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# The photodissociation of water-clustered HNO<sub>3</sub> studied at 193 nm by the LIF method

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#### Abstract

We studied the photodissociation of water-clustered HNO<sub>3</sub> molecules at 193 nm, a process relevant in atmospheric chemistry, by generating water clusters by adiabatic expansion and doping them with HNO<sub>3</sub> using the pickup method. The nascent OH( $X^2\Pi$ ) photofragments were probed by LIF, and the rotational state distribution in the exclusively produced vibrational ground state determined. These experiments were complemented by corresponding ones with argon and methanol clusters. The average cluster size was estimated to be  $\overline{n} \sim 200$ –800 depending on the compound. The results of this study are compared with those previously obtained from the photodissociation of the HNO<sub>3</sub> monomer. © 2001 Elsevier Science B.V. All rights reserved.

## 1. Introduction

The photodissociation of nitric acid HNO<sub>3</sub> has been well investigated, motivated mainly by the importance of this compound for atmospheric chemistry [1–3]. Following excitation into the weak, broad and structureless  $n\pi^*$ -absorption band at 260 nm, the major primary decay process yields OH + NO<sub>2</sub>, both in their electronic ground states with a quantum yield  $\phi$  near unity (channel (1) below) [4,5]. The nascent OH fragments have no vibrational excitation and the rotational state distribution is Boltzmann. With excitation into the strong absorption at 193 nm, the solar photolysis conditions in the stratosphere, where the 190–220 nm region is largely unfiltered by O<sub>3</sub>, O<sub>2</sub> and N<sub>2</sub>, is well reproduced. At this wavelength the primary

$$HNO_3 + hv \rightarrow OH(X) + NO_2 (X^2A_1)$$
  
 $\Delta H^{\circ} = 200 \text{ kJ/mol}$  (1)

$$\rightarrow \mathrm{OH}(\mathrm{X}) + \mathrm{NO}_2 \ (1^2\mathrm{B}_2)$$

$$\Delta H^{\circ} = 315 \text{ kJ/mol} \tag{2}$$

$$\rightarrow O(^1D) + HONO (\tilde{X}^1A')$$

$$\Delta H^{\circ} = 490 \text{ kJ/mol} \tag{3}$$

$$\rightarrow O(^{3}P) + HONO (\tilde{a}^{3}A'')$$
  

$$\Delta H^{\circ} = 550 \text{ kJ/mol}$$
 (4)

$$\rightarrow O(^{3}P) + HONO (\tilde{X}^{1}A')$$
  

$$\Delta H^{\circ} = 300 \text{ kJ/mol}$$
 (5)

The photolysis at 193 nm is dominated by the formation of O+HONO with  $\phi \sim 0.54$  [7] and  $\sim 0.13$  [7] for channels (3) and (4), respectively. A

photodissociation channels, studied by photofragment translational energy spectroscopy [6,7] and LIF [8–10] and REMPI-TOF methods [10], were found to be:

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minor decay path ( $\phi \le 0.075$ ) has tentatively been assigned to (5) [10]. On the other hand, the formation of OH + NO<sub>2</sub> which shows bimodal translational energy [6,7] and rotational state distributions [10] for the OH fragments has been attributed to channel (1) with  $\phi \sim 0.22$  and channel (2) with  $\phi \sim 0.10$  [7].

In view of the potential importance of the photodissociation of HNO<sub>3</sub> attached to ice particles in the stratosphere [11–14], we explored this problem in our laboratory by generating water clusters in a supersonic expansion and attaching HNO<sub>3</sub> to them by a pickup method [15,16]. The subsequent photodissociation was carried out at 193 nm with detection of the nascent OH ( $X^2\Pi$ ) fragments by LIF. These experiments were complemented by corresponding experiments with argon and methanol clusters. The broad cluster size distributions were estimated to have an average size in terms of cluster molecules of  $n \sim 200-800$ depending on the compound. The rotational state distribution and the average translational energy of the OH fragments emerging from the clustered HNO<sub>3</sub> molecules after photodissociation were measured and compared to those previously obtained from HNO<sub>3</sub> monomer photolysis.

