Time-resolved studies of NO2 photoinitiated unimolecular decomposition: step-like variation of $k_{\text{uni}}(E)$

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Time-resolved, subpicosecond-resolution measurements of NO2 photoinitiated unimolecular decay rates are reported for jet-cooled samples in the vicinity of the dissociation threshold. The molecules are excited by 385-400 nm tunable subpicosecond pulses to the $^3B_2$ electronic state which is very strongly mixed with the $^2A_1$ ground electronic state. Subsequent decomposition is probed by a 226 nm subpicosecond pulse which excites LIF in the NO product. When changing the amount of energy in excess of the dissociation threshold, a step-like increase of the reaction rate versus energy is observed.

1. Introduction

The photoinitiated unimolecular reaction of NO2 is presently the only documented case in which a stable triatomic can be excited optically via an allowed one-photon transition to levels that lie above the reaction threshold and that have predominantly ground electronic state character. It is known that near-infrared, visible, and ultraviolet photons excite NO2 into a region of strong mixing between the $^2A_1$ ground state and the $^3B_2$ electronically excited state \[1-12\]. Moreover, excitation between 250 and 398 nm results in NO2 decomposition to NO and O in their electronic ground states, i.e., $^3\Pi_O$ and $^3\Pi_J$, respectively, where $\Omega$ and $J$ label fine structure levels \[5-10\]. Energies above reaction threshold, which is determined to be $D_0 = 25130.6 \text{ cm}^{-1}$ \[7,13\], are spectroscopically accessible over a broad, continuously tunable range, and the NO($^X\Pi$) molecular fragment is particularly amenable to sensitive, state-selective detection.

Recent spectroscopic studies have successfully assigned NO2 vibrational levels of the ground electronic state up to 10000 cm$^{-1}$ \[3\], and the density of vibronic levels on the strongly coupled $^2A_1$ and $^3B_2$ electronic surfaces has been experimentally estimated in the region 16500-18500 cm$^{-1}$ \[4\]. Moreover, spectra of extremely cold ($\approx 1$ K) expansion-cooled samples have been taken in the region just below $D_0$ \[9\]. Such data provide quantitative experimental assessments of vibronic and rovibronic level densities near $D_0$, facilitating comparisons between theoretical predictions and measured reaction rates. These features make NO2 unimolecular decay rates very attractive for scrupulous experimental examinations.

Several groups have examined NO2 photofragmentation in detail, reporting center-of-mass (CM) kinetic energy distributions obtained from time-of-flight (TOF) analyses \[14\], and state selective product detection of O($^3P_J$) \[8,9\] as well as NO($^X\Pi$) \[7,10\]. In a pioneering study by Troe and co-workers, high pressure samples containing NO2 were photolyzed at a number of different ultraviolet wavelengths, yielding the variation of decomposition lifetime versus the photon energy of the photolyzing light \[15\]. These results were analyzed using reaction rate theories, and in fact have been central to the development and refinement of such theories.
On the negative side, interpretations of experimental results on NO$_2$ photochemistry are not always straightforward, due to the molecule's complex electronic structure. There is a conical intersection between the $^2$A$_1$ ground state and the lowest electronically excited $^2$B$_2$ state [11,12] and the use of a single effective potential, in dynamics studies for example, cannot be justified a priori.

Difficulties notwithstanding, NO$_2$ is a scientifically and technologically important species and although several measurements of its photochemistry have been carried out in the region just above $D_0$, there has been no report to date of direct measurements of decomposition lifetimes. Thus, we have carried out time-resolved, subpicosecond resolution measurements of photoinitiated NO$_2$ reaction rates using expansion-cooled samples. These are the first direct measurements of the reaction rates for this system. The rates show an essentially monotonic increase with energy, and are consistent with the step-like variation of $k_{\text{uni}}$ versus $E$ predicted by transition state theories. In this communication, we report the essence of these measurements. These results constitute an important complement to the detailed unimolecular reaction rate studies carried out for NCNO [19] and CH$_2$CO [20].

