Articles

N(²D) Product Velocity Mapped Imaging in the VUV Photolysis of Nitrous Oxide at 118.2 nm

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Resonance-enhanced multiphoton ionization with time-of-flight product imaging of the N(²D) atoms has been used to study the N₂O photodissociation at 118.2 nm and the two-photon dissociation at 268.9 nm. These imaging experiments allowed the determination of the total kinetic energy distributions of the NO(X ²Π) and N(²D_{5/2}) products. The NO(X ²Π) fragments resulting from the photodissociation processes are produced in highly vibrationally excited states. The two-photon photodissociation process yields a broad NO(X ²Π) vibrational energy distribution, while the 118.2 nm dissociation appears to produce a vibrational distribution sharply peaked at NO(X ²Π, ν =14).

Keywords : VUV photolysis, Nitrous oxide, NO vibrational energy distribution, 118.2 nm.

Introduction

Nitrous oxide plays an important role in the global nitrogen cycle and it is an important trace component of the atmosphere.¹ The vacuum ultraviolet (VUV) photolysis of nitrous oxide has been extensively studied.² Dissociation of N₂O in this spectral region has proven to be a good source for various electronically excited states, including O(¹D). O(¹S), N(²P and ²D). N₂(A ³ Σ^+ _u). Much less information is known about either the quantum yields for these excited states² or the energy partitioning in the diatomic fragments. This type of information is needed for a better understanding of the dynamics and the potential energy surfaces of N₂O.

In the 118.2 nm region N₂O has an absorption coefficient of about 1000 atm⁻¹ cm^{-1,3} At this wavelength, dissociation to NO and N fragments primarily produces NO(X ²Π) in coincidence with either N(⁴S). N(²D), or N(²P). The NO(A²Σ⁻) + N(⁴S) channel is also energetically feasible.

Black *et al.*² measured the quantum yields for the production of N₂(A ${}^{3}\Sigma^{-}u$), N(²D), and O(¹S) from the VUV photolysis of N₂O. The quantum yield of N(²D) was measured using the intensity of NO β bands generated by the sequence

N(²D) + N₂O → N₂ + NO(B ² Π_r). NO(B ² Π_r) → NO(X ² Π_r) + h ν (NO β bands).

These authors found that the quantum yield for N(²D) is about 40% in the 118 nm region, but that it drops sharply around 120 nm where the production of O(¹S) + N₂ begins to dominate. This observation is also in agreement with that of Nee *et al.*⁴ who detected O(¹S) metastable atoms produced in the photodissociation of N₂O by observing the fluorescence of the O(¹S \rightarrow ¹D) transition. The van der Waals interaction of Xe buffer gas with the metastable oxygen atoms greatly enhanced the normally weak fluorescence. Nee et al.4 observed a strong O(¹S) fluorescence for wavelengths greater than 120 nm and also measured the absorption spectrum of nitrous oxide over the range 100-140 nm. Above 120 nm. the spectrum is nearly a smooth continuum, whereas below 120 nm, it is dominated by a single peak at 117.8 nm. The authors attributed this peak to a Rydberg state which dissociates to produce only very small amounts of $O(^{1}S)$. In agreement with the previous work by Black *et al.*², this state is observed to produce strong NO β band fluorescence.⁴ The Rydberg state has either ${}^{1}\Sigma$ or ${}^{1}\Pi$ symmetry, and states of either symmetry would correlate to $N(^{2}D, ^{2}P) + NO(X^{2}\Pi)$ or $N_2(X) + O(^1S)$ products.^{4,5,6} Since Nee *et al.*⁴ observed a very small quantum yield for the production of $O(^{1}S)$ atoms, they concluded that the N₂O Rydberg state must be a major factor in the production of metastable N atoms.

That this state does indeed produce metastable N atoms has been shown by the recent work of Umemoto *et al.*,⁷ who investigated the 2-photon photolysis of N₂O between 235 and 240 nm. N(²D, ²P) and NO(v" = 1) products were all detected by resonance-enhanced multiphoton ionization (REMPI) combined with TOF mass spectrometry. Efforts were also made to detect NO(X. v = 0), but the background signal due to the one color MPI process was too great. The authors argued that two processes occur: predissociation of the two-photon excited state of nitrous oxide, and the dissociation of a three-photon excited state into the N(²D) and NO fragments.

