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determine immediately that the spin-orbit splitting is 61.5 ± 1.0 cm⁻¹. Further analysis of the rotational structure of these bands may be expected to refine this, but should not change it by more than two wave numbers.

In addition, a preliminary analysis of the vibrational progressions provides strong evidence that the most intense, highest frequency spin-orbit component of the origin band is actually at 32 306 ± 2 cm⁻², rather than in the region 31 540 cm⁻¹ supposed by other workers. This fact could have easily been missed in previous work because both of the origin components adjoin strong OH lines!

The value of the spin-orbit splitting observed is closer to the value determined from the LMR data⁹ rather than to that of the theoretical prediction.¹³ This theory should be reexamined. Of the three possible theoretical influences on the spin-orbit splitting, we suspect that the orbital quenching in this radical is not as large as supposed. Two pieces of information point to this. First, preliminary analysis of the spectra is consistent with a total Jahn–Teller coupling of about k² = 0.5, which could account for almost all of the reduction from about 140 to 60 cm⁻¹. Second, orbital quenching would require significant delocalization of the valence hole on oxygen onto the hydrogens. The fact that the transition origin is almost exactly at the energy of the analogous transition in OH suggests that the transition is entirely located on the oxygen, with little shift caused by hyperconjugation onto the hydrogens.

We have not yet completed an analysis of the methoxy emission system. A full vibrionic analysis of the spectrum and that of the deuterated species should provide an understanding of the Jahn–Teller effect and the mechanism of orbital quenching or reduction in this simple system.

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Picosecond monitoring of a chemical reaction in molecular beams: Photofragmentation of R–I→R⁺ + I

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Studies of chemical reactions by molecular beam techniques and spectroscopic methods have provided extensive information on the nature of the intermediates involved and on the energy distribution in the products.¹–⁶ Until now, however, there is no report of direct picosecond time resolution of a chemical reaction in molecular beams.

In this communication, we report our first picosecond experimental results on photofragmentation reactions with product state detection. The experiments are performed using the pump–probe picosecond/mass-spectrometry technique.⁷ Two picosecond pulses, delayed in time by a Michelson interferometer, are spatially overlapped in a pulsed molecular beam; the pump pulse initiates the photofragmentation while the probe pulse monitors the dynamics of product state formation using TOF mass spectrometry (see Fig. 1). The reactions studied are those of alkyl iodides, which have been extensively studied by photofragment spectroscopy:⁸⁻¹₀

$R-I \rightarrow R^+ + I$²±$P_{3/2}$ or $P_{1/2}$.

In this report R stands for CH₃ or CF₂⁻I–CF₂. The two spin-orbit states of the iodine atom are probed separately by tun-
ing the wavelength of the probe pulse (two-photon absorption) to 304.0 \( (2P_{1/2}) \) or 304.7 nm \( (2P_{3/2}) \). The results show that for \( \text{CH}_3\text{I} \) the primary C–I bond breakage is occurring within our pulse duration, while for \( \text{CF}_2\text{I–CF}_2\text{I} \) there is a sequential bond breakage; a similar primary and a slower (32 ps) secondary C–I breakage. The scheme for our experiments is illustrated in Fig. 1.

Methyl iodide, whose C–I \( n \rightarrow \sigma^* \) transition is at \( \sim 2800 \) Å, is known to photofragment and form iodine in both the 1/2 and 3/2 states. The \( \text{CH}_3 \) fragment produced is vibrationally excited. Figure 2 shows results obtained by probing \( \text{I} (2P_{1/2}) \) and similar results (not shown) were obtained when probing \( \text{I} (2P_{3/2}) \). In both cases, the iodine signal rises within the pulse and we can fit our data to a buildup time of 0.5 ps. The photodissociation lifetime has been estimated to be on the order of a rotational period, and our results for \( \text{CH}_3\text{I} \) indicate that \( \tau \lesssim 0.5 \) ps.

In Fig. 2, we also show the results for \( \text{I–CF}_2\text{CF}_2\text{I} \). When \( \text{I}^* \) was monitored the signal rose within the pulse similar to the observation made on \( \text{CH}_3\text{I} \). On the other hand, when \( \text{I} \) was monitored a “biexponential” buildup was observed; the least-squares fit gives \( \tau_1 = 0.5 \) and \( \tau_2 = 32 \pm 10 \) ps.