### 2. Experimental

The experimental arrangement is shown schematically in fig. 1. The main parts are: (i) a dual-jet mode-locked dye laser (Coherent 702-1), synchronously pumped by the frequency-doubled output of (ii) an actively mode-locked Nd:YAG laser (Spectron SL 903), (iii) two dye amplifiers pumped by (iv) a regenerative Nd:YAG amplifier (Continuum RGA 69-10), and (v) a vacuum chamber equipped with a pulsed nozzle, quadrupole mass spectrometer and accessories for monitoring LIF. The mode-locked...
Nd: YAG laser with a CAMAC ML 4000 mode-locker generates a train of 100 ps pulses at 76 MHz; typical average power is 20 W at 1064 nm. A 5% portion of this radiation is used for seeding the regenerative amplifier, the remaining 95% is frequency doubled in a 12 mm LBO crystal producing about 2 W of 532 nm radiation for dye laser pumping.

Table 1 presents the combinations of gain and saturable absorber dyes used in the dual-jet dye oscillator and first amplifier. Using a single-plate birefringent filter (BRF) as a tuning element, 250–500 fs pulses tunable from 575 to 645 nm with \( \approx 40–70 \) cm\(^{-1} \) spectral bandwidth are obtained from the mode-locked dye laser. The shortest pulses are observed on the red side of the covered spectral range. Replacing the single-plate BRF by a double-plate BRF results in the bandwidth narrowing to \( \approx 30\% \) of its original value. This narrowing is accompanied by broadening of the pulsewidth. Although 300 mW is obtained at the exit of the dual-jet dye laser, only half of this is used for successive amplification in order to minimize pulse broadening.

The output pulse train from the dye laser is amplified in a three-stage amplifier (Continuum PTA-60). The amplifier is longitudinally pumped at 10 Hz by the second harmonic of the RGA which typically provides 30 mJ, 80 ps pulses at 532 nm. There is a solid state saturable absorber (table 1) positioned between the first and the second amplification stages for reducing pulse broadening. The amplified dye laser pulse, which reaches 2 mJ without noticeable broadening, is divided into two beams by a broadband beam-splitter. The first beam, with \( \approx 80\% \) of the energy, is spatially matched using a telescope and then combined on a dichroic mirror with a 1064 nm, 50 mJ beam from the RGA. These beams are mixed in a BBO crystal producing \( \approx 200 \) \( \mu \)J of sum frequency radiation tunable from 373 to 402 nm. No significant broadening is observed in this near-UV pulse, as ascertained from its cross-correlation with the dye laser fundamental. The cross-correlation is measured by sum-frequency generation in a 150 \( \mu \)m BBO crystal. The near-UV pulse is used to excite NO\(_2\) molecules over the dissociation threshold causing their unimolecular decomposition.

The remaining 20% of the amplified dye laser fundamental is focused by a 10 cm f.l. lens into a \( \text{D}_2\text{O} \) cell producing a white continuum. To avoid boiling, the heavy water is constantly stirred by a magnetic stirrer. Vertically polarized 452 nm radiation (\( \approx 1 \) nJ, 6 nm bandwidth) is selected from the continuum by a Corning 5-57 glass filter followed by a Glan-Taylor polarizer and an interference filter. These components are positioned behind an 8 cm f.l. lens which recollimates the light exiting the \( \text{D}_2\text{O} \) cell. Autocorrelation traces of the filtered continuum show \( \approx 50\% \) temporal broadening of the 452 nm pulses as compared to the dye laser fundamental. We believe that this broadening is due to the interference filter. The 452 nm radiation is amplified in a second three-stage dye amplifier transversely pumped by 45 mJ, 355 nm pulses from the RGA. Coumarin 460/MeOH dye solutions are used in the first and final stages. In order to minimize ASE, a different dye, coumarin 440/MeOH, is used in the second stage. The dye concentrations are adjusted in each stage to satisfy

Table 1
Gain and saturable absorber (S/A) dyes used in the dual-jet dye oscillator and the first dye amplifier.

