Transition State Spectroscopy by Photodetachment of Vibrationally Excited Atoms

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Abstract:

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Technical Summary:

(A) Spectroscopy of the Transition State and of Reactive Intermediates

In these experiments, negative ion photoelectron spectroscopy is used to study the short-lived transition states characteristic of direct reactions or the longer lived reactive
intermediates that occur when the reaction coordinate involves one or more local minima. The idea is to access the neutral transition state or reactive intermediate by photodetachment of a suitable negative ion precursor, and the resulting photoelectron spectrum yields information on the vibrational structure and dissociation dynamics of the neutral complex.

The transition state regions of the OH + OH → O (\(^3P\)) + H\(_2\)O and the OH + H\(_2\)O → H\(_2\)O + OH reactions were studied by photoelectron spectroscopy of the O\(^-\) (H\(_2\)O) and H\(_3\)O\(_2\)\(^-\) anions and their deuterated analogs. The spectra show resolved vibrational progressions attributed to H-atom vibrational motion in the unstable neutral complexes formed by photodetachment. The positions and intensities of the peaks change markedly upon isotopic substitution. One dimensional Franck-Condon calculations using \textit{ab initio} potentials for the anion and neutral are used to interpret the peak spacings and intensities, as well as the strong isotopic effects. The results can be understood in the context of previously obtained transition state spectra for heavy + light-heavy reactions.

The transition state region of the reaction OH + H\(_2\) → H\(_2\)O + H was investigated by photoelectron spectroscopy of the H\(_3\)O\(^-\) and D\(_3\)O\(^-\) anions. The peaks observed in the spectra are from a combination of vibrational progressions and overlapping anion → neutral electronic transitions. The photoelectron angular distributions indicate that two processes contribute the spectra; these are assigned to photodetachment from the H\(^-'\)(H\(_2\)O) and OH\(^-'\)(H\(_2\)) forms of the anion. A comparison of experiments performed in two different laboratories shows that the two forms of the ion readily interconvert, and that the relative populations are determined solely by the temperature of the ions. In order to interpret the spectra, a two-dimensional \textit{ab initio} potential energy surface for the anion was constructed, wavefunctions for the first few vibrational levels were determined, and the photoelectron spectra were simulated using the Walch-Dunning-Schatz-Elgersma surface for the OH + H\(_2\) reaction. A comparison of the experimental and simulated spectra showed that photodetachment from the \(v=0\) level of the anion, which is localized in the H\(^-'\)(H\(_2\)O) well, primarily probes the H + H\(_2\)O exit valley of the neutral surface. The \(v=2\)
level of the anion is the first with significant amplitude in the OH(H₂) well, and photodetachment from this level probes the OH + H₂ transition state region. The simulated spectra are in qualitative agreement with the experimental spectra, but do indicate that the neutral reactive surface needs to be modified.

In order to characterize the N₂O₂ reactive intermediate, photoelectron spectra of the N₂O₂⁻ anion were obtained. The spectra originate from the C₂v isomer of the anion. Vibrationally resolved progressions corresponding to transitions to several electronic states of the previously unobserved N₂O₂ molecule are observed. All of the observed transitions lie above the dissociation asymptotes for N₂ + O₂, NO + NO, and O + N₂O, and several lie above the N + NO₂ and N₂ + O + O asymptotes. \textit{Ab initio} calculations have been carried out for the anion ground state and several singlet and triplet states of neutral N₂O₂. By comparing the observed spectra with Franck-Condon simulations based on these calculations, the lowest bands observed in our spectra were assigned to transitions to the 3A₂ and 3A₁ states (C₂v symmetry) of N₂O₂⁻. These spectra thus represent the first experimental characterization of metastable, high energy forms of N₂O₂. Both the N₂O₂⁻ and the N₂O₂ species are of interest due to their roles as reactive intermediates in the O⁻ + N₂O and N + NO₂ chemical reactions.

Finally, the formyloxyl radical, HCO₂⁺, was experimentally identified for the first time by photoelectron spectroscopy of HCO₂⁻ and DCO₂⁻. Photodetachment accesses the 2A₁, 2B₂, and 2A₂ states of the formyloxyl radical, HCO₂. The 2A₁ state is assigned as the HCO₂ ground state, although it is nearly degenerate with the 2B₂ state (T₀=0.027 eV), and the 2A₂ state lies at T₀=0.536 eV. The electron affinity of HCO₂ is 3.498 ± 0.015 eV. The spectra show partially resolved vibrational features, primarily involving progressions in the CO₂ bending mode. The irregular appearance of the spectra in some regions suggests vibronic coupling between the 2A₁ and 2B₂ states. The possible role of the HCO₂ radical as an intermediate in the OH + CO \rightarrow H + CO₂ reaction and in H + CO₂ inelastic scattering is discussed.
B) Spectroscopy and Electron Detachment Dynamics of Carbon Cluster Anions.

We demonstrated for the first time that stimulated Raman pumping (SRP) can be used to vibrationally excite gas-phase negative ions. In SRP, a molecule or ion is excited by its interaction with two laser pulses whose frequencies differ by a vibrational quantum. This was achieved for \( C_2^- \), in which stimulated Raman pumping excites a rotationally-resolved transition between the \( v=0 \) and \( v=1 \) levels of the ground \( ^2\Sigma_g^- \) state. The resulting vibrational excitation is monitored by resonant two-photon detachment through the excited \( B \ ^2\Sigma_u^+ \) state. At least 35% conversion to the \( v=1 \) level was achieved in this study. It is expected that this method is quite generally applicable and will provide an excellent tool for the study of the vibrational spectroscopy of negative ions.