The above preliminary results indicate the following: First, the primary C–I bond breakage in \( \text{CH}_3\text{I} \) occurs within the time resolution of our picosecond/mass spectrometry when either \( \text{I} \) or \( \text{I}^* \) is monitored. Second, in \( \text{I–CF}_2\text{CF}_2\text{I} \), the buildup time is 32 ps for ground state iodine and very short for the excited iodine. These observations can be understood in terms of a simple sequential bond breakage: a prompt photofragmentation on the repulsive surface (Fig. 1) occurs first, followed by a secondary bond breakage of the other C–I bond. The dynamics of this prompt breakage can be understood using available theoretical techniques. The wave packet approach, as highlighted recently by the work at MIT, indicates that the lifetime is less than a picosecond. The question now: what determines the dynamics of this slower secondary process?

The observation of a 30 ps rise by monitoring I (and not \( \text{I}^* \)) indicates that after the recoil of the fragments in the primary fragmentation the total internal energy in \( \text{R}^2 \) is sufficient for the radical to undergo secondary dissociation and produce I in the ground state. Lee’s group has done a thorough study of the translational energy distribution for similar reactions. From the photon energy (102 kcal/mol) of our experiment, the C–I bond energy (52.5 kcal/mol) and the mean translational energy of similar systems we conclude that the internal energy of the fragment is comparable to the activation energy and that bond breakage of the second io-
dine takes place. The (30 ps)$^{-1}$ represents the average rate for this secondary bond breaking process. This is consistent with the fact that this buildup was only seen when I (and not 1*) was monitored.

The observed "biexponential" buildup of I (Fig. 2) can be related to the internal energy distribution in CF$_3$I–CF$_2$. Assuming a thermal vibrational distribution (width of 6 kcal/mol)$^2$ that peaks near the barrier (taken from thermochemical estimates to be 3–5 kcal/mol), and taking the dependence of the rates on excess energy ($E_x$) to be of the form $Ae^{-α/E_x}$, we obtain a biexponential buildup from this convolution procedure, consistent with our experimental findings. We are currently examining details of the fragmentation pathways, possible surface crossing, and RRKM calculations.

In conclusion, picosecond probing of reactions in beams offers new opportunities for real-time studies of the dynamics of bond breakage. This first report confirms our optimism about the sensitivity of the method which provides excellent time resolution and allows for product state analysis since two color picosecond pulses are used. Extension to other systems, including those studied by Parker and Bernstein,$^{18}$ is currently in progress. We are also extending these studies to the femtosecond time domain.

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1For more reviews, see Refs. 2–5, and the book by R. B. Bernstein, Chemical Dynamics via Molecular Beam and Laser Techniques (Clarendon/Oxford University, Oxford, 1982).


10The absorption spectrum at room temperature shows a broad band at 265 nm. This compound, obtained from PCF Specialty Chemicals, was used without further purification. The GC, UV absorption, and mass spectrum are all consistent with a purity (other than 1) of 98% or better. 14, however, absorbs weakly at λ of interest and has a much lower vapor pressure [Ref. 8(a)].

11In these experiments, the observed fragment is 1 + and there is no detectable 1–, indicating that there is no concerted bond breakage. This is consistent with results on di-iodomethane by Kawasaki et al. [Ref. 12] and Kroger et al. [Ref. 13].

12M. Kawasaki, S. J. Lee, and R. Bersohn, J. Chem. Phys. 63, 809 (1975). These authors presented a clear discussion of the reasons for consecutive bond breakage in di-iodomethane, where resonance interactions between the two C–I bonds is quite strong.


NOTES

Direct measurement of the stop gap for the 31 475 cm$^{-1}$ singlet $a$ exciton of naphthalene$^{a}$

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Recently,$^{1–4}$ we have utilized phase-matched two-photon excitation or polariton fusion$^{5–4}$ to study the spatial dispersion and scattering properties of the polariton associated with the lowest singlet $a$ exciton of naphthalene at $ω_P = 31 475$ cm$^{-1}$. For reasons noted below, these nonlinear results stimulated the present study which deals with the one-photon fluorescence excitation profile from the $a$ exciton.

The $a$ exciton possesses $A_s$ factor group ($C_{2h}$) symmetry and an oscillator strength $F$ of only $2.3 \times 10^{-3}$ (Ref. 4) and lies at the bottom of the band. In the nonlinear experiments,$^{1–4}$ both the SHG (second harmonic generation) and TPA (two-photon absorption) resonant profiles were monitored at a resolution of $\sim 0.06$ cm$^{-1}$. Both processes are weakly allowed via a magnetic dipole–electric dipole mechanism.$^9$ Accurate determinations of the upper and lower polariton branches were made and the curves fit theoretically with $F$ fixed at the above value (determined from one-photon absorption measurements$^4$). The analytic expressions for the dispersion curves were used to show that polaritons with group velocities between $\sim 5 \times 10^6$ and $\sim 5 \times 10^4$ m s$^{-1}$ can be generated by angle tuning of $ab$ plate crystals about the $b$