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Oscillator dyes</th>
<th>Amplifier</th>
<th>solid-state saturable absorber a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>575–595</td>
<td>Rh590(Cl)</td>
<td>DODCI/DQOCI</td>
<td>Rh610</td>
</tr>
<tr>
<td>595–605</td>
<td>Rh590(Cl)</td>
<td>DODCI/DQTCI</td>
<td>kiton red (( \text{H}_2\text{O} )) or SRh640 (MeOH)</td>
</tr>
<tr>
<td>605–618</td>
<td>kiton red</td>
<td>DQTCI</td>
<td>DCM (MeOH) or SRh640 (MeOH)</td>
</tr>
<tr>
<td>618–645</td>
<td>kiton red/SRh640</td>
<td>DQTCI</td>
<td>DCM (MeOH)</td>
</tr>
</tbody>
</table>

a) Solid state saturable absorber: long-pass, Schott-glass filter located at the beam focal point between the first and second stages of amplification to reduce pulse broadening.
two contradictory criteria: maximum power and minimum pulse reshaping due to saturation. The amplified 452 nm beam (≈250 μJ) is frequency doubled in a 150 μm long BBO crystal. The resulting 226 nm laser pulse is used to probe NO via LIF using the γ system.

After passing through a computer-controlled delay stage (Aerotech ATS100) the pump beam is combined with the probe radiation on a dichroic mirror and collinearly focused by a 30 cm f.l. lens into a vacuum chamber pumped to 10⁻⁶ Torr. The chamber is equipped with a pulsed nozzle (250 μs duration, 500 μm dia. orifice), a quadrupole mass spectrometer for monitoring the beam intensity, a PMT (Hamamatsu 943 02) and collecting optics for LIF measurements. The collinear laser beams intersect the supersonic jet ≈40 nozzle diameters downstream from the nozzle. A 3% NO₂/He gas mixture is prepared in situ by blowing 1 atm of He over an NO₂ sample frozen at −28°C. A 760 Torr stagnation pressure is kept throughout the measurements. The NO₂ sample (Matheson, 99.5%) is purified prior to each set of experiments by bubbling O₂ through the liquid at 0°C followed by several freeze–pump–thaw cycles. The prepared sample is kept under 1 atm of oxygen between experimental runs in order to shift the NO₂→NO + ½O₂ equilibrium towards NO₂. To further minimize NO₂ decomposition, the nozzle interior is coated with Teflon and the portion of the gas system between the NO₂ bath and the nozzle is kept at −20°C.

The NO LIF signal is preamplified, digitized and collected in an IBM AT computer simultaneously with both pump and probe pulse energies and a pump–probe cross-correlation signal. The cross correlation is obtained by difference frequency generation in a 100 μm BBO crystal which is positioned shortly after the chamber. The pump, probe and difference frequency radiations are separated in a 60° quartz prism and detected by two UV-grade photodiodes and a PMT, respectively.

3. Results

When the pump radiation wavelength is below 397.9 nm, NO fragments deriving from the photodecomposition of expansion-cooled NO₂ are observed by LIF following the pump pulse. Results obtained with room temperature samples will be described elsewhere. No decomposition is detected at λₚₚₜₜ > 398 nm. This observation is in good agreement with the reported threshold for NO₂ decomposition, 25130.6 cm⁻¹ [7,13]. Typical experimental scans of NO LIF signal versus the delay between the pump and probe pulses are shown in fig. 2. There are typically 300–500 data points in each trace, and each of the points represents an average over 20 laser shots and is normalized to the pump and probe energies. Also shown in the figure are the corresponding pump–probe cross correlations which have been recorded by difference-frequency generation simultaneously with obtaining the LIF signal. We have verified experimentally that the NO LIF signal is a linear function of both pump and probe laser energies and that the experimentally determined decay rates (see below) are independent of the pump energy. Therefore, the observed decomposition of NO₂ results from a one-photon process.

