Theoretical Studies of the Gas-Phase Identity Nucleophilic Substitution Reactions of Cyclopentadienyl Halides

Ikchoon Lee,^{*} Hong Guang Li, Chang Kon Kim, Bon-Su Lee, Chan Kyung Kim, and Hai Whang Lee

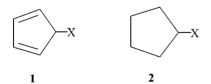
Department of Chemistry Inha University Inchon 402-751, Korea Received February 21, 2003

The gas phase identity nucleophilic substitution reactions of halide anions (X = F, Cl, Br) with cyclopentadienyl halides (1) are investigated at the B3LYP/6-311+G**, MP2/6-311+G** and G2(+)MP2 levels involving five reaction pathways: σ -attack S_N2, β -S_N2'-svn, β -S_N2'-anti, γ -S_N2'-svn and γ -S_N2'-anti paths. In addition, the halide exchange reactions at the saturated analogue, evelopentyl halides (2), and the monohapto circumambulatory halide rearrangements in 1 are also studied at the same three levels of theory. In the σ -attack S_{N2} transition state for 1 weak positive charge develops in the ring with X = F while negative charge develops with X = CI and Br leading to a higher energy barrier with X = F but to lower energy barriers with X = CI and Br than for the corresponding reactions of 2. The π -attack β -S_N2^{*} transition states are stabilized by the strong $n_{\rm C}$ - $\pi^*_{\rm C-C}$ charge transfer interactions, whereas the π -attack γ -S_N2' transition states are stabilized by the strong $n_{C} - \sigma^{2}_{CN}$ interactions. For all types of reaction paths, the energy barriers are lower with X = F than Cl and Br due to the greater bond energy gain in the partial C-X bond formation with X = F. The β -S_N2 paths are favored over the γ -S_N2' paths only with X = F and the reverse holds with X = Cl and Br. The σ -attack S_N2 reaction provides the lowest energy barrier with X = Cl and Br, but that with X = F is the highest energy barrier path. Activation energies for the circumambulatory rearrangement processes are much higher (by more than 18 kcal mol⁻¹) than those for the corresponding $S_N 2$ reaction path. Overall the gas-phase halide exchanges are predicted to proceed by the σ -attack S_N2 path with X = Cl and Br but by the β -S_N2-anti path with X = F. The barriers to the gas-phase halide exchanges increase in the order X = F < Br < CI, which is the same as that found for the gas-phase identity methyl transfer reactions.

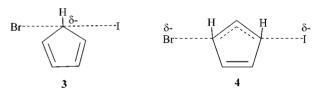
Key Words : Gas-phase, Nucleophilic substitution, Cyclopentadienyl halides, G2(+)MP2

Introduction

The experimental studies of the nucleophilic substitution reactions of cyclopentadienyl halides ((CH)₅X; 1) by Breslow *et al.*¹ with X = Br and I have indicated that cyclopentadi-



envl iodide (1 with X = I) reacts with Br (Bu₄N⁺Br) to give cyclopentadienyl bromide (1 with X = Br) *ca*. 10 times as rapidly as cyclopentyl iodide (2 with X = I) reacts in 1/1 (v/v) CCl₄-CH₃CN at 25.0 °C. The reactivity of cyclopentadienyl bromide (1 with X = Br) with I (Bu₄N⁺I) was found to be even greater. *ca*. 10² times than that of cyclopentyl bromide (2 with X = Br). They invoked (aromatic) cyclopentadienyl anionic character on the carbon atom in S_N2 reactions. **3**, and another possibility of a γ -S_N2² reaction. **4**, to explain the higher reactivity of **1** than **2**. However they were not able to provide satisfactory mechanistic infor-



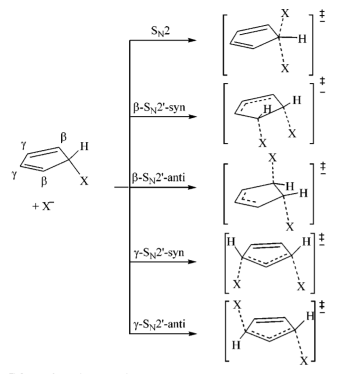
mation for the facts that (i) the reaction of bromide (1 with X = Br) with I is *ca.* 10² times more accelerated than is the reaction of iodide (1 with X = I) with Br under the same reaction condition, and (ii) the cylopentadienyl halides (1) are completely unreactive under S_{VI} solvolytic conditions, but with added halide salts (Bu₄N⁺X) they are significantly more reactive than are the cyclopentyl halides (2).

In order to present plausible answers to these problems by elucidating the detailed mechanism involved in the reactions of halide anions with cyclopentadienyl halides, we have undertaken a high level ab initio molecular orbital studey on the gas-phase identity nucleophilic substitution reactions of 1, eq. 1 with X = F. Cl and Br. The nucleophilic substitution reaction of 1 presents a variety of reaction pathways that

$$(CH)_5 X + X^- \longrightarrow X(CH)_5 + X^-$$
(1)

may involve a ring structure in the transition state (TS) with aromatic (anionic) or antiaromatic (cationic) character.² We have considered altogether 5 types of possible reaction paths

^{*}Corresponding Author, Fax: +82-32-865-4855; E-mail: ilec (\hat{a}) inha.ac.kr



Scheme 1. Various reaction pathways.

shown in Scheme 1. In addition, we have extended the studies (i) to the corresponding S_N2 reactions of the saturated analogues (s- S_N2), cyclopentyl halides. **2**, and (ii) to the sigmatropic 1.5-shifts of halide (circumambulatory rearrangement)³ within the cyclopentadienyl ring.

Computational Methods

Calculations were performed at the B3LYP/6-311+G**. MP2/6-311+G** and G2(+)MP2 levels using the Gaussian 98 programs.⁴ In the G2(+)MP2 level, diffuse functions were added to the standard G2MP2 method in order to get more accurate geometries and thermochemical data. Stationary points were characterized with analytical second derivatives. Enthalpies (ΔH^{τ}) and free energies (ΔG^{τ}) of activation at 298 K were calculated using vibrational contributions calculated with the use of harmonic frequencies computed at the B3LYP/6-311+G** and MP2/6-311+G** levels and scaled by the appropriate factors.5 Activation parameters are reported relative to the separated reactants level. The natural bond orbital (NBO) analyses⁶ were carried out to determine the proximate bond-antibond (which includes $n - \sigma^*$. $n - \pi^*$. σ -p^{*}, σ - σ ^{*}, etc) charge-transfer delocalization energies, $\Delta E_{\sigma=\sigma}^{(1)}$ in eq. 2. in the substrates and transition states (TSs),

$$\Delta E_{\sigma-\sigma^*}^{(2)} = -\sum \frac{2F_{\sigma\sigma}}{\varepsilon_{\sigma^*} - \varepsilon_{\sigma}}$$
(2)

nd natural population analyses (NPA)⁶ were carried out. The percentage bond order changes upon TS formation. $\%\Delta n^{\tau}$, were determined using eq. 3, where r^{τ} , $r_{\rm R}$ and $r_{\rm P}$ are the bond lengths in the TS, reactant and product respectively and

$$\Psi_{0}\Delta n^{*} = \frac{[\exp(-r^{*}/a) - \exp(-r_{\rm R}/a)]}{[\exp(-r_{\rm P}/a) - \exp(-r_{\rm R}/a)]} \times 100$$
(3)

a was set to 0.6 for bond orders less than one and to 0.3 for bond orders greater than one.

