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Citation: The Journal of Chemical Physics **74**, 5093 (1981); doi: 10.1063/1.441716 View online: http://dx.doi.org/10.1063/1.441716 View Table of Contents: http://scitation.aip.org/content/aip/journal/jcp/74/9?ver=pdfcov Published by the AIP Publishing

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Role of impact parameter in branching reactions: Chemical accelerator studies of the reaction $Xe^++CH_4 \rightarrow XeCH_3^++H$

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Integral reaction cross sections and product velocity distributions have been measured for the ion-molecule reaction Xe⁺(CH₄,H)XeCH₃⁺ over the relative reactant translational energy range of 0.7-5.5 eV by chemical accelerator techniques. The kinematic results indicate that reaction proceeds in a direct manner by a rebound mechanism over the energy range studied, suggesting that this substitution reaction occurs predominantly in small impact parameter collisions. This finding contrasts with the results obtained for the competing reaction, Xe⁺(CH₄,CH₃)XeH⁺, where the strong forward scattering of the XeH⁺ product indicates that H-atom abstraction occurs primarily in large impact parameter collisions.

I. INTRODUCTION

An important problem in chemical kinetics is the identification of the factors which determine which products will be formed when a given pair of reactants can form two or more sets of products, as in

$$A+B-C+D \text{ or } E+F$$
, etc. (1)

One approach has been to assume that the reactants form an intermediate complex, (AB)*, and that competition between the various reaction channels is determined by the relative rates of unimolecular decomposition of the complex to form the various sets of products. If it is further assumed that the lifetime of (AB)* is sufficiently long to permit equilibration of the excess energy among the various internal degrees of freedom, then the relative rates of decomposition can be calculated from statistical considerations. The general conclusion of the statistical theory is that the reaction will predominantly follow the most excergic channel.¹

However, statistical theories of unimolecular decomposition would appear to be inapplicable to those reactions which proceed by a direct mechanism rather than by the formation of a long-lived complex because the duration of the collision (usually $\sim 10^{-13}$ sec) in these reactions is insufficient for redistribution of energy within the collision complex. For such reactions it has been postulated² that the reaction path is determined by steric factors (i.e., by the impact parameter and the angle of collision). Until recently, however, evidence in support of this hypothesis has been circumstantial in nature, so that it has been impossible to determine rigorously whether a particular yield pattern was due primarily to steric factors or to other considerations, such as bond energies and inertial effects.³

The first direct evidence to support the hypothesis that factors such as the impact parameter determine the reaction path followed in a given collision was obtained from

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a chemical accelerator study of reactive scattering in the system $Kr^* + CH_4$.⁴ In the hope of obtaining further information concerning the role of steric factors in direct reactions, we have studied the competing ionmolecule reactions

$$Xe^{*} + CH_{4} \rightarrow XeH^{*} + CH_{3}$$
⁽²⁾

$$- XeCH_3^* + H .$$
 (3)

Reaction cross sections and product velocity vector distributions have been measured at collision energies over the range 0.4-8 eV (cm). The data for Reaction (2) have already been published,⁵ so this paper will present data for Reaction (3) only. The results are similar to those obtained for the Kr^{*} + CH₄ system⁴ and thus support the hypothesis that, in direct reactions, factors such as the impact parameter determine which products will be formed.

Ion-molecule reactions (including charge transfer) in xenon-methane mixtures have previously been studied by high-pressure mass spectrometry,⁸ ion cyclotron resonance, ^{7,8} and tandem mass spectrometry.^{9,10}

II. EXPERIMENTAL

A. Apparatus

The instrument used in this study, a single beam, collision chamber type of chemical accelerator (with which product velocity and angular analysis are possible) has been described in detail elsewhere. ^{5,11} It should be pointed out, however, that in the present study the Xe⁺ reactant ions were produced by impact of 33 eV electrons, so that nearly all (98%) of the Xe⁺ ions are in the ^{2}P state and are presumably distributed statistically in a 2:1 ratio between the ${}^{2}P_{3/2}$ ground state and the ${}^{2}P_{1/2}$ excited state, which differ in energy by 1.61 eV.¹²

B. Heats of reaction

Holtz and Beauchamp⁷ have reported a value of 1.87 ± 0.35 eV for the methyl cation affinity of Xe. Assuming values of 4.477 eV for $D_0(H-CH_3)^{13}$ and 9.840 eV for the ionization potential of CH_3 , ¹² one finds that Reaction (3) is endothermic by 0.32 ± 0.35 eV for ${}^{2}P_{3/2}$ ground state

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TABLE I. Integral cross sections for the reaction $Xe^+ + CH_4 \rightarrow XeCH_3^+ + H$.

