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Association reactions at low pressure. V. The CH₃⁺/HCN system. A final word?

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The reaction of the methyl cation with hydrogen cyanide is revisited. We have confidence that we have resolved a long standing apparent contradiction of experimental results. A literature history is presented along with one new experiment and a re-examination of an old experiment. In this present work it is shown that all of the previous studies had made consistent observations. Yet, each of the previous studies failed to observe all of the information present. The methyl cation does react with HCN by radiative association, a fact which had been in doubt. The product ions formed in the two-body and three-body processes react differently with HCN. The collisionally stabilized association product formed by a three-body mechanism, does not react with HCN and is readily detected in the experiments. The radiatively stabilized association product, formed by a slow two-body reaction, is not detected because it reacts with HCN by a fast proton transfer reaction forming the protonated HCN ion. Previous studies either "lost" this product in the extremely large protonated HCN signal that is always present when HCN is used, or discounted it for various reasons. We have been able to show by ion cyclotron resonance (ICR) techniques (both FT-ICR and tandem ICR-dempster-ICR) that the radiative association product does react with the HCN to form the protonated HCN ion. © 1995 American Institute of Physics.

I. INTRODUCTION

The reaction between CH_3^+ and HCN was reported in 1979 (Ref. 1) to have a radiative association mechanism. In that original work two pieces of information were used to make the deduction. It was observed that the number of CH_3^+ ions at 15 Daltons, decreased with time by a second order process with a rate that was proportional to the product of the two concentrations, $[CH_3^+]$ and [HCN]. A bimolecular reaction rate coefficient based on the removal of CH_3^+ of, $k_2 = 2 \times 10^{-10}$ cm³ s⁻¹, was measured at $\sim 10^{-7}$ Torr and at ~ 100 ms. The other piece of information was that at $\sim 10^{-5}$ Torr and after a drift time of ~ 1 ms, the product ion, $C_2H_4N^+$, was identified using double resonance techniques. It was therefore deduced that the reaction removing CH_3^+ could be written as

$$CH_3^+ + HCN \rightarrow CH_3NCH^+ + h\nu. \tag{1}$$

A year later after continued study it was determined that the same reactants also had a competing three-body stabilization mechanism.² It was determined that a second mechanism exhibited third order kinetics with a measured bimolecular reaction rate coefficient that was dependent on the pressure. It was therefore deduced that a second association reaction could be written as

$$CH_3^+ + HCN + M \rightarrow CH_3NCH^+ + M.$$
 (2)

It was assumed in the second study that the double resonance identification of the association product under the three-body conditions reaction (2) also applied to the bimolecular radiative association reaction [reaction (1)].

The reaction rate coefficient for reaction (2) when He was the third-body, M, was reported as $k_3 = 5 \times 10^{-25}$ cm⁶ s⁻¹. This observation is consistent with earlier higher pressure selected ion flow tube (SIFT) observations, 3,4 in which the association reaction was noted to proceed with a two-body reaction rate coefficient of 2×10^{-9} cm³ s⁻¹, at a helium pressure of 0.5 Torr. When M=HCN, the reaction rate coefficient increased to $k_3 = 1.1 \times 10^{-23}$ cm⁶ s^{-1.5} The results for collisional stabilization of (CH₃NCH⁺)* by He were also examined in a variable temperature SIFT-drift study by Smith and Adams⁶ with the similar results to the earlier SIFT study. Elevating the temperature to 580 K resulted in the three-body rate being reduced to $k_3 = 3 \times 10^{-26}$ cm⁶ s⁻¹, when He was the third-body. A similar decrease in the three-body rate was accomplished by increasing the kinetic energy of the methyl ion.⁶