Results and Discussion

Reactants and Reactant Complexes (RCs). The reactant structures are presented in Figures 1-3. In general, bond lengths in the ring are slightly longer with X = F than with Cl and Br. The C-X bond in all cases is shorter than that in the saturated analogue, cyclopentyl halides (2), e.g., for X = F, $d_{C-F} = 1.392$ Å for 1 and 1.409 Å for 2. A stable adduct is formed only in the β -S_N2-anti reaction path with X = F in a triple-well potential energy surface. In all other cases, double-well potential energy profiles are obtained. in which there are two identical ion-dipole complexes (RCs) on either side of the central reaction barrier. The structures of the reactant complexes (RCs) are shown in S1 (Supporting Information). For the γ -S_N2ⁱ paths with X = Cl and Br. the RCs have bifurcated structures with X loosely bonded to two hydrogens on the $\chi\gamma$ -carbons at the B3LYP/6-311+G^{**} level. We failed to locate the RC at the MP2 level for the S_{N2} path with X = F and that shown in S1 is obtained at the RHF/ 6-311+ G^{**} result. The complexation energies (ΔE_{RC}) or well-depths range from -6 to -18 kcal mol⁻¹ as shown in S1.

The σ -attack S_N2 reactions. The halide anion attacks at

Table 1. Activation Parameters' for the Identity Halide Exchange Reactions of Cyclopentadienyl Halide, at the $B3LYP/6-311+G^{**}$ Level (in kcal mol⁻¹)

X	Paths		ΔE^{2}	ΔE^{2} Z.PE	$\Delta H^{\ell h}$	$-T\Delta S^{2b}$	ΔG^{2h}
F	S _A 2		0.46	-0.40	-0.76	8.14	7.38
	$(s-S_N2)^c$		(-2.06)	(-2.58)	(-3.21)	(9.17)	(5.97)
	β -S _A 2-syn		-3.30	-3.84	-4.70	8.88	4.18
	β -S_2-anti	TS	-9.59	-10.10	-10.79	8.28	-2.51
		Inť'	-11.26	-11.55	-12.00	8.31	-3.68
	$\gamma S_{\Lambda}2^{\prime}$ -syn		-4.02	-4.78	-5.43	8.23	2.80
	$\gamma S_N 2^i$ -anti		-2.52	-3.38	-3.96	8.53	4.57
$\overline{C1}$	$S_{\Lambda}2$		2.11	1.45	1.27	7.53	8.79
	$(s-S_{\lambda}2)^{c}$		(2.54)	(1.78)	(1.4.3)	(8.44)	(9.87)
	β -S _A 2-syn		27.69	26.41	26.07	7.33	33.41
	β -S_2-anti		7.84	6.91	6.76	7.25	14.01
	$\gamma S_{\lambda} 2' \text{-syn}$		6.86	6.31	6.00	7.21	13.20
	γ -S $_{\lambda}2$ -anti		5.50	4.91	4.69	7.36	12.05
Br	$S_{N}2$		-0.09	-0.68	-0.83	7.46	6.63
	$(s-S_{\lambda}2)^{c}$		(1.78)	(0.90)	(0.69)	(7.66)	(8.34)
	β -S _A 2-syn		27.28	26.06	25.95	6.44	32.39
	β -S_V2-anti		437	3.44	3.54	7.07	10.40
	$\gamma S_{\lambda} 2^{\prime} \text{-syn}$		4.59	4.13	3.82	7.13	10.95
	γ -S $_{\lambda}$ 2'-anti		3.09	2.56	2.38	7.24	9.61

^aCorrected for zero-point vibrational energy with the scaling factor of 0.9806. ^bAt 298 K. 'For the saturated reactants (*i.e.*, for cyclopentyl halides). ^dFor intermediate.

Table 2. Activation Parameters^{*a*} for the Identity Halide Exchange Reactions of Cyclopentadienyl Halide, at the MP2/6-311+G** Level (in kcal mol⁻¹)

X	Paths	ΔE^{τ}	$\Delta E^{\neq}_{\rm ZPE}$	$\Delta H^{\neq b}$	$-\mathrm{T}\Delta S^{\neqb}$	$\Delta G^{\star b}$
F	S _A 2	6.99	6.19	5.73	8.44	14.17
	$(s-S_{\lambda}2)^{c}$	(3.91)	(3.56)	(2.82)	(9.33)	(12.15)
	β -S _A 2 - syn	-1.88	-1.92	-2.81	8.89	6.08
	β -S _A 2'(A) TS	-5.96	-6.12	-6.84	8.36	1.51
	Int^d	-9.78	-9.71	-10.16	8.19	-1.97
	γ-S _A 2'-syn	-1.73	-2.04	-2.80	8.41	5.62
	γ-S _A 2'-anti	0.40	-0.05	-0.69	8.68	7.96
Cl	S _N 2	8.79	7.71	7.54	7.80	15.34
	$(s-S_N2)^i$	(10.33)	(9.66)	(9.29)	(8.18)	(17.47)
	β -S $_{\lambda}2^{\prime}$ -syn $^{ m e}$	24.28	23.37	22.31	9.43	31.75
	β -S $_{\lambda}2'$ -anti	11.04	10.49	10.04	8.38	18.42
	γ-S _A 2'-syn	11.94	11.81	11.32	7.79	19.11
	γ -S $_{\Lambda}$ 2'-anti	13.35	13.13	12.76	7.96	20.72
Br	S _N 2	5.18	4.29	4.10	8.03	12.13
	$(s-S_N2)^i$	(8.86)	(8.39)	(8.02)	(8.28)	(16.30)
	β -S_2'-syn"	30.64	29.58	28.63	9.40	38.03
	β -S _N 2'-anti	10.76	10.52	10.10	8.45	18.55
	γ-S _A 2-syn	10.24	10.25	9.74	8.07	17.81
	$\gamma S_A 2$ -anti	11.35	11.19	10.84	8.07	18.91

"Thermal data were taken from the MP2 6-31 G" level and zero-point vibrational energies were corrected with the scaling factor of 0.9670. ^bAt 298 K. 'For the saturated reactants, *i.e.*, cyclopentyl halides, "For intermediate, "Two imaginary frequencies (second-order saddle point).