Initial translational energy, E_T (eV)	Relative collection efficiency, K_c/K_A^a	Transmission factor, K(T) ^b	Integral reaction cross sections, σ $(10^{-16} \text{ cm}^2)^c$
0.81	1.44	0.72	0,013
1.08	1.77	0.77	0.033
1.56	2.43	0.54	0.077
2.20	1.53	0.70	0.22
2.39	1.77	0.83	0.23
2.96	1.74	0.75	0.24
4.29	2,45	0.83	0.20
4.38	1.28	0.77	0.23
5.18	1.72	0.83	0.085

^aRatio of collection efficiency correction factors (see Ref. 5, Sec. IIIA).

^bCorrection factor for attenuation of primary beam, calculated from Eq. (4) of Ref. 5.

Calculated from Eq. (4).

Xe⁺ ions but exothermic by 0.99 ± 0.35 eV for the ${}^{2}P_{1/2}$ excited state of Xe⁺.

A similar situation exists for the H-atom abstraction process. Reaction (2) is endothermic by 0.86 ± 0.11 eV for Xe^{*} (²P_{3/2}) but exothermic by 0.45 ± 0.11 eV for Xe^{*} (²P_{1/2}).

III. RESULTS AND DISCUSSION

A. Reaction cross sections

Integral reaction cross sections, σ , for Reaction (3) are calculated from the formula

$$\sigma = \frac{I_C / I_A}{n_B L} K(T) , \qquad (4)$$

where I_c is the total reactively scattered product $(^{132}\text{XeCH}_3^*)$ ion intensity, I_A is the primary $(^{132}\text{Xe}^*)$ ion intensity, n_B is the number density of CH₄ in the collision chamber, L is the collision path length, and K(T) is a factor which corrects the thin target formula, $I_c/I_A n_B L$, for any attenuation of the reactant and product beams.¹⁴ The details of these calculations have been described previously.^{5,14}

Table I presents the integral cross sections calculated from Eq. (4), along with the relative collection efficiencies K_C/K_A and the transmission factor K(T). The uncertainties in the cross sections are estimated to be $\pm 40\%$ relative to each other and $\pm 60\%$ on an absolute basis. The excitation function (σ versus collision energy) is presented in Fig. 1.

As shown in Fig. 1, the excitation function for formation of XeCH₃^{*} rises from an apparent threshold at a relative collision energy (E_T) of 0.7 ± 0.2 eV to a maximum in the region $E_T = 3 - 4$ eV, where $\sigma \sim 0.25$ Å², and then decreases with increasing collision energy. The lack of data at collision energies above 5 eV is due to the complication in the mass spectrum at higher collision energies because of the formation of XeCH₂^{*} and possibly XeCH^{*} with all possible isotopes of Xe. Below 5 eV, XeH^{*} and XeCH₃^{*} are the only detectable products and the data analysis is, therefore, straightforward. The magnitude of the cross section for XeCH₃^{*} formation is somewhat larger than that found for $KrCH_3^*$ formation (about 0.05 Å² at its maximum) in a previous study of the $Kr^* + CH_4$ system, ⁴ but is similar in magnitude to that found for $KrCH_2^*$ formation in that study.

Because the Xe^{*} reactant beam contains ions in both levels of the ²P doublet, it is not possible to deduce from the present data whether the two states differ in the rate at which they produce XeCH₃^{*}. On the one hand, Reaction (3) is exothermic by 0.99 ± 0.35 eV for the Xe^{*}(²P_{1/2}) but endothermic by 0.32 ± 0.35 eV for Xe^{*}(²P_{3/2}). Because most exothermic reactions of positive ions have no activation energy, ¹⁵ the observed threshold energy of 0.7 ± 0.2 eV could be explained by assuming that the ground state Xe^{*}(²P_{3/2}) ion reacts to form XeCH₃ with much greater propensity than does the excited ²P_{1/2} state, so that the threshold is simply a consequence of the endothermicity for the ground state reactants.