We have also studied the electronic spectroscopy of mass-selected carbon cluster anions using resonant multiphoton detachment spectroscopy. The \( C \ ^2\Pi_u \leftarrow X \ ^2\Pi_g \) electronic transition of \( C_4^- \) was studied by both one-color and two-color resonant two-photon detachment (R2PD) spectroscopy. The one-color spectrum reveals vibrational structure in the excited anion state. Transitions due to excitations in one of the symmetric stretching modes as well as the bending modes are observed. Spectral resolution in the one-color experiment is limited by power broadening; using two-color R2PD, rotationally resolved spectra of the origin and \( 2_0^1 \) bands of the \( C \ ^2\Pi_u \leftarrow X \ ^2\Pi_g \) transition are obtained. Molecular constants determined by fitting the rotationally-resolved spectra are generally in good agreement with a recent ab initio calculation by Schmatz and Botschwina. Perturbations in the \( 2_0^1 \) band are attributed to Fermi resonance interactions in the \( C \ ^2\Pi_u \) state.

Resonant multiphoton detachment spectroscopy was also used to obtain vibrationally resolved spectra of the \( C \ ^2\Pi_u \leftarrow X \ ^2\Pi_g \) electronic transitions in \( C_6^- \) and \( C_8^- \). Transitions due to vibrational excitations in the totally symmetric stretching modes as well as the bending modes are observed. The electron detachment dynamics subsequent to multiphoton absorption were studied by measuring the electron emission time profiles and electron kinetic energy distributions. The observation of delayed electron emission combined with the form of the electron kinetic energy
distributions indicates that these species undergo the cluster equivalent of thermionic emission. This interpretation is supported by comparing the experimental results to a microcanonical model for cluster thermionic emission.

C) Studies of Size-selected Weakly-Bound Anion and Neutral Clusters.

These experiments are aimed at understanding how the properties of anions and neutrals are affected by solvation with relatively weakly interacting atoms and molecules. By starting with negative ions, size-selection of the species of interest is straightforward, and photodetachment spectroscopy reveals how binding energies, geometries, and electronic state splittings vary with the number and composition of solvating species.

Photoelectron spectra were measured for the anions $X^-(CO_2)_n$, with $X = I, Br, Cl$ and $F$. The vibrationally resolved spectra show that $I^- (CO_2)_n, Br^- (CO_2)_n$, and $Cl^- (CO_2)_n$ are primarily electrostatically bound clusters, although the charge-quadrupole interaction is strong enough to distort the $CO_2$ molecule by as much as $10^\circ$ (in $Cl^- (CO_2)_n$). *Ab initio* calculations and electrostatic models are used to describe the geometry and bonding of these clusters. The photoelectron spectrum of is qualitatively different and shows transitions to both the $\tilde{X}^2B_2$ ground and the $\tilde{A}^2A_2$ first excited electronic states of the covalently bound $FCO_2$ radical. The previously unobserved $\tilde{A}^2A_2$ state is measured to lie 0.579 eV above the ground state. Vibrational frequencies are assigned with the assistance of *ab initio* calculations. The $FCO_2$ heat of formation is determined to be $\Delta_f H^o_{298} (FCO_2) = -85.2 \pm 2.8$ kcal/mole. While both $FCO_2^-$ and $FCO_2$ are more strongly bound than the other halide-$CO_2$ clusters, the C-F bonds are very weak relative to C-F bonds found in other halocarbon compounds.

These studies of binary clusters were followed by investigations of clusters in which a halide ion is complexed to multiple solvent molecules. Specifically, photoelectron spectra of the $I^- (CO_2)_n, I^-(N_2O)_n$, and $Br^- (CO_2)_n$ clusters were obtained. The spectra provide information about the stepwise solvation of the bromide and iodide anions and about the size of the first solvation shells in these clusters. The data suggest that significantly different solute-solvent
interactions exist in the three sets of clusters studied here. The $X(CO_2)_n$ spectra exhibit resolved progressions which are assigned to in-phase $CO_2$ solvent bending vibrations in the neutral clusters. These vibrations are excited by photodetachment of anion clusters in which the $CO_2$ molecules are distorted from linearity by a charge-quadrupole interaction. The $\Gamma(N_2O)_n$ spectra do not show any vibrational structure, presumable because the weaker ion-solvent interactions are insufficient to distort the $N_2O$ molecules.

Finally, the anion zero electron kinetic energy (ZEKE) spectra of the van der Waals clusters $Ar_{2.3}Br^-$ and $Ar_{2.7}I^-$ were measured, and partially discriminated threshold photodetachment (PDTP) experiments were performed on $Ar_{4.9}Br^-$ and $Ar_{8.19}I^-$. The experiments yield size-dependent adiabatic electron affinities (EAs) and electronic state splittings of the halogen atom in the neutral clusters formed by photodetachment. These results are compared with simulated annealing calculations using model potentials for the anion and neutral clusters, making use of the neutral and anion pair potentials determined from previous work on the diatomic rare gas-halide atom complexes. A simple first-order degenerate perturbation theory model of the neutral cluster potentials was found to agree well with the size-dependent splitting of the halogen $^2P_{3/2}$ state observed in the ZEKE spectra. However, the binding energies calculated from the pair potentials alone were found to be inconsistent with the experimental electron affinities, and it was necessary to include various non-additive terms in the simulated annealing calculations to obtain reasonable agreement with experiment. Many-body induction in the anion clusters was found to be the dominant non-additive effect. The exchange quadrupole effect — i.e., the interaction of the exchange induced electron charge distribution distortion among argon atoms with the halide charge — was also found to be important. This comparison between experiment and theory provides a sensitive probe of the importance of non-additive effects in weakly bound clusters.
PUBLICATIONS:


