Ab initio study of charge-transfer dynamics in collisions of C$_{2}^{+}$ ions with hydrogen chloride

E. Rozsályi, 1 E. Bene, 2 G. J. Halász, 3 Á. Víbók, 1 and M. C. Bacchus-Montabonel 4

1Department of Theoretical Physics, University of Debrecen, P.O. Box 5, H-4010 Debrecen, Hungary
2Institute of Nuclear Research, Hungarian Academy of Sciences, P.O. Box 51, H-4001 Debrecen, Hungary
3Department of Information Technology, University of Debrecen, P.O. Box 12, H-4010 Debrecen, Hungary
4Laboratoire de Spectrométrie Ionique et Moléculaire, Université de Lyon (Lyon I), CNRS-UMR5579, 43 Bd. du 11 Novembre 1918, F-69622 Villeurbanne Cedex, France

(Received 22 March 2011; published 31 May 2011)

**I. INTRODUCTION**

Important experimental and theoretical effort has been devoted recently in order to investigate collisions of slow multiply charged ions with molecular targets. These studies provide detailed analysis on electron capture processes occurring during the collision and on fragmentation dynamics of the ionized species after removal of electrons from the target. A special attention has been focused first on reactions with simple ionized species after removal of electrons from the target. A ring during the collision and on fragmentation dynamics of the multiply charged ions with molecular targets. These studies devoted recently in order to investigate collisions of slow ions with molecular targets. The mechanism has been investigated in detail in connection with avoided crossings between states involved in the reaction. A simple mechanism driven by a strong nonadiabatic coupling matrix element has been pointed out for this process. A comparative analysis with the halogen fluoride target corresponding to a similar electronic configuration shows a quite different charge-transfer mechanism leading to a very different behavior of the cross sections. Such behavior may be correlated to specific nonadiabatic interactions observed in these collision systems.

**Ab initio** quantum chemistry molecular calculations followed by a semiclassical dynamical treatment in the keV collision energy range have been developed for the study of the charge-transfer process in collisions of C$_{2}^{+}$ ions with hydrogen chloride. The mechanism has been investigated in detail in connection with avoided crossings between states involved in the reaction. A simple mechanism driven by a strong nonadiabatic coupling matrix element has been pointed out for this process. A comparative analysis with the halogen fluoride target corresponding to a similar electronic configuration shows a quite different charge-transfer mechanism leading to a very different behavior of the cross sections. Such behavior may be correlated to specific nonadiabatic interactions observed in these collision systems.

DOI: 10.1103/PhysRevA.83.052713 PACS number(s): 34.70.+e
10 core-electron) relativistic pseudopotential has been used for chlorine [18] with the correlation-consistent aug-cc-pVTZ (augmented correlation-consistent polarized valence-triple-zeta) basis sets of Dunning [19]. The same basis set has been chosen for carbon and hydrogen atoms. Calculations have been performed using state-averaged CASSCF-MRCI (complete active space self-consistent field-multireference configuration interaction) methods. The active space includes the 1$s$ orbital of hydrogen, the 1$s$ of chlorine, the core electrons being treated by a pseudopotential. The 1$s$ orbital of carbon has been frozen in the calculation. The geometry of the $1\Sigma^+$ ground state of HCl has been optimized at the CASSCF (complete active space self-consistent field) level of theory leading to an equilibrium distance $r_{eq} = 1.2748$ Å in good agreement with the experimental value 1.2746 Å [20]. The corresponding vertical ionization potential calculated using MRCI (multireference configuration interaction) methods is 12.69 eV, to be compared to the experimental value 12.748 eV obtained from photoelectron spectra measurements [20,21]. The asymptotic energies of the $C^2+$–HCl molecular system may be compared to separated species calculations taking account of experimental ionization potentials and carbon ion levels [20–22] combined with MRCI calculations at optimized equilibrium geometry for the HCl and $C^2+$ ground and excited states. The calculated asymptotic energy difference between the entry channel and the highest $1\Sigma^+ (C(1s^22s^22p)^2D + HCl^+(\Sigma^+))$ exit channel is 2.35 eV, with a discrepancy of 0.06 eV with experimental data assuming a good description of both entry and exit channels. The asymptotic energy difference of the two lowest charge-transfer levels $1\Sigma^+ (C(1s^22s^22p)^2P^o + HCl^+(\Pi^o))$ and $1\Sigma^+ (C(1s^22s^22p)^2P^o + HCl^+(\Sigma^+))$ is 4.15 eV, in agreement with calculated separated species. The spin-orbit effects can be neglected in the collision energy range of interest so electron spin may be conserved in the collision process and only singlet states will be correlated to the $1\Sigma^+$ entry channel.