Table 3. Activation Parameters⁶ for the Identity Halide Exchange Reactions of Cyclopentadienyl Halide, at the $G2(+)MP2//MP2/6-311+G^{**}$ Level (in kcal mol⁻¹)

X	Paths	ΔE^{2}	$\Delta E^{2}_{\rm ZPE}$	ΔH^{2b}	$-\mathrm{T}\Delta S^{2b}$	ΔG^{2b}
F	$S_N 2$	1.32	0.52	0.05	8.44	8.49
	$(s-S_N2)^c$	(-1.25)	(-1.60)	(-2.34)	(9.33)	(6.99)
	β -S _A 2'-syn	-5.08	-5.13	-6.01	8.89	2.88
	β -S _N 2'-anti TS	-8.48	-8.65	-9.37	8.36	-1.02
	Int."	-12.34	-12.27	-12.72	8.19	-4.53
	$\gamma S_{\lambda}2'$ -syn	-4.17	-4.48	-5.24	8.41	3.19
	$\gamma S_{\lambda}2'$ -anti	-2.25	-2.68	-3.32	8.68	5.33
Cl	$S_N 2$	3.55	2.47	2.30	7.80	10,10
	$(s-S_N2)^{\prime}$	(4.52)	(3.84)	(3.47)	(8.18)	(11.66)
	β -S _N 2'-syn ^e	26.14	25.23	24.17	9.43	33.60
	β -S _N 2'-anti	12.81	12.25	11.80	8.38	20.19
	γ-S _A 2'-syn	11.53	11.41	10.92	7.79	18.71
	$\gamma S_{\Lambda}2^{\prime}$ -anti	11.68	11.46	11.09	7.96	19.05
Br	$S_N 2$	1.99	1.09	0.91	8.03	8.94
	$(s-S_N2)^c$	(4.26)	(3.79)	(3.42)	(8.28)	(11.70)
	β -S _A 2-syn ^e	31.23	30.18	29.23	9.40	38.62
	β -S _{NN} 2-anti	10.82	10.59	10.17	8.45	18.62
	$\gamma S_{\Lambda}2'$ -syn	9.80	9.81	9.31	8.07	17.38
	$\gamma S_{\Lambda}2^{i}$ -anti	9.70	9.54	9.18	8.07	17.25

"Thermal data were taken from the MP2 6-31+G^{*} level and zero-point vibrational energies were corrected with the scaling factor of 0.9670. ^bAt 298 K. 'For the saturated reactants, *i.e.*, cyclopentyl halides. ^dFor intermediate, "Two imaginary frequencies (second-order saddle point).

the C_1 atom and a trigonal bipyramidal pentacoordinate (TBP-5C) TS (3) is formed in which the cyclopentadienyl ring provides two equatorial ligands. The reactions proceed through typical double-well potential energy surfaces. As have been pointed out by Breslow *et al.*¹ this TS will be extremely unstable, if positive charge develops in the TS within the ring, due to the antiaromatic character of the cyclopentadienyl cation. By contraries, if negative charge develops in the ring, the TS will be stabilized due to the aromatic character of the cyclopentadienyl anion. The energetics summarized in Tables 1-3 reveals that the activation energies (ΔE^{\star} and ΔG^{\star}) for the σ -attack S_N2 reaction of fluoride anion with cyclopentadienyl fluoride (1 with X = F) are higher by $\delta \Delta G^{\dagger} = \Delta G^{\dagger}(S_{N}2) - \Delta G^{\dagger}(s-S_{N}2) = \pm 1.5$ kcal mol^{-1} at the G2(+) MP2 level than that with cyclopentyl fluoride (s-S_x2; 2 with X = F), while those for the reactions of 1 with X = Cl and Br are lower by $\delta\Delta G^2 = -1.6$ and -2.8kcal mol⁻¹ respectively than those for the corresponding reactions of 2 with Cl and Br at the G2(+)MP2 level. This should indicate that there is an unfavorable electronic effect within the unsaturated ring in the reaction of 1 with X = F, in contrast there is a favorable effect within the ring in the reactions of 1 with Cl and Br, compared with the reactions of 2 with the respective halide. The percentage bond order changes upon TS formation, $\%\Delta n^{27}$ (S2), and the NPA charge⁶ analyses (S3) show that the degrees of bond making and cleavage are approximately 50% in all cases with the NPA charges at the C₁ atom in the TS of $q_1^{\prime} = +0.149$ (X = F), -0.112 (Cl) and -0.159 (Br). These results clearly

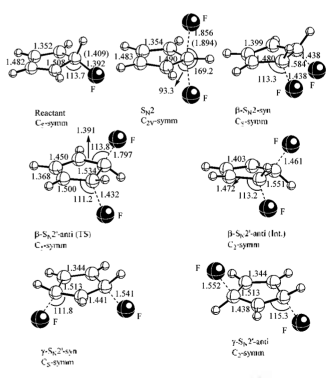


Figure 1. Structures of reactant and transition states for the identity fluoride exchange reactions of cyclopentadienyl fluoride, calculated at the MP2/6-311+G'' level (bond lengths in Å, and angles in degrees). Values in parentheses are for cyclopentyl fluoride.

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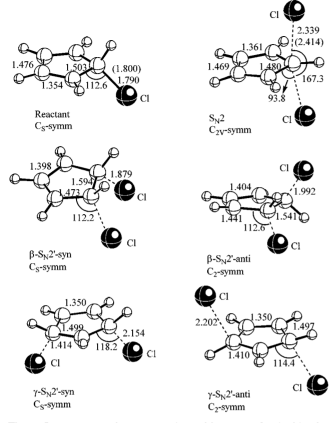


Figure 2. Structures of reactant and transition states for the identity chloride exchange reactions of cyclopentadienyl chloride, calculated at the MP2/6-311+G^{*} level (bond lengths in Å, and angles in degrees). Values in parentheses are for cyclopentyl chloride.

show that the reaction of fluoride (1 with X = F) is unfavorable whereas those of chloride and bromide (1 with X = Cl and Br) are favorable over the corresponding reactions with 2 due to antiaromatic cationic charge (for X = F) and aromatic anionic charge (for X = Cl and Br) development in the TSs, respectively.

The TS structures are shown in Figures 1-3. The C-F bonds in the distorted TBP-5C TS with C_{2T} symmetry for 1 with X = F are longer by 0.038 Å than the corresponding values in the TS for 2 with X = F. In contrast, however, they are much shorter than the corresponding values (d_{C+F}^{τ} = 2.069 Å) in the σ -attack S_N2 TS for cyclopropenyl fluoride (5, X = F), by 0.213 Å.⁸ In the latter, the TS has an "open" structure in a dissociative type S_N2 process so that the cyclopropenium ring has strong delocalized cationic (aromatic) character. In fact all the C-X bonds in the σ -attack S_N2 TSs for the identity exchange of cyclopentadienyl halides (Figures 1-3) are much shorter (by 0.342 Å for X = C1 and 0.395 Å for X = Br at the MP2 level) than the corresponding values for cyclopropenyl halides. 5 ((CH)₃X).⁸



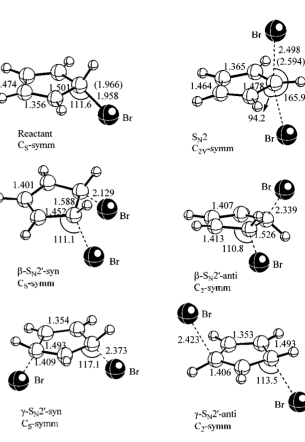


Figure 3. Structures of reactant and transition states for the identity bromide exchange reactions of cyclopentadienyl bromide, calculated at the MP2/6-311+G^{*} level (bond lengths in Å, and angles in degrees). Values in parentheses are for cyclopentyl bromide.

These trends of forming tighter TSs (**S2**) in the $S_N 2$ reactions with 1 than with cyclopropenyl halides (**5**) are of course to preserve as much as possible the (aromatic) anionic ring character in the TSs for the associative type $S_N 2$ reactions of 1. in contrast to the preferred (aromatic) cationic ring character in the TSs for the dissociative type $S_N 2$ reactions of **5**.

