Isotope Effects in the Association Reactions of Methyl and Ethyl Iodide Cations*  
L. Wayne Sieck  
Institute for Materials Research, National Bureau of Standards, Washington, D. C. 20234  
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Rate coefficients for production of stabilized dimeric parent cations at 295 K have been determined in CH$_3$I, CD$_3$I, CH$_2$I—CD$_2$I mixtures, C$_2$H$_5$I, CH$_3$CD$_2$I, CD$_3$CH$_2$I, and C$_2$D$_5$I. These processes are the most rapid reported for association reactions, the various individual values falling within the limits 0.33 X 10$^{-24}$ (CH$_3$I) and 10.1 X 10$^{-24}$ cm$^6$ molecule$^{-2}$ sec$^{-1}$ (C$_2$H$_5$I). The temperature dependence of the stabilization coefficients in CH$_3$I and CD$_3$I was also measured over the range 220 ± 3 to 320 ± 1 K, as well as the efficiencies of other third bodies in the stabilization process. The differences observed for the variously labelled analogues are interpreted in terms of vibrational level (energy) depression upon deuteration, which affects the intrinsic lifetime of the collision complex.

Key words: Alkyl iodides; ion-molecule reactions; isotope effects; mass spectrometry; photo-ionization; rate constants.

1. Introduction

Several years ago [1] we reported a study of parent dimer cation formation in the methyl halides using high pressure photoionization mass spectrometry, which extended earlier work by Hamill and co-workers [2-4] concerned with association ion production in the ethyl and propyl bromides and iodides. The common characteristic of these systems is the high overall efficiency for the third order process  

$$A^+ + 2A \rightarrow (A)_2^+ + A$$  

(1)  

where $A^+$ denotes the parent molecular ion. More recently [5] we completed an investigation of dimer formation in low molecular weight aromatic hydrocarbons in which we found that production of (C$_6$D$_6$)$_2^+$ in C$_6$D$_6$ was a factor of 2.7 times faster than (C$_6$H$_6$)$_2^+$ formation in C$_6$H$_6$ at 295 K. In view of this pronounced isotope effect, we initiated a new photoionization study involving the methyl and ethyl iodides emphasizing the effect of isotopic substitution on the rate coefficient for process 1. These molecules were considered to be ideal systems for detailing some of the general factors influencing stabilization of ion-molecule complexes since >90 percent of the parent ions produced initially via photoionization are in $v = 0$ of the ground electronic state [6] (internal energy effects are minimal), the variously labelled deuterium analogues are readily available, and the room temperature stabilization coefficients are sufficiently high to permit experiments at reduced collision frequencies under controlled conditions.

2. Experimental Procedure

The NBS high pressure photoionization mass spectrometer equipped with a field-free reaction chamber was used for all experiments. Ionization of methyl iodide (I.P. $^7E_{1/2} = 9.54$ eV) and ethyl iodide (I.P. $^7E_{1/2} = 9.34$ eV, I.P. $^7E_{3/2} = 9.93$ eV) [7] was induced via photoabsorption at 123.6 nm (10.0 eV) using a Kr resonance lamp equipped with MgF$_2$ optics. Measurements other than those at 295 K were taken with a variable temperature modification described elsewhere [8].

3. Results

3.1. General Behavior

Typical composite spectra found for methyl and ethyl iodides as a function of pressure are given in figure 1 for CD$_3$I.
and C₃D₅I. In every system the competing reactions

\[ \text{RI}^+ + \text{RI} \rightarrow k \frac{1}{2} (\text{RI})^* \]

followed by

\[ (\text{RI})^* \rightarrow \text{R}_3^+ + \text{I} \]

or

\[ (\text{RI})^* + \text{RI} \rightarrow (\text{RI})^* + \text{RI} \]

where RI = methyl or ethyl iodide, were observed. Process 2 involves bimolecular formation of a transient collision complex (RI)^*, which either dissociates to reform reactants or eliminates an iodine atom (process 3) at lower densities. At higher densities collisional stabilization (process 4) is facilitated, as indicated in figure 1 by the rapid formation of (CD₃)₂⁺ and (CH₃I)⁺ and the concurrent decrease in the residence times of the respective dialkyliodide ions.

