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Electronic spectra and photodissociation of vinyl chloride: A symmetry-adapted cluster configuration interaction study

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The vertical absorption spectrum and photodissociation mechanism of vinyl chloride (VC) were studied by using symmetry-adapted cluster configuration interaction theory. The important vertical $\pi \rightarrow \pi^*$ excitation was intensively examined with various basis sets up to aug-cc-pVTZ augmented with appropriate Rydberg functions. The excitation energy for $\pi \rightarrow \pi^*$ transition obtained in the present study, 6.96 eV, agrees well with the experimental value, 6.7–6.9 eV. Calculated excitation energies along with the oscillator strengths clarify that the main excitation in VC is the $\pi \rightarrow \pi^*$ excitation. Contrary to the earlier theoretical reports, the results obtained here support that the C–Cl bond dissociation takes place through the n_{CI} - $\sigma^*_{\text{C-CI}}$ state. © 2006 American Institute of Physics. [DOI: 10.1063/1.2162539]

I. INTRODUCTION

Vinyl chloride (C₂H₃Cl) (VC) is an atmospheric pollutant. Evidence showed that VC causes cancer in human beings. Because of this, understanding of the photochemistry and spectroscopy of VC is important. There were many studies in this direction in the past; however, detailed studies to get a clear picture about the photochemical behavior of VC are still necessary, because the results obtained in some of the previous studies are ambiguous and speculative. Additionally, the conclusions made in a few recent studies are very much in contradiction with the existing predictions. For example, two recent theoretical studies^{1,2} assigned the π -3s Rydberg state as the optically bright state. It is completely contradictory to the generally agreed fact that for ethylenic compounds the strongest absorption is due to $\pi \rightarrow \pi^*$ transition.³ Another fact is that the VC excited to the bright π - π^* state has generally been thought to be predissociated through repulsive $n_{\rm Cl}$ $\sigma_{\rm C-Cl}^*$ state.^{4,5} Though a reassignment to a π - σ_{C-Cl}^* state has been recently suggested,⁶ it was promptly questioned lately.⁷ Surprisingly, while concluding that the dissociation of fast chlorine atoms occurs through the π - σ_{C-Cl}^* state, the role of the n_{Cl} - σ_{C-Cl}^* state in the dissociation was completely ignored in a recently published report.2

Studies focusing on spectroscopy^{3,4,8–18} and photodissociation mechanism^{4–7,19–24} of VC were made in the past. The bright peak at 6.7–6.9 eV has been considered and agreed to be due to the $\pi \rightarrow \pi^*$ excitation.^{4,8,10–12} Umemoto *et al.*⁵ studied the dissociation mechanism of a few chloroethylenes including VC. They concluded that two processes are competing: one is the transition to a dissociative $n_{\rm CI} - \sigma^*_{\rm C-CI}$ state which crosses the bright π - π^* state in the C-Cl coordinate and the other is the internal conversion to the ground state. The C-Cl bond cleavage resulted in the former process and HCl elimination occurred in the latter. Umemoto et al.⁵ speculated that twisting of the C=C bond leads to mixing of the π - π^* and π - σ^*_{C-Cl} states. Internal conversion to the π - σ_{C-Cl}^{*} state may serve as intermediate state from which internal conversion to the ground state occurs. It is obvious that the π - σ^*_{C-CI} state was considered as a bound state in that study. However, Tonokura et al.⁶ later speculated the π - σ_{C-C1}^* state as a repulsive state. From their calculated vertical excitation energies, they claimed that 193 nm (6.42 eV) photoexcitation is not sufficient to reach the crossing point of the π - π^* and $n_{\rm Cl}$ - $\sigma^*_{\rm C-Cl}$ states and hence they suggested that the π - σ_{C-Cl}^* state is responsible for the C–Cl bond breaking, but Blank et al.⁷ raised doubts over this assignment.