The experimental time dependencies are fit assuming single exponential NO production after the excitation event:

![Fig. 2. NO LIF measured as a function of the pump–probe delay time for the indicated excitation conditions. Also shown are the corresponding cross correlations between the pump and probe pulses. The experimental points are fitted assuming single exponential decay as described in the text.](image)
\[ S(t) \propto \int_{-\infty}^{\infty} ds R(s) \left\{ 1 - \exp[-k_{\text{uni}}(t-s)] \right\}, \]  \( (1) \)

where

\[ R(s) = \int_{-\infty}^{\infty} dt I_{\text{pump}}(t) I_{\text{probe}}(t-s), \]  \( (2) \)

\( R(s) \) is the experimentally measured cross correlation between the pump and probe pulses and \( k_{\text{uni}} \) is the unimolecular reaction rate under consideration. Fits of the experimental data by the above convolution are shown as solid lines in fig. 2. As seen from the figure, the assumption of a single exponential decay provides good agreement with the data. The small discrepancies between the experimental results and the simulations are not serious. We note that the spectral bandwidth of the pump pulse, i.e., 20–80 cm\(^{-1}\), results in excitation of a number of molecular eigenstates, as discussed in the next section. Since the state-to-state dissociation rate is expected to fluctuate considerably \([21]\), modest deviations from single exponential decay are not surprising.

The NO\(_2\) dissociation rates obtained from the experimental delay time dependencies described above are presented in fig. 3. Every point in the figure is an average of 8 to 16 rates, each obtained from independent scans at specific excitation wavelengths. Statistical uncertainties, i.e., standard deviations, are small, generally lying within the circles shown in fig. 3. It should be noted that independent measurements have been carried out for several excitation wavelengths near the reaction threshold with either one or two birefringent filters in the mode-locked dye oscillator. As discussed in the previous section, the single-plate BRF provides better time resolution, i.e., 500 to 900 fs cross correlation fwhm, with lower spectral resolution of \( \approx 40–80 \) cm\(^{-1}\), as compared to the double-plate BRF, which provides \( \approx 1.5 \) ps cross correlations and 20–30 cm\(^{-1}\) spectral resolution. The experimental points obtained with the single- and double-plate BRFs are presented by closed and open symbols, respectively.

One aspect of the results presented in fig. 3 is noteworthy. Namely, \( k_{\text{uni}} \) versus \( h\nu_{\text{pump}} \) is not a smooth ascending function, as is often the case for unimolecular decompositions. For example, the rate rises abruptly to \( \approx 1.6 \times 10^{11} \) s\(^{-1}\) at threshold and then rises only moderately in the range 0–100 cm\(^{-1}\). The rate exhibits another steep rise to \( \approx 4 \times 10^{11} \) s\(^{-1}\) for excess energies near 100 cm\(^{-1}\) and then does not change in the range 120–210 cm\(^{-1}\). At 220 cm\(^{-1}\) the rate climbs to \( \approx 5.5 \times 10^{11} \) s\(^{-1}\) and then flattens again in the range 240–350 cm\(^{-1}\). These step-like rises cannot be dismissed as a S/N problem, i.e. each data point is the result of considerable signal averaging and is reproducible. In similar experiments carried out in our laboratory with room temperature samples \([22]\), a smooth variation of \( k_{\text{uni}} \) versus \( E \) is observed. We also note that the rate on the first step is roughly 1/2 and 1/3 of the rates on the second and third steps, respectively.

### 4. Discussion

The simplest form of microcanonical transition state theory uses eq. (3) to compute the unimolecular decomposition rate, \( k_{\text{uni}}(E) \)

\[ k_{\text{uni}}(E) = \frac{N^*(E-E_0)}{\hbar \rho(E)}, \]  \( (3) \)

where \( N^*(E-E_0) \) is the number of accessible states of the transition state at energy \( E \) without inclusion of the reaction coordinate, \( E_0 \) is the reaction thres-
old, which in the present case is taken to be the same as $D_o$ since rotation is minimal and there is no reverse barrier, and $\rho(E)$ is the density of states of the parent molecule. $N^*(E-E_0)$ can be estimated at a single transition state in the simplest applications of eq. (3), or at transition states obtained variationally to satisfy a minimum flux criterion.

The view embodied in eq. (3) can predict rates with uncanny success. $N^*(E-E_0)$ rises in steps, taking on the value unity just above threshold when only one internal level of the transition state is accessible. With increasing energy, $N^*$ changes from 1 to 2 when two channels become available, and so on. Since $\rho(E)$ changes more slowly than $N^*(E)$, eq. (3) predicts a step-like increase of $k_{uni}$ with energy. For large molecules having many internal degrees of freedom the step-like behavior will generally be too fine-grained to be observed except in the most sophisticated experiments, while for small molecules it may be more easily observed. It is tempting to consider the possibility that the measured $k_{uni}$ values shown in fig. 3 can be rationalized in terms of eq. (3): For example, is the step-like behavior a manifestation of $N^*+1$, 2, etc.?

Although reaction via $S_0$ character is assured, NO$_2$ does not fit comfortably into the niche of standard unimolecular decomposition models. It has a complex electronic structure that derives from the conical intersection between the $^2A_1$ ground and $^2B_2$ excited states; it is a triatomic with a modest density of states and the fact that our measurements do not include a large number of levels, statistical fluctuations in the rates may occur [21]. The above considerations notwithstanding, it is still interesting to compare $k_{uni}(E)$ from eq. (3) to the data.

Fortunately, it is possible to obtain experimentally based estimates of vibronic and rovibronic state densities for NO$_2$ in the region just below $D_o$. For example, Delon et al. recently examined NO$_2$ spectra in great detail in the region 16500–18500 cm$^{-1}$. They observed 159 vibronic $B_2$-symmetry levels of the strongly mixed $^3B_2$ and $^2A_1$ electronic states in this interval [4]. By extrapolating from observed levels in the region 0–10000 cm$^{-1}$ [3], they deduced that there are most likely 210 levels in the 16500–18500 cm$^{-1}$ interval. Taking this level density of $\approx 0.1$ per cm$^{-1}$, and extrapolating to $D_o$ by assuming that the density increases as $(E+E_{00})^2$ [23], where $E_{00}$=1844 cm$^{-1}$ is the NO$_2$ zero-point vibrational energy, yields a level density of $\approx 0.2$ per cm$^{-1}$ near $D_o$. Substituting this into eq. (3) with $N^*=1$ yields 1.5×10$^{11}$ s$^{-1}$ for the threshold rate. Thus, if the vibronic level density measured by Delon et al. is extrapolated to $D_o$, the step-like $k_{uni}$ variation observed experimentally can be assigned to the opening of channels at the transition state.

A more precise evaluation of the near threshold density of NO$_2$ rovibronic states may be obtained from recently reported LIF spectra which have been recorded for 1 K expansion cooled samples having 98% of the NO$_2$ molecules in the lowest rotational level. Under these experimental conditions, Miyawaki et al. observe more than 60 transitions in the 25 cm$^{-1}$ region immediately below $D_o$ [9]. Since $J=1/2$ for the lowest rotational level, the one-photon electric dipole excitation process carries molecules to $J=1/2$ and 3/2, each of which contributes to the observed density of spectral lines. It should be mentioned that the "observed" levels are obtained from visual inspection of a published spectrum and, as a result of limited resolution, there may be additional levels that do not show up; also, some of the small peaks may originate from levels other than $J=1/2$. Other groups have also reported highly structured spectra below $D_o$ [7] which are in general agreement with the Miyawaki et al. results. Above $D_o$, lifetime broadening results in overlap of zero-order levels, making it impractical to recover level densities from absorption spectra.

It is clear from the spectroscopic results of Miyawaki et al. that the observed level density near $D_o$ is higher than that obtained by extrapolation of the vibronic level density. The difference is most likely due to spin-rotation and Coriolis couplings. For example, if $J=1/2$ and 3/2 are produced by one-photon absorption from the $J=1/2$ ground state, we anticipate six rotational sublevels for each vibronic level [22]. Since rotational degeneracies of vibronic levels enter both the numerator and denominator of eq. (3), $\rho(E)$ and $N^*(E-D_o)$ can be interpreted either as the density and number of rovibronic levels
for given \( J \) and \( E \) or as those for vibronic levels, without affecting the final result. Thus, the Miya-
waki et al. data are consistent with a vibronic level density of \( \approx 0.4 \) per \( \text{cm}^{-1} \), which is close to the ex-
trapolated Delon et al. value of \( \approx 0.2 \) per \( \text{cm}^{-1} \). Both estima-
tions are consistent with a simple interpreta-
tion of the steps in \( k_{\text{uni}}(E) \) in terms of transition state levels corresponding to \( N^* = 1,2, \) etc.

