Ion-molecule rate constants and branching ratios for the reaction of $N_3^+ + O_2$
from 120 to 1400 K

The kinetics of $N_3^+$ reacting with $O_2$ has been studied from 120 to 1400 K using both a selected ion flow tube and high-temperature flowing afterglow. The rate constant decreases from 120 K to ~1200 K, then increases slightly to 1400 K, which compares well to most of the previous measurements in the overlapping temperature range. Comparing the results to drift tube data shows little difference between increasing the translational energy available for reaction and increasing the internal energy of the reactants over much of the range, i.e., all types of energies drive the reactivity equally. The reaction produces both $NOO^+$ and $NO_2^+$; the latter is shown to be the higher energy NOO linear isomer. The ratio of $NOO^+$ to $NO^+$ is over 2 at 120 K, decreasing to less than 0.01 at 1400 K because of dissociation of $NOO^+$ at the higher temperatures. This ratio decreases exponentially with increasing temperature. High-level theoretical calculations have also been performed to complement the data. Calculations using multi-reference configuration interaction theory at the MRCCSD(Q)/cc-pVTZ level of theory show that singlet NOO$^+$ is linear and is 4.5 eV higher in energy than NOO$^+$. A barrier of 0.9 eV prevents dissociation into NO$^+$ and O($^3P$); however, crossing to a triplet surface connects to NO$^+$ and O($^3P$) products. A singlet and a triplet potential energy surface leading to products have been determined using coupled cluster theory at the CCSD(T)/aug-cc-pVTZ level on structures optimized at the Becke3-Lee, Yang, and Parr (B3LYP)/aug-cc-pVTZ levels.
Ion-molecule rate constants and branching ratios for the reaction of N₃⁺ + O₂ from 120 to 1400 K

Svetozar Popovic, Anthony J. Midey, Skip Williams, Abel I. Fernandez, and A. A. Viggiano
Air Force Research Laboratory, Space Vehicles Directorate, Hanscom Air Force Base, Massachusetts 01731-3010
Peng Zhang and K. Morokuma
Cherry L. Emerson Center for Scientific Computation and Department of Chemistry, Emory University, Atlanta, Georgia 30322

(Received 7 July 2004; accepted 25 August 2004)

The kinetics of the reaction of N₃⁺ with O₂ has been studied from 120 to 1400 K using both a selected ion flow tube and high-temperature flowing afterglow. The rate constant decreases from 120 K to ~1200 K and then increases slightly up to the maximum temperature studied, 1400 K. The rate constant compares well to most of the previous measurements in the overlapping temperature range. Comparing the results to drift tube data shows that there is not a large difference between increasing the translational energy available for reaction and increasing the internal energy of the reactants over much of the range, i.e., all types of energies drive the reactivity equally. The reaction produces both NO₄⁺ and NO₂⁺, the latter of which is shown to be the higher energy NOO⁺ linear isomer. The ratio of NOO⁺ to NO₄⁺ decreases from a value of over 2 at 120 K to less than 0.01 at 1400 K because of dissociation of NOO⁺ at the higher temperatures. This ratio decreases exponentially with increasing temperature. High-level theoretical calculations have also been performed to compliment the experiments. Calculations using multi-reference configuration interaction theory at the MRCISD(Q)/aug-cc-pVTZ level of theory show that singlet NOO⁺ is linear and is 4.5 eV higher in energy than ONO⁺. A barrier of 0.9 eV prevents dissociation into NO⁺ and O₂(1D); however, a crossing to a triplet surface connects to NO⁺ and O₂(3P) products. A singlet and a triplet potential energy surface leading to products have been determined using coupled cluster theory at the CCSD(T)/aug-cc-pVQZ level on structures optimized at the Becke3-Lee, Yang, and Parr (B3LYP)/aug-cc-pVTZ level of theory. The experimental results and reaction mechanism are evaluated using these surfaces. © 2004 American Institute of Physics. [DOI: 10.1063/1.1807376]

I. INTRODUCTION

Air plasmas at elevated pressures are currently the subject of intensive study for a variety of applications. Typical nonequilibrium plasmas have heavy particle temperatures below 1000 K and average electron energies on the order of a few eV. These conditions are favorable for associative ionization processes and the presence of polyatomic nitrogen ions such as N₃⁺ and N₄⁺ is expected. However, temperature dependent data even for the relatively simple cases involving reactions of these ions with major species are insufficient or nonexistent.