In the TBP-5C TS (Figure 1) an empty p orbital develops as the leaving group. F⁻, departs, and leads to strong electronic charge delocalization from the lone pair orbitals (one p and two sp^3 types) on both F atoms toward the empty p(p') orbital on C₁ (n_F-p' charge transfer interactions)⁹ and the TS is stabilized as much as possible by preventing the formation of antiaromatic cyclopentadienyl cationic ring structure. The $n_{\rm F}$ - p^+ charge transfer energy ($\Sigma \Delta E^{(2)}_{n-p^+} = -295$ kcal mol⁻¹) is the main component of the total increase $(\Sigma \Delta E^{(2)}_{\sigma \sigma^*} = -370 \text{ kcal mol}^{-1}$ in Table 4) in the proximate charge transfer delocalization energies^{6,10} on going from the reactants to the TS. The TS is further stabilizaed by an extra, but partial, C-X bond formation (BE = 102 (X = F), 76 (Cl) and 65 (Br) kcal mol⁻¹).¹¹ Due to the substantially larger gain in the bond energy with X = F(ca, 54%) bond cleavage and 46% bond formation) than with X = Cl and Br (*ca.* 60%) bond cleavage and 40% bond formation) despite the weaker charge transfer stabilization with X = F (Table 5), the barrier height to the σ -attack S_N2 reaction decreases in the order CI

Table 4. Electrostatic (ΔE_{es}) , Deformation (ΔE_{def}) and Charge-Transfer $(\Sigma \Delta E^{(2)}\sigma_{e}\sigma)$ Energies for the Halide Exchange reactions of Cyclopentadienyl Halides at the MP2/6-311+G^{**} Level (in kcal mol⁻¹)

X	Paths	ΔE_{es}	$\Delta E_{ m def}$	$-\delta \Sigma \Delta E^{(2)}{}_{\sigma \cdot \sigma}$ -
F	S _A 2	8	42	-370
	$(s-S_N2)^{\alpha}$	(-12)	(42)	(-131)
	β -S_2-syn	42	32	-235
	β -S _A 2'-anti (TS)	24	8	-218
	β -S _A 2'-anti (Int)	33	28	-238
	γ-S _A 2'-syn	48	29	-148
	γ-S _A 2'-anti	44	29	-145
C1	S _A 2	23	37	-383
	$(s-S_{\Lambda}2)^{a}$	(-8)	(42)	(-111)
	β -S _A 2'-syn	31	.34	-237
	β -S _A 2'-anti	40	25	-294
	γ-S _A 2'-syn	.38	30	-345
	γ -S $_{\Lambda}$ 2'-anti	30	31	-347
Br	$S_{\Lambda}2$	30	31	-379
	$(s-S_{\Lambda}2)^{a}$	(-6)	(42)	(-106)
	β -S _A 2'-syn	36	32	-278
	β -S ₃ 2'-anti	21	27	-438
	γ-S _M 2'-syn	38	27	-444
	γ-S _A 2'-anti	28	28	-454

"Backside attack for the saturated reactants (*i.e.*, cyclopentyl halides). ${}^{b}\delta\Delta E_{es} = \Delta E_{es}(TS) - \Delta E_{es}(Reactant)$. ${}^{c}\delta\Sigma\Delta E^{(2)}_{\sigma\sigma\sigma'} = \Sigma\Delta E^{(2)}_{\sigma\sigma'}(TS) - SDE^{(2)}_{\sigma\sigma'}$ (Reactant).

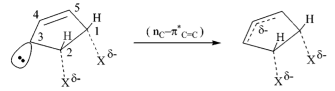
 $(\Delta G^{\tau} = 10.1 \text{ kcal mol}^{-1}) > \text{Br} (8.9) > \text{F} (8.5)$, which is similar to the order of the S_N2 reactions with **2**. Cl ($\Delta G^{\tau} = 11.7 \text{ kcal mol}^{-1}$) $\equiv \text{Br} (11.7) > \text{F} (7.0)$.

Comparisons of activation energies. $\delta \Delta G^{\tau} = \Delta G^{\tau}(S_N 2)$ $-\Delta G^{2}$ (s-S_A2), between those for the reactions of cyclopentadienyl halides $(S_N 2)$ with those corresponding reactions of the saturated, cyclopentyl, halides (s- S_N 2), show that the reactivities of 1 with X = Cl ($\delta\Delta G^{2}$ = -1.6 kcal mol⁻¹) and Br ($\delta\Delta G^{\tau} = -2.8 \text{ kcal mol}^{-1}$) are faster by $k(S_N 2)/k(s-S_N 2) \equiv$ 10 and 10² at 298 K respectively, but is slower ($\delta\Delta G^{2}$ = +1.5 kcal mol⁻¹) by 10^{-1} with X = F. The G2(+)MP2 reactivity ratio of 10^2 is in good agreement with the experimental reactivity ratio, $k(S_N 2)/k(s-S_N 2) \equiv 10^2$ for X = Br⁺ considering the fact that our theoretical values are for the gas-phase reactions whereas the experimental data are for the reactions of cyclopentadienyl ($S_N 2$) and cyclopentyl (s- $S_N 2$) bromides with $I^{-}(Bu_4N^{+}I^{-})$ in CCl₄-CH₃CN mixtures at 298 K.⁺ The lower reactivity of cyclopentadienvl fluoride $(k(S_N2)/$ $k(s-S_N 2) \equiv 10^{-1}$) than cyclopentyl fluoride is most probably due to the weak antiaromatic cyclopentadienyl cationic character of the TS with X = F, while the higher reactivity of cyclopentadienyl chloride and bromide than the corresponding evelopentyl halides are due to the aromatic cyclopentadienyl anionic character of the TSs involved, as discussed above.

In general there are several factors that are important for the TS stability, and hence for the reactivity. in the S_N2 reactions: (i) The deformation energy, ΔE_{det} , that is needed to supply for the structural deformation of the reactant to achieve the TS structure. The major component of ΔE_{def} is the stretching of the leaving group in the TS.¹² (ii) The bond energy (BE) gain in the newly formed ((partial) bond by the nucleophile. (iii) Changes in the proximate charge transfer energies (eq. 2) upon TS formation. $\delta\Delta E_{\sigma,\sigma}^{(2)} = \Delta E_{\sigma,\sigma\sigma}^{(2)}$ (TS) – $\Delta E_{\sigma,\sigma}^{(2)}$ (Reactants). The charge transfer interaction takes place between vicinal (and / or geminal) bond (n, π , σ , etc) and vacant antibond (π . σ , etc) orbitals, and the interaction energy can be calculated as the second-order perturbation energy by eq. 2.^{6,9} (iv) Changes in non-chargetransfer energies.⁶ $\delta\Delta E_{\rm NCT} = \Delta E_{\rm NCT}(\rm TS) - \Delta E_{\rm NCT}(\rm Reactants).$ upon TS formation. Non-charge-transfer term ($\Delta E_{\rm NCT}$) includes electrostatic ($\Delta E_{\rm es}$) and steric interaction energies ($\Delta E_{\rm st}$) and corresponds essentially to the energy associated with the Lewis structure ($\Delta E_{\rm Lew}$).¹³

