The decomposition of NO\(_2\) on Rh(111): product NO velocity and angular intensity distributions

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Abstract

The velocity and angular distributions of NO produced from the decomposition of NO\(_2\) on Rh(111) under both reducing and oxidizing conditions have been measured at surface temperatures between 500 K and 1000 K. When a concurrent H\(_2\) beam is used, which keeps the surface free of oxygen, the NO product has much more translational energy than expected for equilibration at the surface temperature, but is dependent on \(T_s\). There is total energy scaling; the translational energy is independent of final angle. A small amount of N\(_2\) is also produced. When the H\(_2\) beam is turned off, oxygen builds up on the surface. Under this oxidizing condition, the NO product has an almost Maxwell–Boltzmann velocity distribution at the surface temperature. © 1999 Elsevier Science B.V. All rights reserved.

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1. Introduction

Rhodium, a component of three-way automotive exhaust catalysts, is important because it selectively reduces NO to N\(_2\) [1,2]. Platinum and palladium also catalyze the reduction, but produce large quantities of undesirable NH\(_3\) and N\(_2\)O. In our laboratory, we have been systematically investigating the dynamics of the NO\(_x\) reduction on Rh(111). In a previous paper, we studied the NO + H\(_2\)/Rh(111) reaction [3]. In this paper, we report our results for the decomposition of NO\(_2\). On many transition metal surfaces, NO\(_2\) readily decomposes. It is a very effective agent for quickly depositing large quantities of oxygen, particularly under ultra-high vacuum (UHV) conditions [4-6]. There have been studies investigating the adsorption and desorption of NO\(_2\) from Pt [7,8], Pd(111) [9], and Ru(001) [10]. These involved dosing the crystals at cryogenic temperatures. On Pt(111), the co-adsorption of NO\(_2\) and oxygen was also studied [11]. When the Pt(111) crystal is exposed to NO\(_2\) at 100 K and then heated, the NO\(_2\) completely decomposes to NO and oxygen. At larger coverages, some of the NO\(_2\) desorbs. High-resolution electron energy loss spectroscopy (HREELS) showed that the NO\(_2\) is adsorbed as a \(\mu\)-N,O-nitrito surface complex (bound to the surface through the N and one O atoms) [11]. When the surface is pre-covered with oxygen atoms, most of the NO\(_2\) is reversibly adsorbed. HREELS spectra also indicated that the NO\(_2\) is nitro-bonded (bound to the surface through the N atom only). On an initially clean Pd(111) surface, the behavior is similar to that of Pt(111), and the NO\(_2\) is also

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adsorbed as a \( \mu \cdot N_2O \cdot \text{nitrito complex}. \) On Ru(001), some of the NO\(_2\) dissociates to NO and O even at 80 K, and any NO\(_2\) in the first layer are adsorbed only through the N atom. Also, the initial decomposition of the NO\(_2\) inhibits any further decomposition. The trend is that the presence of oxygen on the surface changes the adsorption geometry and suppresses the decomposition of NO\(_2\).

In our laboratory, the kinetics of NO\(_2\) decomposition on Rh(111) have been studied [12]. When a Rh(111) crystal [surface temperature \( T_s = 525 \) K] is exposed to a room temperature NO\(_2\) beam, oxygen is very quickly deposited on the surface, \( \sim 50 \) times faster than with an equivalent O\(_2\) beam. The reaction products are principally gas phase NO and adsorbed (and absorbed) oxygen. Temperature programmed desorption (TPD) experiments show that for equivalent quantities of deposited oxygen, the O\(_2\) desorption peak due to absorbed oxygen occurs at a higher temperature when the surface was dosed with NO\(_2\) rather than O\(_2\) [13].

Using our ability to measure the velocity of reaction products under steady-state conditions, we studied the dynamics of NO\(_2\) decomposition on the Rh(111) surface. The reaction was studied at elevated temperatures, where practical NO\(_2\) reduction catalysts operate. This contrasts with previous investigations where reactants were typically deposited at cryogenic surface temperatures, and the reactions studied under non-steady state conditions during which the catalyst’s temperature was subquently ramped up to reach threshold temperatures suitable for reaction. We performed experiments both when there was a concurrent H\(_2\) beam (reduc- ing conditions) and when the oxygen was allowed to accumulate on the surface.

2. Experimental

These experiments were performed in a scattering machine thoroughly described elsewhere [14,15]; only the essential features will be mentioned here. The UHV scattering chamber has a differentially pumped quadrupole mass spectrometer with an angular resolution of \( 1^\circ \). Attached to the detector housing, and rotating with it, is a double-differentially pumped housing containing an AC hysteresis motor for driving a post-crystal chopper, which allows us to make time-of-flight (TOF) measurements of desorbing molecules. This has a 511 channel cross-correlation pattern, and was rotated at 195.69 Hz (10 \( \mu \)s/channel). The chopper interrupts the flux of molecules leaving the Rh(111) surface at a distance of 10.5 cm from the electron impact ionizer.

It is possible to make three independent molecular beams in our machine. The center beam is in the plane defined by the detector rotation, the other two beams define a perpendicular plane. They are canted at \( 15^\circ \) to the plane of the detector rotation, so that all of the beams overlap at the position of the crystal. The crystal can be lowered out of the beam path, and the detector rotated so that the incident velocity distribution and intensity of the center beam can be directly measured. The Rh crystal was cut to within 1\( ^\circ \) of the (111) plane, determined by Laue X-ray back-reflection. The crystal was cleaned by Ar\(^+\) ion sputtering (\( \sim 5 \mu\)A at 900 K), followed by annealing at 1300 K. Carbon was removed from the sample by exposing the 900 K surface to an O\(_2\) beam followed by annealing at 1300 K for 3 min, which results in the removal of both surface and sub-surface oxygen. Cleanliness and surface flatness could be checked by Auger electron spectroscopy and He atom scattering.

During these experiments, we used a continuous beam of NO\(_2\) (Matheson, 99.5\%) and, when we wanted the surface free of oxygen, a continuous beam of H\(_2\). By controlling the relative flux of the two beams, we were able to keep the surface nearly oxygen free during the course of the experiments. Oxygen coverage could be checked by titrating off adsorbed oxygen with an H\(_2\) beam at surface temperatures \( T_s \geq 650 \) K while monitoring the signal of desorbing H\(_2\)O. This result could be calibrated by comparison with the H\(_2\)O from a saturation coverage of oxygen deposited using O\(_2\) dosing. 0.5 monolayers (ML). For the NO\(_2\)+H\(_2\) experiments reported in this paper, the oxygen coverage was less than 0.04 ML.

To determine the velocity distributions for the NO produced at the surface, the measured signal
3. Results

Fig. 1 shows \( m/e = 30 \) TOF spectra for the \( \text{NO}_2 + \text{H}_2 \) reaction at \( T_g = 800 \text{ K} \). The circles are the result of deconvolving the raw data from the cross-correlation pattern. The lines are the result of using a nonlinear least-squares fitting routine, convolving shifted Maxwell-Boltzmann velocity distributions:

\[
f(v|x) \propto \exp \left[ -\left( \frac{v-v_0}{\Delta v} \right)^2 \right]
\]

(1)

with the flight time broadening due to the finite length of the ionizer (\( v \) is the velocity, and the three fitting parameters are the intensity, \( v_0 \), and \( \Delta v \)). The solid line is the sum of all of the contributions to the \( m/e = 30 \) TOF spectra. To arrive at these results, a spectrum with the mass spectrometer tuned to \( m/e = 46 \) had to be taken at each angle. This was fit with the sum of two shifted Maxwell-Boltzmann velocity distributions; the results are the two broken lines with the lowest intensities. (As mentioned, we have used a 1:1 \( \text{NO} : \text{NO}_2 \) fragmentation). The long dashed line is the computed result for the product \( \text{NO} \) from the surface reaction using the same values of \( v_0 \) and \( \Delta v \) for all of the spectra. The average values are 315 meV for \( \langle E \rangle \) and 1360 m/s for \( \langle v \rangle \). For comparison, a Maxwell-Boltzmann velocity distribution \((v_0=0)\) and \( \Delta v = 2k_bT_g/m_g \), where \( k_b \) is Boltzmann’s constant, \( T_g \) is the temperature of the gas, and \( m_g \) is the mass of the gas molecule) at \( T_g = 800 \text{ K} \), what you might expect for desorbing molecules equilibrated to the surface temperature has \( \langle E \rangle = 138 \text{ meV} \).