Chan *et al.*⁸ obtained the electronic excitation spectrum of nitrous oxide along with the corresponding oscillator strengths. In the energy range of 10.36-10.64 eV they find an integrated optical oscillator strength of 0.0523. The photoabsorption spectrum of N₂O in the regions 10-12 and 12-21 eV consists mainly of transitions from the 2π . 7σ . 1π and 6σ orbitals to

Rydberg levels. At 117.8 nm, which corresponds to 10.5 eV. the transition is assigned to a $3p\pi \leftarrow 2\pi$, the first member of a Rydberg series leading to an ionization threshold of 10.54 eV.

In this paper we present results for the VUV photodissociation of N₂O at 118.2 nm, which is nearly resonant with the transition $(3p\pi \leftarrow 2\pi)$ to a Rydberg state. The VUV radiation was produced by non-resonantly tripling 354.6 nm laser light in a rare gas cell. We detect the N(²D_{5/2}) product following the primary VUV photolysis:

$$N_2O + 118.2 \text{ nm} \rightarrow N(^2D_{5/2}) + NO(X)$$
 (1)

using the velocity-mapped, ion imaging technique. The focused ionization laser produces a background of $N(^2D_{5:2})$ atoms by a two-photon process:

$$N_2O + 2 \times (268.9 \text{ nm}) \rightarrow N(^2D_{5/2}) + NO(X)$$
 (2)

which contributes to the image. A third process produces a background of N^+ signal in a multi-photon process involving the 118.2 nm light, and possibly the 354.6 nm laser light:

$$N_2O + 118.2 \text{ nm}/354.6 \text{ nm} \rightarrow N^- + NO(X)$$
 (3)

By making use of the spatial properties of the imaging technique, we were able to distinguish the primary process of interest from the background contributions.

Experimental Section

The technique of ion imaging has been described in more detail elsewhere.⁹ Neat N₂O (Matheson) and 10%NO/He (Matheson) were used from the cylinder without further purification. The gases were expanded at 17 psia through a pulsed 0.5 mm diameter nozzle and collimated by a 0.5 mm diameter skimmer mounted 0.5-1.0 cm from the nozzle orifice. Further downstream, the molecular beam was crossed at right angles by two counterpropagating laser beams. one used to dissociate the molecules and the other to probe the resulting $N(^{2}D_{52})$ fragments using (2+1) resonanceenhanced multiphoton ionization (REMPI) via the 3p²S_{1/2} state at 268.9 nm.⁷ Due to the extremely low $N(^{2}D_{5/2})$ signal in these experiments, the ionization laser was not scanned over the $N(^2D_{5/2})$ Doppler profile, however the resonance was monitored periodically by detecting $N(^2D_{5/2})$ fragments from the two-photon dissociation of NO at 268.9 nm.

The dissociation laser radiation at 118.2 nm was produced by nonresonant tripling in Xe of 354.6 nm from a Nd:YAG laser. The production of VUV was monitored by detecting the NO⁻ signal from the ionization of NO at 118.2 nm. Power dependence studies showed that the NO⁻ signal levels increased non-linearly with 354.6 nm pulse energies, with saturation occurring at pulse energies above 45 mJ. Lower laser energies in the range of 25-30 mJ/pulse were used in the experiments with N₂O in order to minimize a background signal caused by the 118.2 nm light. The 354.6 nm laser light was focused into a cell containing 30 Torr of Xe using a quartz lens of 7 cm focal length. The output VUV radiation along with the 354.6 nm light were collimated by a LiF lens which served as the exit window of the Xe cell. The 118.2 nm beam was then focused onto the molecular beam by a 12 cm focal length LiF lens located inside the chamber. The 354.6 nm light was focused some 6 cm beyond the interaction region.

The tunable light needed to probe the $N(^{2}D_{5/2})$ fragments at 268.9 nm was generated by a 354.6 nm pumped SCANMATE (Lambda Physik) with Coumarin 540A dye using second harmonic generation in a BBO I crystal. Typical powers were 0.5 mJ/pulse with a pulse duration of 8-10 ns. The probe beam was focused using a MgF₂ lens with a focal length of 46 cm. The polarization directions of the dissociation and the ionization lasers were parallel and perpendicular to the plane of the detector, respectively.

The imaging technique uses an electrostatic immersion lens which serves to extract the ionized $N(^2D_{5/2})$ fragments from the interaction region and to focus ions with equal velocity vectors to the same point on the detector.¹⁰ The magnification factor for this electrostatic lens was measured to be 1.21 ± 0.03 by dissociating NO using 2 photons of 268.9 nm and probing the $N(^2D_{5/2})$ fragment using the same REMPI scheme as described above. The magnification factor was obtained by comparison of the single ring in the image with the known energetics of NO dissociating to $N(^2D)$ and $O(^3P_I)$.

The ionized fragments were accelerated into a field-free flight tube along the axis of the molecular beam. The ions were imaged by a position sensitive detector consisting of a chevron-mounted, double microchannel plate (MCP) assembly (Galileo) coupled to a fast phosphor screen. The image on the screen was recorded by a 640 × 480 pixel CCD camera (Xybion). Both the MCP and the camera were electronically gated to collect signal corresponding to only the mass of the N($^{2}D_{5/2}$) fragment. Signal levels from the 118.2 nm photodissociation of N₂O were very low, on the order of 10-20 ions per frame. We demonstrated that the signal was VUV dependent by pumping out the Xe gas cell and observing that the signal disappeared. There was no contribution to the N($^{2}D_{5/2}$) signal solely from the 354.6 nm light.

Large background levels were encountered from both the photolysis wavelength (118.2 nm) and the probe wavelength (268.9 nm). In order to account for these backgrounds the imaging acquisition consisted of 3 separate toggle states in which the timing between the photolysis and the probe was changed. The first toggle state ensured that the probe laser was delayed from the photolysis by 20 ns. This gave us the desired N(²D_{5/2}) distribution from the photolysis of N₂O at 118.2 nm and subsequent ionization of the fragments. The second toggle state accounted for the background caused by N₂O two-photon absorption of the 268.9 nm probe laser followed by REMPI ionization of the $N(^2D_{5/2})$. We recorded this background by moving the photolysis laser pulse to a timing 1 μ s after the probe laser pulse. Similarly, the third toggle state accounted for the 118.2 nm background by removing the probe laser from the time frame of relevance. Each data set was collected for up to a total of 30.000 laser shots.



Figure 1. Velocity-mapped ion image of the $N(^{2}D_{5/2})$ fragment detected by (2+1) REMPI at 268.9 nm approximately 20 ns after photodissociation of N₂O at 118.2 nm (toggle state 1). The image is the sum of nine datasets totaling 136,980 laser shots.

Results and Discussion

In these experiments, we have observed signal at mass 14 from a variety of sources. These different sources were distinguished by acquiring images with both photolysis and probe lasers together and each separately, by toggling the digital delay generator controlling the laser timing. The unique spatial appearance of the images taken with each laser separately (toggle states two and three: TS2, TS3), provides the basis by which to understand the image acquired with both lasers together, (toggle state one: TS1), shown in figure 1. The signal acquired in each of the three toggle states depended on the presence of the molecular beam of N_2O . Only a very small background of ions was produced in the absence of the N_2O due to interaction of the focused lasers, presumably, with traces of pump oil.

The focused probe laser produced a strong background signal due to detection of the N(²D) product from a twophoton dissociation of N₂O at the (2+1) REMPI resonance wavelength of 268.9 nm. A power dependence experiment demonstrated a quartic dependence of the N(²D) signal on the laser pulse energy below 1.4 mJ/pulse in keeping with the 2 photon dissociation and 2 photon resonance in the REMPI step, and a saturated ionization step. At higher pulse energies, the power dependence was quadratic, indicating that one of the two photon processes became saturated. This background image was acquired with the second toggle state (TS2), when mass 14 signal could only arise from the probe laser. The intensity in the TS2 image is concentrated in a central disc approximately 150 pixels in diameter, and tails off rapidly in pixels beyond this region.

Significant background signal was also produced by the photolysis laser. This background image was acquired with the third toggle state (TS3) in which mass 14 signal could only arise due to the 354.6 nm/118.2 nm laser radiation. Although a TOF spectrum showed peaks due to hydrocarbons from interaction with traces of pump oil, the dominant features

Bull. Korean Chem. Soc. 2002, Vol. 23, No. 2 181

corresponded to N⁺. NO⁻. and N₂O⁺. This background signal disappeared upon removal of the VUV tripling medium. by pumping out the Xe, indicating a dependence on the 118.2 nm light. Multiphoton absorption processes would be required to ionize N₂O and its dissociation products: N(⁴S. ²D, ²P) and NO. The TS3 background N⁻ image uniformly covers the detector surface suggesting that there is a very broad range of translational energies available to these dissociation products. consistent with a multiphoton absorption scheme.

The TS1 image acquired with both lasers is a composite of the background signals arising from each laser interacting with the molecular beam of N_2O and the additional $N(^2D)$ signal from the single photon dissociation of N₂O at 118.2 nm. The TS1 image thus shows a bright central region and a diffuse, uniform outer coverage extending to the edge of the detector. The additional signal arising from both lasers together is seen as a brighter image intensity in the central region for TS1 than TS2. Unlike the background contributions, this additional signal is crucially dependent upon the spatial overlap between the two focused laser beams. The overlap was demonstrated when the central region of the N⁺ image increased significantly more than the sum of either laser background at a particular (x.y) position of a laser focusing lens, but dropped to background levels at different lens positions to the left or right, up or down.

The extra signal due to N_2O dissociation at 118.2 nm was weak and was not apparent on every laser shot. It was practical to acquire only about 9000 laser shots in each of the three toggle states before the overlap deteriorated. Individual images were shifted to a common center, then added to make up the image seen in Figure 1, as well as the images corresponding to the TS2 and TS3 toggle states. Matlab routines were used to perform an inverse Abel transform so as to obtain the 3-dimensional velocity distribution. Separation of the velocity distribution into angular and speed components

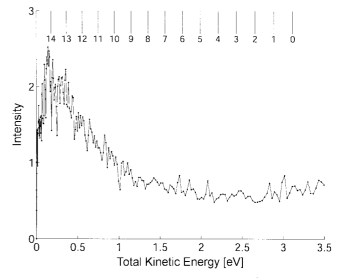


Figure 2. Total kinetic energy distribution of $N(^2D_{5/2}) + NO(X)$ obtained after the inverse Abel transformation of the image displayed in Figure 1, which was recorded in toggle state 1 (both lasers present).

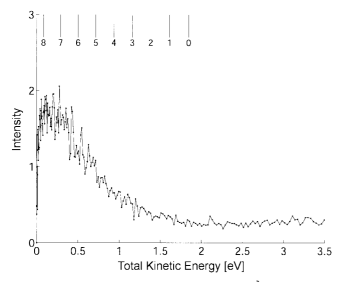


Figure 3. Total kinetic energy distribution of $N(^2D_{5/2}) + NO(X)$ obtained after the inverse Abel transformation of the background image recorded in toggle state 2 (268.9 nm REMPI probe laser only).

was achieved by integrating over speed intervals for each angle. and similarly integrating over all angles for each speed. The distribution of total kinetic energy released for the dissociation channel leading to N and NO cofragments was obtained by using the conservation laws of energy and momentum for each toggle state, and correcting for the magnification factor of our ion collection optics.

The total kinetic energy distributions derived from the summed images for each toggle state are shown in Figure 2 (TS1), Figure 3 (TS2), and Figure 4 (TS3). The broad, flat appearance of the kinetic energy distribution for the photolysis laser background (TS3), reflects the uniform image intensity extending to the perimeter of the detector surface.

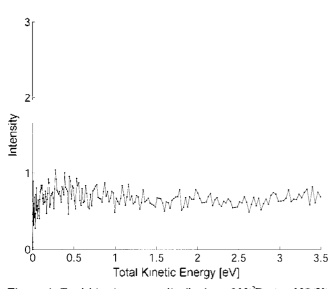


Figure 4. Total kinetic energy distribution of $N(^{2}D_{5/2}) + NO(X)$ obtained after the inverse Abel transformation of the background image recorded in toggle state 3 (118.2 nm/354.6 nm photolysis laser only).

This background from the photolysis laser contributes a pedestal-like appearance to the kinetic energy distribution derived from the image acquired with the photolysis and probe lasers (TS1). This is most apparent in Figure 2 as the non-zero baseline at kinetic energies above 1.5 eV. Since the kinetic energy distributions for TS1 and TS3 are obviously the same in this region, it is reasonable to assume that the separation of the photolysis laser background contribution to TS1 should be a simple matter of subtraction.