# 2. Experiment

The apparatus used for this work is described elsewhere [17,18]. A pulsed molecular beam of

HNO<sub>3</sub> seeded in a noble gas was generated with a specially designed corrosion resistant valve driven by a piezoelectric translator [19]. Fuming nitric acid (Merck, labeled 100%) was degassed to remove air and traces of NO<sub>2</sub> until it was colorless. In order to generate the cluster beams, two kind of cluster preparations were employed. In the first case the components were premixed and then subjected to supersonic expansion, in the second case a pickup technique was utilized.

Following premixing of 1% HNO<sub>3</sub> in Ar at a backing pressure of  $\leq 2.4$  bar or 1% HNO<sub>3</sub> with 250 mbar CH<sub>3</sub>OH in ≤ 2 bar He, expansion was conducted through a pulsed valve equipped with a conical nozzle (aperture d = 1 mm, length 10 mm, opening angle  $2\theta = 20^{\circ}$ ) which favors cluster formation. In contrast, for cluster production by pickup the setup shown in Fig. 1 was used. First a pulsed supersonic cluster beam of either pure argon, methanol or water was generated which then was crossed by a pulsed HNO<sub>3</sub> monomer beam at right angle. The HNO<sub>3</sub>-doped clusters formed in the interaction region were extracted through a skimmer into a second chamber with an additional differential pump stage. For production of the pure cluster beams we used the valve described above and 2 bar Ar, or in the case of methanol and water, 2 bar He which was guided into a flask filled with CH<sub>3</sub>OH or H<sub>2</sub>O at a temperature of 35 and 60 °C, respectively. The outgoing gas mixtures, possessing partial pressures of  $p(CH_3OH) = 300$ mbar or  $p(H_2O) = 250$  mbar, were then passed

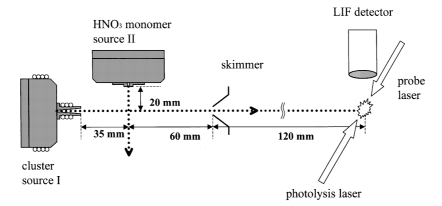


Fig. 1. Experimental setup for cluster production by the pickup method and cluster photodissociation with LIF fragment detection. The conditions for the pulsed cluster (I) and monomer (II) beams are given in the text.

through a heated tube to the pulsed valve. The latter was also heated to prevent condensation of CH<sub>3</sub>OH or H<sub>2</sub>O. On the other hand, the 'soft' HNO<sub>3</sub> monomer beam was produced by expanding a mixture of 4% HNO<sub>3</sub> in 300 mbar He through a pinhole aperture (d = 0.5 mm) attached to the piezoelectric pulsed valve. The carrier gas He will not form clusters and the low stagnation pressure prevents HNO<sub>3</sub> from clustering. The pulse still provides a well localized, relatively highly concentrated HNO<sub>3</sub> environment. In order to optimize the mixed cluster formation the distance between nozzle I and the crossing of the HNO<sub>3</sub> monomer beam (Fig. 1) could be adjusted (to usually about 35 mm) as well as the time delay between the opening of the pulsed valves I and II.

In our work there is no direct way to measure the cluster size distributions but a rough estimate can be given based on the work of Buck and Krohne [20] and Hagena [21]. For a pure argon expansion under the conditions used to produce  $Ar_nHNO_3$  clusters, we obtain an average size in terms of  $\overline{n} \sim 200$ –400. In the cases of  $(CH_3OH)_nHNO_3$  and  $(H_2O)_nHNO_3$   $\overline{n}$  is expected to be in the range of  $\sim 300$ –600 and  $\sim 400$ –800, respectively. We expect the mixed clusters to contain only one (pickup) or a few (premixing)  $HNO_3$  molecules; associations of the latter were not found to be important below about 8%  $HNO_3$  concentration.