A very different result was presented by Kemper, Bass, and Bowers⁷ in 1985. They followed the three-body stabilization reaction from 1×10^{-4} Torr to 1×10^{-5} Torr and found only an upper limit of $k_2 = 5\times10^{-12}$ cm³ s⁻¹ for the radiative association channel. The reaction rate coefficient they measured for the three-body reaction with He was $k_3 = 2.2 \times 10^{-25}$ cm⁶ s⁻¹. This set of experiments was carried out in a tandem ICR-dempster-ICR spectrometer where the ion source was separated from the reaction region and only the reaction region contained HCN. The authors concluded that⁷ "The $CH_3^+ + HCN \rightarrow [CH_3^+ - HCN]$ association reaction has been re-examined by using both tandem ion cyclotron resonance (ICR) and drift ICR spectrometers. It appears that the fast radiative stabilization channel reported previously was due to interfering bimolecular reactions and that radiative stabilization does not occur to a significant extent in this system; i.e., any low-pressure bimolecular rate

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coefficient must be less than $\sim 5 \times 10^{-12}$ cm³ s⁻¹."

All the three-body reaction studies at low pressures ($<10^{-3}$ Torr) used both the parent ion abundance, CH_3^+ , and the product ion abundance, $C_2H_4N^+$, to determine the reaction rate coefficient. This differs from the radiative association rate constant measurements measured in the trapped-mode ICR experiments, where only the CH_3^+ ion was followed. In point of fact at no time has the presence of a product ion for the "radiative association" reaction been accounted for in the ICR trapped mode experiments.

A MIKES-CID study⁸ was able to show that at high pressure (10^{-4} Torr) the low energy collisionally stabilized products had the structure of CH_3CNH^+ . At lower pressures ($<10^{-5}$ Torr) the adduct had a structure like that of CH_3NCH^+ .

Gilbert and McEwan⁹ and Smith *et al.*¹⁰ using RRKM methods, modeled the CH₃⁺/HCN system. The model was found to be very sensitive to the structure of the collision complex and the transition state back to reactants. They concluded that the CH₃NCH⁺ is the structure of the association product that best fits the kinetic data of both the ICR and SIFT, even though the CH₃CNH⁺ structure has the lower heat of formation.

Results of four different experiments will be presented and discussed. Two will be the results of the low-pressure trapped-mode ICR and FT-ICR experiments between 10^{-7} and 10^{-6} Torr. The other two results will be higher pressure results from drift-mode ICR and tandem ICR-dempster-ICR experiments between 10^{-5} and 10^{-3} Torr. An interpretation will be presented and applied to the observations from the four different experiments.

The new experiment is the FT-ICR. A reexamination of the tandem ICR-dempster-ICR experiment will also be presented.

II. EXPERIMENT

The trapped-mode ICR and drift-mode ICR experiments were not performed as part of this work. Descriptions of these experiments can be found in the references cited.

A. FT-ICR

The FT-ICR experiments were performed using an OMEGA 50 IonSpec (Ref. 11) Fourier transform mass spectrometer. Briefly, the instrument utilized computer controlled digital FT-ICR technology. Special features of this instrumentation included sequential multiple double resonance ejection, pulsed gas inlet system, and a 10 in. Walker Scientific electromagnet. The magnetic field was typically 1.1 T. The cell was a single 5.0 cm cube. Electron impact ionization was used to initiate ionization. The cell was pumped by a Balzer 330 ℓ s⁻¹ turbomolecular pump.

B. Tandem ICR-dempster-ICR

This instrument was built at UCSB and has been described previously. 12 It has been relocated to the JPL (see Acknowledgments).

The design is an adaptation of that used by Smith and Futrell. ¹³ Instead of the standard dempster source an ICR cell

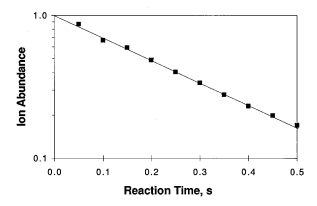


FIG. 1. Semilog plot of the methyl ion decay with time. HCN present at 6.0×10^{-7} Torr. From trapped-mode ICR experiments.

was used as the ion source. Ions generated in this cell are accelerated to typically 3000 V and bent through 180°. A second ICR cell is located 4.74 cm from the first, the 15 Dalton ions are guided into this second ICR by adjusting the magnetic field. The ions are decelerated and introduced into the second ICR through a 0.5 mm thick Wein filter which is 3.8 mm long. This design only allows ions with less than 0.3 eV of translational energy to enter the detection cell. A Wronka bridge detection circuit¹⁴ was used to measure the ion abundances within the detection cell.