Chang and Chen¹ studied few low-lying electronic states of VC by using complete active space self-consistent field (CASSCF) and multireference configuration interaction (MRCI) theories. Their calculated vertical excitation energies for the $\pi \rightarrow \pi^*$ transition which is well established as responsible for the bright peak in the absorption spectrum are around 0.6-0.8 eV higher than the experimental values (strong peak). However, the authors assigned the π -3s Rydberg state as the optically bright state on the basis of their calculated adiabatic excitation energies. Through configuration interaction singles calculation,¹ a long tail observed at $45\ 000-54\ 000\ cm^{-1}\ (5.58-6.70\ eV)$ in the absorption spectrum⁴ was speculated due to the $1 {}^{3}A'' \leftarrow 1 {}^{1}A'$ and $2^{3}A'' \leftarrow 1^{1}A'$ transitions via spin-orbit coupling. As a result, Chang and Chen¹ also speculated that upon the excitation of VC at 193 nm (6.42 eV), the $1^{3}A''$ or $2^{3}A''$ excited state,

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instead of π - π^* , is initially prepared for the photodissociation processes. These speculations were contradictory to the existing reports. However, Chang² studied triplet excited states of VC using CASSCF and MRCI and from the calculated values he abandoned the earlier speculations that (1) the second and third triplet excited states might be responsible for the absorption spectra at $45\,000-54\,000\,\mathrm{cm}^{-1}$ (5.58-6.70 eV) and (2) the $1^{3}A''$ or $2^{3}A''$ state might involve in the photodissociation process. But, he maintains that the strong peak in the absorption spectra is due to the Rydberg $\pi \rightarrow 3s$ excitation. Chang² also studied the potentialenergy surface of the first singlet excited state of VC and concluded that the chlorine dissociation occurs via π - $\sigma^{*}_{\mathrm{C-Cl}}$ state followed by a crossing between π -3s and π - σ^*_{C-CI} states. These conclusions are inconsistent with the existing speculations.

In these circumstances, we made the present study on VC by using symmetry-adapted cluster²⁵/configuration interaction^{26–28} (SAC-CI) theory to clear the existing contradictions. Considering the importance of the photochemistry of chloroethylenes, comprehensive and accurate knowledge about vinyl chloride, the smallest molecule in the chloroethylene's family, is vital. The main focus of this paper is two-fold: (1) studying the electronic spectrum of VC by using SAC-CI to assign the excited states including the valence π - σ_{C-CI}^* , π - π^* , n_{CI} - σ_{C-CI}^* , π , and the Rydberg states and (2) understanding the C–Cl bond breaking mechanism in VC by studying the potential-energy surfaces (PESs) of its excited states including the important n_{CI} - σ_{C-CI}^* state.

II. COMPUTATIONAL DETAILS

All computations were performed with GAUSSIAN 03 suite of programs.²⁹ The vertical absorption spectra were obtained using SAC-CI singles and doubles (SD)-R method utilizing many different basis sets ranging from Dunning's double zeta [D95(d,p)] (Ref. 30) to augmented triple zeta type (aug-cc-pVTZ).³¹ The influence of polarization and diffuse functions was explored. Notations, for example, cc-pVTZ(-d) for H representing the basis set cc-pVTZ without *d* polarization function on H atoms and aug-cc-pVTZ(-f) for C and Cl atoms representing the basis set aug-cc-pVTZ without f diffuse function on C and Cl atoms are used here. Calculations were made using these basis sets which were augmented with two sets of Rydberg spd functions for carbon ($\zeta_s = 0.0437$ and 0.017 25, ζ_p =0.0399 and 0.015 75, and ζ_d =0.0285 and 0.011 25) and two sets for chlorine ($\zeta_s = 0.0475$ and 0.018 75, $\zeta_p = 0.0380$ and 0.0150, and $\zeta_d = 0.0285$ and 0.011 25). Chlorine Rydberg functions were placed on chlorine atom in all the calculations while carbon Rydberg functions were used in two ways: placing the functions (1) on both carbons and (2) on C=Cbond center (BC). The preliminary PES of several low-lying excited states along the C–Cl bond and C= CH_2 twisting coordinates were obtained mainly by using D95(d,p) basis set in this study. Electronic spectra calculations were done within C_s symmetry, with the molecule placed in the xyplane.

III. RESULTS AND DISCUSSIONS

Before going to the calculations on VC, we studied $\pi \rightarrow \pi^*$ transition of ethylene to examine the basis sets to study VC. Since VC is similar to ethylene and there were many theoretical studies on ethylene, such examination is meaningful. The strong peak in the excitation spectrum of ethylene is the broadband with the maximum at 7.65 (Ref. 32) or 7.66 eV,³³ which arises from the $\pi \rightarrow \pi^*$ transition. This broadband is overlapped by several Rydberg states. While the exact vertical excitation energy of this transition has not been obtained experimentally, high-level theoretical studies have converged to a value of 8 eV.³⁴ Finally, this energy has been estimated to fall in between 7.90 and 7.95 eV on the basis of sophisticated theoretical calculations.^{35,36}

The results obtained for the $\pi \rightarrow \pi^*$ transition of ethylene in the present study were tabulated in Table I. The table clearly shows that the SAC-CI excitation energies are very accurate and are in close agreement with the available theoretical results. Preliminary calculations were performed on VC to obtain the excitation energies for the $\pi \rightarrow \pi^*$ transition. The results were presented in Table II. These calculations conclude the following two points: (1) Though the results obtained by placing the Rydberg diffuse functions either on the carbon atoms or on the C=C bond center do not have much differences, the former has a slight edge over the latter and (2) aug-cc-pVTZ basis set for heavy atoms seems very essential to obtain accurate energies, while cc-pVTZ basis set, even without *d*-type polarization function, is enough for hydrogen atoms. By keeping these points in mind, we performed the calculations of the vertical excitation spectrum in the wide energy region of VC.

Table III shows the vertical excitation energies and oscillator strengths calculated using SAC-CI with LevelThree, along with the results obtained in the previous theoretical and experimental studies for the singlet excited states in each (A' and A'') symmetry. The lowest excited state can be better denoted as π -3s/ σ_{C-CI}^{*} since a small contribution from $\pi \rightarrow \sigma_{C-CI}^{*}$ transition is also involved in this Rydberg excitation.

The second excited state represents $\pi \rightarrow \pi^*$ excitation. The excitation energy, 6.96 eV, obtained for this transition is close to the experimental values, 6.7-6.9 eV. On the other hand, the values derived from MRCI (Refs. 1 and 6) were 0.5-0.76 eV higher than the experimental values. Tonokura et al.⁶ used cc-pVTZ basis set with a set of s and p diffuse functions on chlorine in their MRCI calculations, while Chang and Chen¹ adopted 6-311+G(d,p) basis set with a diffuse sp function on each carbon. The present calculations reveal that the basis sets used in those two MRCI studies were not sufficient to properly describe $\pi \rightarrow \pi^*$ excitations. Using the basis set by Chang and Chen,¹ we performed the SAC-CI calculations on ethylene and VC. The excitation energy for $\pi \rightarrow \pi^*$ transition in VC was calculated to be 7.32 eV. Though this value is better than that of MRCI with the same basis set (7.46 eV), it is far from our best SAC-CI value (6.96 eV). On the basis of their calculation, Chang and Chen¹ assigned the maximum at 6.71 eV in the absorption spectrum is due to the $\pi \rightarrow 3s$ Rydberg transition. They proposed that the observed peaks at 6.82 and 6.88 eV are pos-

TABLE I. Ethylene $\pi \rightarrow \pi^*$ vertical excitation energies (ΔE in eV) using the SAC-CI method with various basis sets at experimental geometry (Ref. 37).

	Ba	sis set	Rydberg function		
Method	Н	С	(two sets of <i>spd</i>)	ΔE	
	DZP	DZP	On BC	8.195	
	cc-pVTZ	cc-pVTZ	On BC	8.027	
	cc-pVTZ	aug-cc-pVTZ	On BC	7.949	
	cc-pVTZ	aug-cc-pVQZ	On BC	7.922	
	DZP	DZP	On C	8.142	
SAC-CI	cc-pVTZ	cc-pVTZ	On C	8.098	
	cc-pVTZ(-pd)	aug-cc-pVTZ(-f)	On C	8.029	
	cc-pVTZ(-pd)	aug-cc-pVTZ	On C	7.981	
	cc-pVTZ(-d)	aug-cc-pVTZ	On C	7.959	
	cc-pVTZ	aug-cc-pVTZ(-f)	On C	7.967	
	cc-pVTZ	aug-cc-pVTZ	On C	7.943	
CASPT2				8.00^{a}	
Best theory estimate				7.90–7.95 ^b	
Expt.				7.65, ^c 7.66 ^d	

^aReference 34.

^bReferences 35 and 36.

^cReference 32.

^dReference 33.

sible candidates for the $\pi \rightarrow \pi^*$ excitation. They used their calculated adiabatic excitation energy, 6.82 eV, to propose the π - π^* state assignment. This may be due to the fact that their calculated vertical excitation energy, 7.46 eV, of the same state is far away from the experimental values. However, Chang² raised doubts over this calculated adiabatic excitation energy of π - π^* state.¹ He found that the π - π^* state has the lowest energy at 90° (completely different from their first report), however, he neither mentioned the exact adiabatic energy nor did he make a reassignment for $\pi \rightarrow \pi^*$ excitation. According to that study,² the adiabatic energy of π - π^* state was around 5 eV!

As mentioned earlier, the π - π^* vertical excitation energy

obtained in the present study (6.96 eV) agrees very well with the experimental value (6.7-6.9 eV). Our results show that the oscillator strength is mainly carried by π - π^* transition in the vertical region (Table III). These results clearly support the fact that the maximum in the absorption spectrum is due to the $\pi \rightarrow \pi^*$ excitation. However, the oscillator strengths obtained in the previous study using CASSCF level of theory for $\pi \rightarrow 3s$ and $\pi \rightarrow \pi^*$ transitions are close to each other (0.0515 and 0.0818).¹

Walsh³⁹ and Merer and Mulliken⁴⁰ analyzed the π $\rightarrow \pi^*$ transition in ethylene and concluded that the π - π^* state has an equilibrium geometry which is twisted 90° about the C=C bond with the CH_2 groups deformed. Walsh³⁹ pre-

TABLE II. Vinly chloride $\pi \rightarrow \pi^*$ vertical excitation energies (ΔE in eV) using the SAC-CI method with various basis sets at experimental geometry (Ref. 38).

	Basis set		Rydberg function	
Method	Н	C and Cl	(two sets of <i>spd</i>)	ΔE
	DZP	DZP	On BC and on Cl	7.432
	cc-pVTZ	cc-pVTZ	On BC and on Cl	7.211
	cc-pVTZ	aug-cc-pVTZ(-f)	On BC and on Cl	7.116
	DZP	DZP	On C and Cl	7.351
SAC-CI	cc-pVTZ	cc-pVTZ	On C and Cl	7.155
	cc-pVTZ(-pd)	aug-cc-pVTZ(-f)	On C and Cl	7.134
	cc-pVTZ(-pd)	aug-cc-pVTZ	On C and Cl	7.065
	cc-pVTZ(-d)	aug-cc-pVTZ	On C and Cl	7.028
	cc-pVTZ	aug-cc-pVTZ(-f)	On C and Cl	7.068
	cc-pVTZ	aug-cc-pVTZ	On C and Cl	7.103
Expt.				6.70, ^{a,b} 6.90, ^c
-				6.72, ^d 6.74 ^e

^cReference 10.

^dReference 11.

^eReference 12.

TABLE III. Several lowest-lying singlet excited states of C₂H₃Cl obtained using SAC-Cl with cc-pVTZ(-d) for H and aug-cc-pVTZ for C and Cl at experimental geometry (Ref. 38). Two sets of Rydberg *spd* functions for both carbons and for chlorine were added. Excitation energies (ΔE), oscillator strengths (f), and second moments ($\langle r^2 \rangle$). Note that the Rydberg 3*s*, 3*p*, and 3*d* orbitals are predominantly attributed to carbons while the Rydberg 4*s* and 4*p* orbitals are attributed to chlorine.