The σ -attack S_N2 reactions of cyclopentadienvl halides (1) have relatively tight associative $S_N 2$ type TS structures in which the bond formation (40-45%) and cleavage (54-60%) have progressed approximately 50% with negative charge development at the reaction center carbon (except with X = F). The major stabilization energy is provided by the n_X-p^+ interaction (-295 (F): -309 (Cl); -299 kcal mol⁻¹ (Br)). In addition to this charge delocalization toward the developing empty $p(p^{\dagger})$ orbital at C₁, small but significant $\pi_{C,C}$ -p charge transfer (-50 (F); -68 (Cl); -74 kcal mol⁻¹ (Br)) occurs and the cyclopentadienyl ring acquires delocalized anionic (aromatic) character. For X = F, these charge delocalizations are not sufficient to gain aromatic character but is supplemented by the strong bond energy of a newly, but partially, formed C-F bond to stabilize the TS. Due to the slightly cationic antiaromatic character with X =F, the reaction barrier becomes higher than that for the saturated, cyclopentyl fluoride, in which there is no delocalization within the (saturated) ring. This means that other factors ((i) and (iv) above) differ very little and contribute similarly to the TSs for the reactions of 1 and 2. Quite interestingly, the stabilizing factors in the TSs for 1 form a striking contrast to those in the TSs for cyclopropenvl halides (5). For the halide exchanges in 5, the TSs are of the dissociative $S_N 2$ type and have an "open" (loose) structure with low degree of bond making (22-33%) and extensive bond cleavage (67-78%).8 The cylopropenyl ring becomes strongly positive $(+0.67 \sim +0.76)$ and achieves strong aromatic cyclopropenyl cation structure. The major charge delocalization comes from the $\pi_{C=C}-p^{+}$ interaction (-307 ~ -367 kcal mol⁻¹) with small stabilizing electrostatic ($-61 \sim$ -86 kcal mol⁻¹) and destabilizing deformation (35-46 kcal mol⁻¹) energies. Here due to the extensive stretching of the C-X bonds (very small overlap between the two orbitals leads to negligible Fock matrix element $F_{p,p}$) there is no n_X -p⁺ type charge transfer interaction.

The π -attack β -S_N2' reactions. The nucleophile can attack at a neighboring β -carbon either with syn (β -S_N2'-syn) or anti (β -S_N2'-anti) orientation to the existing C-X bond forming a 1.2-dihalocyclopentenyl anion like TS. The anti 1.2-dihalo adduct is stable enough to exist as an intermediate only for X = F with a well-depth of -3.5 kcal mol⁻¹ at the G2(+)MP2 level, and the β -S_N2'-anti reaction with X = F



Scheme 2. β -S_A2'-syn TS.

exhibits a triple-well potential energy profile. The stable 1,2dihalo adducts are not formed with X = CI and Br. so that β - $S_{\lambda}2^{\prime}$ reactions with X = Cl and Br proceed by a double-well potential energy surfaces. The energetics in Tables 1-3 reveals that the B3LYP activation energies are lower than those corresponding values at the G2(+)MP2 level, excepting for the two cases of β -S_N2'-syn TS and β -S_N2'-anti adduct with X = F for which the reverse holds. *i.e.*, the B3LYP ΔG^{\prime} values are higher. This suggests that the DFT method overestimates electron correlation effects in general for the delocalized TS.^{14,15} but underestimates interorbital pair correlation energies between neighboring lone pairs on the two F atoms¹⁶ of the vicinal C-F bonds which are much shorter ($d^{\tau}_{CF} = 1.438$ Å) than those of C-Cl (1.879 Å) and C-Br (2.129 Å) in the β -S_N2'-syn TS. The β -S_N2' TSs are relatively tight (S2) with large degree of bond formation (75-93%) and small extent of bond cleavage (7-25%) in the syn TSs compared with the corresponding values of 53-71% and 6-47% in the anti TSs.

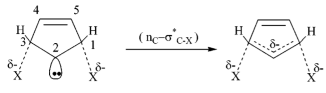
On the other hand, the MP2 ΔG^{\dagger} values for the β -S_N2ⁱ paths with X = Cl and Br are lower than those corresponding values at the G2(+)MP2 level. This indicates that, in contrast to underestimation of electron correlation effects in general at the MP2 level, they are overestimated for the delocalized TSs^{14,15} of the β -S_N2' paths with X = Cl and Br. The NBO analyses indicate that in both the syn and anti TSs of the β -S_N2' paths. an incipient lone pair formed on C₃ is delocalized toward the C = C bond by a strong $n_C - \pi c_C$ charge transfer interaction and the delocalized allyl anionic structure is formed over C3-C1-C5 molety as shown in Scheme 2. The lone pair on C_3 is a p type in the anti path, whereas it is an sp^3 type in the syn path. This $n_C - \pi^*_{C,C}$ charge transfer delocalization provides major part (> 80%) of the overall proximate charge transfer energies listed in Table 4. There are also weak vicinal n_{C} - σ^{*}_{C-X} interactions in all of the β -S_N2' TSs. This means that electronic delocalization extends partly to C_{2s} *i.e.*, over C_{2} - C_{5} moiety (Scheme 2). The overall proximate charge transfer energies $(-\delta\Delta E_{\sigma-\sigma}^{(2)})$ summarized in Table 4 show that the anti TSs (compare the adduct value for the anti path with X = F) have stronger stabilization than the syn TSs due to the proximate charge transfer delocalization leading to lower activation energies for the anti β -S_N2' reactions. The β -S_N2'-svn TSs (C_s svmmetry) for X = CI and Br are characterized by two imaginary frequencies at the MP2/6-311+G** level indicating that they are second-order saddle points with a maximum with respect to two mutually perpendicular directions.¹⁷ Such a TS may be considered as a transient structure in the inversion profile of the β -S_A2'-anti TSs (C₂ symmetry) and has an energy

higher by 13-20 kcal mol⁻¹ at the G2(+)MP2 level than the C_2 structure. The higher ΔG^{τ} values for the syn than anti path are in most cases due to higher steric repulsion ($\Delta E_{cs} > 0$) as evident from more negative ΔS^{τ} value by *ca.* 2-4 e.u for the syn path. and also due to higher deformation energies (ΔE_{def}) for the syn (Table 4) than the anti path.

For the β -S_N2 reaction paths also the fluoride exchanges have considerably lower activation energies despite low proximate charge transfer stabilizations, *e.g.*, for the anti path, $\Delta G^{\tau} = -1.0$ (F). 20.2 (Cl) and 18.6 kcal mol⁻¹ (Br). Since stretching of the C-F bond in the TS is smaller than that of the C-Cl and C-Br bond the DE_{def} value is smaller for X = F by *ca.* 17-19 kcal mol⁻¹ than for X = Cl and Br. We also invoke the bond energy factor (factor (ii). *vide supra*) to rationalize the considerably lower ΔG^{τ} value with X = F since bond energy of the C-F bond is much stronger than those of the C-Cl and C-Br bonds.¹¹ This interpretation was also applicable to the lower ΔG^{τ} values with X = F for the halide exchanges in cyclopropenyl halides.⁸

The activation energies (ΔG^{*}) for the β -S_N2[•] processes of halide exchanges in cyclopropenyl halides, (5), are substantially lower than the corresponding reactions with 1, especially with X = C1 [lower by *ca*. 20 ((syn) and 14 kcal mol⁻¹ (anti)] and with Br [by *ca*. 25 (syn) and 15 kcal mol⁻¹ (anti)]. A possible origin of this activation energy differences could be the more delocalized (covering whole cyclopropene ring) 1,2-dihalocyclopropyl anion-like TSs than the partially delocalized (covering only three carbon atoms out of five of the ring) 1,2-dihalocyclopentenyl anion-like TSs. In both cases, there is no involvement of aromatic vs antiaromatic character of small rings, since in the former the TS is cyclopropyl (not cyclopropenyl) anion like, whereas in the latter the TS has cyclopentenyl (not cyclopentadienyl) anion like structure.