On the other hand, translational energy thresholds have been found for the exothermic reactions $N_2^*(CH_4, CH_3)N_2H^{*16}$ and $Ar^*(CH_4, CH_3)ArH^{*14}$.¹⁷ Although the thresholds found for these reactions ($\approx 0.1 \text{ eV}$) are con-



FIG. 1. Integral reaction cross section σ vs initial relative translational energy E_T for the reaction Xe⁺+CH₄ \rightarrow XeCH₃⁺+H (solid circles). The values of the cross sections, calculated from Eq. (4), are listed in Table I. Estimated experimental uncertainties in σ are indicated by error bars. The solid line represents an empirical fit to the experimental data.



FIG. 2. Product probability distribution for Reaction (3) at the relative collision energy $E_T = 1.91$ eV. The product ion intensities, normalized to 100 at the position of maximum intensity, are shown relative to the Cartesian system, P_C . Arrows represent scattering angle with respect to the center of mass (marked C.M.).

siderably lower than the threshold found for Reaction (3), it is clear that the observation of a translational energy threshold is not sufficient to prove that one is dealing with an endothermic reaction.

B. Kinematics

As described previously,⁵ the measured product translational energy distributions at various laboratory scattering angles were transformed to probabilities in Cartesian velocity space.¹⁸ The results are presented as contour maps showing the relative yields of XeCH₃⁺ as functions $P_C(u, \theta)$ of the c.m. velocity u and scattering angle θ .

Such product velocity vector distributions for XeCH^{*}₃ produced at relative collision energies E_T of 1.91 and 4.29 eV are shown in Figs. 2 and 3, respectively. At both energies the XeCH₃^{*} product is scattered almost entirely in the backward direction (i.e., opposite to the direction of the incident Xe* ion). This asymmetry about the $\theta = \pm 90^{\circ}$ plane is a clear indication that Reaction (3) proceeds in a direct manner (i.e., on a time scale shorter than a rotational period, about 10⁻¹² sec). As reported previously,⁵ the competing H-atom abstraction, Reaction (2), was also found to be dominated by a direct mechanism in this range of collision energies, although a transition to a mechanism involving the formation of a long-lived XeCH₄ collision complex was observed in the previous study at relative collision energies less than 0.5 eV.

Additional information on the reaction mechanism can be obtained from the product c.m. angular distributions, $I(\theta)$, derived from the product velocity vector distributions (Figs. 2 and 3) by the relation

$$I(\theta) = \int_0^\infty u^2 P_C(u, \theta) \, du \, . \tag{5}$$

The results, presented as differential polar cross sec-

tions, $d\sigma/d\theta = 2\pi I(\theta) \sin \theta$, are shown in Fig. 4. As can be seen, the displacement product XeCH₃^{*} is almost entirely scattered through angles greater than 90°. This finding contrasts sharply with that for the abstraction process, where the XeH⁺ product is found almost entirely at scattering angles less than 90° at comparable collision energies. (Compare Fig. 4 with Fig. 9 of Ref. 5.) To the extent that a correlation exists in direct reactions between impact parameter and scattering angle, ¹⁹ we can conclude that the displacement reaction producing XeCH^{*}₃ occurs principally in close (i.e., small impact parameter) collisions, while the abstraction reaction producing XeH⁺ occurs predominantly in glancing (i.e., large impact parameter) collisions. Thus, over the energy range considered here, competition between the two reaction channels is apparently governed by the collision geometry.

Similar results have been reported for the analogous reactions of Kr^* with CH_4 .⁴ As discussed in that paper, the principle of conservation of angular momentum provides a qualitative explanation for this dependence of reaction path upon impact parameter. Reaction will be limited to low angular momentum (i.e., small impact parameter) collisions *if* (1) the products reduced mass (with respect to translation) is much less than that of the

$$Xe^+ + CH_4 \longrightarrow XeCH_3^+ + H$$

E_= = 4.29 eV



FIG. 3. Product probability distribution for Reaction (3) at $E_T = 4.29$ eV.