The points shown in Fig. 2 are the average energies for the surface produced \( \text{NO} \) at the surface normal and \( T_g \) from 500 K to 1000 K. The line through the points is simply a cubic fit to aid the eye. Also shown is a line for what would be expected for the barrierless desorption of the molecules if they were equilibrated to the surface temperature; the measured energies are much higher. Interestingly, for the lower surface temperatures, the slope of the energy versus \( T_g \) function is greater than \( 2k_bT_g \), indicating a stronger dependence of the final translational energy than expected for surface accommodation. A common method for characterizing the width of a velocity distribution is the speed ratio:

\[
\text{SR} = \sqrt{\left( \frac{\langle v^2 \rangle}{\langle v \rangle^2} - 1 \right) / \left( \frac{32}{9\pi} - 1 \right)}.
\]

(2)
which compares the width of the measured velocity distribution with that of a Maxwell–Boltzmann distribution with the same average energy. For the results shown in Fig. 2, this ratio varies from about 0.83 to 0.92, narrower than a Maxwell–Boltzmann distribution measured for higher surface temperatures having on average a larger speed ratio.

Fig. 3 shows the intensity of the surface produced NO as a function of detector angle, normalized to the value at the surface normal, for three values of $T_s$. The solid line is for a $\cos(\theta_f)$ distribution, and the dashed line is for an intensity distribution that varies as $0.43 \cos^2(\theta_f) + 0.57 \cos(\theta_f)$, derived from a fit to the data taken at $T_s=600$ K, which appears to be slightly peaked towards normal. For the higher temperatures, the angular dependence of the intensities is cosine within our error. At both $T_s=600$ K and 800 K, the measured velocity distributions are independent of final angle.

The hydrogen prevents oxygen from accumulating on the surface, and H$_2$O is produced on the surface. We looked at the H$_2$O product TOF spectrum at $T_s=800$ K. The result is fit by a Maxwell–Boltzmann velocity distribution cooler (733 K) than $T_s$, which agrees with our previous results for the H$_2$+O$_2$ reaction on Rh(111) [19]. NO also decomposes to nitrogen and oxygen on Rh(111). At $\theta_f=0^\circ$ and $T_s=800$ K, we were able to detect a very small amount of very high energy N$_2$, but too little for decent statistics to be accumu-
Fig. 2. Average energies for NO produced from the NO$_2$+H$_2$ reaction as a function of surface temperature. The solid line through the points is a cubic fit to aid the eye, the other solid line is what would be expected if the NO had equilibrated and desorbed with a thermal Maxwell–Boltzmann velocity distribution at the surface temperature.

Fig. 3. Relative NO product intensity, normalized by the value at the surface normal, from the NO$_2$+H$_2$ reaction as a function of final angle probed for three different surface temperatures. The solid line is $\cos(h_f)$, and the dashed line is $0.43\cos(h_f)+0.57\cos^2(h_f)$.

Fig. 4a shows an $m/e=30$ TOF spectrum measured at the crystal normal and with $T_s=800$ K when there was no H$_2$ beam. For these data, the crystal was exposed to NO$_2$ for ~2 min before starting data collection, allowing oxygen to build up. We did not measure the amount of oxygen at the end of the experiment, but from our knowledge of oxygen desorption, there is at least ~0.5 ML of adsorbed and absorbed oxygen at this temperature. The integrated intensity, as measured by a

Fig. 4. NO TOF spectrum at $h_f=0^\circ$ for the decomposition of NO$_2$ on Rh(111) after oxygen has been allowed to accumulate taken at $h_i=45^\circ$ and $T_s=800$ K. The circles are the data after deconvolving from the cross-correlation pattern. (a) The lowest intensity broken line is a fit to the signal from NO$_2$ fragmentation; the other dashed line is the estimate for the surface produced NO; the solid line is their sum. (b) The same data, compared to the fit of the spectrum for the NO$_2$+H$_2$ reaction taken at $h_f=0^\circ$ and $T_s=800$ K.

be due to the decomposition of NO in the incident NO$_2$ beam.

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counting ratemeter, did not vary during the course of the experiment, so the dynamics were essentially invariant. Again, we had to measure a spectrum at \( m/e = 46 \), but the results are well represented by a single shifted Maxwell-Boltzmann velocity distribution. The result is shown as the lower intensity broken line in Fig. 4a, with the intensity ratio for NO:NO\(_2\) = 1. Relative to the experiments taken with reducing conditions, there is much more scattered NO\(_2\) away from specular under these oxidizing conditions. The second dashed line is the expected result for a Maxwell-Boltzmann velocity distribution at the surface temperature (\( \langle E \rangle = 138 \) meV). The solid line is then the sum of the two contributions. The product NO is notably less energetic than for NO produced on the oxygen-free surface. Fig. 4b clearly shows this; the dashed line is the fit to the spectrum taken at \( h_f = 0 \)°, shown in Fig. 1. Within experimental uncertainties, the product NO energy is independent of \( T_s \), and the angular intensity distribution is essentially cosine, or only slightly peaked towards normal.

