Reactions of Oxygen Atoms with Van der Waals Complexes: The Effect of Complex Formation on the Internal Energy Distribution in the Products

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Reactions of oxygen atoms with van der Waals complexes: The effect of complex formation on the internal energy distribution in the products

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Reactions of atomic oxygen with complexes containing HCl are investigated and the OH product state distributions are compared to those observed for the corresponding reactions of HCl monomers. In previous studies of reactions of O(3P) with HCl and hydrocarbon complexes, rotationally colder OH product state distributions were observed, when compared to the corresponding reactions of monomers. In contrast, we find that reactions of O(1D) with HCl clusters yield OH rotational distributions that are unaffected by the incorporation of HCl into a van der Waals complex. Quasiclassical trajectories are run on collisions of oxygen with HCl and Ar·HCl at 1 eV collision energies to investigate the differences in the dynamics of the O(1D) and O(3P) reactions. It is found that when the van der Waals complex is longer lived than the collision complex, rotational and vibrational cooling are observed. In contrast, when the dissociation of the van der Waals complex is prompt, compared to the collision complex lifetime, the effects of complex formation on the internal energy of the OH product become negligible. © 1998 American Institute of Physics. [S0021-9606(98)01623-7]

I. INTRODUCTION

Over the past 25 years, experimental and theoretical approaches have been developed to the point that a variety of atom plus diatom (A + BC) reactions have been mapped out very accurately.1-3 While much can be learned from these studies, most chemistry occurs in solvated environments. Therefore, understanding how changes to the environment affect reaction dynamics and product state distributions has become an important question in chemical physics. One way to begin to understand these effects is through studies of reactions in which one or both of the reacting species are incorporated into a weakly bound cluster. A variety of reactions have been investigated in this manner, either by photo-initiating a reaction inside a cluster,4-11 or by studying the bimolecular reaction of a weakly bound complex with an atom or molecule directly.9,12-19

In the present study, we focus on reactions of atomic oxygen in its ground (3P) or first excited (1D) electronic state with complexes containing HCl. Because of their importance in fundamental reaction dynamics20,21 and the significant role reactions of O(1D) play in atmospheric chemistry,22 the reaction of atomic oxygen with HCl has been studied extensively both theoretically23,24 and experimentally.25,26 However, much less work has probed reactions of atomic oxygen with complexes of HCl.17

The specific question that we will address is how the product state distributions resulting from reactions in clusters differ from those obtained from the corresponding reactions of isolated molecules. In many cases, it has been found that incorporating one or both of the reactants into a van der Waals complex leads to products that contain less internal energy compared to the products of the corresponding reaction of isolated atoms and molecules. This effect is particularly profound for the angular momentum distributions. The simple-minded explanation for the observed rotational cooling assumes that the "third body," namely the spectator to the reaction, shares angular momentum with the reacting species. As a result, less angular momentum is expressed in the reaction products. When the issue was studied in detail,27,28 it was found that in some cases the rotational cooling exceeds that predicted by a simple statistical model. This can be explained by assuming that during the very slow dissociation of the collision complex, there is efficient transfer of rotational energy to translational energy.29

The reaction of atomic oxygen with HCl provides a good laboratory in which we can investigate the mechanisms for the above processes. From our previous experimental studies of the reactions of O(3P) with HCl, (HCl)2, and Ar·HCl, we know that the addition of the complexing atom or molecule leads to longer lived reaction complexes than is observed for the reaction of isolated HCl molecules.17 This observation is consistent with observed colder angular momentum distributions for the OH products. By contrast, in the present work, we find that in the O(1D) + HCl reaction rotational cooling of the OH products cannot be observed experimentally when the HCl is complexed with Ar or other HCl molecules.

In order to understand the experimental findings described above, we use classical mechanics30,31 to simulate reactions of O(1D) and O(3P) with HCl and Ar·HCl. We obtain identical rotational distributions for the OH products from the reactions of O(1D) with HCl and Ar·HCl, whereas there is measurable cooling in the OH products of the O(3P) reaction. We explain the observed differences in
the ability of the complexing atom or molecule to act as a "cooling element" for the OH products in terms of the lifet ime of the van der Waals complex compared to that of the collision complex. In the case of the \( O(^{1}D) + HCl \) reaction, the HOCI reaction complex is strongly bound and can exist for several picoseconds before dissociating into products whereas the dissociation of the Ar·OHCl complex is prompt. In contrast, for the \( O(3P) + HCl \) reaction, the OHCl transition state complex lives for no more than one or two vibrational periods of the HCl stretch whereas the Ar·OH complex survives for several hundred femtoseconds.

The remainder of the paper is organized as follows. In Sec. II the experimental methods and theoretical approaches are discussed. Results are presented and discussed in Sec. III and the results are summarized in Sec. IV.

II. METHODS

A. Experiment

The experimental setup for the investigation of the reactions of \( O(1D) \) with HCl clusters has been described previously. Therefore, the details will not be repeated here. For the present study, a crossed molecular beam setup is used in which a beam of HCl collides with fast \( O(1D) \) atoms, produced by photodissociation of \( N_2O \) at 193 nm in the region where the two beams intersect. The photodissociation laser is a ArF excimer laser (Lambda Physik, Compex 102). A frequency doubled Nd:Yag pumped dye laser (Spectra-Physics) is used to create the tunable UV light for the laser-induced fluorescence (LIF) measurements of the OH product.

B. Theory

In order to understand the experimental OH product state distributions, we have run several ensembles of classical trajectories. We focus on four separate systems that model reactions of ground state atomic oxygen \( [O(3P)] \) and oxygen in its first excited electronic state \( [O(1D)] \) with HCl and Ar·HCl complexes. For this comparison, all four sets of trajectories are run at 1 eV collision energies with zero-point energy in all degrees of freedom.

1. Initial conditions and propagations

The initial conditions of the trajectories are sampled from a long time propagation of intermolecular dynamics of Ar·HCl with zero-point energy in both of the intermolecular modes, the HCl in its minimum energy configuration and total angular momentum \( J = 0 \). The dynamics is initiated with the oxygen atom at least 10.6 Å from the center of mass of HCl or Ar·HCl. The relative kinetic energy of the collision partners is 1 eV. Impact parameters and the orientation of reactants are chosen randomly using a Monte Carlo sampling procedure. For \( O(1D) + HCl \) collisions, \( b_{max} = 7.94 \) Å, while \( b_{max} = 3.97 \) Å for the \( O(3P) + HCl \) collisions. Finally, to facilitate comparison of the reactivity of HCl and Ar·HCl, the same initial conditions are used for both sets of propagations.

The trajectories are propagated using a Gear algorithm and all trajectories are run until the distances between the product species are larger than 8 Å or the trajectory has been run for longer than 1.2 ps for collisions with HCl reactions and 2.4 ps for collisions with Ar·HCl. For all reactions except the \( O(3P) + Ar + HCl \) reaction, 10 000 trajectories are run. In the case of the \( O(3P) + Ar + HCl \) reaction 20 000 trajectories are required to obtain meaningful statistics.

2. Potential energy surfaces

For the present study, we approximate the intermolecular O+Ar·HCl potential as a sum of pairwise interactions. Many of the contributing interactions have been characterized by fitting a combination of scattering data, the observed spectra, and \textit{ab initio} points to simple functional forms. The most important ingredient in the global surface is the underlying O+HCl potential. In the present study, we use the \( O(1D) + HCl \) potential of Schinke and the \( O(3P) + HCl \) potential of Koizumi et al. Both of these potentials were fit to \textit{ab initio} as well as scattering data and have been used in a variety of classical and quantum mechanical studies of the O+HCl reaction.

The interactions with the argon atom are more difficult to construct. In the reactant channel, the argon atom sees an oxygen atom, either \( O(3P) \) or \( O(1D) \), and an HCl molecule. In the product channels, the argon atom is in the presence of either Cl+OH or H+OCl. While potential surfaces that describe most of the necessary intermolecular potentials have been characterized, the interaction potentials between an argon atom and the reaction intermediates have not. In the absence of this information, we model the intermolecular potential as the sum of atom–atom interactions. The advantage of this approach, which has been previously employed in studies of reactions of van der Waals dimers, is that the resulting potentials are necessarily smooth everywhere.