III. RESULTS AND DISCUSSION

A. Trapped-mode ICR

Typical results of experiments reported earlier from our laboratory^{1,2} are shown in Fig. 1. The experiment consisted of recording CH_3^+ ion densities at different trapping times for known pressures of HCN in the ICR cell. The reaction rate coefficients for the reaction of CH_3^+ with HCN were found from the slope of the semilog plot of CH_3^+ abundance against the trapping time. Analysis of many such decays resulted in the reaction rate coefficient of $2\times10^{-10}\pm10\%$ cm³ s⁻¹. A typical mass spectrum of the ions in the ICR cell is shows ions present at masses 15, 16, 17, 26, 27, 28, 29, 38, 39, 40, 41, 42, and 43 Daltons. These correspond to the ions, CH_3^+ , CH_4^+ , CH_5^+ , CH_3^+ , CH_5^+ , C

$$CH_{2}^{+}+HCN\rightarrow CH_{2}CN^{+}+H$$
 (3)
 $CH_{2}^{+}+CH_{4}\rightarrow C_{2}H_{2}^{+}+2H_{2}$
 $\rightarrow C_{2}H_{3}^{+}+H_{2}+H$
 $\rightarrow C_{2}H_{4}^{+}+H_{2}$
 $\rightarrow C_{2}H_{5}^{+}+H$ (4)

$$CH_3^+ + CH_4 \rightarrow C_2H_5^+ + H_2$$
 (5)

$$CH_4^+ + CH_4 \rightarrow CH_5^+ + CH_3$$
 (6)

$$CH_4^+ + HCN \rightarrow HCNH^+ + CH_3$$

 $\rightarrow CH_3CNH^+ + H$ (7)

$$CH_5^+ + HCN \rightarrow HCNH^+ + CH_4$$
 (8)

$$C_2H_2^+ + CH_4 \rightarrow C_3H_4^+ + H_2$$

$$\rightarrow C_3 H_5^+ + H \tag{9}$$

 $C_2H_2^+$ +HCN \rightarrow HCNH $^+$ + C_2H

$$\rightarrow$$
CHCCNH⁺+H (10)

$$C_2H_3^+ + CH_4 \rightarrow C_3H_5^+ + H_2$$
 (11)

$$C_2H_3^+ + HCN \rightarrow HCNH^+ + C_2H_2$$
 (12)

$$C_2H_5^+ + CH_4 \rightarrow C_3H_7^+ + H_2$$
 (13)

$$C_2H_5^+ + HCN \rightarrow HCNH^+ + C_2H_4$$
 (14)

$$CN^{+}+CH_{4}\rightarrow CH_{3}^{+}+HCN$$

$$\rightarrow CH_{4}^{+}+CN$$

$$\rightarrow HCN^{+}+CH_{3}$$

$$\rightarrow HCNH^{+}+CH_{2}$$

$$\rightarrow CH_{2}CN^{+}+H_{2}$$
(15)

$$CN^+ + HCN \rightarrow HCN^+ + CN$$

$$\rightarrow C_2 N_2^+ + H \tag{16}$$

$$HCN^+ + CH_4 \rightarrow HCNH^+ + CH_3$$

$$\rightarrow C_2 H_3^+ + NH_2 \tag{17}$$

$$HCN^{+} + HCN \rightarrow HCNH^{+} + CN.$$
 (18)

Of the ions produced in the ICR cell by electron impact and the reaction sequence (3)–(18), the protonated hydrogen cyanide ion, HCNH⁺ is the most abundant. Using the double resonance technique all of reactions (3)–(18) could be verified, but the product ion of reaction (1) could not be confirmed. On several occasions double resonance experiments indicated a small fraction of the HCNH⁺ ion was derived from CH₃⁺, but this was not reproducible. A problem inherent in establishing a double resonance link between CH₃⁺ and HCNH⁺ is that the 28 Daltons signal is very large due to primary ionization of HCN and subsequent proton transfer to HCN.

