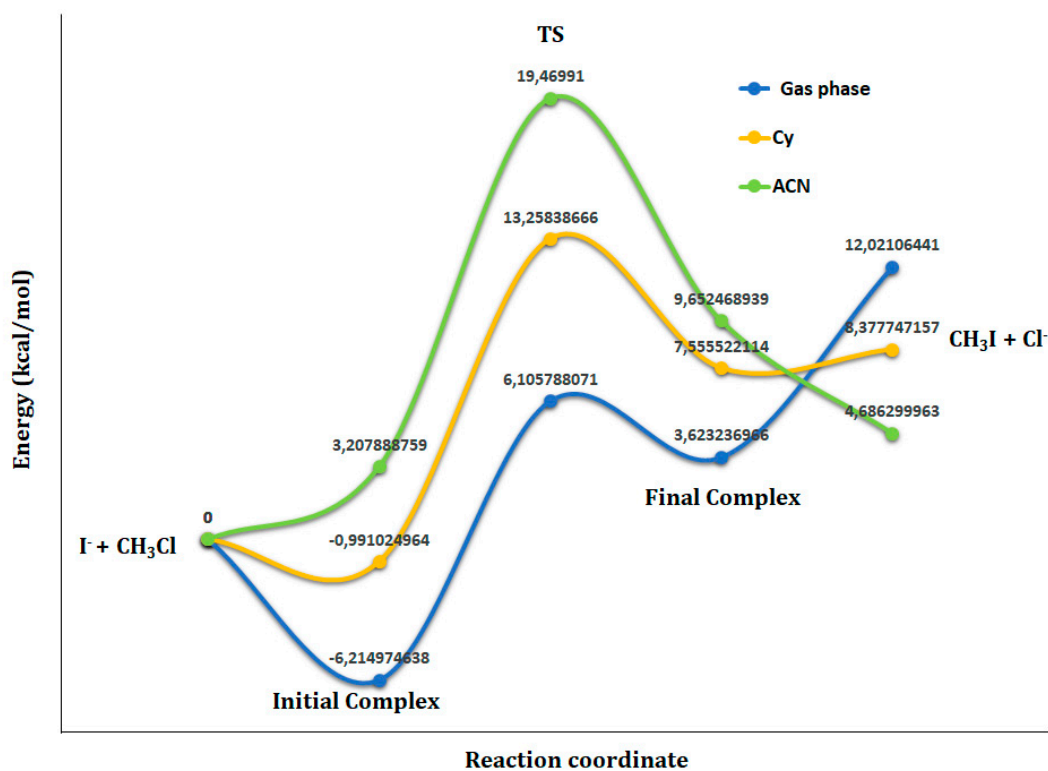


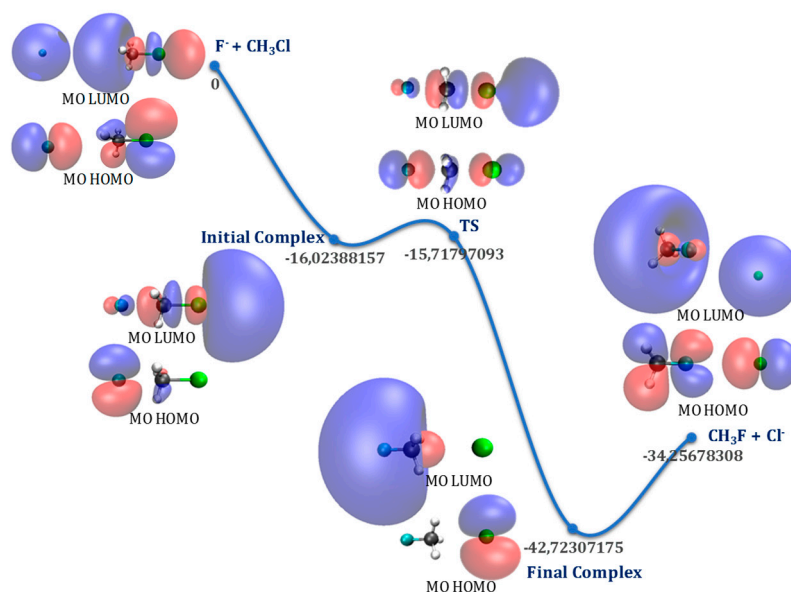
Figure 5. Solvent effect on the PES for the  $S_N2$  reaction of  $I^-$  and  $CH_3Cl$ .