In the present study, the Ar·Cl and Ar·O(1D) potentials are modeled by a Lennard-Jones \((12,6)\) function,

\[
V(r) = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} \right],
\]

with \( \epsilon = 0.0144 \) eV and \( \sigma = 3.47 \) Å. The Ar·H interaction is given by the Slater–Buckingham \( \exp(a,6) \) function,

\[
V(r) = \frac{\epsilon}{1 - 6/\alpha} \left[ \frac{6}{\alpha} \exp(1-r_m/r) - \left( \frac{r_m}{r} \right)^6 \right],
\]

where \( \alpha = 14.23 \), \( \epsilon = 38.717 \) cm\(^{-1}\), and \( r_m = 2.85 \) Å. Finally, the Ar·O(3P) is given by a Morse function,

\[
V(r) = D \left[ 1 - e^{-\beta(r-r_c)} \right]^2,
\]

with \( D = 0.010401 \) 5 eV, \( r_c = 3.45 \) Å, and \( \beta = 1.748 \) Å\(^{-1}\).

While the above pairwise additive potentials will not describe the Ar·HCl or the Ar·OH intermolecular potentials with spectroscopic accuracy, they provide good descriptions of these complexes. When HCl is in its equilibrium geometry, the dissociation energy of Ar·HCl is 150 cm\(^{-1}\), which is in good agreement with the 175 cm\(^{-1}\) value computed from Hutson’s H6(3) potential. A second minimum on the Ar·HCl surface is found for the Ar·Cl–H geometry. In this geometry, the system is predicted to be bound by 140 cm\(^{-1}\).
FIG. 1. The laser-induced fluorescence spectrum of the nascent OH products formed from the reaction of O(1D) with a beam of HCl. The Q1 branch of the A-X transition is shown. In the upper panel the HCl pressure behind the nozzle was 1000 mbar and the lower panel was obtained at a pressure of 300 mbar.

according to the spectroscopic potential, while the pairwise additive potential yields a value of 120 cm\(^{-1}\). The variation in the minimum Ar−Cl distance as a function of the Ar··HCl angle is larger in our pairwise approximation to the intermolecular potential than in the spectroscopic surface, but in both cases, the average value is approximately 4 Å. The Ar··OH surface is also well represented by the pairwise additive approximation. For example, the sum of the Ar··O(3P) and Ar··H surfaces has a minimum at the Ar−H−O geometry with \(D_e = 108\) cm\(^{-1}\), and \(R_e = 3.6\) Å. These values are in good agreement with the reported \(ab\) \initio \Ar−\OH(X \(2\Sigma^+\)) surface: 100 cm\(^{-1}\) and 3.8 Å. \(^{39}\)

III. RESULTS

A. Experiment

The OH rotational distributions, resulting from the reaction of O(1D) with HCl, have been investigated as a function of the HCl pressure behind the nozzle for \(v_{\text{OH}} = 0,1\). Part of the resulting LIF spectrum, taken at two different HCl pressures, is shown in Fig. 1. The spectrum was obtained by monitoring the A−X \(Q_1\) transition of OH(\(v = 0\)), generated by the reaction of O(1D) with a molecular beam of HCl. The spectrum in the lower panel of Fig. 1 was obtained when the pressure behind the nozzle was 300 mbar of neat HCl, while the spectrum in the upper panel was obtained at 1000 mbar. The rotational population obtained for the two HCl pressures show no noticeable differences. Further, the distributions remain basically identical when the pressure is varied from approximately 100 to 2000 mbar. Under the same conditions dramatic changes in the OH rotational distributions have been reported for reactions of O(3P) and HCl \(^{17}\) and O(1D) and CH\(_4\). \(^{40}\)

We conclude that for the reaction of O(1D) with HCl complexes, the rotational energy distribution is not sensitive to cluster formation. To understand the origin of this apparent insensitivity, we look at the results of the classical trajectory studies of the O(3P)+HCl and O(1D)+HCl reactions.