The *n*-attack γ -S_N2 reactions. The nucleophile can substitute at a y-carbon with a double well potential energy surface leading either to a syn (γ -S_N2'-syn) or anti (γ -S_N2'anti) orientation to the C-X bond in the TS. No stable γ adduct was found at all levels of theory. Reference to Tables 1-3 reveals that the B3LYP ΔG^2 (ΔE^2) values for the γ -S_N2'syn paths with X = Cl and Br are higher than those for the γ - S_N2 '-anti paths in contrast to the lower $G2(+)MP2 \ \Delta G^{\tau}$ values (except for X = Br. $\Delta G^2(syn) \cong \Delta G^2(anti)$) indicating that the interorbital pair electron correlation effects between the lone pairs on the two halogen atoms of the eclipsing C-X bonds (and also lone pairs on C_2) in the γ -S_N2'-svn TSs are underestimated as in the β -S_N2'-syn paths (*vide supra*). The MP2 ΔG^{τ} values (Table 2) are all higher than those at the G2(+)MP2 level (Table 3) so that electron correlation effects for the γ -S_N2[•] TSs are underestimated at the MP2 level. This is in contrast to the overestimation of electron correlation effects at the MP2 level for the β -S_N2^o TSs due to strong electronic charge delocalized structures.^{14,15} This means that the electronic charge delocalization in the γ -S_N2 TSs is weaker than that in the β -S_N2[•] TSs in which delocalization occurs partially over four carbon atoms. C₂-C₅ (Scheme 2), as discussed above. The TS structures are shown in Figures

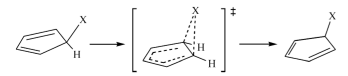


Scheme 3, γ -S_N2'-syn TS,

1-3. A lone pair develops at the β -(C₂) position in the γ -S_N2' TSs, scheme 3, a p type for the anti and an sp^3 type for the syn, and relatively strong n_{C} - σ_{C-X}^* vicinal charge transfer interactions occur and the C1-C2-C3 molety becomes a delocalized allyl anion. There are also very weak $\pi_{C-C} - \sigma^*_{C-X}$ charge transfer interactions; the $\Delta E_{ps}^{(2)}$ values range -8 (X - F) ~ -16 (X - Br) kcal mol⁻¹ whereas the $\Delta E_{ns}^{(2)}$ values are much greater than these ranging $-67 (X - F) \sim -219 (X$ - Br) kcal mol⁻¹. These latter $\Delta E_{n \sigma}^{(2)}$ charge transfer energies in the $\gamma S_{\lambda}2$ TSs are, however, considerably lower than the n_C- π^*_{C-C} interaction energies $(\Delta E_{n,\pi^*}^{(2)} - -198 \sim$ -334 kcal mol⁻¹) in the β -S₃2' TSs (Scheme 1) mainly due to wider energy gaps, $(\Delta \in - \in \sigma^* - \in n \text{ in eq. } 2)$ with the lower n levels $|e,g_{n}| \in [n] - [0.022 \text{ (syn)} \text{ and } 0.019 \text{ a.u. (anti) in}$ the γ -S_N2' TS $vs \in n - 0.025$ (syn) and 0.026 a.u. (anti) in the β -S_N2' TSs for X – CI and the higher σ^*_{C-F} than π^*_{C-C} levels. The $\gamma S_{\lambda}2$ TSs are somewhat looser with lower degree of bond formation and larger degree of bond cleavage than the β -S_{λ}2' TSs. For example all the C-X bonds for the γ - $S_{\lambda}2'$ in Figures 1-3 are longer than those for the β - $S_{\lambda}2'$ TSs. The negative charges on C_2 in the $\gamma S_N 2^{\circ}$ TSs are stronger (e.g., for X – Cl q_2 – -0.402 (syn) and -0.397 (anti)) than those on C₄ in the β -S_A2' TSs (e.g., for X – Cl, q₄ – -0.186 (syn) and -0.161 (anti)). This means that the lone pairs on C₂ in the γ -S_X2^{*} TSs are less delocalized (or more localized) than those on C₄ in the β -S_N2' TSs.

The proximate charge transfer stabilization $(-\delta \Sigma \Delta E_{\sigma \sigma}^{(2)})^{-1}$ in Table 4) of the $\gamma S_{\lambda}2'$ TS increases in the order X – F < Cl < Br. However, the barrier height decreases in the order Cl $(\Delta G^{+} \cong 19 \text{ kcal mol}^{-1}) > \text{Br}(17) > \text{F}(3-5);$ in other words the halide exchanges of fluoride have the lowest proximate charge transfer TS stabilization but have the highest reactivity. These trends of the lowest charge transfer delocalization with the highest reactivity for X – F in the γ $S_{\lambda}2'$ processes are similar to those found in the β - $S_{\lambda}2'$ processes. We interpret this highest reactivity with X – F as the stronger gain of the C-F bond energy relative to C-Cl and C-Br bonds in the β - as well as $\gamma S_{\lambda}2'$ TSs.

Although the differences in the ΔG^2 values between γ S_N2'-syn and anti processes are almost insignificant for X – Cl and Br, there is *ca*. 2 keal mol⁻¹ difference in favor of the γ S_N2'-syn for X – F. This appears to be caused by the

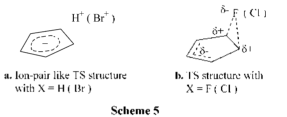


Scheme 4. 1.5-Sigmatropic rearrangements.

Table 5. Energetics⁶ for Circumambulatory Rearrangements of Cyclopentadienyl Halides. Calculated at Various Methods with $6-311+G^{**}$ Basis Sets (in keal mol⁻¹)

Method	Х	ΔE^{2}	$\Delta E^{2}_{ m ZPE}$	$\Delta H^{\ell h}$	$-T\Delta S^{\neq b}$	$\Delta G^{\neq h}$
B3LYP	П	27.52	25.47	25.26	-0.06	25.20
	F	44.87	43,43	43.34	0.25	43.59
	CI	27.83	26.88	26.70	0.39	27.10
	Br	21,91	21,11	20.87	0.49	21,36
MP2	H	25.32	23,19	22,99	-0.06	22,93
	F	52.84	51.66	51,51	0.37	51,88
	CI	34.37	33.79	33.52	0.61	34,14
	Br	26.93	26.36	26.08	0.65	26,72
G2(+)MP2 ^c	A	28.58	26.45	26.25	-0.06	26,19
	F	50.85	49.67	49,52	0.37	49,89
	CI	33.60	33.03	32.76	0.61	33.37
	Br	26.95	26.38	26.09	0.65	26,74

"Corrected for zero-point vibrational energy with appropriate scaling factors: 0.9806 for the B3LYP method, and 0.9748 for the MP2 method. "At 298 K. "The MP2 geometries were used, and thermal data were taken form the MP2/6-311+G" level.



stronger n_c - σ^*_{C-F} interaction in the syn (-68.8 kcal mol⁻¹) than anti (-66.7 kcal mol⁻¹) leading to stronger delocalized allyl anionic moiety (C₁-C₂-C₃) as evidenced by the lower negative charge on C₂ for syn (-0.609) than anti (-0.619). In fact the lower activation energy for the syn than anti paths by 2 kcal mol⁻¹ is the same as the difference in the n_c - σ^*_{C-F} charge transfer energy difference.