In light of these issues, the kinetics of the reaction of N₃⁺ with O₂ has been studied in more detail. The reaction has two observed product channels:

\[ \text{N}_3^+ + \text{O}_2 \rightarrow \text{NO}_2^+ + (\text{N}_2\text{O}) + 466 \ \text{kJ mol}^{-1} \]  

\[ \rightarrow \text{NO}_2^+ + \text{N}_2 + 124 \ \text{kJ mol}^{-1}, \]  

where the N₂O product may be dissociated, i.e., N₂ + O, reducing the exothermicity of the reaction to 299 kJ mol⁻¹, or production of N + NO₂, which would make the reaction endothermic by 14 kJ mol⁻¹. The heat of formation of N₃⁺ has been obtained by combining the average of the experimental heats of formation of N₃ from the work of Continetti et al. and Martin et al.² with the ionization potential from the work of Dyke et al.,³ which are in good agreement with the recent theoretical values of Dixon et al.⁴ For the NO₂⁺ thermochemistry, we use the present ab initio calculations for the NOO⁺ isomer (see below). All of the other values are from the NIST Webbook.⁵ Note also that the NIST Webbook recommended value for the heat of formation of N₃ is 52 kJ mol⁻¹ lower than the values used here.

In a flowing afterglow, Dunkin et al.⁶ measured the first experimental value for the total rate constant for reaction (1) at 200 K and obtained a value of \( k = 1 \times 10^{-10} \text{cm}^3 \text{s}^{-1} \). Smith et al.⁷ have determined a rate constant of 5.1 \( \times 10^{-11} \text{cm}^3 \text{s}^{-1} \) for the reaction at 300 K in a selected ion flow tube (SIFT), which is an order of magnitude less than the Langevin collision rate constant of 7.0 \( \times 10^{-10} \text{cm}^3 \text{s}^{-1} \). In addition, Lindinger⁸ has measured the rate constants for the N₃⁺ reaction with O₂ in a flow drift tube (FDT) at 298 K...
over a wide range of center-of-mass kinetic energies at low pressures. A rate constant of $6 \times 10^{-11}$ cm$^3$ s$^{-1}$ has been reported by Lindinger at zero field (0.038 eV), which decreases to a minimum at 0.25 eV kinetic energy and then increases at higher kinetic energy. The rate constant has been observed to be independent of pressure over the limited pressure range 0.139–0.253 Torr studied. Hiraoka$^2$ measured the rate constant for reaction (1) from 64 to 552 K using a high pressure mass spectrometer (HPMS) at ca. 3 Torr and found that the rate constant decreases monotonically with increasing temperature over the entire temperature range. Matsuoka et al.$^{10}$ have also found a total reaction rate constant at 294 K for reaction (1) of $7.1 \times 10^{-11}$ cm$^3$ s$^{-1}$ using a time-resolved atmospheric pressure ionization mass spectrometer (TRAPI).$^{10}$

There is good agreement among the room temperature thermal rate constants reported by Smith et al., Lindinger, and Matsuoka et al.

Product formation has also been studied. Smith et al. have determined that the product ions formed in reaction (1) are 70% NO$^+$ and 30% NO$_2^+$ at 300 K.$^7$ By contrast, the drift tube results of McCrumb and Warneck$^{11}$ and the TRAPI measurements of Matsuoka et al.$^{12}$ indicate that the majority product is NO$_2^+$. Dunkin et al.$^6$ suggest that an additional “switching” reaction channel results in a third product channel giving O$_2^+$, proceeding through an intermediate excited state of a (N$^+$ + O$_2$)$^*$ complex that can decay into NO$^+$ + O and O$_2$ + N products, or can be stabilized collisionally or radiatively to form NO$_2^+$. Matsuoka et al.$^{10}$ have also reported that the NO$_2^+$ ions formed are rapidly lost in collisions with N$_2$ at atmospheric pressure with an effective decay constant of $3.0 \times 10^8$ s$^{-1}$, carefully ruling out any interfering reactions of NO$_2^-$. Hiraoka$^9$ measured the relative branching ratios of the NO$_2^+$ and NO$^+$ products from 64 to 552 K and found that $[\text{NO}_2^+]/[\text{NO}^+]$ decreases monotonically with increasing temperature which is in general agreement with the values of Matsuoka et al. from 303 to 384 K.$^{12}$

Matsuoka et al.$^{12}$ further addressed the problem of thermal stability of the NO$_2^+$ product of reaction (1). They determined that the decay of NO$_2^+$ into NO$^+$ has a strong temperature dependence, increasing exponentially with increasing temperature. Furthermore, they have found that NO$_2^-$ from reaction (1) is reactive and thermally dissociates, while the NO$_2^+$ formed in the reaction of O$_2^+$ with NO$_2$ is thermally stable and unreactive. Therefore, they conclude that the two NO$_2^+$ ions do not have the same structure. These authors further argue that the atomic arrangement in the NO$_2^+$ from reaction (1) should be the higher energy NOO$^+$ linear isomer.$^{12}$