B. Reactions of O(3P) and HCl and Ar·HCl

There is one energetically accessible product channel for an O(3P)+HCl collision at 1 eV: formation of OH+Cl and dissociation to the starting materials. As shown in Table I, given our choice of initial conditions only 0.6% of the trajectories form OH.

When an argon atom is introduced, additional product channels become accessible. The observed outcomes are listed in Table I. Although the argon atom could form a complex with the OH products, the Ar··OH interaction potential is sufficiently weak, \(D_e \approx 2.5\) of the available energy, such that these complexes are extremely rare. The Ar··HCl binding energy is also two orders of magnitude smaller than the collision energy, explaining why the Ar··HCl complex dissociates in approximately 40% of the trajectories.

As the results in Table I show, the cross sections, \(\sigma\), decrease by 30% when the argon atom is introduced while \(b_{\text{max}}\) for the reactive process increases by 1.6 Å. The primary source of the increase in \(b_{\text{max}}\) for the collision of O(3P) and Ar··HCl comes from the definition of this quantity. We have defined \(b\) to be the impact parameter measured between the centers of mass of the colliding species, either HCl or Ar··HCl. If instead we compare \(b_{\text{max}}\), measured from the center of mass of HCl we obtain \(b_{\text{max}}^{\text{O+HCl}} = 1.5\) Å and \(b_{\text{max}}^{\text{O+Ar+HCl}} = 1.77\) Å.

The observed decrease in reaction cross section, \(\sigma\), for the

<table>
<thead>
<tr>
<th>Products</th>
<th>(N_i)</th>
<th>(\sigma/\text{Å}^2)</th>
<th>(b_{\text{max}}/\text{Å})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl+OH(^b)</td>
<td>62</td>
<td>0.14(0.02)</td>
<td>1.50</td>
</tr>
<tr>
<td>O+Ar+HCl(^c)</td>
<td>7969</td>
<td>15.8(0.2)</td>
<td>3.93</td>
</tr>
<tr>
<td>Cl+Ar+OH</td>
<td>68</td>
<td>0.11(0.01)</td>
<td>3.13</td>
</tr>
</tbody>
</table>

\(^a\)The reported uncertainties, in Parentheses, represent one standard deviation from the reported cross sections.

\(^b\)Based on 10 000 trajectories run at 1.0 eV collision energies with \(b_{\text{max}} = 3.97\) Å.

\(^c\)Based on 20 000 trajectories run at 1.0 eV collision energies with \(b_{\text{max}} = 3.97\) Å.
reaction also can be easily understood. It reflects the fact that even when we average over the zero-point vibrations of the Ar·HCl complex, our initial configurations are dominated by near collinear Cl—H—Ar geometries, as this represents the minimum energy geometry of Ar·HCl. Because the O+HCl reactive collisions occur with a near linear O—H—Cl transition state geometry, the argon atom in Ar·HCl effectively blocks the formation of the transition state complexes.

To clarify this point, a pair of representative trajectories are plotted in Fig. 2. Here, the panels on the left-hand side show a trajectory that represents an O+HCl collision while the ones on the right give an O+Ar·HCl trajectory. In the top panels, the distances between the hydrogen, oxygen, and chlorine atoms are plotted with thick lines. In both cases the oxygen and chlorine atoms undergo an impulsive collision, forming an O—H—Cl transition state complex that lives for approximately two vibrational periods of the HCl stretch before dissociating into the Cl+OH products. In the lower panels, the thick line provides cos θ_{HCl}. We find that this value is approximately −1 when the complex is formed, indicating a near linear O—H—Cl transition state geometry. The thin lines in Figs. 2(c) and 2(d) provide information about the location of the argon atom during this O+Ar·HCl collision. Focusing on Fig. 2(d), in which the thin line gives cos θ_{ArHCl}, we find that initially the complex is in a near linear Ar—H—Cl geometry. As the oxygen atom approaches the Ar·HCl complex, θ_{ArHCl} approaches 90°, enabling the formation of the OHC1 complex. Finally, the thin line in Fig. 2(b) shows that as the complex dissociates, the Cl—O distance increases faster than the Ar—O distance.

