

-ittle: Polyatomic Photodissociation Rynamic 394 Principal Investigator: Stephen J. Riley Assistant Professor (DFinal rept. Department of Chemistry 20 Yale University AD A O 5 8 0 8 New Haven, Conn. August 6, 1975 Polyatomic Photodissociation Dynamics . AUG 25 1978 6 Una During the contract period puotofragment spectroscopy studies of several small halide molecules were carried out. In two cases evidence for novel free radical dissociation processes was found. 1) Methylene iodide (CII_2I_2) . When photolyzed at a wavelength of 266nm direct evidence for the following dissociation pathway was seen:

 $CH_2I_2 + hv \Rightarrow CH_2I + I.$



The fragments were found to carry a very high fraction (85-90%) of excess energy in internal excitation, most likely in the form of rovibrational excitation of the CH₂I species. While this result is consistent with a simple statistical model for energy partitioning, the fragment angular distribution indicates parent molecule breakup occurs very quickly on a rotational timescale, so that a statistical decay is unlikely. A simple direct inpulsive model, which treats the carbon atom as loosely bound to the rest of the radical and requires momentum censervation as the carbon atom recoils from the I atom, predicts an internal excitation of 84% of available energy, in good agreement with experiment.

A second dissociation process was observed, yielding CH₂ fragments with surprisingly high translational energy. The lowest energy assumption, that the process is

$CH_2I_2 + hv + CH_2 + I_2$,

was found to be inconsistent with the generally accepted value (91kcal mole⁻¹) of the heat of formation of CH_2 . If the above process were occurring, the observed photofragment energetics would place an <u>upper</u> limit to the heat of formation of CH_2 of 40 kcal mole⁻¹. Two possible explanations were considered: photodissociation of a dimeric species in the molecular beam and a two photon dissociation process. The former was ruled out by experiments at elevated molecular beam temperatures, the latter was presumably excluded by the observed linear dependence of CH₂ signal on laser intensity. However, if the second photon absorption

DISTRIBUTION STATEMENT A Approved for public release Distribution Unlimited

382 475 B

has a very large extinction coefficient, this dependence would appear linear except at very low laser intensities, where poor signal-to-noise ratios make accurate measurements impossible. The existence of a second absorption was proven by measuring CH_2I signal as a function of laser intensity. This dependence shows a strong saturation at relatively low light levels, indicating some photon-dependent depletion process is occurring. Thus there must be a sequential two photon absorption, in which an initially formed CH_2I radical absorbs a second photon and dissociates to $CH_2 + I$. The detailed dynamics of this radical dissociation process are still not understood. Theories of two photon angular distributions will be applied to the data to determine the extent of correlated dynamics in the three-body dissociation system.

2) Acetyl iodide (CH₃COI). As expected, the primary photodissociation process in acetyl iodide at 266nm is C-I bond breakage and recoil of an I atom fragment. However, the excess available energy above that needed to break the C-I bond is sufficient to further decompose the acetyl fragment to CO and CH₃. In fact, no evidence for a recoiling acetyl fragment could be found. Photofragment spectra of both CO and CH₃ were observed, but could conceivably have arisen from dissociation of an acetyl fragment in the detector's electron bombardment ionizer. To test this, the spectra can be transformed to recoil translational energy distributions. If both CH₃ and CO signal originated from a recoiling CH₃CO radical, their energy distributions would be identical, and would match that observed for the I atom fragment. This was not the case. Also, the angular distributions for CO and CH₃ are more isotropic than for I, indicating something more complex than a simple two-body breakup is occurring.

Acetyl iodide thus provides another example of a radical dissociation process, initiated by the impulsive energy release during C-I bond breakage. The simplest dynamical model might predict the primary initial excitation of the acetyl radical to be in the CCO bending mode, yet dissociation must occur from a radical highly excited in the C-C stretching vibration. Thus the radical might live for many vibrational periods, undergoing statistical redistribution of its excess energy prior to decomposing via a classic unimolecular decay process. Further theoretical work on three body kinematics will be needed to tell if this, rather than instantnaeous decomposition, is happening. The results should prove useful to an understanding of intramolecular energy transfer processes.

3) Tin tetrabromide (SnBr₄). A clear example of statistical unimolecular decay of photoexcited species was found in the ultraviolet photodissociation of SnBr.. Fragment recoil translational energy distributions peak at or near zero and have an exponential fall-off, exactly the behavior expected for classical unimolecular decomposition. Statistical decay is the heart of many reaction rate theories, yet experimental verification of its theoretical aspects has not been easy. Crossed molecular beam reaction and ionic dissociation studies have provided some input, but have been complicated by large and sometimes ill-defined distributions in kinematic parameters such as angular momentum and excess internal energy. Photodissociation, on the other hand, offers a "cleaner" experimental test, since the angular momentum of the complex is known (just the thermal momentum of the parent molecule) and the excess energy (photon energy minus bond strength) is known, has a narrow distribution, and can be systematically varied via wavelength selection. Analysis of photofragment spectra should provide insight into such basic aspects of unimolecular decay theories as determining the effective number of oscillators and the role of zero point energy.

Watte Section atrs. 202 SMAR DER TH arsteinetion / AVAILABILITY CODES Sist AVAIL and/or SPESIAL