Charge-transfer processes are driven mainly by nonadiabatic interactions in the vicinity of avoided crossings [23,24], in particular between the entry channel and charge-transfer levels. The radial coupling matrix elements between all pairs of states of the same symmetry have thus been calculated by means of the finite difference technique [25]:

$$g_{KL} (R) = \langle \psi_K | \partial / \partial R | \psi_L \rangle = \langle \psi_K (R) | \lim_{\Delta \to 0} \frac{1}{\Delta} [\psi_L (R + \Delta) - \psi_L (R)] \rangle ,$$

which, taking account of the orthogonality of the eigenfunctions $|\psi_K (R)\rangle$ and $|\psi_L (R)\rangle$ for $K \neq L$, reduces to

$$g_{KL} (R) = \langle \psi_K | \partial / \partial R | \psi_L \rangle = \lim_{\Delta \to 0} \frac{1}{\Delta} \langle \psi_K (R) | \psi_L (R + \Delta) \rangle .$$

The parameter $\Delta$ has been tested and the value $\Delta = 0.0012$ a.u. has been chosen as in previous calculations [26] using the three-point numerical differentiation method for reasons of numerical accuracy.

The interaction between $1\Sigma^+$ and $1\Pi$ states by means of rotational coupling has also been taken into account. The rotational coupling matrix elements $\langle \psi_K (R) | L_z | \psi_L (R) \rangle$ between states of angular moment $\Delta \Lambda = \pm 1$ have been calculated directly from the quadrupole moment tensor from the expression $i L_z = x \partial / \partial z - z \partial / \partial x$ with the center of mass of the system as the origin of electronic coordinates [27].

Taking account of the $1\Sigma^+$ symmetry of the $C^2+(1s^22s^2)1S + HCl(1\Sigma^+)$ entry channel, only $C^+(2P^o)$ or $C^+(2D)$ states could be involved in the collision process. Effectively, $C^+(2P)$ states could lead only to triplet and quintet states, which cannot be correlated to the entry channel as spin-orbit coupling is negligible. With regard to the different excited states of HCl, there are thus four $1\Sigma^+$ states, which can be correlated by means of radial coupling, the entry channel $4\Sigma^+\{C^+(1s^22s^22p)^2D + HCl(1\Sigma^+)\}$ and three charge-transfer levels $1\Sigma^+\{C^+(1s^22s^22p)^2P^o + HCl(1\Pi^o)\}$, $2\Sigma^+\{C^+(1s^22s^22p)^2P^o + HCl(2\Sigma^+)\}$, $3\Sigma^+\{C^+(1s^22s^22p)^2D + HCl(2\Pi^o)\}$. We have also to take into account the $1\Pi$ states, which can be correlated to the $1\Sigma^+$ entry channel by rotational coupling interaction; all charge-transfer levels may be considered: $1\Pi\{C^+(1s^22s^22p)^2P^o + HCl(2\Pi^o)\}$, $2\Pi\{C^+(1s^22s^22p)^2P^o + HCl(2\Sigma^+)\}$, $3\Pi\{C^+(1s^22s^22p)^2D + HCl(2\Pi^o)\}$. The corresponding potential-energy curves have been calculated in the [2.0-14.0] a.u. internuclear distance range. They are presented in Fig. 2 for the equilibrium distance in the linear geometry. The main feature is a very strong avoided crossing between the $1\Sigma^+$ entry channel and the $3\Sigma^+\{C^+(1s^22s^22p)^2D + HCl(2\Pi)\}$ charge-transfer level around $R = 11$ a.u. This avoided crossing appears to be the leading nonadiabatic interaction in the present $C^2+$–HCl collision system. The other avoided crossings, between $1\Sigma^+\{C^+(1s^22s^22p)^2P^o + HCl(2\Pi^o)\}$ and $2\Sigma^+\{C^+(1s^22s^22p)^2P^o + HCl(2\Pi^o)\}$ exit channels, or between the $2\Sigma^+\{C^+(1s^22s^22p)^2P^o + HCl(2\Sigma^+)\}$ and $3\Sigma^+\{C^+(1s^22s^22p)^2D + HCl(2\Pi^o)\}$ levels, are significantly smoother and correspond to large energy gaps. They could certainly not be determinant in the process. Such a strong interaction between the entry channel and one charge-transfer level was not present in the $C^2+$–HF collision system. Effectively, in that case, the $3\Sigma^+\{C^+(1s^22s^22p)^2D + HFC(2\Pi^o)\}$ level was higher in energy than the entry channel with regard to the ionization
FIG. 2. (Color online) : Potential-energy curves for the $^1\Sigma^+$ (full line) and $^1\Pi$ (broken line) states of the C$_2$+–HCl molecular system at equilibrium, $\theta = 0^\circ$. 1: $C^+(1s^22s^22p)^2P^o + \text{HCl}^+(2\Sigma^+)$; 2: $C^+(1s^22s^22p)^2P^o + \text{HCl}^+(2\Sigma^+)$; 3: $C^+(1s^22s^22p)^2D + \text{HCl}^+(2\Pi)$; 4: $C^+(1s^22s^22p)^2S + \text{HCl}(^2\Sigma^+)$ entry channel.