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FIG. 4. Polar differential cross sections, $2\pi I(\theta) \sin\theta$, vs c.m. scattering angle θ for XeCH₃⁺ produced in Reaction (3) at the relative collision energies of 1.91 and 4.29 eV. These cross sections were obtained by integrating over the product probability distributions shown in Fig. 2 and 3, using Eq. (5) of the text, and then scaled so that the area under each curve equaled the measured integral reaction cross section at the energy, in accord with the relation $\sigma = 2\pi \int_0^{\pi} I(\theta) \sin\theta \, d\theta$.

reactants, and (2) the intermolecular potential lacks anisotropic terms which could convert reactant orbital to product rotational angular momentum. In the present case $\mu = 14.3$ amu, while $\mu' = 13.5$ amu for the abstraction products, but only 0.99 amu for the displacement products. For abstraction, $\mu' \simeq \mu$, and therefore even large amounts of reactant orbital angular momentum can be taken up by the products' orbital rather than rotational motion. Consequently, we can observe XeH* formation from a strippinglike mechanism occurring in glancing collisions. On the other hand, $\mu' \ll \mu$ for displacement. XeCH₃^{*} formation in large impact parameter collisions would be possible only if large amounts of the reactants' orbital angular momentum could be converted to rotational angular momentum in the products. Evidently this is difficult, because we see XeCH₃ produced by a rebound (large scattering angle) mechanism, with an integral cross section much smaller than that for XeH* production. Both observations indicate that displacement is possible only when the impact parameter (and hence the reactants' orbital angular momentum) is small.

Product velocity distributions, P(u), or relative translational energy distributions, $P(E'_T)$, can also be derived from Figs. 2 and 3 by appropriate integration over the scattering angle. Unfortunately, in the present case the mass ratio between the detected (XeCH₃) product and the undetected (H) product is so large that the recoil velocity u of the detected product is only a very small fraction $(\frac{1}{148})$ of the final relative velocity, u'_R . Because of this very unfavorable kinematic factor, even moderately wide spreads in u'_R arising from a distribution in E'_T will result in distributions in u that are much narrower than the experimental resolution.

An approximate one-dimensional treatment of the problem demonstrates that nearly all of the observed

spread in XeCH₃^{*} velocity is caused by the spread in the initial relative velocities arising from the velocity spread of the Xe^{*} reactant beam and the thermal motion of the CH₄ target molecules.

As Fig. 5 shows, the distribution $P_1(v_1)$ of reactant Xe^{*} laboratory velocities in the 0° direction can be fairly well approximated by a Gaussian distribution centered around the most probable Xe^{*} laboratory velocity, v_1^0 . The distribution of centroids, P(V), along the 0° direction, will then also be a Gaussian distribution centered around V_0 , the nominal velocity of the c.m. [see Eq. (A5) of the Appendix].

To determine the extent to which this spread in V contributes to the width of the observed XeCH₃^{*} c.m. velocity distribution, P(u), we have assumed that, at a given collision energy, all XeCH₃^{*} product ions actually have the same c.m. velocity, w_0 . As shown in the Appendix, this one-dimensional approximate treatment then predicts that the observed XeCH₃^{*} c.m. velocity distribution should be of the form

$$P(u) \propto \exp\left[-\left(\frac{u-w_0}{\sigma_{\rm cm}}\right)^2\right] \quad . \tag{6}$$

Figures 6 and 7 compare the experimental data at E_T =1.91 and 4.29 eV, respectively, with the distributions predicted by Eq. (6), using the parameters given in Table II. As Fig. 6 shows, agreement between the measured distribution and the predicted distribution is excellent at a c.m. scattering angle of 180°, indicating that the shape of the observed distribution is a consequence of the distribution of centroids rather than actual dynamics of the reaction. The only measured quantity carrying any dynamical information, then, is the most probable recoil velocity, which equals (in this approximate treatment) w_0 , the true recoil velocity of XeCH₃^{*}. The experimental distributions at smaller angles are skewed