However, the HPMS measurements of Hiraoka indicate that ONO$^+$ is the isomer formed by reaction (1), based on the lack of thermal unimolecular dissociation of NO$_2^+$ at 465.7 K and $ab$ initio calculations of the successive binding energies of N$_2$ ligands in NO$_2^+(N_2)_n$ clusters containing ONO$^+$. The calculated energies agree with the measured values, whereas the preliminary calculations of Hiraoka and Yamabe show that NOO$^+$ is highly unstable.$^{13}$ Differences in the nucleophilic properties of N$_2$ vs O$_2$ further imply that the observed product is ONO$^+$.$^9$

In order to elucidate the kinetics of the N$_3^+$ + O$_2$ system, the rate constants and product ion branching ratios for reaction (1) were measured from 120 to 1400 K using two complementary instruments. The discrepancy concerning the NO$_2^+$ product isomer was investigated by experimental and theoretical analyses of the thermal stability, reactivity, energetics, and structure of the various isomeric forms of NO$_2^+$ (including barriers to rearrangement). The combination of a SIFT and high-temperature flowing afterglow (HTFA) allows this wide temperature range to be probed. Comparing the current rate constants to the previous measurements as a function of temperature$^7$ and kinetic energy$^9$ shows the effects of different types of energies on the reactivity. A potential energy profile has been calculated to elucidate the reaction mechanism.

II. EXPERIMENT/COMPUTATION

A. Kinetics measurements

An experimental study of the N$_3^+$ + O$_2$ reaction was performed using two separate techniques in order to encompass the temperature range from 120 to 1400 K. The SIFT has been used in the temperature range between 120 and 500 K, while the HTFA measurements have spanned the 350–1400 K range. Details of the two systems are described elsewhere.$^{14-16}$ All of the reagent gases have been used without further purification.

In the SIFT, N$_3^+$ ions are generated from N$_2$ in an effusive source in a high-pressure source region. High purity molecular nitrogen is taken from the boil-off of a high-pressure liquid nitrogen dewar. Two mechanisms for N$_3^+$ generation are operative: three-body association of N$^+$ with N$_2$ and Penning ionization of N$_2$, the latter of which is shown in Eq. (2).

$$\text{N}_3^+ + \text{N}_2 \rightarrow \text{N}_2^+ + \text{N}. \quad (2)$$

Mass selection of N$_3^+$ using a quadrupole mass filter before injection into the fast flow of He buffer gas (AGA, 99.995%) produces N$_3^+$ in the flow tube in >99% purity. After reaction, the ions are sampled through a blunt nose cone aperture and are analyzed by a quadrupole mass analyzer, and then detected by an electron multiplier. Rate constant measurements are performed at two reaction distances, 36 and 52 cm, and the data do not vary between the two reaction lengths within the experimental uncertainty. Measurement of the exponential decay of the primary ion signal as a function of the O$_2$ concentration (Massachusetts Oxygen, 99.999%) under pseudo-first-order conditions, coupled with the previously measured ion velocities, yields the rate constant. The average relative error in the rate constants is ±15% with absolute errors of ±25%.$^{15}$

The product ion branching ratios are determined in the SIFT by measuring the relative magnitude of the NO$^+$ and NO$_2^+$ signals obtained from extrapolating to zero O$_2$ concentration:

$$\%[\text{NO}^+] = 100 \times [\text{NO}^+]/([\text{NO}_2^+] + [\text{NO}^+]). \quad (3)$$

In addition, the ratio of NO$_2^+$ to NO$^+$, $r$, is obtained as shown in Eq. (4),

$$r = \frac{[\text{NO}_2^+]}{[\text{NO}^+]}.$$
This ratio has been measured over a relatively high range of oxygen concentrations, approaching 0.2% of the total gas flow. No secondary chemistry is observed and the percentage of NO₂⁺ is independent of O₂ flow. The relative uncertainty of the branching ratios is ±10% of the major product peaks.¹⁷

In the HTFA, the ions are generated in an ion source region positioned orthogonal to the quartz reaction tube.¹⁴ The source has a 0.5 in. diameter exit aperture for these experiments, so that the source pressure is higher than that found in the flow tube. Nitrogen is introduced downstream from a thoriated iridium filament biased to 100–150 V with 0.1–4 mA emission current. The same two reaction processes generate N⁺ ions as in the SIFT. A helium buffer flow up to 30 000 SCCM is introduced through the source region and carries the ions into the flow tube where a commercial furnace controls the temperature (SCCM denotes cubic centimeter per minute at STP). Typical operating pressures in the flow tube are between 0.8 and 1.5 Torr. O₂ is added 52.7 cm upstream from the nose cone sampling plate in the region where the ions have reached thermal equilibrium. A small fraction of the flow is again sampled and the ions are analyzed by a quadrupole mass analyzer, and then detected by an electron multiplier. The N₂⁺ ion decay rate does not change with the N₂ source gas flow at rates of 125–500 SCCM, indicating that the production of the ions is complete before the reactant inlet. Residual water contaminant in the helium buffer is reduced using a liquid nitrogen cooled sieve trap.