### 3.2. Reaction Rates

Rate coefficients for process 3 at 295 K were derived from the initial slopes of the growth curves found for R₃⁺ when the respective fractional yields were plotted in semilogarithmic form versus RI concentration. Residence times (\(\tau\)'s) for various RI⁺ ions at the zero pressure limit were approximated by first evaluating average \(\tau\)'s for several ions of differing m/e ratios at 295 K under molecular flow conditions (<1-2 millitorr)² using calibration reactions [9] and assuming the usual (mass)\(^{1/2}\) dependence for relative \(\tau\)'s. Calibration reactions included \(i-C_4H_9^+ + CH_3NH_2 \rightarrow CH_2NH_2^+ (k = 1.30 \pm 0.15 \times 10^{-10}),\) \(C_2H_5^+ + i-C_4H_9 \rightarrow C_4H_9^+ (k = 4.2 \pm 0.2 \times 10^{-19}),\) \(C_2H_5^+ + CH_3H \rightarrow \text{products} (k = 8.0 \pm 0.5 \times 10^{-19})\), and \(NH_2^+ + NH_2 \rightarrow NH_4^+ (k = 1.9 \pm 0.2 \times 10^{-9})\).

Third order stabilization coefficients (process 4) were calculated from the initial slopes of the logarithm of the fractional intensities of (RI)^* versus (concentrations) using the same relative \(\tau\)'s for the various RI⁺ ions. Residence times at other temperatures were calculated by applying the appropriate velocity corrections.

### 3.3. Methyl Iodides

Rate coefficients for production of dialkyliodide ions in CH₃I, CD₃I, and CH₃CD₂I (1:1) mixtures at 295 K are given in table 1, along with values for CH₄I and CD₄I at 220 ± 3 and 320 ± 1 K. Within the indicated reproducibility limits, the coefficients for (CH₃)₂⁺ formation in CH₃I and (CD₃)₂⁺ in CD₃I were invariant at 2.4 ± 0.1 and 2.25 ± 0.1 \(\times 10^{-11}\), respectively, over this temperature range. The overall 295 K coefficient for production of (CH₃)₂⁺ in CH₃I—CD₃I (1:1) mixtures was found to be \(2.3 \pm 0.2 \times 10^{-11}\). Products indicating isotopic scrambling such as (CH₂D)(CD₃)⁺, (CD₂H)(CH₃)⁺, etc., were not observed; the only second order products detected were (CH₃)₂⁺, (CD₃)₂⁺, and (CD₃)⁺ with an essentially statistical distribution (0.26:0.50:0.24).

In contrast to the near equivalency of the halogen elimination rates, substantial isotope effects were found in the third order coefficients for collisional stabilization of (CX₃I)₂⁺ (X = H or D). Data for CH₃I, CD₃I, and CH₃—CD₃I (1:1) mixtures are shown in figure 2, which gives the appropriate (CX₃I)₂⁺/(CX₃I)⁺ ratios versus CXI concentration. Relative stabilization efficiencies are given by the slope ratios after correction for the slight differences found in the production rates for (CX₃I)⁺, and were determined to be 1.0, 1.8(3), and 2.8(0) in CH₃I—CD₃I (1:1), and CD₃I respectively. Absolute values, as well as those determined at 320 ± 1 K in CH₃I and CD₃I, are listed in table 1. The isotopic compositions of the (CX₃I)₂⁺ dimer ions formed in (1:1) CH₃I—CD₃I mixture as a function of concentration and percent reaction of (CX₃I)⁺ are given in figure 3. Again no evidence was found for hydrogen-scrambled species such as (CH₂D)(CD₃)⁺, etc.; i.e., the methyl groups retain their isotopic integrity during stabilization.

A number of measurements were also taken to estimate the stabilization efficiencies for other third bodies relative to methyl iodide itself. These results, which were obtained in mixtures containing ~5 percent CH₃I or CH₄I in the bulk additive, are shown in figure 4. Approximate relative values in CH₃I were derived from the additive concentration at which the intensity of (CH₃I)₂⁺ was equal to that for (CH₃)₂⁺.