FIG. 5. Distribution $P(v_1)$ of the reactant Xe⁺ ion laboratory velocities along the 0° axis vs $v_1 - v_1^0$, the deviation of the Xe⁺ laboratory velocity from its nominal value, v_1^0 . In panel (a), $E_T \approx 1.91 \text{ eV}$ and $v_1^0 \approx 5.09 \times 10^5 \text{ cm/sec}$; in (b) $E_T = 4.29 \text{ eV}$ and $v_1^0 \approx 7.63 \times 10^5 \text{ cm/sec}$. The solid circles represent the measured distribution, and the solid line represents the assumed Gaussian distribution given by Eq. (A4).



FIG. 6. Distribution $P(u; \theta)$ of the product XeCH₃^{*} ion c.m. velocities at various c.m. scattering angles for the experiment at a relative collision energy of 1.91 eV. The solid circles represent the experimental data, and the solid lines represent the product velocity distributions predicted by Eq. (6).

somewhat to higher velocities than Eq. (6) predicts, but this may well be a consequence of the angular divergence of the Xe^{*}, a factor which was not included in this approximate, one-dimensional treatment.

As shown in Fig. 7, the measured product velocity' distributions at E_T =4.29 eV are somewhat broader than would be expected from just the distribution of centroids,



FIG. 7. Distribution $P(u; \theta)$ of the product XeCH₃^{*} ion c.m. velocities at various c.m. scattering angles at $E_T = 4.29$ eV. The solid circles represent the experimental data, and the solid lines represent the product velocity distributions predicted by Eq. (6).

TABLE II. Parameters for calculating product velocity distribution, P(u).

	$E_T = 1.91 \text{ eV}$	$E_{T} = 4.29 \text{ eV}$	
$\sigma_i (10^4 \text{ cm/sec})$	0.54	0.35	
$\sigma_{\rm cm}$ (10 ⁴ cm/sec)	0.81	0.72	
γ	0.892		
α	$6.03 \times 10^4 \text{ cm/sec}$		

even at $\theta = 180^{\circ}$. This indicates that our approximation of the true distribution, $P_3(w)$, by a δ function is no longer valid. Nevertheless, it can be seen that the distribution of centroids is still the dominant factor influencing the shape of the observed distribution, thereby making it very difficult to deduce $P_3(w)$ from P(u). Again, the poorer agreement between experiment and Eq. (6) at smaller scattering angles is presumably a consequence of the approximate treatment employed here.

As the preceding considerations disclose, the peak in the observed product velocity distribution relates directly to the actual reaction dynamics $(u_{mp} = w_{mp})$, while the shape of P(u) is predominantly a result of the shape of P(V) rather than $P_3(w)$. Consequently, as an alternative to the construction of a complete contour map and the subsequent derivation of complete product translational energy distributions, $P(E'_T)$, we have simply determined the velocity of maximum product ion intensity in a number of experiments by measuring the product ion translational energy distribution at 0° in the laboratory and then converting this distribution to the corresponding Cartesian velocity spectrum by multiplying the intensity at each energy by the overall Jacobian factor of 1/v. The velocity corresponding to the peak in this velocity spectrum was then used to calculate the most probable value of the translational excergicity, $Q_{mp} = E'_{T,mp}$ $-E_{T,mp}$. The values so obtained, as well as those derived from that data shown in Figs. 2 and 3, are plotted versus E_T in Fig. 8. For comparison, values of Q_{mp} taken from Ref. 5 for the abstraction process yielding XeH⁺ are also shown in Fig. 8.

As can be seen, Q_{mp} for XeCH^{*}₃ is more negative than that for XeH^{*} at the same collision energy. Since, by conservation of energy, the internal energy of the products is given by

$$E'_{I} = E_{I} + (-\Delta E_{0}) - Q , \qquad (7)$$

we see that displacement products are more excited internally than the abstraction products formed at the same collision energy, and that the internal excitation of the former increases more rapidly with increasing E_T than that of the latter. This is just what one would expect if, as suggested earlier, abstraction occurs predominantly in glancing collisions while displacement requires a more nearly head-on collision. The larger fraction of E_T directed along the line of centers in the latter situation could be expected to excite more effectively the internal modes of the products.

