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Evidence for the Production of N_4 via the N_2 $A^3\Sigma_u^+$ + N_2 $A^3\Sigma_u^+$ Energy Pooling Reaction

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We report mass spectrometric evidence supporting our proposed mechanistic pathway for the production of N_4 through the energy pooling reaction N_2 $A^3\Sigma_u^+ + N_2$ $A^3\Sigma_u^+$. N_2 $A^3\Sigma_u^+$ is generated from the quenching of resonantly excited xenon in a mixture of xenon, $^{15}N_2$, and $^{14}N_2$ that is illuminated with xenon resonant lamps (147 nm). Mass spectra are periodically taken of the mixture. Over time, we observe significant isotopic scrambling of the $^{15}N_2$ and $^{14}N_2$, generating $^{15}N^{14}N$ in concentrations approaching 10% (\sim 2 Torr) of the initial $^{15}N_2$ concentration. Though we do not observe the direct formation of N_4 , the isotopic ratios indicate that an excited complex ($^{15}N_2^{14}N_2$) exists long enough so that scrambling of the nitrogen atoms can occur, offering a possible route to the formation of tetrahedral nitrogen (17 _d N_4).

Introduction

Tetrahedral nitrogen (tetraazatetrahedrane, N_4 1T_d), was first predicted as a stable isomer of nitrogen in the early 1970s. Since that time, a number of theoretical calculations have been performed at increasing levels of sophistication that have supported these first calculations. Interest in the experimental confirmation of these calculations increased over the years, culminating with the funding of programs to pursue the synthesis of N_4 as well as other allotropes of nitrogen. The interest lies in the fact that all of the predicted allotropes of nitrogen are energetic and would form the basis for a new set of environmentally friendly, or "green", energetic materials. At that time, we proposed a concept of photosensitization of N_2 for the production of N_4 , analogous to mercury photosensitization of hydrocarbons. Early on, it became clear that a simple scheme, such as

$$N_2 X^1 \Sigma_g^+ + M^* (M = Kr, Xe) \rightarrow N_2^*$$
(state not specified)
 $N_2^* + N_2 X^1 \Sigma_g^+ + M \rightarrow {}^1 T_d N_4 + M$ (1)

is not, practically speaking, feasible, given the known energetics and orbital diagrams of these species. However, in the 1995 paper by Dunn and Morokuma,⁴ it was shown that the longrange N_2 fragments correlated (at large molecular separation) with *both* fragments electronically excited, each having an electronic configuration of $\pi^3\pi^{*1}$. A review of the comprehensive spectroscopic data set of N_2 revealed that the lowest excited state to contain the required symmetry is the $A^3\Sigma_u^+$ state of N_2 ,⁵ which, coincidentally, has a 1.9 s fluorescence lifetime, with this exceptionally long lifetime critical to our proposed mechanism for the formation of N_4 (M is a third body):

$$N_2 A^3 \Sigma_u^+ + N_2 A^3 \Sigma_u^+ + M \rightarrow N_4^1 T_d + M$$
 (2)

Reaction 2 is a possible singlet-coupled channel of the well-known energy pooling reaction of $N_2~A^3\Sigma_u^{+}+N_2~A^3\Sigma_u^{+},$ first observed by Stedman and Setser in 1969^6 and later by Hays and Oksam in $1973.^7$ The detailed study of this reaction has provided rate constants and hence "branching" ratios for the energy pooling of the A-state as follows: $^{8-10}$