	Nature	SAC-CI			ΔE (eV)	
State		ΔE (eV)	f	$\langle r^2 \rangle$	Other theory	Expt.
$1^{1}A'$	Ground state			245.4		
$ ^{1}A''$	π -3s/ $\sigma^*_{\rm C-Cl}$	6.81	0.0111	274.4	6.68 ^a	
2 A'	π - π^*	6.96	0.3274	257.1	7.46 ^a	6.7, ^{b,c} 6.9, ^d
					7.40 ^e	6.72, ^f 6.74 ^g
$^{1}A''$	$\pi - \sigma^*_{C-CI}/3s$	6.99	0.0035	268.0	7.27 ^a	
	0-01				7.26 ^e	
$^{1}A''$	π -3 $p\sigma_x$	7.48	0.0006	297.6	7.48^{a}	
$^{1}A''$	π -3 $p\sigma_v$	7.70	0.0102	312.6	7.56 ^a	
${}^{1}A''$	$n_{\rm Cl} - \pi^*$	7.82	0.0005	251.6	7.9 ^e	
$^{1}A'$	π -3 $p\pi$	7.89	0.0098	309.7		7.817, ^h 7.82 ⁱ
$^{1}A'$	$n_{\rm Cl} - \sigma^*_{\rm C-Cl}$	8.20	0.0012	256.2	8.50 ^e	
${}^{1}A''$	π -3 $d\sigma/4s$	8.42	0.0152	334.8		8.33 ^{h,i}
${}^{1}A''$	π -3 $d\sigma/4p\sigma_x$	8.46	0.0001	354.4		$8.38^{h,i}$
$^{1}A'$	π -3 $d\pi/4p\pi$	8.57	0.0050	362.2		8.52 ^h
$^{1}A'$	$n_{\rm Cl}-3s$	8.60	0.0465	293.0		8.51 ^{h,i}

^aReference 1.

^bReference 4.

^cReference 8. ^dReference 10.

^eReference 6.

^fReference 11.

^gReference 12.

^hReference 15.

ⁱReference 18.

dicted that the Raman spectrum of ethylene would show activity in the C=C bond stretching mode, the C=C twist, and various CH₂ bends and deformations. These predictions were confirmed by Ziegler and Hudson⁴¹ and Sension and Hudson.³³ Like ethylene, the emission spectra of VC obtained by Browning *et al.*¹⁶ also indicated C=C stretching and twisting motion as well as CH₂ bending or deformation, verifying that excitation has $\pi \rightarrow \pi^*$ character. All these facts support that the strong peak in the absorption spectrum is due to the $\pi \rightarrow \pi^*$ excitation.

Excitation energies obtained for other states also agree very well with the available experimental values (Table III). The π - σ_{C-Cl}^* , n_{Cl} - π^* , and n_{Cl} - σ_{C-Cl}^* states are assigned at 6.99, 7.82, and 8.20 eV, respectively.

We also investigated a few low-lying triplet states by using SAC-CI with the same basis set used to study the singlet states. It is worth to mention here that the vertical excitation energy of 4.12 eV obtained for ${}^{3}(\pi - \pi^{*})$ state is very close to the experimental value, 4.13 eV,¹² whereas it is 4.25 eV in MRCI calculation.² The detailed results regarding excitations to the triplet states will be published in a forthcoming article.

Chang² studied the potential-energy surface of the first excited state of VC by using CASSCF. It showed that departing from the planar equilibrium structure through a barrier, the first excited state is mainly of π - σ^*_{C-Cl} character as the C-Cl distance increases, whereas it is of π - π^* character as the twisting angle increases. This is as a result of the forma-

tion of avoided crossings between π -3s which has been assigned as the bright state by Chang and Chen¹ and π - σ_{C-Cl}^* states at planar geometries (both have the same *A''* symmetry) and between π -3s and π - π^* states at nonplanar structures (all states have the same *A* symmetry). Accordingly, Chang² concluded that the dissociation of Cl takes place via two pathways: one is through π - σ_{C-Cl}^* at planar leading to fast Cl and the other through π - π^* at twisted geometries from which internal conversion to the ground state and subsequent dissociation produces slow Cl.

Our results, however, show some interesting differences. The PES of some excited states along the C–Cl bond length and C==C bond twisting were depicted in Figs. 1 and 2, respectively. For this purpose, single point calculations on different C–Cl bond lengths and twisting angles were performed at SAC-CI/d95(d, p) level with all other internal coordinates were kept at the ground-state values optimized at SAC/d95(d) level. The orbital characters given in Table IV show that σ^*_{C-Cl} has an antibonding nature along C–Cl bond. The nature of the π - σ^*_{C-Cl} state was in debate in literature^{5,6} as mentioned in the Introduction section. The present study clearly shows that this state is a repulsive state as shown in Fig. 1(a).

The PES along the C–Cl bond length clearly reveals the formation of an avoided crossing between the bright π - π^* and n_{Cl} - $\sigma^*_{\text{C-Cl}}$ states (both have the same symmetry A') at around 1.85 Å. This reveals that the excited VC transfers

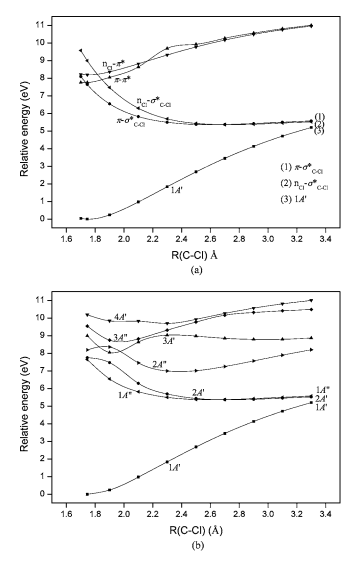


FIG. 1. PES along the C–Cl bond with the other geometrical parameters optimized for the ground state: (a) diabatic PES of π - σ_{C-Cl}^* , π - π^* , n_{Cl} - π^* , and n_{Cl} - σ_{C-Cl}^* states and (b) adiabatic PES of several low-lying excited states.

from its bound π - π^* state to the dissociative n_{Cl} - $\sigma^*_{\text{C-Cl}}$ state through level crossing. So dissociation can take place through this repulsive n_{Cl} - $\sigma^*_{\text{C-Cl}}$ state.

The PES along torsion shows that the low-lying two states π - σ^*_{C-Cl} and π - π^* start mixing from the very beginning of torsion. As the twisting angle increases, the mixing becomes stronger. As a result, the π - π^* character of the first excited state increases and simultaneously the π - σ_{C-Cl}^* character of the second and third states dominates at higher twisting angles. It reflects the bound nature of the π - σ_{C-C1}^* state and the dissociation of VC could occur through the well of π - π^* state or through the ground state followed by internal conversion from π - π^* to the ground state. However, the mixing of the π - π^* and π - σ^*_{C-C1} states due to twisting could possibly lead into fast dissociation of chlorine atoms on π - σ_{C-Cl}^{*} state which has a repulsive nature along C–Cl bond coordinate. So there is a possibility for the C-Cl bond dissociation through π - σ_{C-Cl}^{*} state in the C–Cl bond coordinate after it started mixing with π - π^* state.

Earlier experimental studies^{6,42} on VC concluded that the

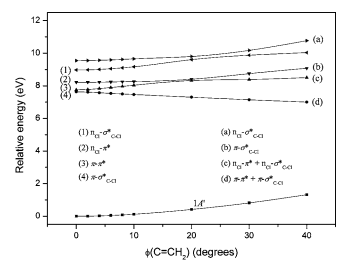


FIG. 2. PES of several low-lying excited states along C = C twisting with the other geometrical parameters optimized for the ground state.

dissociation from the excited state is much faster $(40 \pm 10 \text{ fs})$ than from the ground state (110 ps). That means the fast dissociation of chlorine atoms resulted from the repulsive excited state and not from the ground state followed by twisting and internal conversion. So it is obvious that the dissociation from the ground state has less importance.

The forgoing facts reveal the following: The main excitation is the $\pi \rightarrow \pi^*$ transition and at planar geometries along C-Cl bond coordinate, the bright π - π^* state forms an avoided crossing with the $n_{\rm Cl}$ - $\sigma^*_{\rm C-Cl}$ state which leads to fast C-Cl bond dissociation. Twisting causes mixing of π - π^* state with π - $\sigma^*_{\mathrm{C-Cl}}$ state, however, dissociation from the π - σ^*_{C-Cl} state after initial twisting may not have much importance because the fast dissociation takes place at the planar structure and the dissociation takes place within a time much shorter than molecular twisting.^{6,7,42} But there may be a small possibility for this π - σ_{C-CI}^* state to be involved in the C-Cl dissociation. There is a small transition from the ground state to the Rydberg 3s state (see Table III) in the excitation. This state, π -3s, could form an avoided crossing with the dissociative π - σ_{C-C1}^* state in the planar structure (both have the same A'' symmetry), as mentioned by Chang,¹ which could lead to fast dissociation like that from the repulsive $n_{\rm Cl}$ - $\sigma^*_{\rm C-Cl}$ state. However, the oscillator strength for π $\rightarrow 3s$ transition is very small compared with that for π $\rightarrow \pi^*$ transition and hence it is obvious that the dissociation through this π - σ_{C-Cl}^* state has a minor contribution. Again, the C-Cl bond dissociation through the ground state does not have much importance. All these facts suggest that the fast dissociation of the C-Cl bond takes places mainly through the repulsive $n_{\rm Cl}$ - $\sigma_{\rm C-Cl}$ state.

TABLE IV. Nature of π , π^* , σ^*_{C-Cl} , and n_{Cl} orbitals. (d) Dominant.

Nature	C=C	C–Cl
π	Bonding (d)	Antibonding
π^{*}	Antibonding (d)	Antibonding
$\sigma^*_{ ext{C-Cl}}$	Antibonding	Antibonding (d)
$n_{\rm Cl}$	Antibonding	Bonding (d)

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Chang² studied the first singlet excited state to understand the dissociation process, since presumably the dissociation could take place through this state. It should be mentioned here that our results show that the first two states π - σ^*_{C-Cl} and n_{Cl} - σ^*_{C-Cl} are energetically very close once the C-Cl bond length reaches at around 2.4 Å and, in fact, the n_{Cl} - σ^*_{C-Cl} state becomes the first excited state when the C-Cl bond length increases to around 3 Å (see Fig. 1).

The present PES calculation does not include Rydberg states; however, we believe that the inclusion of Rydberg states in the PES will not have any adverse effect beyond the Frank-Condon region and we expect that the C–Cl bond dissociation would occur on valence states. Results of excitations to the higher Rydberg states and the detailed two-dimensional (2D) PES of low-lying excited states, including Rydberg states, incorporating both C–Cl bond length and C==C twisting at the excited-state geometries will be presented in a future publication.

IV. CONCLUDING REMARKS

The vertical excitation energies obtained using SAC-CI with the necessary basis sets agree very well with the experimental values. The present study not only clarified the existing ambiguity in the assignment for the optically bright state of VC but also gave the confidence that the same level of theory could be used to obtain the theoretical electronic spectra of other chlorinated ethylenes, which would be useful to assign the observed spectra of these important atmospheric pollutants.

In the case of photodissociation, the optically bright π - π^* state forms an avoided crossing with n_{Cl} - $\sigma^*_{\text{C-Cl}}$ state which leads to fast C–Cl bond dissociation. Though π - $\sigma^*_{\text{C-Cl}}$ is a repulsive state, its participation in the dissociation is expected to be small.

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- ¹J.-L. Chang and Y.-T. Chen, J. Chem. Phys. **116**, 7518 (2002).
- ²J.-L. Chang, J. Chem. Phys. **122**, 194321 (2005).
- ³M. B. Robin, *Higher Excited States of Polyatomic Molecules* (Academic, New York, 1975), Vol. 2, p. 56.
- ⁴M. J. Berry, J. Chem. Phys. **61**, 3114 (1974).
- ⁵M. Umemoto, K. Seki, H. Shinohara, U. Nagashima, N. Nishi, M. Kinoshita, and R. Shimada, J. Chem. Phys. **83**, 1657 (1985).