![Table 1. Rate coefficients for second and third order reactions in methyl and ethyl iodide.](image-url)

<table>
<thead>
<tr>
<th>System</th>
<th>Temp. (K)</th>
<th>k(RI)⁺⁺</th>
<th>k(RI)⁺⁺⁺</th>
<th>Relative k(RI)⁺⁺⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₃I</td>
<td>295 ± 1</td>
<td>2.37 ± 0.12 (\times 10^{-11})</td>
<td>0.53 (\times 10^{-14})</td>
<td>1.0</td>
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<tr>
<td>CH₃I</td>
<td>220 ± 3</td>
<td>2.36 ± 0.25 (\times 10^{-11})</td>
<td>0.45 (\times 10^{-14})</td>
<td>1.0</td>
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<td>CH₄I</td>
<td>320 ± 1</td>
<td>2.37 ± 0.05 (\times 10^{-11})</td>
<td>0.54 (\times 10^{-14})</td>
<td>1.0</td>
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<tr>
<td>CD₃I</td>
<td>295 ± 1</td>
<td>2.23 ± 0.05 (\times 10^{-11})</td>
<td>1.04 (\times 10^{-14})</td>
<td>2.8</td>
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<tr>
<td>CD₃I</td>
<td>220 ± 1</td>
<td>2.04 ± 0.12 (\times 10^{-11})</td>
<td>1.26 (\times 10^{-14})</td>
<td>2.8</td>
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<tr>
<td>CD₄I</td>
<td>320 ± 1</td>
<td>2.16 ± 0.12 (\times 10^{-11})</td>
<td>0.67 (\times 10^{-14})</td>
<td>2.8</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.3 ± 0.1 (\times 10^{-11})</td>
<td>0.61 (\times 10^{-14})</td>
<td>1.8</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>285 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>4.1 (\times 10^{-14})</td>
<td>1.0</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>3.3 (\times 10^{-14})</td>
<td>1.4</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>7.4 (\times 10^{-14})</td>
<td>2.79</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>10.1 (\times 10^{-14})</td>
<td>4.44</td>
</tr>
<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>0.194 (\times 10^{-14})</td>
<td>1.0</td>
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<tr>
<td>CH₃I—CD₃I (1:1)</td>
<td>295 ± 1</td>
<td>2.0 ± 0.1 (\times 10^{-11})</td>
<td>0.355 (\times 10^{-14})</td>
<td>2.71</td>
</tr>
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</table>

² Units are cm² molecule⁻¹ s⁻¹.
³ Values taken from Reference 5.
3.4. Ethyl iodides

Measurements in C₂H₅I, CH₃CD₂I, CD₃CH₂I, and C₂D₅I were taken at 295 K only. The rate coefficients for production of diethyliodonium ions, (C₂H₅I)₂⁺, were determined to be 2.0 ± 0.15 × 10⁻¹¹ for all analogues. Stabilization rates (process 4) were again found to increase with increasing deuteration in the order C₂H₅I = 1.0, CH₃CD₂I = 1.28, CD₃CH₂I = 1.79, and C₂D₅I = 2.44. On an absolute scale, the third order coefficient for stabilization of (C₂H₅I)₂⁺ was deter-
mained to be $4.1 \times 10^{-24}$, or a factor of 12 higher than that observed in CH$_3$I (see table 1).

Biomolecular formation of long-lived dimeric ions was also directly detected in the ethyl iodides. Figure 5, which gives plots of $(C_2X_3I)_2^+/(C_2X_3I)_1^+$ versus concentration, indicates non-zero intercepts for all of the analogues, including C$_2$H$_2$I. Observation of these entities corresponds to lifetimes for detected $(C_2X_3I)_2^+$ on the order of several milliseconds, which is the total transit time in our apparatus for ions of these approximate masses. The plots also exhibit downward curvature at higher concentrations (higher densities), which is ascribed to formation of undetected trimeric ions having masses beyond the operational range of the quadrupole mass filter used in these experiments (~m/e 400). Association ions such as [(C$_2$H$_5$)$_2$I·C$_2$H$_5$I]+, etc., which correspond to solvation of dialkylidionium ions, were not detected in either the ethyl or methyl iodides at pressures up to 0.1 torr, which was the maximum in the present study.

![Figure 5](image)

**Figure 5.** $(C_2X_3I)_2^+/(C_2X_3I)_1^+$ ratio (X = H or D) as a function of the extent reaction of CH$_3$I, CH$_3$CD$_2$I, CD$_3$CH$_2$I, and C$_2$I$_3$.