The above differences between the lifetimes of the OHC1 collision complex and the Ar·OH van der Waals complex are found to represent a general trend for the reactive collisions. This can be seen by comparing the lifetime distributions of the OHC1 collision complex formed in O+HCl and O+Ar·HCl collisions to the distribution of Ar·OH lifetimes, plotted in Fig. 3. The shift in the distribution in Fig. 3(b) relative to that in Fig. 3(a) is a result of the difference between the relative velocity of O(^3 P)+HCl, compared to that for O(^3 P)+Ar·HCl at the same collision energy.

Based on the masses of the two systems, the difference in the time of the collision, resulting from the change in the relative velocities, is expected to be approximately 0.15 ps, and this is what is seen. In addition, in Fig. 3(b), some of the collision energy is converted into kinetic energy of the departing argon atom. This second effect explains the broadening of the OCI lifetime distribution in Fig. 3(b) compared to that in Fig. 3(a). Finally, comparison of Figs. 3(b) and 3(c) shows that the argon atom remains in a complex with the OH product for as much as 0.5 ps after the OHC1 complex has dissociated.

One consequence of the existence of short-lived Ar·OH van der Waals complexes comes in lowering the internal energy of the OH product, plotted in Fig. 4. Here the distributions for the OH formed from O+HCl collisions are plotted with thin lines and those distributions obtained from O+Ar·HCl collisions are plotted with broad lines. Comparing the distributions, we find that the rotational and vibrational distributions are shifted to lower energies. These trends are consistent with the short-lived Ar·OH intermediates.

![FIG. 2](image2.png)

**FIG. 2.** Representative trajectory for O(^3 P)+HCl [(a) and (b)] and Ar·HCl [(c) and (d)] collisions that results in the formation of OH+Cl. In (a) and (c) the OH (dashed line), OCI (solid line), and HCl (dotted line) distances are plotted as a function of time. In (b) and (d) cos(θ_{HCl}) is plotted. The thin lines in (c) and (d) provide the ArO distance and cos θ_{ArHCl}, respectively. Finally, the vertical dotted line is provided to show when the collision complex is initially formed.

![FIG. 3](image3.png)

**FIG. 3.** Lifetime distributions for (a) the time when r_{OCI} exceeds 2.7 Å for O+HCl reactive collisions, (b) the time when r_{OCI} exceeds 2.7 Å for O+Ar·HCl reactive collisions, and (c) the time when r_{ArH} exceeds 5.3 Å for O+Ar·H Cl reactive collisions.

![FIG. 4](image4.png)

**FIG. 4.** Internal energy distributions for OH formed from reactions of O(^3 P) with HCl (thin line) and Ar·HCl (broad line). In (a), the vibrational energy distributions of the OH products are plotted. Each point represents results that have vibrational energies between E_{vib}−ω_{OH}2 cm⁻¹ and E_{vib}+ω_{OH}2 cm⁻¹. The uncertainties represent one standard deviation. The angular momentum distributions are plotted in (b). Here each point represents all trajectories that have OH angular momentum between J_{OH}−2 and J_{OH}+2. All plots are normalized so that the sum of the relative cross sections is unity.
TABLE II. Outcomes of the O+(1D)+HCl collisions. Results are based on 10,000 trajectories run at 1.0 eV collision energies with b_{max}=7.94 Å.

<table>
<thead>
<tr>
<th>Products</th>
<th>N_i</th>
<th>σ/Å²</th>
<th>b_{max}/Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl+OH</td>
<td>914</td>
<td>9.5(0.3)</td>
<td>2.95</td>
</tr>
<tr>
<td>H+OCl</td>
<td>31</td>
<td>0.32(0.06)</td>
<td>1.65</td>
</tr>
<tr>
<td>HOCb</td>
<td>29</td>
<td>0.34(0.06)</td>
<td>2.54</td>
</tr>
</tbody>
</table>

aThe reported uncertainties, in parentheses, represent one standard deviation from the reported cross sections.

bThe atoms remain in an HOCI complex when the trajectory is terminated after 1.2 ps.