**Table 2.** ESP-charges for the stationary points at PES in the gas phase, Cy and ACN (B3LYP/6-31+G\* and 6-311G\* for I, PCM = solvent).

Medium	Initial Complex	Initial Complex2	Columna2	TS	TS2	Columna 4	Final Complex	Final Complex2	Columna 6
	F	C	Cl	F	C	Cl	F	C	Cl
Gas phase	-0.919226	0.350280	-0.431054	-0.855653	0.419605	-0.563953	-0.386110	0.366005	-0.979895
Cyclohexane	-0.969862	0.295338	-0.325476	-0.834176	0.457668	-0.623492	-0.359097	0.351721	-0.992624
Acetonitrile	-0.995115	0.218417	-0.223702	-0.827332	0.494112	-0.666780	-0.329067	0.324878	-0.995811
	Cl	C	Cl	Cl	C	Cl	Cl	C	Cl
Gas phase	-0.966727	0.288636	-0.321905	-0.721680	0.440154	-0.718474	-0.321905	0.288636	-0.966727
Cyclohexane	-0.989683	0.265494	-0.275810	-0.728953	0.455967	-0.727014	-0.275810	0.265494	-0.989683
Acetonitrile	-0.996887	0.221818	-0.224931	-0.742535	0.483993	-0.741457	-0.224931	0.221818	-0.996887
	Br	C	Cl	Br	C	Cl	Br	C	Cl
Gas phase	-0.977789	0.287873	-0.310084	-0.714824	0.420334	-0.705510	-0.313878	0.259218	-0.945340
Cyclohexane	-0.998460	0.269990	-0.271531	-0.718476	0.419629	-0.701153	-0.250189	0.228794	-0.978605
Acetonitrile	-1.013396	0.247397	-0.234001	-0.737449	0.447250	-0.709801	-0.191500	0.188741	-0.997241
	I	C	Cl	I	C	Cl	I	C	Cl
Gas phase	-0.940664	0.245309	-0.304644	-0.612063	0.351473	-0.739410	-0.348560	0.275338	-0.926778
Cyclohexane	-0.969899	0.245269	-0.275370	-0.642339	0.371812	-0.729473	-0.244679	0.223227	-0.978548
Acetonitrile	-0.988262	0.213118	-0.224856	-0.673620	0.405243	-0.731623	-0.149619	0.144718	-0.995099

According to the Hammond postulate, when two consecutive states in a potential energy surface have similar energies, their interconversion will involve only a small reorganization of the molecular structure and so most exothermic reaction should be having an early TS. This happens in the  $S_N2$  with  $F^-$  as nucleophile, where the C-F bond distance is relatively large at the transition state, similar to the distance C-F at initial complex, as we already mentioned in gas phase, where the reaction is more exothermic (Table 3). The opposing happens in the  $S_N2$  with  $Br^-$  and  $I^-$  as nucleophiles, which are endothermic in gas phase and, in agreement, have a late TS.

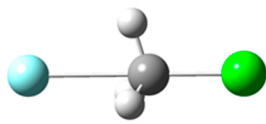

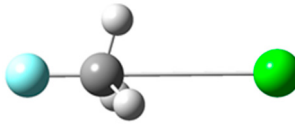
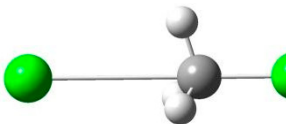
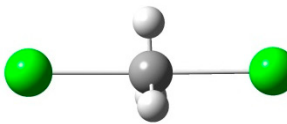
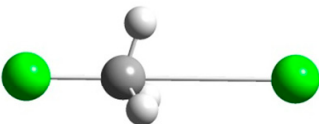
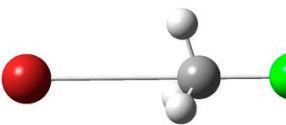
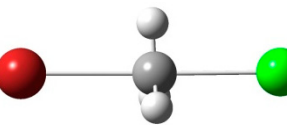
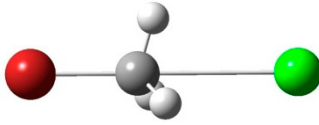
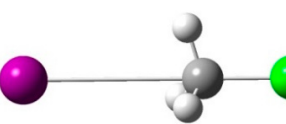
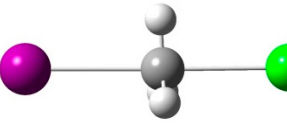
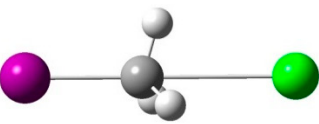
On the other hand, we carry out a reactivity analysis from the point of view of molecular orbitals. As can be seen from Figures 6–9, we graphed the MOs HOMO and the MOs LUMO of the different stationary points at the PES, for the  $S_N2$  reaction between methyl chloride and fluoride and bromide as nucleophiles, both in a gas phase and in acetonitrile as the solvent. If we observe the MO LUMO of the substrate, it is an  $\sigma^*$  orbital, antibonding between C-Cl, indicating that this bond will cleave during the attack of the nucleophile. This is also observed at the initial complex, where the orbital on the Cl is larger, indicating that the leaving group is less attached to the C. It is also observed from the shape of the MOs, that the TS is very similar to the initial complex in the case of the reaction with  $F^-$ , indicating an early TS, in the gas phase and in acetonitrile.