At higher temperatures, alkali ions thermionically emitted from the quartz flow tube are inevitable, but these ions are nonreactive background species in the mass spectrum. Invariance of the alkali ion signal as a function of O₂ flow has been used as an indicator that the flow tube chemistry has properly equilibrated as demonstrated previously.¹⁸ At 1400 K, the level of the K⁺ signal was comparable to the level of the N₂⁺ signal (m/z = 42), and the mass resolution of the quadrupole filter was adjusted so that the two mass peaks are separated. Average relative errors of the N₂⁺ decay rate are ±12% with absolute errors of ±25%.¹⁶

Unfortunately, a pure N₂⁺ signal is not possible in the HTFA because of the lack of upstream mass selection and the mechanism for N₂⁺ ion generation. Due to the complexity of the reactions involving N⁺, N⁺, N⁺, N⁺, and NO⁺ also present in the flow tube, it is impossible to determine the branching ratios for the two reaction channels by simultaneously following the two products, especially since NO⁺ is also a product of a number of background reactions, e.g., N⁺ + O₂. Fortunately, none of the contaminant ions react with O₂ to produce an ion at m/z=46, the mass of NO₂⁺. Therefore, the NO₂⁺ fraction at a given concentration of oxygen has been evaluated in the HTFA from Eq. (5) below,

\[
\%[\text{NO}_2^+] = 100\left(\frac{[\text{NO}_2^+]-[\text{NO}_2^+]_0}{([\text{N}_2^+]_0-[\text{N}_2^+]_0)}\right),
\]

where the differences between the ion signals at a given O₂ concentration and zero O₂ concentration are used. The relative uncertainty in the product distributions determined this way is also 10%. This uncertainty is determined from the agreement with the SIFT results shown below.

B. Ab initio calculations

Energies of the various isomers of the NOO⁺ species on the ground state singlet potential energy surface have been calculated at different levels of theory. Here we present only the energetics of the NOO⁺ isomer optimized using multireference configuration interaction at the MRCISD(Q)(15e,12o)/cc-pVTZ level. The detailed theoretical results will be presented in a separate paper.

A potential energy profile has also been calculated. Geometries were optimized using density functional theory (DFT) at the Becke3-Lee, Yang, and Parr (B3LYP)/aug-cc-pVTZ level, where aug-cc-pVTZ denotes Dunning’s correlation consistent polarized valence triple-zeta basis set augmented with diffuse functions.¹⁹ Frequency calculations have been performed at the same level as the geometry optimization to determine whether the species are stable molecules or transition states. Intrinsic reaction coordinate (IRC) calculations have been carried out to confirm the connectivity of every optimized transition state at the same level. To obtain better energetics, single point energy calculations using coupled cluster theory at the CCSD(T)/aug-cc-pVQZ level have been performed on the B3LYP/aug-cc-pVTZ optimized geometries, where aug-cc-pVQZ denotes Dunning’s correlation consistent polarized valence quadruple zeta basis set augmented with diffuse functions.¹⁹

III. RESULTS AND DISCUSSION

A. Rate constants

Figure 1 shows the rate constants of the N₂⁺ + O₂ reaction as a function of temperature as measured in the SIFT and HTFA covering the 120–1400 K range; error bars denote our ±15% experimental uncertainty. The rate constants decrease substantially with temperature up to 1200 K, reaching...
a minimum at this temperature and then increase up to 1400 K. Data from the two instruments, the SIFT and the HTFA, agree in the overlapping temperature region within the experimental uncertainty.

Measurements from the other temperature dependent experiments discussed in the Introduction are also shown in Fig. 1 and agree well with the present work, if the rate constants measured below ~180 K in the HPMS are excluded. The previous flow drift tube results are discussed separately below. These values are not shown because clusters of the type \( N_3^+ (N_2)_n \) are formed and because the rate constants do not correspond purely to \( N_3^+ \) as the reactant ion. Below 180 K, the HPMS rate constants show a steeper increase with decreasing temperature, indicating that the \( N_3^+ (N_2)_n \) ions react faster than does bare \( N_3^+ \). This interference is not a problem in the current experiments because no \( N_2 \) is present in the flow tube of the SIFT and clustering to \( N_3^+ \) does not occur at high temperature in the HTFA. The room temperature value measured by Matsuoka et al. at atmospheric pressure is slightly larger than the rest of the measurements and may have been slightly influenced by clustering. However, the atmospheric pressure value does agree within the experimental error. All of the rate constants shown are considerably smaller than the Langevin collision rate constant of \( 7.0 \times 10^{-10} \text{ cm}^3 \text{s}^{-1} \).