Photolysis at 193 nm was carried out with an ArF excimer laser (Lambda Physik EMG 101 MSC). The softly focussed laser beam crossed the molecular beam at right angles (Fig. 1). A counterpropagating probe laser beam recorded the nascent OH photofragments on the  $A^2\Sigma^+ \leftarrow X^2\Pi$ system (vibrational bands with  $\Delta v = 0$ ) by the LIF method. The probe laser (Lambda Physik dye laser FL2002) was pumped by an XeCl excimer laser (Lambda Physik LPX200) operated with Sulforhodamine B dye. After frequency doubling the linewidth of the laser light was 0.4 cm<sup>-1</sup> between 306 and 311 nm. Fluorescence from the OH fragments was measured at right angles to the plane of the molecular and laser beams by a Hamamatsu R928 photomultiplier tube equipped with an appropriate cutoff filter. The signal was fed to a digital boxcar integrator (Stanford Research SR 250), and photodiodes monitored the photolysis

and probe laser intensities to correct for shot to shot fluctuations. Since the LIF signal was found to be linear when the dissociation laser was operated in the range 3–16 mJ/cm<sup>2</sup>, we used 5 mJ/cm<sup>2</sup> to safely avoid multiphoton processes in dissociation. The probe laser power density was kept at 0.1 mJ/cm<sup>2</sup> thus avoiding saturation effects. The Doppler profile measurements were carried out by operating the probe laser with an intracavity etalon which provided  $\Delta v = 0.08 \pm 0.01$  cm<sup>-1</sup>.

#### 3. Results

#### 3.1. Clusters by premixing

The unpolarized LIF spectra of OH ( $A^2\Sigma^+ \leftarrow$  $X^2\Pi$ ,  $\Delta v = 0$ ) fragments upon 193 nm photodissociation of Ar, HNO<sub>3</sub> and (CH<sub>3</sub>OH), HNO<sub>3</sub> clusters generated by premixing of the components are shown in Fig. 2. No signal was detected when the wavelength of the probe laser was tuned to the position of the OH (1,1) band indicating a negligible population of OH (v = 1) and very probably also of higher vibrational states. From the LIF spectra we obtained the rotational state populations of the OH fragments given in Fig. 3 using standard procedures [22]. A comparison between the LIF spectra of Fig. 2a,b to that recorded from HNO<sub>3</sub> monomer dissociation [10] shows that the intensity of high rotational lines is considerably smaller in the former indicating rotational state relaxation in cluster dissociation.

The rotational state distribution of OH (v = 0) from Ar<sub>n</sub>HNO<sub>3</sub> clusters (Fig. 3a), showing a maximum at N = 2 and extending to N = 10, is well described by a Boltzmann distribution (solid line) with a temperature  $T_{\rm rot} = 400 \pm 100$  K. That of the (CH<sub>3</sub>OH)<sub>n</sub>HNO<sub>3</sub> clusters has a maximum at N = 1 followed by a population decrease to N = 8 and fits a Boltzmann distribution with a temperature  $T_{\rm rot} = 250 \pm 150$  K (solid line in Fig. 3b). The resulting mean rotational energy  $\langle E_{\rm rot} \rangle = \sum_J P(J)E_{\rm rot}(J)$ , where P(J) is the state distribution and  $E_{\rm rot}$  the energy of a given rotational state, was found to be 290 cm<sup>-1</sup> (3.5 kJ/mol) and 175 cm<sup>-1</sup> (2.1 kJ/mol) for the Ar<sub>n</sub>HNO<sub>3</sub> and (CH<sub>3</sub>OH)<sub>n</sub> HNO<sub>3</sub> clusters, respectively.

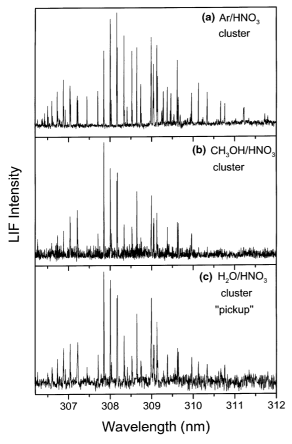


Fig. 2. Unpolarized LIF spectra of nascent OH  $(A^2\Sigma^+ \leftarrow X^2\Pi)$  (0,0) fragments after 193 nm photolysis of HNO<sub>3</sub> (a) in argon clusters (premixed), (b) in methanol clusters (premixed) and (c) in water clusters (pickup).