$$N_2 A^3 \Sigma_u^{\ +} + N_2 A^3 \Sigma_u^{\ +} \rightarrow N_2 B^3 \Pi_g$$

$$k \sim 1 \times 10^{-10} \text{ cm}^3/\text{molecule/s (3)}$$

$$N_2 A^3 \Sigma_u^{\ +} + N_2 A^3 \Sigma_u^{\ +} \rightarrow N_2 C^3 \Pi_u$$

 $k \sim 1.5 \times 10^{-10} \text{ cm}^3/\text{molecule/s} \ (4)$

$$N_2 A^3 \Sigma_u^{\ +} + N_2 A^3 \Sigma_u^{\ +} \rightarrow N_2 C'^5 \Pi$$

 $k \sim 2.5 \times 10^{-11} \text{ cm}^3/\text{molecule/s (5)}$

$$N_2 A^3 \Sigma_u^+ + N_2 A^3 \Sigma_u^+ \Rightarrow N_2 C''^5 \Pi$$

 $k \sim 9 \times 10^{-11} \text{ cm}^3/\text{molecule/s (6)}$

We note that reaction of two molecules, each triplet coupled, will lead to potential surfaces that are singlet, triplet, and quintuplet coupled. As identified in reactions 3-6, the triplet and quintuplet branches are open and available for the energy pooling reactions. However, the singlet channel has never been observed. The absence of the observation of a singlet channel may result from (a) no reaction, (b) a dark channel leading to the production of ground state nitrogen molecules, or (c) a dark channel producing highly excited N_4 (excess energy of $\sim 20~000~cm^{-1}$; $\sim 36~500~cm^{-1}$ above the ground rovibrational state of N_4) that would dissociate in the absence of quenching collisions. Thus, the singlet coupled surface may provide a pathway for generating singlet N_4 (highly vibrationally excited: either tetrahedral⁴ or rectangular¹¹) that eventually could be quenched to ground-state N_4 .

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To attempt a practical realization of our scheme, we used known quenching pathways, which allowed us to develop the following mechanism and energy flow system:¹²

$$Xe + hv (147 nm) \rightarrow Xe [3/2]_1$$
 (7)

$$Xe [3/2]_1 + N_2 X^1 \Sigma_g^+ \rightarrow Xe + N_2 B^3 \Pi_g$$
 (8)

$$N_2 B^3 \Pi_g \rightarrow N_2 A^3 \Sigma_u^+ + h\nu$$
 (9)

$$^{15}N_{2} A^{3}\Sigma_{u}^{+} + {}^{14}N_{2} A^{3}\Sigma_{u}^{+} \rightarrow {}^{15}N_{2}^{14}N_{2} \rightarrow {}^{15}N_{2} + \\ ^{14}N_{2} \text{ or } 2^{15}N^{14}N \text{ (10)}$$

where reaction 10 is assumed to occur in the absence of quenching collisions. Thus, if the A-state of nitrogen could be made in high enough concentration (which is feasible given the 1.9 s lifetime), reactions 3-6 and 10 could occur with good efficiency. A more detailed kinetic scheme using all of the known kinetic parameters and fluorescent lifetimes available in the literature, and a rate constant for N₄ creation equivalent to those measured by Piper for the triplet and quintuplet channels, was used to determine an approximate optimal nitrogen pressure for this reaction.¹³ This optimal pressure is less than 50 Torr and results from the fact that the B-state in reaction 8 can be depleted through self-quenching by groundstate nitrogen as identified by Piper.9 Thus, by illuminating a mixture of xenon, ¹⁵N₂, and ¹⁴N₂ with xenon resonance radiation, we expect to see scrambling of the isotopes if the excited-state intermediate of reaction 10 results in either a single bonded nitrogen structure (tetrahedral N₄) or through a double bonded resonance structure (via rectangular N₄)¹¹ and lives long enough for the atoms to lose memory of their initial atomic partner.

Other groups have proposed alternative schemes and have conducted experiments in an attempt to form N₄. Ostmark and co-workers have proposed a mechanism similar to reaction 1: the difference is that the excited N₄ would be generated through excitation of nitrogen using laser radiation, ion bombardment, or in hollow cathode discharge. 14 They have not observed the formation of N₄ to date. Radziszewski and co-workers have conducted a number of experiments in which a microwave discharge of pure nitrogen is generated and the resulting fragments and species are collected on a helium cooled sapphire window. 15 The resulting nitrogen "ice" gave rise to the observation of a weak peak at the expected N₄ IR active frequency of \sim 936 cm⁻¹. However, substitution to the ¹⁵N₂ starting material led to an isotopic shift that was inconsistent with that expected for this molecule. ¹⁶ Finally, Cacace et al. have observed an N₄⁺ cation in a neutralization-reionization experiment that they have attributed to the weakly bound open chain form of N₄. ¹⁷

Experimental Section

A mixture of 2 Torr Xe (Air Products 99.995%), 20 Torr $^{15}\rm{N}_2$ (Cambridge Isotopes, 98%), and 20 Torr "standard" \rm{N}_2 (UHP 99.999%; 99.635% $^{14}\rm{N})$ was prepared in a small vacuum chamber consisting of a double-sided $2^3/_4$ in. knife-edge flange (MDC vacuum) mated to two double-sided knife-edge flange LiF windows (St. Gobain), as shown in Figure 1. The gases were additionally purified to below ppb impurity levels by passage through SAES PS11 series ambient temperature gas purifiers before mixing in the cell. Xenon resonance lamps (10: 1, 147 nm:129.5 nm; COREX of Russia; $\sim\!\!20$ mW average power) were mounted in water-cooled stainless nipples, which were back-filled with UHP argon (620 Torr 99.999%) and bolted