- ⁶K. Tonokura, L. B. Daniels, T. Suzuki, and K. Yamashita, J. Phys. Chem. A **101**, 7754 (1997).
- ⁷D. A. Blank, W. Sun, A. G. Suits, Y. T. Lee, S. W. North, and G. E. Hall, J. Chem. Phys. **108**, 5414 (1998).
- ⁸A. D. Walsh, Trans. Faraday Soc. **41**, 35 (1945).
- ⁹S. P. Sood and K. Watanabe, J. Chem. Phys. 45, 2913 (1966).
- ¹⁰ J. H. Moore, Jr., J. Phys. Chem. **76**, 1130 (1972).
- ¹¹A. Kuppermann, W. M. Flicker, and O. A. Mosher, Chem. Rev. (Washington, D.C.) **79**, 77 (1979).
- ¹²C. F. Koerting, K. N. Walzl, and A. Kuppermann, Chem. Phys. Lett. 109, 140 (1984).
- ¹³K. H. Sze, C. E. Brion, A. Katrib, and B. El-Issa, Chem. Phys. **137**, 369 (1989).
- ¹⁴ R. Locht, B. Leyh, K. Hottmann, and H. Baumgartel, Chem. Phys. 220, 207 (1997).
- ¹⁵B. A. Williams and T. A. Cool, J. Phys. Chem. **97**, 1270 (1993).
- ¹⁶ P. W. Browning, D. C. Kitchen, M. F. Arendt, and L. J. Butler, J. Phys. Chem. **100**, 7765 (1996).
- ¹⁷J.-L. Chang, J.-C. Shieh, J.-C. Wu, R. Li, and Y.-T. Chen, Chem. Phys. Lett. **325**, 369 (2000).
- ¹⁸J.-L. Chang, R. Li, J.-C. Wu, J.-C. Shieh, and Y.-T. Chen, J. Chem. Phys. **115**, 5925 (2001).
- ¹⁹A. Fahr and A. H. Laufer, J. Phys. Chem. **89**, 2906 (1985).
- ²⁰ Y. Mo, K. Tonokura, Y. Matsumi et al., J. Chem. Phys. 97, 4815 (1992).
- ²¹ Y. Huang, Y.-A. Yang, G.-X. He, and R. J. Gordon, J. Chem. Phys. **99**, 2752 (1993).
- ²² Y. Huang, Y.-A. Yang, G. He, S. Hashimoto, and R. J. Gordon, J. Chem. Phys. **103**, 5476 (1995).
- ²³ T. L. Myers, D. C. Kitchen, B. Hu, and L. J. Butler, J. Chem. Phys. **104**, 5446 (1996).
- ²⁴K. Sato, S. Tsunashima, T. Takayanagi, G. Fujisawa, and A. Yokoyama, J. Chem. Phys. **106**, 10123 (1997).
- ²⁵H. Nakatsuji and K. Hirao, J. Chem. Phys. **68**, 2053 (1978).
- ²⁶H. Nakatsuji, Chem. Phys. Lett. **59**, 362 (1978).
- ²⁷H. Nakatsuji, Chem. Phys. Lett. **67**, 329 (1979); **67**, 334 (1979).
- ²⁸H. Nakatsuji, Acta Chim. Hung. **129**, 719 (1992); *Computational Chemistry—Review of Current Trends* (World Scientific, Singapore, 1997), Vol. 2, pp. 62–124.
- ²⁹ M. J. Frisch, G. W. Trucks, H. B. Schlegel *et al.*, GAUSSIAN 03, Revision B.01, Gaussian, Inc., Pittsburgh, PA, 2003.
- ³⁰ S. Huzinaga, J. Chem. Phys. **42**, 1293 (1965); T. H. Dunning, Jr., *ibid.* **53**, 2823 (1970).
- ³¹ T. H. Dunning, Jr., J. Chem. Phys. **90**, 1007 (1989); D. E. Woon and T. H. Durning, Jr., *ibid.* **98**, 1358 (1993).
- ³² A. J. Merer and R. S. Mulliken, Chem. Rev. (Washington, D.C.) **69**, 639 (1969); R. S. Mulliken, J. Chem. Phys. **66**, 2448 (1977); **71**, 556 (1979).
- ³³R. J. Sension and B. S. Hudson, J. Chem. Phys. **90**, 1377 (1989).
- ³⁴R. Lindh and B. O. Roos, Int. J. Quantum Chem. **35**, 813 (1989).
- ³⁵ J. D. Watts, S. R. Gwaltney, and R. J. Bartlett, J. Chem. Phys. **105**, 6979 (1996).
- ³⁶S. Krebs and R. J. Buenker, J. Chem. Phys. **106**, 7208 (1997).
- ³⁷E. Hirota, Y. Endo, S. Saito, K. Yoshida, I. Yamaguchi, and K. Machida, J. Mol. Spectrosc. **89**, 223 (1981).
- ³⁸I. Merke, L. Poteau, G. Wlodarczak, A. Bouddou, and J. Demaison, J. Mol. Spectrosc. **177**, 232 (1996).
- ³⁹ A. D. Walsh, J. Chem. Soc. **1953**, 2353.
- ⁴⁰ A. J. Merer and R. S. Mulliken, J. Chem. Phys. **50**, 1026 (1969).
- ⁴¹L. D. Ziegler and B. S. Hudson, J. Chem. Phys. **79**, 1197 (1983).
- ⁴² P. Farmanara, V. Stert, and W. Radloff, Chem. Phys. Lett. **288**, 518 (1998).