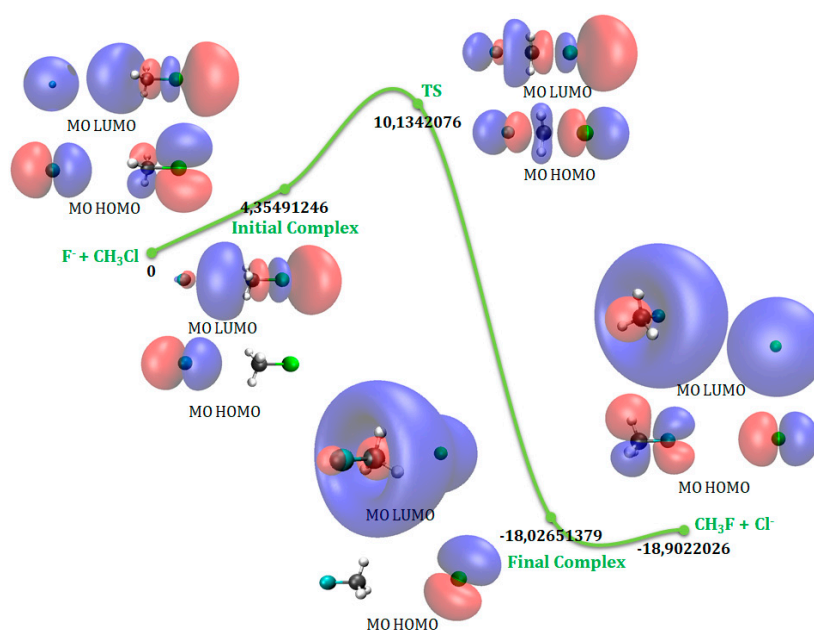


**Figure 6.** MOs HOMO and MOs LUMO for the stationary points at PES for the  $S_N2$  reaction of  $F^-$  and  $CH_3Cl$  in the gas phase.