create where the argon atom acts as an effective cooling element. Similar results have been reported for classical studies of O(3P)+hydrocarbon reactions.19

C. Reactions of O(1D) and HCl and Ar·HCl

The above results should be contrasted with those for the O(1D)+HCl reaction. Again the reaction is studied for collision energies of 1 eV. At this total energy, there are four possible outcomes of an O(1D)+HCl inelastic collision. As before, OH+Cl can be formed or the oxygen atom can give up energy to the HCl molecule. Because there is an additional 2.28 eV of energy in this system compared to the O(3P)+HCl reaction, the OCl+H product channel is opened. This channel is approximately 2.3 eV higher in energy than the O(3P)+HCl and OH+Cl channels. Finally, a long-lived, metastable HOCI molecule may be formed. When an argon atom is introduced, all of the above channels remain energetically accessible. In addition, the Ar atom may form a complex with either one of the product species. The cross sections for all observed processes are reported in Tables II and III.

In contrast to the results for the O(3P) system, here the introduction of the argon atom increases the cross sections. Further, in contrast to the O(3P) reaction which proceeds through a linear O–H–Cl transition state geometry, in the O(1D)+HCl reaction, the oxygen atom inserts into the HCl bond, forming a strongly bound HOCI molecular intermediate. Therefore, the introduction of an Ar atom in a near linear Ar–H–Cl complex geometry will have little effect on the dynamics of this process. This can be seen in the representative O(1D)+HCl trajectories plotted in Fig. 5. In Figs. 5(a) and 5(c), we find that when the collision complex is formed, the OHCl angle flips from 180° to 0° indicating that the oxygen atom is now between the hydrogen and chlorine atoms. Further examination of the argon atom dynamics in Figs. 5(c) and 5(d) shows that it begins to separate from the HOCI complex approximately 50 fs after this complex is formed and long before the complex falls apart. Not only does the argon atom leave the collision complex shortly after formation, but, comparing Figs. 2(c) and 5(c), we find that the argon atom leaves the reaction complex formed from the O(1D)+HCl collision with more kinetic energy.

While the above observations are based on individual trajectories, they translate into several general results, as can be seen in the average lifetime distributions, plotted in Fig. 6. Comparing Figs. 6(a) and 6(b), we find that the trajectories for O(1D)+Ar·HCl lead to longer-lived complexes than when the Ar atom is not present. Further, by comparing Figs. 6(b) and 6(c), we find that for many of the trajectories the Ar·HOCI complex dissociates before the HOCI complex falls apart.

An important consequence of these relative lifetimes comes in the resulting internal energy distributions of the OH products, plotted in Fig. 7. In this case, the angular momentum distribution is unaffected by the presence of the argon atom. The vibrational energy distribution is shifted to lower energies. This trend can be rationalized by the fact that the argon atom removes internal energy from the HOCI complex, leading to the observed shift in the vibrational energy distributions. The prompt dissociation of the van der Waals complex results in the observation that there is no difference

TABLE III. Outcomes of the O(1D)+Ar·HCl collisions. Results are based on 10,000 trajectories run at 1.0 eV collision energies with b_{max}=7.94 Å.

<table>
<thead>
<tr>
<th>Products</th>
<th>N_i</th>
<th>σ/Å²</th>
<th>b_{max}/Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>O+Ar+HCl</td>
<td>3020</td>
<td>37.3(0.7)</td>
<td>7.76</td>
</tr>
<tr>
<td>Cl+Ar·OH</td>
<td>2</td>
<td>0.02(0.01)</td>
<td>2.02</td>
</tr>
<tr>
<td>Cl+Ar+OH</td>
<td>890</td>
<td>9.8(0.3)</td>
<td>4.90</td>
</tr>
<tr>
<td>Ar·Cl+OH</td>
<td>11</td>
<td>0.12(0.04)</td>
<td>3.67</td>
</tr>
<tr>
<td>H+Ar+OCl</td>
<td>59</td>
<td>0.60(0.08)</td>
<td>3.79</td>
</tr>
<tr>
<td>Ar+HOCb</td>
<td>36</td>
<td>0.38(0.06)</td>
<td>4.03</td>
</tr>
</tbody>
</table>

aThe reported uncertainties, in parentheses, represent one standard deviation from the reported cross sections.

bThe atoms remain in an HOCI complex when the trajectory is terminated after 2.4 ps.