### 4. Discussion

As expected [12] for association reactions, a negative temperature coefficient was observed for the third order stabilization of $(C_3X_3I)_2^+$ in CH$_3$I and CD$_2$I (see table 1). However, the variation with temperature does not yield a straight line when $k_{(stab.)}$ is plotted versus $T^{-n}$ for any values of $n$ from 0.1 to 5, although unique fits of this form have been obtained for other systems. The lack of a unique functional form or profound temperature effect in methyl iodide is ascribed to the fact that the stabilization coefficients are extremely high over the range studied, so that no substantial dependence would be anticipated except at much higher temperatures (other studies [12] have been restricted to association reactions having 300 K stabilization coefficients 2-5 orders of magnitude slower than those characteristic of the alkyl iodides). In addition, the ratios of the third order coefficients in CD$_3$I and CH$_3$I at any temperature are constant at 2.8, which would be expected since the physical factors involved in the stabilization mechanism are presumably the same for both analogues.

Consideration of the relative stabilization efficiencies for the labelled ethyl iodides (table 1) indicates that deuteration on either the methyl or methylene carbon facilitates stabilization (comparison of CH$_3$CD$_2$I and CD$_3$CH$_2$I with C$_2$H$_5$I or C$_2$D$_5$I). Based on the relative values, attempts were made to calculate the effect of D substitution on both carbons choosing various trial isotope effects at each site. However, no unique self-consistent set of relative or individual contributions compatible with the experimental ratios could be computer-generated due to the fact that all models predicted a substantially higher relative value for CH$_2$CD$_2$I ($\geq$15 percent higher, which was considered outside the possible error in measurement). The only common result of all calculations was that substitution of D for H on the methylene carbon has a slightly lower effect (~10 percent) than exchanging a D for an H on the methyl group.

Among all of the active vibrational fundamentals in the alkyl halides, the methyl deformation and the methylene rocking modes (ethyl iodide), which lie substantially lower in energy than the C—H stretch, are the most easily excited. The effect of deuteration is to reduce the C—H stretching, CH$_3$ deformation, and CH$_2$ rocking frequencies by a factor of ~0.3, while the C—I stretching frequency remains essentially unaffected [13]. The large differences observed in the third order stabilization coefficients for the variously labeled iodides are therefore ascribed to a depression of the deformation and rocking frequencies (shift to lower energies and higher state densities) upon deuteration substitution. This depression is manifested by an increased dissociative lifetime for the unstable initial ion-molecule collision complex, as evidenced by the fact that the relative yields of the biomolecular produced ethyl iodide dimer ions which survived ion transit (extrapolated zero conversion intercept in figure 4) also appear to increase with increasing deuteration. The lack of isotopic scrambling also indicates that both reacting partners retain their structural integrity within the excited collision complex, facilitating a favorable energy level match between the third body and the excited dimeric ion in the pure halide systems. This favorable condition would account for the magnitude of the stabilization coefficients, which are the highest recorded for third order association reactions at 295 K.

With respect to the measured stabilization efficiencies of other third bodies relative to methyl iodide itself, it is somewhat surprising that methane and Xe exhibit the same relative value (~0.5) since Xe can only deactivate via vibrational to translational energy conversion. Also, CH$_4$ and CD$_4$ are found to be equally effective even though there is a 30 percent decrease in the vibrational frequencies for CD$_4$ relative to CH$_4$, which should increase the probability for energy transfer. The low overall efficiency found for methane may be due either to the fact that the duration of the collision between excited dimeric ions and methane is less than the duration with Xe, SF$_6$, or CH$_3$I (based on relative velocity considerations), or that the active vibrational frequencies in
the excited dimers are substantially lower than those associated with the methane fundamentals (≥ ~1000 cm⁻¹). Sulfur hexafluoride has both a high mass (146 a.m.u.) and several very low-lying vibrational modes (<800 cm⁻¹) which accounts for the observation that it is slightly more effective than CH₃I itself as a deactivator for (CH₃I)₂.

In conclusion we would like to emphasize the intriguing fact that the stabilization efficiencies for CD₃I, C₂D₅I, and C₆D₆ (see table 1) are all 2.6 ± 0.2 times higher than those for the perprotonated analogues. Whether this equivalence is fortuitous, is due to common physical property of the excited dimer cation in these particular systems, or is generally characteristic of polyatomic organic ion-molecule collision complexes which have no highly exothermic or favorable fragmentation channels, deserves further study.

5. References


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