potential of HF and could not be populated directly. So the only exit channel which could be accessible from the entry channel was the $^1\Sigma^+$ states for the C$_2$+–HCl molecular system at equilibrium, $\theta = 0^\circ$. Same labels as in Fig. 2.

FIG. 3. (Color online) Radial coupling matrix elements between $^1\Sigma^+$ states of the C$_2$+–HCl molecular system at equilibrium, $\theta = 0^\circ$. Same labels as in Fig. 2.

trajectories are satisfying [28] and semiclassical approaches may be applied with a good accuracy. The EIKONX code based on an efficient propagation method has been used [29]. In the collision process, electronic transitions are much faster than vibration and rotation motion so that the sudden approximation can be applied. The cross sections, corresponding to purely electronic transitions, are thus determined by solving the impact-parameter equation as in the usual ion-atom approach, considering the internuclear distance of the molecular target fixed in a given geometry. Simple model has been widely used in the field of ion-molecule collisions and leads to reliable results in the keV energy range we are dealing with [30,31].

The coupled equations have been solved taking account of all the molecular states involved in the process by means of the transitions driven by radial and rotational coupling matrix elements. Translation factors may be introduced so that cross

FIG. 4. (Color online) Rotational coupling matrix elements between $^1\Sigma^+$ and $^1\Pi$ states for the C$_2$+–HCl molecular system at equilibrium, $\theta = 0^\circ$. Same labels as in Fig. 2. $\text{rot11} = \langle 1^1\Pi|l|1^1\Sigma^+\rangle$; $\text{rot22} = \langle 2^1\Pi|l|2^1\Sigma^+\rangle$; $\text{rot32} = \langle 3^1\Pi|l|2^1\Sigma^+\rangle$; $\text{rot34} = \langle 3^1\Pi|l|4^1\Sigma^+\rangle$.

III. COLLISION DYNAMICS

The collision treatment has been performed in the keV energy range. For energies higher than 10 eV/amu, straight-line
sections would be independent of the origin of coordinates and to avoid spurious coupling terms at long range. They may be evaluated in the approximation of the common translation factors \[32,33\]. The radial and rotational coupling matrix elements are thus transformed respectively into

\[
\langle \psi_K | \partial/\partial R - (\varepsilon_K - \varepsilon_L) z^2/2R | \psi_L \rangle
\]

and

\[
\langle \psi_K | i L_\gamma + (\varepsilon_K - \varepsilon_L) z \chi | \psi_L \rangle,
\]

Where \(\varepsilon_K\) and \(\varepsilon_L\) are the electronic energies of states \(\ket{\psi_K}\) and \(\ket{\psi_L}\), \(z^2\) and \(z\chi\), the components of the quadrupole moment tensor. Such effect has been tested for the C\(^{2+}\) + HF collision system in the [3–108]-keV laboratory energy range ([0.1–0.6]-a.u. collision velocity range). The corresponding cross sections are presented in Fig. 5. The introduction of translation factors induces a very small variation on the total cross sections, less than 3% at \(E_{\text{lab}} = 100\) keV. The effect decreases at lower collision energies and is completely negligible below about 30 keV. Such effect depends of course of the collision system, but, in a first approach, it can be considered with a good approximation to be weak in the energy range we are dealing with and has not been taken into account in the C\(^{2+}\) + HF collision treatment.

The partial and total cross sections are presented in Fig. 6 and Table I for the linear C-H-Cl geometry \((\theta = 0^\circ)\). The total cross section presents a peak around \(v_{\text{coll}} = 0.1\) a.u. \((E_{\text{lab}} = 3\) keV\) and then decreases at higher collision energy. Such a peak is mainly due to the contribution of the corresponding peak of the partial cross section sec43. As pointed out from the potential-energy curves, the charge-transfer process appears clearly dominated by one nonadiabatic in-

\[
\text{FIG. 6. (Color online) Total and partial charge-transfer cross sections for the C}^{2+} - \text{HCl collision system at equilibrium, } \theta = 0^\circ. \text{ Full line: transition to } 1\Sigma^+ \text{ states; broken line: transition to } 1\Pi \text{ states. sectot: total cross section; sec43: partial cross section on } \{C^{(+1s^22s^22p)^2}P + \text{HCl}(^2\Sigma^+)\}; \text{ sec42: partial cross section on } \{C^{(+1s^22s^22p)^2}P + \text{HCl}(^2\Sigma^+)\}; \text{ sec41: partial cross section on } 1\Sigma^+(C^{(+1s^22s^22p)^2}P + \text{HCl}(^2\Sigma^+)\).
\]

}\]

As already pointed out for C\(^{2+}\) + HF, rotational effects may be observed also for this system as charge-transfer channels may be all correlated to the entry channel by means of rotational coupling. In particular, a hump is shown on the partial cross section on the \(2\Pi\{C^{(+1s^22s^22p)^2}P + \text{HCl}(^2\Sigma^+)\}\) exit channel in connection with the nonadiabatic interaction between the \(2\Pi\{C^{(+1s^22s^22p)^2}P + \text{HCl}(^2\Sigma^+)\}\) and \(3\Pi\{C^{(+1s^22s^22p)^2}D + \text{HCl}(^2\Pi)\}\) charge-transfer levels pointed out in the molecular calculations. However, the
contribution of $^1\Pi$ exit channels decreases at higher collision energies, and is almost of the same order of magnitude as the contribution of corresponding $^1\Sigma^+$ charge-transfer states coupled by radial coupling (Table I). The mechanism of the $^1\Sigma^+$ HCl charge transfer is clearly dominated by the nonadiabatic radial coupling interaction rad34. On the contrary, rotational effects remain significant in $^2\Sigma^+$ at higher collision energies, in particular for the $^2\Pi\{C^+(1s^22s^22p)^2P^0 + HF^+ (2\Sigma^+)\}$ channel, and contribute significantly to the total cross section.