B. FT-ICR

These results are new. We were able to observe the reaction of CH₃⁺ with HCN using FT-ICR technology on an IonSpec FT-ICR mass spectrometer. ¹¹ The methyl ion was generated by electron impact on methane in a one cell instrument with HCN present during the whole experiment. A pulsed valve was used to introduce the methane into the spectrometer. The valve was open for 2 ms. After a delay of 80 ms, a 5 ms pulse of electrons was used to ionize the gases. In a resultant mass spectrum ions were identified at 15, 16, 17, 27, 28, 29, and 42 Daltons. Minor peaks at 18 and 19

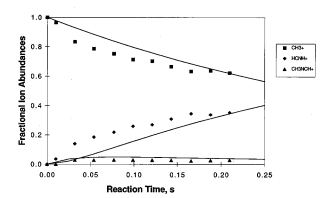


FIG. 2. The reaction of the methyl ion with hydrogen cyanide. Using the IonSpec FT-ICR. The data points are experimental results. The solid lines are from model calculations.

Daltons were also present. The mass spectrum at this stage was identical to the trapped-mode ICR experimental results. All ions were then ejected from the cell by sequential double resonance ejection except the CH₃⁺ ion at 15 Daltons. The ejection sequence started 15 ms after the electron pulse and lasted for about 16 ms. The reaction of CH₃⁺ and HCN was then allowed to proceed and the ions in the cell were monitored for the next 210 ms as they reacted. Figure 2 shows the results of one of these experiments as data points. It is noted that the methyl ion concentration decreases exponentially with time, while the protonated HCN species increases. The protonated methyl isocyanide product, representing the collision complex and the radiative association product ion, stays at a steady state level of a few percent. We also found the HCNH⁺ ion signal decreased when the CH₃⁺ ion was ejected using a double resonance rf field as it also did when the 42 Dalton ion was irradiated. This second observation of a decrease in HCNH⁺ upon double resonance ejection of (CH₃NCH⁺)*, indicates some HCNH⁺ ions are derived from CH₃NCH⁺, via reaction (19),

$$CH_3NCH^+ + HCN \rightarrow HCNH^+ + CH_3NC.$$
 (19)

Although reaction (19) is endothermic by 1.48 kJ/mol for reactants in their ground states, the formation process for CH₃NCH⁺ in reaction (1) is so exothermic that the CH₃NCH⁺ ion should have sufficient excess internal energy to drive the reaction. Several other reactions were considered linking the CH₃⁺ ion to the HCNH⁺ ion, but reaction (19) was the least endothermic option.

These observations allow us to present the following mechanism to represent the reaction sequence in the system:

$$CH_3^+ + HCN \leftrightarrow (CH_3NCH^+)**$$
 (20)

$$(CH3NCH+)** \rightarrow (CH3NCH+)* + h\nu$$
 (21)

$$(CH3NCH+)**+M\rightarrow CH3CNH++M$$
 (22)

$$(CH3NCH+)*+HCN\rightarrow HCNH++CH3NC$$
 (23)

$$CH_3CNH^+ + HCN \rightarrow no reaction.$$
 (24)

A model based on reactions (20)–(24) is plotted in Fig. 2 as a set of solid lines. We note that in this model the collision

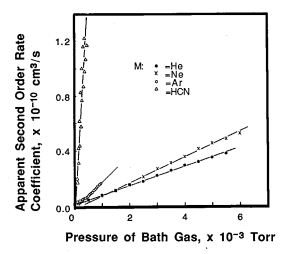


FIG. 3. Plot of the observed 2nd order reaction rate coefficient vs bath gas pressure for the reaction of the methyl ion with hydrogen cyanide. Taken from Ref. 7. Using the drift-mode ICR experiment.

complex (CH₃NCH⁺)** can be stabilized by either radiative association or collision stabilization. The collisionally stabilized CH₃CNH⁺ is not reactive with HCN, while the radiatively stabilized (CH₃NCH⁺)* is. The results of two further experiments were examined to test the proposed mechanism.