One of several types of collision models that could

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FIG. 8. Most probable translational excergicity, Q_{mp} , vs relative collision energy E_T for the reactions Xe⁺+CH₄-XeH⁺+CH₃ (open circles) and Xe⁺+CH₄-XeCH₃⁺+H (solid circles). The dashed line represents the prediction of the ideal knockout model as given by Eq. (9) for the reaction yielding XeCH₃⁺.

produce the back-scattering we see here for $XeCH_3^*$ is the billard-ball or knockout (IK) process.²⁰⁻²³ In this model the Xe^{*} projectile ion would collide impulsively with one hydrogen atom, ejecting it from the CH₄ molecule. The Xe^{*} ion, its velocity diminished, would then pick up the remaining CH₃ group to form XeCH₃^{*}. Such a process will presumably occur with greatest probability when the axis of the C-H bond is perpendicular to the velocity vector of the projectile ion and the impact parameter is approximately equal to one-half the C-H bond distance.

For the general reaction $X^* + YZ - XY^* + Z$, the kinematics of the IK model predicts that, for 180° scattering, the translational excergicity should be the negative of the translational energy of the reactant ion X^* relative to the abstracted group Y after the ion has collided elastically with the ejected species Z:

$$Q_{\rm IK} = - \frac{(Y)(X-Z)^2(X+YZ)}{(X+Y)(X+Z)^2(YZ)} E_T , \qquad (8)$$

where (Y) represents the mass of Y, etc. For Reaction (3), with the masses of X, Y, and Z equal to 132, 15, and 1 amu, respectively, the IK model predicts $Q_{IK} = -0.916E_T$. This prediction is shown by the dashed line in Fig. 8.

We find that the IK model overestimates product internal excitation (i.e., Q_{mp} is less negative than Q_{IK}) at relative collision energies less than 4 eV. Previous applications of the IK model to the T-CH₄, ^{2b} O^{*}-H₂, ²³ and N^{*}₂-H₂ (Ref. 24) systems have also been unsatisfactory, and reasons for such failure have been discussed. ²³ At collision energies above 4 eV, however, the IK model much more nearly predicts the observed Q_{mp} . The improved agreement at high energy would seem to be in accord with one's chemical intuition. At high collision energies, the reaction dynamics should be dominated more by the steep repulsive part of the system's potential energy surface and be less affected by the attractive part.

IV. SUMMARY AND CONCLUSIONS

(1) In the Xe⁺-CH₄ system, the displacement reaction producing XeCH₃⁺ is only a minor reaction channel. The reaction cross section has an apparent translational energy threshold of 0.7±0.2 eV, passes through a maximum of about 0.25 Å² in the region $E_T = 3-4$ eV, and then decreases with increasing collision energy.

(2) The XeCH^{*}₃ velocity vector distributions at E_T = 1.91 and 4.29 eV are asymmetric about the c.m., indicating that reaction proceeds by a direct mechanism over this energy range, as was found⁵ for the abstraction product XeH^{*} at relative collision energies above 0.5 eV.

(3) The XeCH^{*}₃ product is scattered through large angles with high internal excitation. In contrast, the XeH^{*} product is scattered through small angles with relatively low product excitation. This implies that the opacity functions (reaction probability as a function of impact parameter) are significantly different for the two reactions, with abstraction being favored by moderately large impact parameters and displacement being most likely in small impact parameter collisions. This interpretation is consistent with the need to conserve angular momentum and the difficulty of doing so when the reduced mass of the products is much less than that of the reactants.

ACKNOWLEDGMENTS

Acknowledgment is made to the University of Kansas General Research Fund for support of this research. We wish to thank Professor R. Sampson and the Kansas Geological Survey for providing the computer program used to construct the intensity contour diagrams.