Circumambulatory rearrangements. The halogen atom in the cyclopentadienyl halides can undergo sigmatropic rearrangement by 1,5-shift and migrates around the periphery of the five-membered conjugated ring in a so-called monohapto (1,5-suprafacial) circumambulatory (merry-go-round) rearrangement reaction,3 Scheme 4. We have used four migrants (X - H, F, Cl and Br) and the activation energies calculated at the B3LYP, MP2 and G2(+)MP2 levels are summarized in Table 5. The ΔG^{2} values at the B3LYP level are lower than those corresponding values at the G2(+)MP2 level, due to the overestimation of electron correlation effects in the TSs. The barrier heights increase in the order $X - H \le Br \le Cl \le F$ at the MP2 and G2(+)MP2 levels, but the barrier height for X - H rises higher than that for X - Brat B3LYP level. Moreover, the ΔG^2 value for X – H (and Br) at the MP2 level is lower than that at the G2(+)MP2 level. The origin of this lower ΔG^2 values at the MP2 than G2(+)MP2 level for X – H is again the overestimation of electron correlation effects for the relatively strong charge delocalized TS structure^{14,15} with X - H at the MP2 level. The percentage bond order changes, $\%\Delta n^2$ (S4), reveal that

the TS with X = H has a particularly tighter structure (65%) bond formation; 35% bond cleavage) than those with other migrants (for X = F. Cl and Br. bond formation is *ca.* 40% and bond cleavage is ca. 60%). The NPA charges (S5) at the NBO-MP2/6-311+G** level show that negative charge delocalization over the ring is the strongest with X = H $(q_1 = q_2 = -0.340, q_3 = q_5 = -0.252 \text{ and } q_4 = -0.236)$ and the weakest with X = F ($q_1 = q_2 = +0.035$, $q_3 = q_5 = -0.365$ and $q_4 = -0.024$). Furthermore, the formal charge of the migrant in the TS (S5) is positive with $X = H (q_X = +0.341)$ and X = Br ($q_X = +0.033$), while it is negative for X = Fand Cl. These NPA charge analyses (S5) indicate that the TS structure with X = H (and to a smaller extent with X = Br) is nearly an ion pair type, (Scheme 5a) with charge delocalized cyclopentadienyl anionic (aromatic) ring. On the other hand, the TS with X = F (and to a lesser degree with X = CI) has a quadrupole structure (Scheme 5b) with little charge delocalization within the ring. The ion pair like TS with X = H (and with Br) is stabilized by an aromatic cyclopentadienyl anion like ring structure. This is in sharp contrast to the strong antiaromatic negative charge delocalized ring structure found in the sigantropic 1.3-shift for cyclopropene (X = H) with extremely high activation energy. $\Delta G^{2} = 88.5$ kcal mol⁻¹ at the G2(+)MP2 level.⁸ These trends of extremely high energy barrier to the 1,3-sigmatropic hydride shift on the cyclopropene ring in contrast to much lower barrier to the 1,5-sigmatropic hydride shift on the cyclopentadiene ring ($\Delta G^{\dagger} = 26.2$ kical mol⁻¹ at the G2(+)-MP2 level) are consistent with the experimental results; thermally induced 1.3-hydride shifts have not been reported for simply substituted cyclopropenes.^{3,18} whereas the corresponding 1.5-sigmatropic hydride shifts for cyclopentadiene are known to be facile at ca. 60 °C with a barrier height of $\Delta G^{\tau} = 24.3$ kcal mol^{-1,19} In fact circumambulation around evelopentadiene ring is not limited to hydride as the migrating group.³ A wide variety of other types of groups can undergo comparable thermally induced migration. Alkyl groups shift in general with substantially higher activation energies (e.g., methyl group shifts at ca. 200 °C with ΔG^{τ} = 41.8 kcal mol^{-1})²⁰ than hydride.

We have correlated the activation energy (ΔG^2 at the G2 (+)MP2 level) for the circumambulatory rearrangement of X

$$\Delta G^{2} = (0.29 \pm 0.03) \text{BE} + (11.79 \pm 0.59) \chi - (26.54 \pm 2.40)$$

(r = 0.999) (4)

(= H, F, Cl and Br) with bond energy (BE) of the C-X bond and electronegativity (χ) of X. eq. 4. For the cyclopentadienyl ring, both the BE and electronegativity (χ)²¹ contribute to elevate the barrier height.⁴ *i.e.*, the stronger the BE and the electronegativity, the barrier becomes higher. Alternatively, a migrant X with low electronegativity and weak C-X bond will readily form a ((C₅H₅)⁻ + X⁺) like TS (Scheme 5), which is stabilized by an aromatic cyclopentadienyl anion ring. By contrast, the same type of correlation with ΔG^{τ} (G2(+)MP2) values⁸ for circumambulatory rearrangement of X (= H, F, Cl and Br) in the cyclopropenvl ring, eq. 5, leads to a negative coefficient of χ . This means

$$\Delta G^{2} = (1.34 \pm 0.09) \text{BE} - (41.38 \pm 2.36) \chi + (36.16 \pm 8.76) (-r = 0.997)$$
(5)

that an aromatic cyclopropenyl cation- X^- ion pair like⁵ TS. (6). formation is facilitated by a strong electronegativity of X.



We therefore conclude that the lower activation energies with X = H and Br than with X = F and Cl are due to strong (delocalized) negative charge development in the ring (as evidenced by the overestimation of electron correlation effects at the MP2 level) in the TS leading to aromatic cyclopentadienyl anion ring structure.

Relative reactivities. For the gas-phase identity nucleophilic reaction pathways of fluoride anion with cyclopentadienyl fluoride, the *reactivity* increases at the G2(+)MP2 level in the order $S_N 2 \le \gamma \cdot S_N 2^*$ -anti $\le \gamma \cdot S_N 2^*$ -syn $\le \beta \cdot S_N 2^*$ -anti. The lowest reactivity of $S_N 2$ path with X = F is due to the weak antiaromatic cationic ring structure in the TS while the highest reactivity with $\beta \cdot S_N 2^*$ -anti can be attributed to the formation of a stable 1,2-difluorocyclopentenyl anion, which in turn is stabilized by the strong $n_C - \pi^* \in C$ charge transfer interaction with a delocalized allylic anion structure. The strong C-F bond energy in the adduct also contributes to its stability. The sigmatropic 1.5-shift of F has much higher activation barrier with 49.9 kcal mol⁻¹ than other fluoride exchange reaction paths so that can not interfere or compete with other reaction path.