**IV. CONCLUDING REMARKS**

We have presented a theoretical treatment of charge-transfer processes induced by collision of the $^2\Sigma^+$ projectile on hydrogen chloride. This reaction may be compared to the $^1\Sigma^+$ HF collision system involving a quite similar molecular target. Different features may be observed for these collision systems. They do not appear to rely on possible steric effects, but more likely to the electronic interactions in the vicinity of avoided crossings. In $^2\Sigma^+$ + HCl, the charge transfer is driven essentially by the nonadiabatic radial coupling interaction between the entry channel and the highest $3^1\Sigma^+\{C^+(1s^22s^22p)^2D + HF^+ (2\Pi)\}$ charge-transfer channel. The total cross section presents a maximum of $16.6 \times 10^{-16}$ cm$^2$ around $E_{\text{lab}} = 3$ keV and decreases at higher energies. The mechanism is completely different for the $^1\Sigma^+$ + HF collision system. In that case, the $\{C^+(1s^22s^22p)^2D + HF^+ (2\Pi)\}$ charge-transfer channel cannot be accessible directly from the entry channel. A two crossing mechanism is observed, mainly driven by the interaction at shorter range with the $\{C^+(1s^22s^22p)^2P^0 + HF^+ (2\Sigma^+)\}$ exit channel leading to important values of the total cross section at higher collision energy by contribution of radial coupling with a significant rotational effect [9]. The charge transfer is globally more efficient with the HF molecular target with total cross sections from $14.2 \times 10^{-16}$ cm$^2$ at $E_{\text{lab}} = 3$ keV to about $22.0 \times 10^{-16}$ cm$^2$ at $E_{\text{lab}} = 100$ keV. These comparative results show clearly that the charge-transfer mechanism is fundamentally dependent of the specific nonadiabatic interactions involved in each collision system. General conclusions for a series of molecular targets, even of similar electronic configurations, have to be handled with care and the analysis of the behavior of each molecular system is essential.

**ACKNOWLEDGMENTS**

This work was granted access to the HPC resources of [CCRT/CINES/IDRIS under the allocation 2011-1201181655] made by GENCI (Grand Equipement National de Calcul Intensif) as well as from COST actions CM0702 CUSPFEL and MP1002 Nano-IBCT.

### Table I. Charge-transfer cross sections for the C$_2$H + HCl collision system (in 10$^{-16}$ cm$^2$). Comparison with the C$_2$H + HF collision system. Same labels as in Fig. 2.

<table>
<thead>
<tr>
<th>Velocity (a.u.)</th>
<th>$E_{\text{lab}}$ (keV)</th>
<th>sec43 $^4\Sigma^+{3\Sigma^+}$</th>
<th>secpi43 $^4\Sigma^+{3\Pi}$</th>
<th>sec42 $^4\Sigma^+{2\Sigma^+}$</th>
<th>secpi42 $^4\Sigma^+{2\Pi}$</th>
<th>sec41 $^4\Sigma^+{1\Sigma^+}$</th>
<th>secpi41 $^4\Sigma^+{1\Pi}$</th>
<th>Sectot $^2\Sigma^+$+HCl</th>
<th>Sectot $^2\Sigma^+$+HF</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>0.75</td>
<td>6.99</td>
<td>1.78</td>
<td>0.14</td>
<td>0.22</td>
<td>0.17</td>
<td>0.09</td>
<td>9.38</td>
<td>10.54</td>
</tr>
<tr>
<td>0.1</td>
<td>3</td>
<td>12.22</td>
<td>2.80</td>
<td>0.39</td>
<td>0.77</td>
<td>0.14</td>
<td>0.27</td>
<td>14.26</td>
<td>16.10</td>
</tr>
<tr>
<td>0.15</td>
<td>6.75</td>
<td>7.48</td>
<td>4.78</td>
<td>0.36</td>
<td>1.14</td>
<td>0.27</td>
<td>0.23</td>
<td>14.26</td>
<td>16.10</td>
</tr>
<tr>
<td>0.2</td>
<td>12</td>
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<td>4.99</td>
<td>1.80</td>
<td>1.38</td>
<td>0.73</td>
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<tr>
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<td>2.36</td>
<td>2.91</td>
<td>3.91</td>
<td>0.95</td>
<td>0.77</td>
<td>0.81</td>
<td>11.70</td>
<td>19.10</td>
</tr>
<tr>
<td>0.35</td>
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<td>2.14</td>
<td>2.34</td>
<td>3.43</td>
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<td>0.66</td>
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<td>2.08</td>
<td>2.03</td>
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<tr>
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<td>0.67</td>
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<td>7.37</td>
<td>22.23</td>
</tr>
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</table>

[17] MOLPRO (version 2009.1) is a package of ab initio programs written by H.-J. Werner et al.