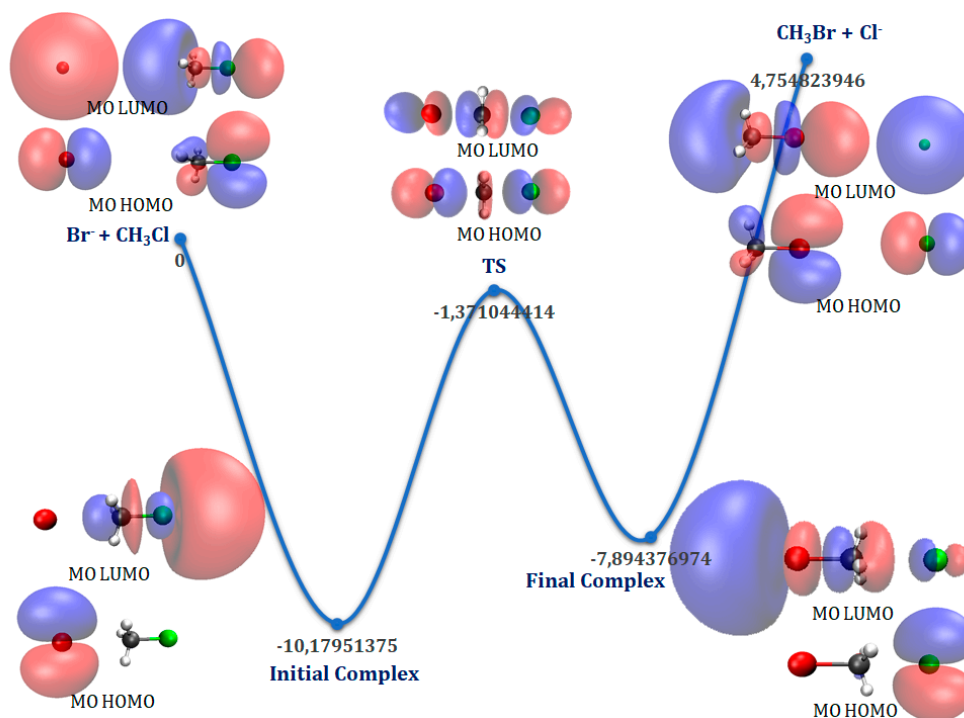


**Table 3.** C-X distances (Å) for the stationary points at PES in the gas phase (top value), Cy (middle value) and ACN (bottom value), (B3LYP/6-31+G\* and 6-311G\* for I, PCM = solvent).

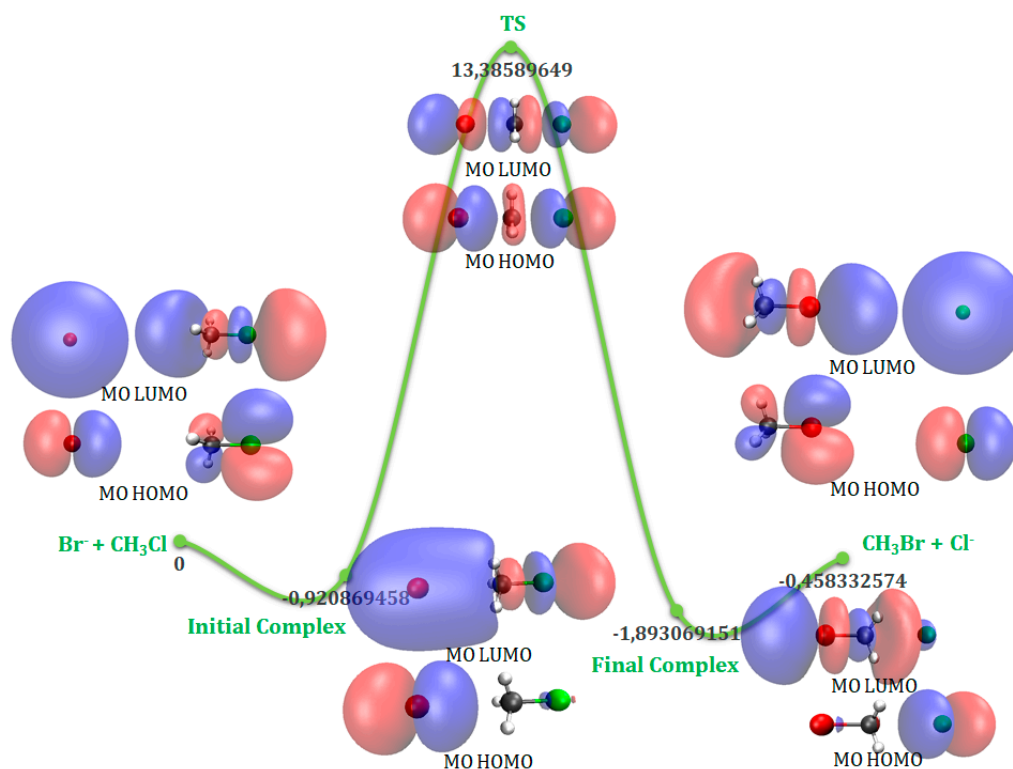
	Initial Complex2		TS2		Final Complex2	
F <sup>-</sup>	2.431	1.933	2.194	2.074	1.432	3.286
	2.681	1.863	2.097	2.157	1.423	3.506
	3.966	1.818	2.038	2.213	1.413	5.373
						
Cl <sup>-</sup>	3.203	1.856	2.364	2.370	1.856	3.203
	3.436	1.835	2.363	2.367	1.835	3.436
	5.610	1.817	2.365	2.367	1.817	5.610
						
Br <sup>-</sup>	3.296	1.855	2.483	2.366	2.022	3.072
	3.445	1.837	2.494	2.351	1.994	3.240
	3.677	1.824	2.506	2.339	1.975	3.543
						
I <sup>-</sup>	3.651	1.847	2.331	2.495	2.274	3.051
	3.854	1.831	2.672	2.442	2.212	3.365
	4.872	1.817	2.705	2.408	2.185	5.533
						



**Figure 7.** MOs HOMO and MOs LUMO for the stationary points at PES for the  $S_N2$  reaction of  $F^-$  and  $CH_3Cl$  in ACN.