Further analysis of the spectra reveal that the spin-orbit states  $\Pi_{3/2}$  and  $\Pi_{1/2}$  spaced by 126 cm<sup>-1</sup> [23] are equally populated for both cluster systems which is also the case for the almost degenerate lambda doublet states  $\Pi(A')$  and  $\Pi(A'')$  as obtained from the intensities of the R and Q branches. The translational energy of the OH fragment was estimated from the Doppler width of the spectral lines N=2 and N=6, depicted as inserts in Fig. 3. After deconvolution with the probe laser profile, we found the Doppler width to be  $0.11\pm0.01~{\rm cm^{-1}}$  for the argon clusters and  $0.09\pm0.01~{\rm cm^{-1}}$  for the methanol clusters which corresponds to a mean translational energy  $E_{\rm trans}({\rm OH})=540~{\rm cm^{-1}}$  (6.5 kJ/mol or  $T_{\rm trans}\sim500$ 

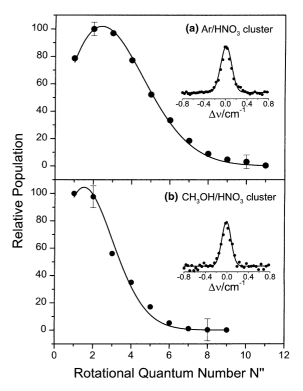


Fig. 3. Rotational state distributions of nascent OH  $(X^2\Pi, v=0)$  fragments from 193 nm photolysis of HNO<sub>3</sub> in (a) argon and (b) methanol clusters (premixed). The solid lines represent the best fits for a Boltzmann distribution with  $T_{\rm rot}=400$  and 250 K, respectively. The inserts show the Doppler profiles for the  $P_1$  (2) rotational state.

K) and 350 cm<sup>-1</sup> (4.2 kJ/mol or  $T_{\rm trans} \sim 350$  K), respectively.

In order to study the formation of the  $Ar_nHNO_3$  and  $(CH_3OH)_nHNO_3$  clusters, we monitored the intensity of a high rotational line of the OH fragment, which is prominent in the monomer and absent in the cluster dissociation, as a function of the stagnation pressure  $p_0$  [18,24]. Thus we selected the OH (v = 0)  $Q_1(13)$  rotational line at the probe wavelength of 311.02 nm and obtained the results shown in Fig. 4. Changing  $p_0$ from 0.3 to 2.4 bar of a mixture of 1% HNO<sub>3</sub> in Ar carrier gas results in the intensity of this rotational line (Fig. 4a) decreasing abruptly at  $\sim 0.8$  bar, and assuming a constant minimum value at  $\sim 1.3$  bar. This behavior is typical for a 'phase transition' associated with cluster formation [18,25,26]. A similar behavior (Fig. 4b) is observed for the

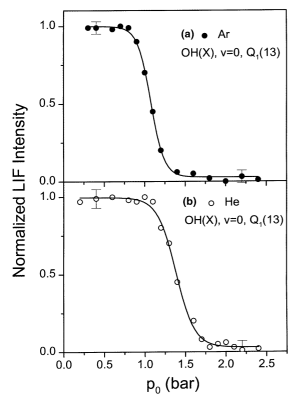


Fig. 4. Cluster formation ('phase transition') as a function of the stagnation pressure  $p_0$  monitored by the intensity of the  $\mathrm{OH}(v=0)$ ,  $Q_1(13)$  rotational transition which is essentially absent following cluster photolysis. The expansion conditions were (a) 1% HNO<sub>3</sub> in argon and (b) 1% HNO<sub>3</sub> with 250 mbar CH<sub>3</sub> OH in helium.

 $(CH_3OH)_n$ HNO<sub>3</sub> system using a mixture of 1% HNO<sub>3</sub> in He and keeping the partial pressure of CH<sub>3</sub>OH constant at 200 mbar. The phase transition occurs here between ~1.1 and 1.7 bar. Thus the stagnation pressures applied in the above measurements guarantee a high cluster production.