FIG. 5. Same as Fig. 2 for a typical O(1D)+HCl and Ar·HCl collisions.

FIG. 6. Internal energy distributions for OH formed from reactions of O(1D) with HCl (thin line) and Ar–HCl (broad line). In (a), the vibrational energy distributions of the OH products are plotted. Each point represents results that have vibrational energies between E_{vb}–q_{vib}2 cm⁻¹ and E_{vb}+q_{vib}2 cm⁻¹. The uncertainties represent one standard deviation. The angular momentum distributions are plotted in (b). Here each point represents all trajectories that have OH angular momentum between J_{OH}–1 and J_{OH}+1. All plots are normalized so that the sum of the relative cross sections is unity.
between the angular momentum distributions of the OH formed from O+Ar-HCl and O+HCl collisions.

D. The Ar atom as a “cooling element”

Comparing the dynamics of O(3P) and O(1D) collisions with HCl and Ar-OH there are several significant differences. First, for the O(3P) + Ar-OH collisions, the lifetime of the collision complex is short compared to the lifetime of the Ar-OH complex. This difference is manifested in the observation that the rotational and vibrational energies of the OH products are lower for reactions of Ar-OH than they are for the HCl reactions. In contrast, in O(1D)+Ar-OH collisions, the HOCl collision complex survives for as long as several picoseconds. Due to the large energy difference between the reactants and the products, the argon atom leaves promptly thereby having a minor effect on the energy distributions of the OH products. In this case the OH product state distribution will be determined by the lifetime of the HOCl complex.

These results point to the conditions under which the complexing atom or molecule can act as an effective “cooling element.” If the collision complex lifetime is short relative the time required for the van der Waals complex to fall apart, then rotational cooling takes place. However, if the collision complex is long lived relative to the lifetime of the van der Waals complex, then the product state energy distribution will be similar to that observed for the corresponding reaction of uncomplexed species.

To test the above hypothesis we look at the second product channel that is populated by 1 eV collisions of O(1D) with HCl: the H+OCl products. To obtain improved statistics for this comparison, we ran these trajectories with a maximum impact parameter of 3.97 Å. This yields 139 and 171 reactive trajectories for the collisions of O(1D) with HCl and Ar-OH, respectively.

In Fig. 8 we compare the lifetimes of the collision and van der Waals complexes, as we did for the OH channel in Fig. 6. Here we define the HOCl complex to dissociate when the OH and HCl distances both exceed 2.17 Å, and the van der Waals complex is dissociated when $r_{Ar-OCl}$ becomes larger than 5.29 Å. In this case, the HOCl lifetimes, plotted in Figs. 8(a) and 8(b), are similar, and the Ar-OCl complex has a longer lifetime, on average. In Fig. 9 we plot the OCl rotational and vibrational distributions and, as expected by our model, the distributions are colder for the O(1D)+Ar-OH reactions than for the O(1D)+HCl.

IV. SUMMARY AND CONCLUSIONS

To summarize, we have found that for the reaction of O(1D) with HCl the rotational distribution of the OH products is insensitive to complex formation. This result is in stark contrast to previous work on O(3P) collisions with hydrocarbon molecules complexed with rare gas atoms, where the product state distribution of the OH was significantly cooler than in the reactions with HCl monomer.40 We have rationalized the above findings by running classical simulations of the atomic oxygen + HCl reactions for O(1D) and O(3P) and compare the OH rotational distributions to those obtained when HCl is in a van der Waals complex. Our results are consistent with the experimental findings. By studying the trajectories, we find that the effectiveness of argon as a “cooling element” is closely related to the life-
times of the collision and van der Waals complexes. This is further verified by comparing the OCI product state distributions for O(1D) collisions with HCl and Ar···HCl. Here the lifetime of the complexes of Ar with OCI are longer lived than the OHCl complex and the corresponding OCI angular momentum and vibrational energy distributions are colder.

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