C. Drift-mode ICR

The drift-mode ICR results we have called on were extracted from literature sources.^{2,5,7} The experiments consist of observing the parent ion and product ions as a function of the cell pressure. The CH₃⁺/HCN system has been examined in this way for the parent neutral, HCN, as well as other third bodies like He, Ne, and Ar. The analysis performed was to measure the peak heights of both the reactant and the products and using the power absorption equations to determine an effective second order reaction rate coefficient. The effective second order reaction rate coefficients were then plotted against the third body pressure. Figure 3 shows these results and reveals that the effective second order reaction rate coefficients increase linearly with pressure in this range and have an apparent zero intercept. This observation appears at first to indicate that the association reaction is third order and has no measurable second order reaction rate coefficient. This was in fact the conclusion of Kemper, Bass, and Bowers⁷ on viewing their results. The drift-mode operation of the ICR results in the reaction sequence (3)–(18) competing simultaneously with reactions (20)–(24). A mass spectrum taken in the drift-mode reviels the multiple ion problem observed in the trapped-mode.

To confirm their predictions, Kemper, Bass, and Bowers⁷ used the tandem ICR-dempster-ICR instrument which avoids complications of multiple many ions and neutrals that occur in the reaction region of a single cell instruments.

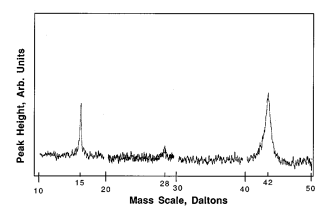


FIG. 4. Mass spectrum of a typical ion concentration in the tandem ICR-dempster-ICR. Reaction time \sim 1 ms and HCN pressure \sim 2×10⁻⁴ Torr.

D. Tandem ICR-dempster-ICR

The results we present from this instrument are from one previous study⁷ plus some new results using the same instrument as in the earlier study. The literature results from the tandem instrument were entirely consistent with the driftmode ICR results, in that collisional stabilization of the association complex is found to be very efficient. These earlier results which are characteristic of both the drift-mode ICR and tandem instruments, are shown in Fig. 3 and present the variation in effective second order reaction rate coefficient with the third body pressure. There is an important distinction between the drift-mode ICR and the tandem instruments. In the tandem, the ion source is completely separate from the reaction region. Methyl ions are generated in the source ICR cell from either methane or methyl bromide. The dempster section transfers the methyl ions from the source cell into the ICR reaction-detection cell. HCN at a known pressure is added into the reaction-detection ICR cell and the reactant and product ions are all monitored.

With this configuration it becomes possible to detect very low densities of the product ions that have more than one source of production. The model that we proposed in reactions (20)–(24) for the association of CH₃⁺ and HCN, as well as the results of the experiments with the FT-ICR instrument, suggest that HCNH⁺ is the product of the proton transfer reaction between the radiatively stabilized collision complex (CH₃NCH⁺)* and HCN. As we have noted earlier, other sources of HCNH⁺ in single cell instruments obscure this reaction product. Our new experiments with the tandem did show low concentrations of HCNH⁺ as predicted by the model. A mass spectrum of the ions seen in the tandem experiment is shown in Fig. 4. The abundances of ions at 15, 28, and 42 Daltons were recorded at different HCN pressures and their measured abundances are compared with calculations based on the model presented in reactions (20)–(24) in Fig. 5. The instrumental signal at 42 Daltons includes both the collision stabilized adduct and the radiative stabilized adduct. The points are experimental and the lines are the model calculations. Finally we have shown in Fig. 6, the relative amounts of CH₃NCH⁺ predicted by the model arising from association from complexes stabilized by radiation