Also shown in Fig. 1 are rate constants from a flow drift tube experiment (FDT) taken as a function of center-of-mass kinetic energy. The kinetic energy is converted to a translational temperature by \( \langle K E_{\text{c.m.}} \rangle = 3/2 k T_{\text{eff}} \). The agreement with the drift tube data is good at low temperature, but at temperatures around 800 K and higher, the data diverge for reasons to be discussed below. In addition, the current results suggest that the rate constants begin to increase after reaching a minimum at a translational temperature lower than that found in the drift tube.

In order to compare the role of various types of energies, the data are replotted as a function of average total energy in Fig. 2. The rotational and vibrational temperatures of the reactant ions in the FDT experiments are determined from the center-of-mass collision energy for ion-buffer gas collisions \( \langle (K E_{\text{buffer}}) = 3/2 k T_{\text{rot}} \rangle \). Note that a \( N_2 \) buffer was used in the drift tube experiments. The average vibrational energy is calculated as an ensemble average over a Boltzmann distribution of harmonic vibrational frequencies of the reactants. At low temperatures, \( N_3^+ \) vibrations predominantly contribute, while the \( O_2 \) vibrations contribute only at temperatures over about 1000 K.

The rate constants from all of the data sets decrease with increasing energy and reach a small plateau in the 0.2–0.4 eV range before decreasing again to a minimum at 0.7 eV. Above this energy, the rate constants increase with increasing energy. The temperature and kinetic energy data are indistinguishable, indicating that all forms of energy that are excited in this range control reactivity approximately the same. Of course, cancellation of effects is possible, but past experience indicates this occurrence is unlikely. The equivalency of translational and rotational energies is a typical behavior for most ion-molecule reactions. Vibrational excitation often has a different effect than the other forms of energy.

**B. Branching ratios**

Previous results on the branching ratios for reaction (1) are few in number and contradictory. A SIFT experiment by Smith et al. yields a value of \( r = [\text{NO}_2^+]/[\text{NO}_4^+] = 0.43 \) at 300 K. However, a preference for reaction channel (1b) \( r = 1.33 \) was observed in the drift tube measurement by McCrumb and Warneck at the same temperature. Matsuoka et al. reported that \( r \) decreases from \( r = 1.72 \) at 303 K to \( r = 1 \) at 384 K in a TRAPI experiment, which suggests that reaction (1b) is preferred at lower temperatures and that this preference may be reversed at higher temperatures, i.e., \( r < 1 \). Thus, the results from an earlier SIFT experiment are the opposite of the results from the drift tube\textsuperscript{1} and the TRAPI experiment.\textsuperscript{1} The present work attempts to (a) address the discrepancy in the value of \( r \), (b) verify its temperature dependence, and (c) elucidate the nature of the \( \text{NO}_2^+ \) product.

Branching ratios as a function of temperature obtained from the current experiments are shown in Fig. 3, given as...
the ratio $r$ defined in Eq. (4). The most striking feature of the present results is a monotonic exponential decrease of $r$ with increasing temperature over several orders of magnitude, i.e., a substantial decrease in the amount NO$_2^+$ observed for reaction (1) at high temperature. This downward trend is in excellent agreement with the HPMS temperature dependent data up to ca. 450 K, indicating that the crossover to $r<1$ predicted by Matsuoka$^{12}$ does occur. The present data further indicate that the trend continues until the highest temperatures. There are no obvious explanations for (a) the exponential nature of the decline and (b) for the disagreement of the O- with the crossover to $r<1$.

The other anomalous point on the branching ratio plot is from the earlier SIFT study.$^7$ In contrast to the present study where N$_2^+$ is injected cleanly into a helium buffer, the previous SIFT value was obtained by injecting N$^+$ into a mixed He (0.45 Torr)/N$_2$ (0.2 Torr) buffer and letting N$_2^+$ cluster with N$_2$ to form N$_2^+$$. A potential problem would occur if some N$^+$ had survived into the reaction zone (but not necessarily to the mass spectrometer) since the reaction of N$^+$ with O$_2$ produces NO$^+$ but not NO$_2^+$. In addition, Matsuoka$^{12}$ et al. have shown that the NO$_2^+$ formed in reaction (1) subsequently reacts with N$_2$ with a rate constant of $1.2 \times 10^{-15}$ cm$^3$ s$^{-1}$ to form NO$^+$. This reaction would partially convert NO$_2^+$ to NO$^+$ in the previous SIFT experiment and skew the branching ratio to a value less than 1.