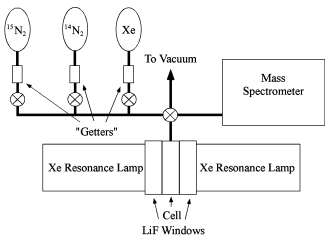


Figure 1. Schematic of the experimental apparatus.

to the LiF window flanges. The VUV radiation passed through LiF windows into the $\sim\!30~\text{cm}^3$ volume cell. Mass spectra were taken periodically over several weeks using a Ominstar GSD 300 (Pfeiffer Vacuum). Calibration and normalization was accomplished by scaling the mass spectral intensities to maintain a constant xenon signal, which should not change over time. Two replica experiments were performed to demonstrate reproducibility.

Results

The Xe signal in the mass spectra is used for an internal standard, and all mass spectra are Xe peak area normalized such that the area under the curve from $126 \ m/z$ to $137 \ m/z$ is equal to 1. Prior to normalization, a simple baseline correction is performed using the best fit line through a highly linear region of the spectra from $50 \ m/z$ to $60 \ m/z$ and from $70 \ m/z$ to $120 \ m/z$ (Xe⁺² peaks are omitted) and this line is subtracted from each spectrum. Without this baseline correction, the peak area of the Xe species would be overestimated by as much as 1.2%. Following the baseline correction and peak area normalization, the peaks (or clusters of peaks) of interest are fit with commercially available software (PeakFit) to separate overlapping peaks, e.g., 28, 29, and $30 \ m/z$.

Baseline-corrected, Xe+-normalized mass spectra are illustrated in Figure 2 for the range 24-34 m/z. This series of spectra shows the time evolution throughout the first experiment. From t = 0 days to t = 24 days, there is a clear increase in the $29 \, m/z \, \text{peak} \, (^{14}\text{N}^{15}\text{N})$. During this time period, the reaction cell is illuminated with the lamps. After the 24th day, the lamps are switched off. While the lamps are switched off, there is no appreciable change to the 29 m/z signal. Given that the 29 m/zpeak increases under VUV irradiation and remains constant under conditions of no illumination suggest that isotope exchange only takes place (to any measurable extent) when the reaction is provided energy via the VUV photons and that, once exchanged, the ¹⁴N¹⁵N molecules are stable in the absence of VUV, as to be expected. These two observations provide strong evidence that our proposed reaction mechanism may be correct and could form the basis for N₄ synthesis. The increase in 29 m/z peak and simultaneous decrease in 30 m/z peak seen in our mass spectra indicate that an excited complex of N₄ with the proper bonding and sufficient lifetime to allow for this exchange to occur. Shown in Figure 3 are the time-dependent mass spectra for the second experimental run, demonstrating both reproducibility and an increase in the 29 m/z peak to \sim 10% of the 30 m/z peak.

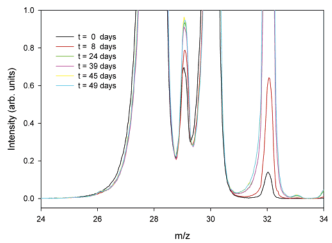


Figure 2. Mass spectra from 24 m/z to 34 m/z over the course of 49 days. The VUV lamps are illuminating the cell for the first 24 days only. During this period, there is a significant increase of the 29 m/zpeak. After the lamps are switched off, the change of the 29 m/z peak is insignificant.

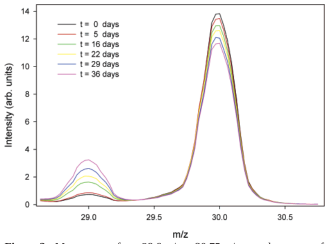


Figure 3. Mass spectra from 28.6 m/z to 30.75 m/z over the course of 36 days. The VUV lamps are illuminating the cell for the entire time. During this period, there is a significant increase of the 29 m/z peak, with a corresponding decrease in the 30 m/z peak.