APPENDIX

If v_1 and v_2 are the laboratory velocities of the ionic and neutral reactants, respectively, the velocity V of the center of mass is given by

$$V = \gamma v_1 + (1 - \gamma) v_2 , \qquad (A1)$$

where $\gamma = m_1/(m_1 + m_2)$, m_1 and m_2 being the masses of the ionic and neutral reactants, respectively.

If $P_1(v_1)$ and $P_2(v_2)$ represent the distributions in ionic and neutral reactant velocities, the distribution in V is found by convoluting $P_1(v_1)$ with $P_2(v_2)$, subject to the condition that $v_2 = (V - \gamma v_1)/(1 - \gamma)$:

$$P(V) = \int_{-\infty}^{\infty} P_1(v_1) \cdot P_2\left(\frac{V - \gamma v_1}{1 - \gamma}\right) dv_1 .$$
 (A2)

In the approximate one-dimensional treatment employed here, the distribution of target velocities along the 0° direction is given by the one-dimensional Maxwellian distribution function,

$$P_2(v_2) \propto \exp\left[-\left(\frac{v_2}{\alpha}\right)^2\right]$$
, (A3)

where $\alpha = (2kT/m_2)^{1/2}$, k being the Boltzmann constant and T the target gas temperature. If the distribution of reactant ion velocities in the 0° direction is assumed to be a Gaussian distribution centered around the most probable ionic velocity, v_1^0 , we have

$$P_1(v_1) \propto \exp\left[-\left(\frac{v_1 - v_1^0}{\sigma_1}\right)^2\right],$$
 (A4)

where σ_1 is related to the measured full width of the ionic velocity distribution at half its maximum (FWHM) by $\sigma_1 = FWHM/2\sqrt{\ln 2}$.

Substitution of Eqs. (A3) and (A4) into (A2), followed by integration of the resulting formula, produces the result

$$P(V) \propto \exp\left[-\left(\frac{V-V_0}{\sigma_{\rm em}}\right)^2\right] , \qquad (A5)$$

where V_0 , the nominal velocity of the c.m., is given by γv_1^0 , and

$$\sigma_{\rm cm}^2 = \gamma^2 \, \sigma_1^2 + (1 - \gamma)^2 \, \alpha^2 \, . \tag{A6}$$

If a reactant pair whose c.m. has velocity V produces an ionic product whose true recoil velocity in the c.m. system is w, this product will be detected at a laboratory velocity $v_3 = V + w$. If, in interpreting the data, we assume that the velocity of the c.m. is simply V_0 for all reactant pairs, we would attribute a c.m. recoil velocity $u = v_3 - V_0$ to the ionic product detected at laboratory velocity v_3 . Thus, the assumed recoil velocity u is related to the true recoil velocity w by

$$u = w + (V - V_0)$$
 (A7)

Since any combination of w and V satisfying Eq. (A7) for a given value of u would contribute to the measured product intensity at u, the observed product c.m. velocity distribution P(u) is the result of convoluting the true (but unknown) product velocity distribution $P_3(w)$ with the distribution of centroids, P(V), subject to the condition expressed in Eq. (A7):

$$P(u) = \int_{-\infty}^{\infty} P_3(w) \cdot P(V = V_0 + u - w) \, dw \, . \tag{A8}$$

It is hoped that P(V) is much narrower than $P_3(w)$. In the limit that P(V) is a δ function located at $V = V_0$, we have the happy result that the observed and the true distributions are identical, $P(w) = P_3(w)$.

Unfortunately, the unfavorable kinematic factor in the present situation probably causes $P_3(w)$ to be much narrower than P(V). The limit in this case would be a situation where all the ionic products have the same c.m. recoil velocity w_0 . Then

$$P_{3}(w) = \delta(w_{0}) = 1$$
 if $w = w_{0}$
0 if $w \neq w_{0}$. (A9)

Substitution of this assumed product velocity distribution into Eq. (A8) yields

$$P(u) = P(V = V_0 + u - w_0)$$

$$\propto \exp\left[-\left(\frac{u - w_0}{\sigma_{\rm em}}\right)^2\right]$$
(A10)

That is, the observed c.m. velocity distribution for the ionic product would be a Gaussian distribution centered around the true c.m. recoil velocity w_0 and having a width equal to that of the distribution of centroids.

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