The kinetic energy distribution corresponding to TS2 provides information about the internal energy distribution of the NO co-fragment from the two-photon dissociation of N_2O at 268.9 nm. The difference between the two-photon energy of the laser radiation and the N2O dissociation energy to give $N(^{2}D) + NO$ products is the available energy to be partitioned between the kinetic energy of the recoiling cofragments and the internal energy of the NO diatom. The comb appearing above the data plotted in Figure 3 indicates the NO vibrational level corresponding to the difference between the known available energy and the observed kinetic energy. The right hand side of the comb marks the maximum energy available to translational motion and thus correlates with vibrationless. rotationless NO(v=0). In agreement with the energetics, the observed kinetic energy distribution has little intensity past the maximum allowed energy. The inferred vibrational distribution is inverted, peaking at $NO(\nu=7)$. Previous two-photon dissociation studies at shorter wavelengths between 235 and 240 nm did not provide vibrational distributions because only NO(ν =1) was detected.⁷

The contribution of the probe laser induced background is not so easily separated from the two-laser contribution in TS1 as was the photolysis laser background. A simple consideration of the relative intensities in Figures 2-4 will show that the kinetic energy distributions corresponding to TS2 and TS1-TS3 are comparable in magnitude and that further subtraction of TS2 would result in near zero or even negative intensities. This is likely a consequence of the competitive nature of the two multi-photon processes involving the probe laser, and we should expect that the probe laser background determined without the photolysis laser (TS2), will be reduced somewhat in the presence of the competing two-laser pump/probe process (TS1). By constraining the kinetic energy distributions TS1-f*TS2 to be equal to TS3, we could estimate the factor "f", by which the probe laser background should be reduced, and thus obtain the kinetic energy distribution for the two laser process alone.

Even without explicitly correcting for the probe laser background, it is clear from a comparison of the kinetic energy distributions (see Figures 2-4), that there is a significant peak near 0.14 eV which is present only in the case of TS1. This is a reproducible feature in the data sets acquired from day to day, and which emerges more clearly after summing several images. The comb of NO vibrational levels in Figure 2, which correlate with the kinetic energy available to the N(²D) and NO products, indicates that the peak corresponds to NO(v=14). Although there are no available calculations or experimentally derived bond lengths for the 10.5

N₂O Photolysis at 118.2 nm

eV Rydberg states of N_2O , it appears likely that the NN--O bond is severely compressed or extended in the excited state of N_2O in order to create such high vibrational excitation in the NO photofragment.

The peak value of the total kinetic energy of 0.14 eV corresponds to a lab frame velocity of 1133 ms⁻¹ for the probed N(²D) fragment. The Doppler width (full width at half maximum) for the REMPI transition under these conditions works out to be about 0.47 cm⁻¹. The nominal bandwidth of the dye laser is quoted by the manufacturer to be 0.15 cm^{-1} in the fundamental. It is our experience that the bandwidth after second harmonic generation is increased by a factor of two. Since the REMPI transition is a two-photon absorption in the resonance step, the laser bandwidth will further increase by a factor of $\sqrt{2}$. The effective laser bandwidth in the VUV may thus be estimated as 0.42 cm^{-1} . It is clear from these bandwidth considerations that our experiment was somewhat biased towards less efficient detection of N(²D) photofragments with velocities faster than at the peak of the kinetic energy distribution. Due to the weak signal, however, and the long accumulation times, it was deemed more practical to keep the probe laser wavelength on resonance rather than to scan over the full Doppler profile.

The very nearly isotropic appearance of the images indicates that the beta parameter is small, with analysis giving a value on the order of 0.1 for the peak of the kinetic energy distribution derived from TS1. This nearly zero anisotropy parameter indicates that the timescale for dissociation must be comparable to or slower than the rotational period. The shape of the angular distribution for the two photon dissociation of N₂O at 268.9 nm showed a significant contribution from the P₄(cos θ) Legendre polynomial, as is expected for a two photon absorption.

Conclusion

 $N(^{2}D_{5/2})$ fragments have been detected from the photo-

dissociation of N₂O at 118.2 nm and from the two-photon dissociation at 268.9 nm using the velocity-mapped, ion imaging technique. The total kinetic energy distributions for the products N(²D) and NO(X), derived from analysis of the images and the conservation laws. indicate that the NO(X) co-fragments appear to have inverted vibrational distributions for both processes. Remarkably, the 118.2 nm photodissociation produces a dominant peak in the total kinetic energy distribution corresponding to NO(v=14). In the 2-photon dissociation at 268.9 nm, the NO product was observed to have a broader vibrational distribution spread over several levels near to the energy limit.

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