**Figure 8.** MOs HOMO and MOs LUMO for the stationary points at PES for the  $S_N2$  reaction of  $Br^-$  and  $CH_3Cl$  in the gas phase.



**Figure 9.** MOs HOMO and MOs LUMO for the stationary points at PES for the  $S_N2$  reaction of  $Br^-$  and  $CH_3Cl$  in ACN.

#### 4. Conclusions

We performed a computational analysis for the ionic  $S_N2$  reaction, where the nucleophile charged ( $X^-$ ;  $X=F, Cl, Br, I$ ) attacks the carbon atom of the substrate ( $CH_3Cl$ ) through a backside pathway and simultaneously the leaving group is displaced ( $Cl^-$ ). All the structures were shown to



be more stable energetically under solvated conditions than the ones in vacuum. As is known, fluoride was the best nucleophile in the gas phase, and bromide was the best in ACN as the solvent. Despite the notable difference in the solvation capacity between acetonitrile and cyclohexane, an increase in polarity solvent produces a small decrease on the rates of the reaction, due to the dispersal of charges in the transition state. The values obtained for the atomic charges (ESP) and the bond distances C-X are in accord with the Hammond postulated for the system, that is, an early TS is observed in the reaction with fluoride, and a late TS is observed when the bromide is the nucleophile. The MOs analyses are in agreement with the concepts mentioned above.

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## References

1. James, B. Foresman and Aeleen Frisch. In *Exploring Chemistry with Electronic Structure Methods*, 2nd ed.; Gaussian Inc.: Pittsburgh, PA, USA, 1996.
2. Hamlin, T.A.; van Beek, B.; Wolters, L.P.; Bickelhaup, F.M. Nucleophilic Substitution in Solution: Activation Strain Analysis of Weak and Strong Solvent Effects. *Chem. Eur. J.* **2018**, *24*, 5927–5938.
3. Hamlin, T.A.; Swart, M.; Bickelhaup, F.M. Nucleophilic Substitution (SN2): Dependence on Nucleophile, Leaving Group, Central Atom, Substituents, and Solvent. *ChemPhysChem* **2018**, *19*, 1315–1330.
4. Kohn, W.; Sham, L.J. Self-consistent equations including exchange and correlation effects. *Phys. Rev.* **1965**, *140*, A1133–A1138.
5. 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 09, Revision C.01*; Gaussian, Inc.: Wallingford, CT, USA, 2010.
6. Lee, C.; Yang, W.; Parr, R.G. Development of the Colle-Salvetti correlation-energy formula into a functional of the electron density. *Phys. Rev. B* **1988**, *37*, 785–789.
7. Becke, A.D. Density-functional exchange-energy approximation with correct asymptotic behavior. *Phys. Rev. A* **1988**, *38*, 3098–3100.
8. Miehlich, E.; Savin, A.; Stoll, H.; Preuss, H. Results obtained with the correlation energy density functionals of Becke and Lee, Yang and Parr. *Chem. Phys. Lett.* **1989**, *157*, 200.
9. Miertus, S.; Scrocco, E.; Tomasi, J. Electrostatic interaction of a solute with a continuum. A direct utilization of AB initio molecular potentials for the prevision of solvent effects. *Chem. Phys.* **1981**, *55*, 117.
10. Miertus, S.; Tomasi, J. Approximate evaluations of the electrostatic free energy and internal energy changes in solution processes. *Chem. Phys.* **1982**, *65*, 239.
11. Cossi, M.; Barone, V.; Cammi, R.; Tomasi, J. Ab initio study of solvated molecules: A new implementation of the polarizable continuum model. *Chem. Phys. Lett.* **1996**, *255*, 327.
12. Cox, S.R.; Williams, D.E. Representation of the molecular electrostatic potential by a net atomic charge model. *J. Comp. Chem.* **1981**, *2*, 304.



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