### C. NO$_2^+$ structures and unimolecular reactions

The structure of the NO$_2^+$ ions produced in reaction (1) has also been the subject of debate. Matsuoka and co-workers$^{10,12}$ have reported that the NO$_2^+$ ions formed by reaction (1b) in the TRAPI are lost in collisions with N$_2$ at atmospheric pressure with an effective decay constant of $3.0 \times 10^7 s^{-1}$ or a bimolecular rate constant of $\sim 3 \times 10^{-16}$ cm$^3$ s$^{-1}$. Furthermore, they observed that this decay (a) produces NO$^+$, (b) is independent of the O$_2$ and N$_2$ content in their reaction gas mixture, and (c) is temperature dependent. In their earlier paper,$^{10}$ they assigned the reactivity to

$$\text{NO}_2^+ + \text{N}_2 \rightarrow \text{NO}^+ + \text{N}_2 \text{O} \quad (6)$$

with a bimolecular rate of $1.2 \times 10^{-15}$ cm$^3$ s$^{-1}$. In their subsequent paper, they attribute the loss of NO$_2^+$ to thermal decomposition of an isomer having a NOO$^+$ structure, mainly because the rate did not change with buffer gas density. The observed decay rate of NO$_2^+$ measured over a relatively narrow range between 294 and 384 K using TRAPI increases exponentially with increasing temperature, giving a rate of $(5.6 \times 10^7)e^{[-4500/RT]}$ s$^{-1}$. Furthermore, they have found that NO$_2^+$ from reaction (1) reacts strongly with C$_2$H$_6$ to form C$_2$H$_5$. However, the NO$_2^+$ formed in the reaction of O$_2^+$ with NO$_2$ does not react with C$_2$H$_6$. Therefore, they concluded that the two NO$_2^+$ ions do not have the same structure, also supporting the thermal decomposition hypothesis.

In contrast, the HPMS measurements of Hiraoka indicate that ONO$^+$ is the isomer formed.$^9$ Loss of NO$_2^+$ is not observed at 465.7 K in the HPMS. Ab initio calculations of the successive binding energies of N$_2$ ligands in NO$_2^+$ (N$_2$) clusters with ONO$^+$ agree with the measured values, whereas their preliminary calculations show that NOO$^+$ is highly unstable.$^{13}$ In the experiments of Matsuoka et al. at atmospheric pressure,$^{10,12}$ the single exponential decay of the NO$_2^+$ signal indicates that only one isomer is formed.

To address this issue, both high-level theoretical calculations and an experimental test of the reactivity of the NO$_2^+$ produced in reaction (1) have been performed. The details of the calculations are presented in a separate paper, with only the most relevant results at the highest level of calculation, MRCISD(Q)/cc-pVTZ, presented here in Fig. 4. The calculations show that the NOO$^+$ isomer is stable, with an energy of 4.51 eV above the ground state linear ONO$^+$ isomer. The NOO$^+$($^1\Sigma^+$) isomer is linear with a N--O bond distance of 1.146 Å and an O--O bond distance of 1.233 Å. A barrier of 0.89 eV prevents the isomer from dissociating to NO$^+$($^1\Sigma^+$)+O($^1D$), which is only 0.060 eV endothermic. Production of NO$^+$($^1\Sigma^+$) and O(g$^3P$) from NOO$^+$($^1\Sigma^+$) is exothermic by almost 2 eV, correlating to three triplet surfaces accessed by curve crossings. Two minima of the singlet-triplet seam-of-crossing (MSX) have been located: MSX$^1\Sigma^+/3\Pi$ ca. 0.07 eV below NOO$^+$-TS and MSX$^3\Sigma^+/3\Sigma$ ca. 0.02 eV below NOO$^+$-TS, as shown in Fig. 4. Spin-orbit calculations indicate that both of them have
very strong interactions: 65.5 cm\(^{-1}\) for each of the \(^3\Sigma^+\) components with \(^1\Sigma^+\) and 35.7 cm\(^{-1}\) for the MSX \(^3\Sigma^+\) isomer, suggesting that predissociation could easily occur. On the singlet ground state, there is also a bent \((\angle_{\text{ONO}})^\text{is}TS)\) isomer of ONO\(^+\) with \(\angle_{\text{ONO}}\) of 80°. This bent form lies 4.03 eV above the most stable linear ONO\(^+\), and a small barrier of 0.23 eV relative to the bent isomer links them. This barrier is relatively small and thermal isomerization in the presence of buffer gas should be fast enough to prevent this species from being observed in our apparatus.\(^1\) However, the barrier between linear NOO\(^+\) and bent ONO\(^+\) isomers, NOO\(^+\) isoTS, is found to be quite large, 1.36 eV from the linear NOO\(^+\) isomer. The dissociation processes of ONO\(^+\) isomers are also investigated. Two corresponding transition states, ONO\(^+\) disTS1 and ONO\(^+\) disTS2, connecting the linear and bent isomers are found to be 4.77 and 1.01 eV above the corresponding isomers, respectively. Note that the activation energy found by Matsuoka et al.\(^10,12\) for NO\(_2\) reacting with N\(_2\) is much smaller than the barrier found here for dissociation to NO\(^+\), indicating that the process observed is probably not thermal dissociation. (Note also that the new energetics for N\(_3\), the production of N\(_2\) + O(\(^1\)D) + NO\(^+\) is 0.99 eV exothermic.) Therefore, all of the transition states and isomers except the NOO\(^+\) isoTS discussed above lie below the entrance channel energy for N\(_2\) + O\(_2\) at 300 K as shown in Fig. 4.