#### 3.2. Clusters by pickup

The formation of water clusters doped with HNO<sub>3</sub> molecules could not be obtained by premixing the components and co-expansion through the same source. Reactions of HNO<sub>3</sub> with H<sub>2</sub>O prevented application of this simple method. We therefore had to use the pickup arrangement

shown in Fig. 1 where the initial preparation of very cold water clusters before the pickup of HNO<sub>3</sub> reduced the reactivity by hindered restructuring. This method in combination with LIF probing of the OH fragments was also applied for Ar, HNO<sub>3</sub> and (CH<sub>3</sub>OH), HNO<sub>3</sub> allowing us to directly compare the results of the pickup and premixing methods. Fig. 2(c) depicts the unpolarized LIF spectra recorded for OH ( $A^2\Sigma^+ \leftarrow$  $X^{2}\Pi$ ,  $\Delta v = 0$ ) from dissociation of the water cluster. Again no evidence for the population of excited vibrational states was found. The rotational state populations of OH obtained from the line strengths in Fig. 2 are displayed in Fig. 5. The results could be well fitted by a Boltzmann distribution (solid line) with a rotational temperature of 600, 330, and 300 K for the  $Ar_nHNO_3$ ,

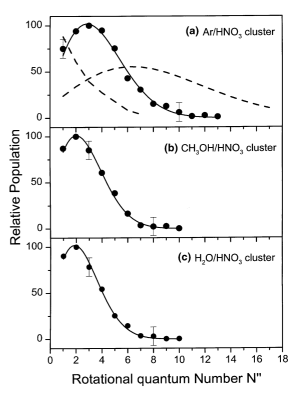


Fig. 5. Rotational state distributions of OH ( $X^2\Pi$ , v=0) from 193 nm photolysis of HNO<sub>3</sub> (a) in argon, (b) methanol and (c) water clusters produced by pickup. The solid lines represent best fits by Boltzmann distributions with  $T_{\rm rot}=600$ , 330 and 300 K, respectively. The broken lines in (a) show the two rotational distributions from HNO<sub>3</sub> monomer dissociation [10].

Reaction (2)         Reaction (1)         Ar,HNO <sub>3</sub> (CH <sub>3</sub> OH),HNO <sub>3</sub> Ar,HNO <sub>3</sub> (CH <sub>3</sub> OH),HNO <sub>3</sub> Non-thermal         250         400         250         600         330           120         1700         290         175         445         260           9700         4000         540         350         830         -           1 3+0 1         1 3+0 1         1 0+0 1         1 0+0 2         -         -	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Cluster by pickup		
Non-thermal         250         400         250         600         330           120         1700         290         175         445         260           9700         4000         540         350         830         -           13+01         13+01         10+01         10+02         -         -	Non-thermal 2500 400 2 120 1700 290 1 9700 4000 540 3 1.3±0.1 1.0±0.1 1		(CH <sub>3</sub> OH),,HNO <sub>3</sub>	$(H_2O)_nHNC$
120     1700     290     175     445     260       9700     4000     540     350     830     -       13+01     13+01     10+01     10+02     -     -	120 1700 290 1 9700 4000 540 3 1.3±0.1 1.3±0.1 1.0±0.1 1	009	330	300
9700 4000 540 350 13+01 13+01 10+01	9700 4000 540 3 1.3±0.1 1.3±0.1 1.0±0.1 1		260	220
13+01 13+01 10+01	$1.3\pm0.1$ $1.3\pm0.1$ $1.0\pm0.1$ 1	830	1	1
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(CH<sub>3</sub>OH)<sub>n</sub>HNO<sub>3</sub> and (H<sub>2</sub>O)<sub>n</sub>HNO<sub>3</sub> clusters, respectively. The mean rotational energy  $\langle E_{\rm rot} \rangle$  was found to be 445, 260 and 220 cm<sup>-1</sup> (i.e, 5.3, 3.1 and 2.6 kJ/mol) for the Ar<sub>n</sub>HNO<sub>3</sub>, (CH<sub>3</sub>OH)<sub>n</sub>HNO<sub>3</sub> and (H<sub>2</sub>O)<sub>n</sub>HNO<sub>3</sub> clusters, respectively. In the case of the argon cluster the Doppler profile of OH rotational lines were also examined. The width was found to be  $0.14 \pm 0.02$  cm<sup>-1</sup> which corresponds to a mean translational energy  $E_{\rm trans}(OH) = 830$  cm<sup>-1</sup>( $\sim 10$  kJ/mol or  $T_{\rm trans} \sim 800$  K). These findings are summarized in Table 1 and compared with previously reported results from the photodissociation of HNO<sub>3</sub> monomer [10].