For the reactions with X = CI, the σ -attack $S_N 2$ reaction path is the most favored and has the strongest reactivity; the reactivity increases in the order at the G2(+)MP2 level, β -S_N2'-syn $\leq \beta$ -S_N2'-anti $\leq \gamma$ -S_N2'-syn $\leq \gamma$ -S_N2'-anti \leq S_N2. The strongest reactivity of the σ -attack S_N2 path is ascribed to the weak aromatic cyclopentadienyl anion like TS. For X = CI, the two γ -S_N2ⁱ paths, *i.e.*, syn and anti, have similar reaction barriers. ca. 19 kcal mol⁻¹, which are lower than those for the β -S_N2' paths. The high energy barrier with β - S_{N2} -syn is due to smaller proximate charge transfer stabilization (Table 4). The signatropic 1,5-shift of chlorine atom has also a high energy barrier, 33.4 kcal mol⁻¹ at the G2(+)MP2 level, which is considerably higher than those for all halide exchange reaction paths except for the β -S_N2'-syn path which has almost the same barrier height of 33.6 kcal mol⁻¹. Thus the circumanbulatory rearrangement reaction of chlorine can not compete with halide exchange reactions in general, in particular with the σ -attack S_N2 path (ΔG^{τ} = 10.1 kcal mol^{-1}) which is favorable by more than 20 kcal mol^{-1} over the signatropic 1.5-shift.

For the reactions of X = Br, the reactivity trends are similar to those of X = Cl above: the σ -attack $S_N 2$ path is the most favored due to the aromatic cyclopentadienyl anion like TS structure. Again the two $\gamma S_N 2^{\prime}$ paths have similar energy barriers, *ca*. 17 kcal mol⁻¹. The β -S_N2'-syn path has the highest energy barrier (36.6 kcal mol⁻¹), which is even higher than the signatropic 1,5-shift of the bromine atom (26.7 kcal mol⁻¹). The much higher barrier for β -S_N2'-syn (by 20 kcal mol⁻¹) than for β -S₃/2 anti results from much lower proximate charge transfer energies [-278 (syn) vs -438 (anti) kcal mol⁻¹. Table 4]. For β -S_N2ⁱ bromide exchanges. the energy gaps between n_c and $\sigma_{\text{C-Br}}$ levels are 0.30 (syn) and 0.21 a.u. (anti) so that the $n_{\rm C}$ - $\sigma^{*}_{\rm C-Bt}$ charge delocalization energies are -57 (svn) vs -178 (anti) kcal mol⁻¹, and lead to large overall proximate charge transfer energy difference cited above. The activation barrier to the circumambulatory bromine migration is low with 26.7 kcal mol⁻¹, but is still higher than all the halide exchange paths except for β -S_N2'syn path. Therefore the Br migration can not compete with the halide exchanges, especially with the σ -attack S_N2 path. The reactivity increases in the order 1.5-shift $\leq \beta$ -S_N2'-svn \leq β -S_N2'-anti < γ -S_N2'-svn $\cong \gamma$ -S_N2'-anti < S_N2.

For the reactions with X = F, the β -S_N2' paths are the most preferred and the σ -attack S_N2 path is the most disfavored. In contrast, however, for the reactions with X = CI and Br, the most preferred path is the σ -attack S_N2 and the S_N2' paths are much less favored (by more than 8 kcal mol^{-1}) relative to the $S_{\lambda}2$ process. We therefore conclude that in the gas phase the chloride and bromide exchanges in the corresponding cvclopentadienvl halides (1) proceed by the σ -attack S_N2 path and the $S_N 2'$ processes should be difficult to compete and will constitute the minor reaction pathways. This conclusion supports the experimentally deduced S_N2 mechanism of Breslow et al. for the halide exchanges in cyclopentadienyl halides.¹ However, the possibility of competing the γ -S_N2' path (as well as the β -S_N2' path) with the $S_{\Lambda}2$ reaction raised by them can be disregarded.¹ Our results also show that reactions involving displacement on the halogen atom to form a dihalogen molecule leading to cyclopentadienvl anion formation is not possible in the gas phase.1

The gas phase fluoride exchange reaction exhibits entirely different trends compared to chloride and bromide exchanges. The TSs in the fluoride exchanges are mostly stabilized by an extra, albeit partial, strong C-F bond energy, which seems to provide much lower energy barriers to β -as well as γ -S_N2^{*i*} reactions but not in the S_N2 path since F is bound much looser (d^{*}_{C-F} = 1.856 Å vs d^{*}_{C-F} 1.438 and 1.541 Å in the β -and γ -S_N2^{*i*} TSs respectively).

The barrier heights at the G2(+)MP2 level increase in the order X = F < Br < C1 for all the gas-phase identity halide exchanges in the cyclopentadienyl halides. This order is the same as that found for the gas-phase identity methyl transfer reactions²² and also for the π -attack S_A2' paths of the halide exchange reactions of cyclopropenyl halides (5) at the G2(+) level of theory.⁸ In the latter reactions of 5, the barrier heights of the σ -attack S_A2 paths are strongly dependent on the development of positive charge in the cyclopropenyl ring, an aromatic cyclopropenium ion character, in the TS,

which decreases in the order X = Br > Cl > F so that the barrier heights increase in the order X = Br < Cl < F. The lowest degree of positive charge development with X = F in this case is the results of rather tighter TS with larger degree of bond formation and lower degree of bond cleavage than those with X = Cl and Br. Thus there is a compromise between a greater degree of aromatic ring formation (the looser the better) and the stronger bond energy gain (the tighter the better) for the $S_N 2$ TS with X = F.

Finally it is appropriate here to summarize the relationship between electron correlation effects calculated and delocalization within a TS or an intermediate. (i) Electron correlation enhances delocalization. Various resonance delocalized forms of a molecular species are already present at the uncorrelated MO level, but electron correlation provides larger weights to the resonance delocalized forms rather than new types of resonance delocalization. This is why the activation barriers to reactions with a delocalized TS become lower with correlated than uncorrelated MOs. Ample examples are presented by Weinhold and coworkers.^{6b,c} (ii) The electron correlation effects are in general underestimated at the MP2 level, but for electron delocalized structures the electron correlation effects are overestimated.^{14,15} We encounter often a reaction with delocalized TS, for which the electron correlation effects are overestimated and hence the activation energy is underestimated at the MP2 level relative to a higher level (e.g., G2 level) result. Examples: The lower β -S_N2^{*} reaction barriers with X = Cl and Br at the MP2 level (Table 2) than the corresponding values at the G2(+)MP2 level (Table 3). (iii) The electron correlation effects are overestimated in the DFT (e.g., B3LYP) energetics especially for a delocalized structure so that for reactions with a delocalized TS the activation energies are underestimated.¹⁵ However. the interorbital pair correlation energies between lone pairs on neighboring atoms are underestimated, and hence if such neighboring lone-pairs are present in the TS, the activation energy is overestimated. Examples: The higher DFT ΔG^{2} (ΔE^{2}) values (Table 1) than the corresponding values at the G2(+)MP2 level (Table 3) for β -S_N2'-syn (TS) and β -S_N2'anti (intermediate) processes.

These trends of accounting for electron correlation effects at lower levels of theory can therefore help understanding TS structures as the examples given above demonstrate.

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