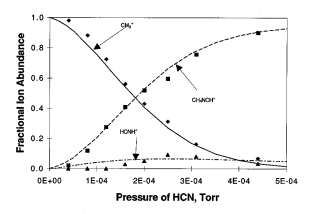


FIG. 5. Data from the tandem ICR-dempster-ICR for the methyl ion reaction with hydrogen cyanide. Data points are notes. Lines are from model calculations

compared to those stabilized by collision with a third body.

We have noted that the new tandem experiments confirm the predictions of the model, in that a small steady state concentration of HCNH⁺ was observed in the 10⁻⁴ Torr range of HCN. The earlier work on the tandem instrument on this system makes no mention of a product at 28 Daltons.⁷ Private communication with the authors on this work revealed no evidence that this peak was observed or even looked for. The very low concentrations of the HCNH⁺ ion would however have made it very easy to have been overlooked. The observation of a very small but nevertheless significant density of HCNH⁺ is vital for a complete understanding of the stabilization mechanism of the association complex in the CH₃⁺/HCN system.

IV. CONCLUSION

We have amalgamated measurements from four different techniques in order to understand the association mechanism between CH₃⁺ and HCN. When used in isolation, the conclusions based on evidence from a single technique can be interpreted quite differently than conclusions based on the results of the four techniques taken together. It is the inclusion and extrapolation of results from isolated experimental methods that have led to conflicting statements about the association mechanism in the CH₃⁺/HCN system. The situation is

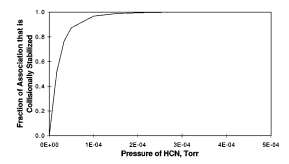


FIG. 6. Showing the fraction of the association product formed from collisional stabilization.

very similar to the Indian fable¹⁶ in which an elephant is examined by six blind men. Each individual touches a different part of an elephant and each reaches a different conclusion as to the nature of the beast.

The amalgamation in this work of the results from all four techniques shows that the simplest mechanism that can explain all the observations is the one given in reactions (20)–(24). In brief, CH₃⁺ does react with HCN by a radiative association channel, with a reaction rate coefficient of $k=2\times10^{-10}$ cm³ s⁻¹. The association complex also undergoes very efficient collisional stabilization. At the lower pressure ICR experiments (e.g., Trapping-mode), reaction times are 100 times longer than the higher pressure ICR experiments (e.g., drift-mode). Quite different outcomes of the collision complex can eventuate in the different pressure regimes making it difficult to extrapolate the results from one pressure regime to the other. The drift-mode and the tandem results which gave effective zero intercepts on their k_2 vs pressure plots (Fig. 3) cannot be interpreted as evidence that radiative association is unimportant in the CH₃⁺/HCN system. Rather, it simply reflects the fact that the time between collisions is shorter than the collision complex lifetimes to radiative stabilization. At all pressures higher than 8×10^{-5} Torr (M=HCN), the collision stabilization process will be the major complex stabilization mechanism. This results from the fact that the radiative relaxation step is too slow to be competitive with the more probable stabilizing collision.

Finally, we note that the differences in reactivity between radiative stabilization complex (CH₃NCH⁺)* and the collisionally stabilized complex (CH₃CNH⁺) are consistent with earlier structural analysis of the products. As noted in the Introduction the high pressure MIKE-CID results⁸ indicate the collision stabilized product has the CH₃CNH⁺ structure. On the other hand Smith *et al.*¹⁰ found the CH₃CNH⁺ structure to be inconsistent with the transition state requirements. It seems reasonable to assume therefore, that the initial stabilized structure is the CH₃NCH⁺ ion by both stabilization channels which is reactive in the initial energy state towards HCN. Collisions with a third body rapidly isomerizes this to the more stable CH₃CNH⁺ ion which is unreactive towards HCN.

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