#### 4. Discussion

The rich photochemistry of the nitric acid monomer after excitation at 193 nm is summarized by the five primary dissociation channels given in Section 1. Two of these decay processes lead to the products OH + NO<sub>2</sub> and three to O + HONO involving different electronic states. Because we exclusively probed the OH fragment in the present cluster study, we are confined to investigating channels (1) and (2). These two OH channels were found to differ not only in their translational energy distributions [6,7] but also in their rotational state distributions [10]. Thus the OH fragments with the electronically excited and stable counterfragments  $NO_2(1^2B_2)$  of channel (2) possess a non-statistical rotational state distribution with low excitation (N < 7) while OH with  $NO_2(X^2A_1)$ in the ground state (1) shows a broad Boltzmann distribution (N < 18) with  $T_{\text{rot}} = 2500$  K, as reproduced in Fig. 5a by broken lines.

HNO<sub>3</sub> photodissociation in a cluster environment reveals for both preparation modes a single and statistical OH rotational distribution described by a rotational temperature between 250 and 600 K (Figs. 3 and 5 and Table 1) depending on the nature of the cluster. Again only OH fragments in the vibrational ground state are formed, but in contrast to the monomer dissociation [9,10], the lambda doublet states are equally populated. In general the rotational as well as the translational excitation of OH are strongly relaxed in our cluster dissociation as compared to that of

the monomer [6,7,10,27]. The clusters prepared by pickup show a higher  $T_{\rm rot}$  than those generated by premixing (Ar<sub>n</sub>HNO<sub>3</sub>: 600 and 400 K, (CH<sub>3</sub>OH)<sub>n</sub>HNO<sub>3</sub>: 330 and 250 K, respectively). Since the stagnation pressure is lower and the chamber pressure higher for beam II than for beam I (Fig. 1) it is conceivable that this effect is caused by a slight warm up of the clusters after pickup thus assuming the interaction between the HNO<sub>3</sub> molecule and the cluster to be temperature dependent. Alternatively, it could arise from a less constrained surface site of HNO3 after pickup than that associated with the premixing preparation. The photodissociation of strongly 'bonded' and even partly or fully caged HNO3 molecules has been found to lead to substantially relaxed and often thermalized fragments, particularly with respect to the rotational and translational degrees of freedom [17]. In this context it might not be surprising that among the pickup  $(H_2O)_nHNO_3$  yields the lowest  $T_{rot}$  (Table 1). The binding forces between HNO<sub>3</sub> molecules and the water cluster, being dipole-dipole interaction and hydrogen bonding, are expected to be the strongest among the present cluster systems.

While decay paths (1) and (2) proceeding to OH products show different rotational state distributions in the monomer, the single thermalized distribution observed in all our cluster systems might suggest that only one OH channel, probably decay path (1) with the lowest ΔH° value, is important in cluster dissociation. However, the broad cluster size distribution inherent in our preparation techniques gives rise to different environments for the HNO<sub>3</sub> molecules and thus to a distribution of interaction strengths between HNO<sub>3</sub> and the cluster. A distinction of different rotational state distribution of the ejected photoproduct OH may under these condition no longer be feasible.

### 5. Conclusion

The photochemistry of the HNO<sub>3</sub> molecule, an important source of OH radicals and NO<sub>2</sub> in the stratosphere, has been well studied at 193 nm and five reaction channels have been identified. The

main goal of the present investigation was to extend this work to HNO<sub>3</sub> in or on water clusters, a system relevant to atmospheric chemistry. For the first time we produced these clusters using a pickup source and photolysed the mixed clusters at 193 nm detecting the OH ( $X^2\Pi$ ) fragments by LIF. Complementary investigations were carried out with argon and methanol clusters, where also the preparation method by premixing of the components was applied. In contrast to the monomer, the cluster photodissociation was found to generate OH in only a single and thermalized rotational state distribution and the rotational as well as translational fragment excitation is greatly relaxed. Thus the reduced internal energy of this fragment, which is also expected for its counterfragment NO2, makes consecutive reactions of these species involving an activation energy less probable.

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