Because the peak at 29 m/z is expected to be a reaction product of atom exchange between $^{14}N_2$ (28 m/z) and $^{15}N_2$ (30 m/z), tracking the temporal evolution of these and other species provide clues as to what is taking place in the reaction cell with the lamps on and off. The relative peak areas for peaks at 18, 28, 29, 30, 32, and 44 m/z are plotted in Figure 4. During the portion of the experiment when the reaction cell is irradiated with VUV, there is a dramatic increase in the 29 m/z peak and a concomitant decrease in the 28 m/z and 30 m/z peaks. The decrease in these two peaks is not unexpected, as the gain of ¹⁴N¹⁵N must come with the loss of both ¹⁴N₂ and ¹⁵N₂. During the period of no illumination, the peak at $30 \, m/z$, much like the peak at 29 m/z, remains relatively constant. There is, however, an increasing trend in the 28 m/z peak. There are several possible explanations. First, ¹²C¹⁶O and ¹⁴N₂ are isobaric interferences and any ¹²C¹⁶O in the system will appear as ¹⁴N₂. The CO may arise as a decomposition product of CO₂, due to reactions taking place either in the reaction cell or during the ionization and detection event in the mass spectrometer. Additionally, the reaction cell is not actively pumped once it is filled with the reactant gases to a total pressure far less than atmospheric pressure. Therefore, it is possible that N2, CO, and CO2 are leaking into the cell.

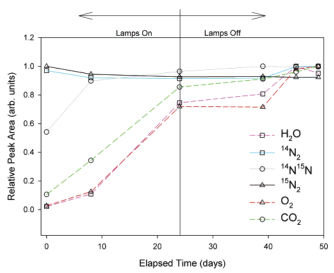


Figure 4. Relative mass spectra peak areas as a function of time. Several peaks in the mass spectra are tracked as a function of time to understand what is taking place in the reaction cell. Peaks for ¹⁴N₂ (28 m/z) [solid line, squares], for $^{14}N^{15}N$ (29 m/z) [solid line, circles], and for $^{15}N_2$ (30 m/z) [solid line, triangles] are directly related to the isotope exchange reaction. Peaks for H_2O (18 m/z) [dashed line, squares], for O_2 (32 m/z) [dashed line, triangles], and for CO_2 (44 m/z) [dashed line, circles] are indicators of an atmospheric leak into the cell or are indicators of unintentional reactions.

Evidence for atmospheric contaminants leaking into the cell is provided by tracking the evolution of the peaks at 18 m/z (H_2O) , at 32 m/z (O_2) , and at 44 m/z (CO_2) . All these tend to increase as time is elapsed. The rate of increase is most dramatic while the lamps are illuminating the reaction cell but continue to increase after the lamps are switched off. Two likely explanations are (1) the heat from the lamps stresses the flanges, making a leak more probable and (2) the VUV is stimulating the desorption of $C_xO_yH_z$ species directly (or indirectly with H₂O, CO, and CO₂ as major decomposition products) from the stainless steel vacuum walls.

Despite the certain interference at 28 m/z, the peaks at 29 m/z and at 30 m/z are not appreciably affected by the leak and or stimulated desorption, because of the extremely low isotopic abundances of ¹³C (1.07%), of ¹⁷O (0.038%), and of ¹⁸O (0.205%). The contributions of $^{13}C^{16}O$ and of $^{12}C^{17}O$ to 29 m/zare negligible. Likewise, the contribution of ${}^{13}C^{17}O$ to 30 m/zis insignificant. Therefore, the 29 m/z and 30 m/z peaks are better indicators of the reaction progress due to the smaller contributions from interferences. In fact, the 30 m/z peak behaves as expected on the basis of the behavior of the 29 m/z peak. The 30 m/z peak decreases while the lamps are on with the concomitant increase in the 29 m/z peak, and while the lamps are off, the 30 m/z peak remains largely unchanged, just as the 29 m/z peak.

Conclusion

Here, we present results of our ongoing efforts to synthesize tetrahedral nitrogen, N₄. Mass spectrometric observations show that resonant energy transfer from Xe* to N2 has allowed for isotopic exchange to occur between ¹⁴N₂ and ¹⁵N₂, creating the mixed-isotopic species ¹⁴N¹⁵N. Although N₄ has yet to be observed directly, this is a promising development due to the inherent stability in the nitrogen-nitrogen triple bond. The increase in 29 m/z peak and simultaneous decrease in 30 m/zpeak seen in our mass spectra indicate the existence of an excited complex of N₄ with the proper bonding and sufficient lifetime to allow for this exchange to occur. We note that an alternate possibility exists, which is that the energy pooling reaction leads to dissociation of either the $^{14}\mathrm{N}_2$ and $^{15}\mathrm{N}_2$, yielding atomic nitrogen that subsequently recombines forming $^{14}\mathrm{N}^{15}\mathrm{N}$. We believe that this is unlikely, because the excitation energy from both molecules must end up in one molecule to yield dissociation, which seems improbable given the known orbital relationships (see ref 11). Experiments are underway in an effort to quench some portion of the N_4 complex to either the theoretically stable tetrahedral or rectangular species.

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