It is relatively easy to determine if a high energy form of NO\(_2\) is generated in the present experiments. A sufficiently large amount of O\(_2\) can be added upstream in the flow tube of the SIFT to drive reaction (1) to completion before a second gas is added downstream to test the chemistry of the product ions. What is required is a gas that does not react with the ground state ONO\(^+\) isomer, but will react with the NOO\(^+\) isomer (assuming that the bent ONO\(^+\) species, if formed, thermally isomerizes to the linear ground state). Xenon is a good candidate since it is a generally unreactive gas. For this test, Xe (Air Products, 99.995%) is introduced into the SIFT at a distance 30 cm downstream from the O\(_2\) inlet. In a separate experiment for comparison purposes, ground state ONO\(^+\) is generated in the source region of the SIFT by electron impact on nitrogen dioxide (Matheson, 99.5%). These ONO\(^+\) ions are mass-selected and then reacted with Xe in the normal manner. NO\(_2\) injected into the flow tube from the ionization of nitrogen dioxide in the source does not react with Xe. By contrast, the 46 amu ion generated by reaction (1) is totally consumed when Xe is added to the SIFT. The reaction turns out to be more complicated than expected, giving two products:

\[
\text{NOO}^+ + \text{Xe} \rightarrow \text{NO}^+ + (\text{Xe} + \text{O}) + 356 \text{ kJ mol}^{-1} \quad (6a)
\]

\[
\rightarrow \text{XeO}^+ + \text{NO} + 108 \text{ kJ mol}^{-1}. \quad (6b)
\]

The bulk of the reaction produces NO\(^+\). However, the reaction is sufficiently exothermic so that the neutral product could be either XeO or Xe+O. This difference in reactivity confirms that the NO\(_2\) made in reaction (1) is a high-energy isomer and is not in the linear ONO\(^+\) form, in agreement with the TRAPI results.\(^12\) Production of XeO\(^+\) is rare, but it has been observed previously.\(^24\) We have also confirmed that C\(_2\)H\(_6\) reacts with NOO\(^+\) from reaction (1) to produce C\(_2\)H\(_5\) and that linear ONO\(^+\) does not.

**D. Potential energy surface and reaction mechanism**

Lindinger suggests that the decrease of the rate constant with kinetic energy (or temperature) is due to the lifetime of the complex intermediate decreasing with the increasing relative velocity of the reactants.\(^8\) This assumes that product formation is slow compared to that for dissociation into reactants and that no barrier is above the energy of the reactants. This assertion is probably qualitatively correct but somewhat simplistic for a quantitative description. The increase at higher temperatures/energies is probably attributable to a new mechanism that becomes available.

In order to elucidate the probable mechanisms, CCSD(T) calculations of the potential energy profile have been made as mentioned in the Experiment/Computation. The resulting

---

**FIG. 5.** Singlet potential energy surface for the reaction of N\(_2\) + O\(_2\) calculated at the CCSD(T)/aug-cc-pVQZ//B3LYP/aug-cc-pVTZ level of theory. On the structures, numbers are bond distances and angles (in angstrom and degree). For the energetics, ZPE correction has been applied.
The reaction of \( \text{N}_2^+ + \text{O}_2 \)

Singlet and triplet potential surfaces are shown in Figs. 5 and 6, respectively. The reactants are triplets and correlate to singlet and triplet products. The connectivity of the transition states has also been confirmed as discussed in the Experiment/Computation.

On the singlet surface shown in Fig. 5, two intermediates are formed directly from the reactants. The first intermediate (INT_1) has the \( \text{O}_2 \) axis nearly parallel to the quasilinear \( \text{N}_2^+ \) \((\angle \text{N-N-N}=168^\circ)\) axis, with \( \angle \text{N-N-O}=105^\circ \) and \( \angle \text{N-O-O}=115^\circ \). This “trans” structure lies 0.51 eV below the energy of the reactants. The other “cis” intermediate (INT_2) is more v shaped and lies 0.56 eV below the reactants. A transition state (TS_4) separates the two intermediates and is 1.20 eV above INT_1. INT_1 correlates only to the NOO+ product, but the reaction must pass through TS_1 with a barrier 0.17 eV above the reactants. Therefore, it is not important at low energies or temperatures. INT_2 is connected to both NO+ and NOO+ through separate transition states (TS_3 and TS_2) that are 0.15 and 0.01 eV, respectively, above the energy of INT_2.