In experiments carried out by Moore and co-work-
ers using expansion-cooled \( \text{CH}_2\text{CO} \), step-like rises in
the yield for the singlet channel were observed with
energy increments corresponding to \( \text{CO rotational} \) levels \[24,25\]. These steps may be due to com-
petition between singlet and triplet exit channels past
a rate-limiting transition state. With our experimen-
tal resolution, steps of such small increments could
not be resolved even if the \( \text{NO}_2 \) spectral features were
more continuous. However, if such small steps were
present, they would manifest themselves at our res-
olution as a monotonic rise in \( k_{\text{uni}} \) versus \( E \), which
is not the case. Zewail and co-workers, on the other
hand, measured rates for \( \text{CH}_2\text{CO} \) decomposition
[20] which were reconciled using the variational
RRKM approach espoused by Marcus and co-work-
ers with transition state parameters that include low
frequency vibrations [26]. Also, in the experiments
of Zewail and co-workers on \( \text{NCNO} \) decomposi-
tion, it was shown that the rates increased smoothly with
energy except near \( 320 \) \( \text{cm}^{-1} \), where structure in the
\( k(E) \) versus \( E \) curve was clearly discerned [19];
again, a variational RRKM analysis employed low-
frequency vibrations at the transition state [27].

The observed spacings of \( \approx 100 \) \( \text{cm}^{-1} \) for \( \text{NO}_2 \)
are a bit high in light of the above detailed studies of
small molecules decomposing via loose transition states [19,20,24-27]. These spacings may corre-
spond either to transition state levels that correlate
with atomic oxygen \( ^3\text{P}_2, ^3\text{P}_1, \) and \( ^3\text{P}_1 \) fine-structure
levels, which are at 0, 158 and 227 \( \text{cm}^{-1} \), respec-
tively, in the free atom [28]; or alternatively to a bending frequency at the transition state. Since the first interpretation implies a low frequency bend or rotor at the transition state, resulting in a monotonic rise which is not observed, we favor the latter inter-
pretation. The conical intersection between \( ^2\text{A}_1 \) and
\( ^2\text{B}_2 \) may also play a role, since it lies near the exit
channel and zeroth-order \( ^2\text{B}_2 \) does not dissociate to
ground electronic state products. Thus, the \( ^2\text{B}_2 \) part
of the wavefunction lends “tight transition state”
character to the overall wavefunction, so the steps
may reflect passage through the region of conical in-
tersection. Such passage from one region of bound
potential to another has been clearly demonstrated by
Zewail and co-workers in the \( \text{NaI} \) system [31].

At the energies accessed in the present ex-
periments, the transition state involves only one “vibra-
tion” since the \( \text{NO}-\text{stretich} \) is too energetic, i.e. 1876
\( \text{cm}^{-1} \) for free \( \text{NO} \) [29]. Thus, evaluating
\( N^*(E-E_0) \) is a matter of choosing the bend fre-
quency and counting open channels. The rate in the
region \( 0-100 \) \( \text{cm}^{-1} \) can then be attributed to a vi-
breationless transition state having \( N^* = 1 \). The rise
near \( 100 \) \( \text{cm}^{-1} \) is associated with \( v = 1 \) of the bend,
and likewise the step near \( 200 \) \( \text{cm}^{-1} \) is associated with
\( v = 2 \). This is consistent with the approximate 1:2:3
ratio of the relative rates on the first, second and third
steps. Finally, we note that steps are only evident to
\( \approx 350 \) \( \text{cm}^{-1} \). Two explanations may be offered: (i) deteriorating experimental accuracy past \( \approx 350 \) \( \text{cm}^{-1} \)
hides details of the \( k_{\text{uni}}(E) \) dependence, and (ii) the
pattern may change, for example because of a trans-
ition from bending to internal rotation. Such ques-
tions are left for further investigations.

In summary, \( \text{NO}_2 \) photoexcited above \( D_0 \) under-
goes unimolecular decomposition due to its ground
electronic state character. With expansion-cooled
samples, \( k_{\text{uni}} \) versus \( E \) is not smooth, but shows step-
like behavior that can be reconciled with RRKM
theory, using experimental densities of states. Much
remains to be done with this most unusual system.

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References