On the triplet surface shown in Fig. 6, a V-shaped intermediate (INT_3) is 0.54 eV below the reactants and connects to NO+ and \( \text{N}_2\text{O}_2 \) singlet products through transition state (TS_6) only 0.04 eV above INT_3. However, INT_3 can only be reached through a cyclic \( \text{N}_2\text{O}_2^+ \) transition state complex (TS_5) that is 2.09 eV higher in energy. Another transition state (TS_7) lies 0.53 eV above the reactants and simultaneously produces three fragments: NO+ (\( ^1\Sigma^+ \)), \( \text{N}_2 \), and O(\( ^3P \)).

The features of the surfaces illustrated in Figs. 5 and 6 at least qualitatively explain the reactivity. At low temperatures, the reaction occurs at an appreciable fraction of the collision rate and decreases with increasing energy and temperature. That trend is typical for ion-molecule reactions involving surfaces with stable intermediates followed by barriers to reaction. The simplest explanation is that decay of the intermediates into reactants involves a loose transition state and into products involves a tight one. This picture is a more complete description of the mechanism than proposed by Lindinger.

At low energies, reaction (1) occurs on the singlet surface in Fig. 5. Once INT_2 forms, it can decay back into reactants or into either product channel, NOO+ or NO+. NOO+ has the lower barrier and is thus favored at low energy. However, the barrier to NO+ production is not much higher and the data show that NO+ is favored by 400 K. Any molecules that form intermediate INT_1 must dissociate back to reactants unless there is enough energy to overcome the barrier. However, sufficient energy is not available until \(-800 \text{ K (~0.7 eV)}\).

The upturn in the rate constants occurs at a total energy of \(-0.7 \text{ eV}, \text{Fig. 2} \). Two reaction paths may contribute to this trend at higher energies. First, TS_4 on the single surface connecting the two intermediates INT_1 and INT_2 has a barrier of 0.69 eV, approximately coinciding with the energy at the upturn. Therefore, this additional mechanism leading to products may begin to contribute to the overall reactivity above 0.7 eV, causing an increase in the total rate constant. Second, reaction also becomes possible on the triplet surface in Fig. 6 where the 0.54 eV barrier to TS_7 can be overcome, opening up the channel that produces three fragments including NO+. In addition, a further reduction in the NOO+ product at higher energies can come from unimolecular decomposition of NOO+ giving NO+. The multireference configuration interaction calculation results shown in Fig. 4 indicate that a crossing from the singlet NOO+ surface to a triplet surface at MSX \( ^1\Sigma^+ / \Pi \) occurs about 0.8 eV above NOO+ minimum that allows predissociation to NO+ (\( ^1\Sigma^+ \)) + O(\( ^3P \)) that is 2 eV lower in energy. This pathway is consistent with the NOO+ unimolecular decomposition results of Guilhaus et al. showing that NOO+ dissociation produces NO+ fragments with 2 eV kinetic energy.\(^{23}\) The present calculations also indicate that a barrier of 0.89 eV exists to dissociate NOO+ into NO+ and O(\( ^1D \)). Since the products...
The rate constants to rate constants are found to decrease from 120 to 1200 K. IV. CONCLUSIONS

The kinetics of the reaction of N$_3^+$ with O$_2$ has been studied over a wide temperature range, up to 1400 K. The rate constants are found to decrease from 120 to 1200 K before increasing up to 1400 K and agree with previous measurements taken over limited temperature ranges. Comparing the rate constants to those obtained in a drift tube shows that all forms of energy behave similarly.

Two products are observed, namely, NO$_2^+$ and NO$^+$. The former is favored at low temperature and the latter at higher temperature. The ratio $r$ of NO$_2^+$ to NO$^+$ decreases exponentially with temperature from $r=2$ at 120 K to $r<0.01$ at 1400 K. In this paper, a straightforward test of the reactivity on the NO$_2^+$ product ion was made that shows conclusively that this product is a high energy isomer, most likely NOO$^+$. Theoretical calculations confirm that this is a stable isomer, with a barrier to dissociation of about 0.9 eV. The energy of the NOO$^+$ isomer is determined to be 4.5 eV above that of the most stable isomer, linear ONO$^+$. The calculated potential energy surfaces for the reaction giving singlet products explain most of the current data.

ACKNOWLEDGMENTS

The authors thank the Air Force Office of Scientific Research (AFOSR) for its continued support of this laboratory. A.J.M. is under contract (Grant No. F19628-99-C-0069) to Visidyne, Inc., Burlington, MA. A.I.F. acknowledges support under the NRC Research Associate program. S.P. also acknowledges support under the NRC Summer Faculty Program. The work done at Emory University was supported in part by a grant from AFOSR (Grant No. FA9550-04-1-0080). A part of the computer time was made available by a grant from AFOSR under the DoD High Performance Computing Program.