Fig. 5 shows examples of the signal due to NO\(_2\) scattered from the surface. Fig. 5a shows the results when the surface is exposed to NO\(_2\)+H\(_2\) with \( h_f = 60 \)°, and Fig. 5b shows results for just NO\(_2\) at \( h_f = 0 \)°, both measured at \( T_s = 800 \) K. Incident NO\(_2\) beam conditions, including the flux, were the same. As already mentioned, the amount of NO\(_2\) scattered at angles away from specular is much greater after oxygen has been deposited on the surface. The spectrum shown in Fig. 5a was fit with the sum of two shifted Maxwell-Boltzmann velocity distributions while the spectrum in Fig. 5b was fit with only one. The low energy (long flight time) feature in Fig. 5a (\( \langle E \rangle = 76 \) meV, \( \langle v \rangle = 540 \) m/s) and the feature in Fig. 5b (\( \langle E \rangle = 74 \) meV, \( \langle v \rangle = 520 \) m/s) have nearly identical velocity distributions. For the NO\(_2\)+H\(_2\) reaction, where we took data at several surface temperatures, the velocity distributions had average values of about 65-85 meV for \( T_s \) from 600 K to 1000 K, with the energies being slightly greater at the higher surface temperatures. In all cases, this is much lower than expected for accommodation to the surface (2\( k_B T_s \) varies from 103 meV to 172 meV for \( T_s \) varying from 600 K to 1000 K). In fact, the energy is close to that of the incident beam, and the signal is probably due to both elastic and inelastic scattering of the incident molecules.

The high energy feature in Fig. 5a is also fit with a shifted Maxwell-Boltzmann velocity distribution, but with \( \langle E \rangle = 392 \) meV and \( \langle v \rangle = 1245 \) m/s. For the data at \( T_s = 800 \) K and 1000 K, the \( \langle E \rangle \approx 370 \) meV, independent of the final angle investigated, and with the intensity somewhat peaked towards normal. For the data at \( T_s = 600 \) K, \( \langle E \rangle \approx 250 \) meV. However, the signal at all surface temperatures was weak and so these results are very approximate, though the difference...
between \( T_s = 600 \) K data and the higher surface temperatures is real. At the present time, we have no explanation for this high energy feature, which occurs only under reducing conditions.

4. Discussion and conclusions

It is quite common that the products of surface reactions have high translational energies, and this is certainly true for the NO product from the \( \text{NO}_2 + \text{H}_2 \) reaction. In previous papers, we have used surprisal analysis in an attempt to discover the minimum number of parameters necessary to describe the dynamics \cite{3,15}. For NO production at 800 K and 1000 K, where the angular variation of the intensity is cosine, the results can be reasonably described with the form:

\[
\int(v) = \int \left[ v^3 \cos(\phi_v) \exp\left(-\frac{v^2}{2\sigma_T^2}\right) \right] \exp(iz) \int(v) \int_T.
\]

The first term in brackets, the prior distribution, is a Maxwell–Boltzmann velocity distribution at the surface temperature. The second term contains the single surprisal parameter, \( \lambda \). This form gives nearly as good a fit (but a consistently larger value of \( \chi^2 \) from the least squares analysis) as the shifted Maxwell–Boltzmann distribution, and with one less parameter \( (\lambda = 3.62 \times 10^{-3} \text{ s/cm at } T_s = 800 \text{ K}) \). However, this way of examining the data adds little apparent insight. To attempt to interpret the data by detailed balance is more difficult, because the NO not only desorbs, but also dissociates. The second term in Eq. (3) can be viewed as a barrier to desorption, but the small value of \( \lambda \) makes it very steep. It might be that the energy comes totally from the breaking of the \( \text{N}=\text{O} \) bond, but this is hard to rationalize with the strong surface temperature dependence of the translational energy.

Without an H\(_2\) beam to remove the adsorbed oxygen from the decomposition of the NO\(_2\), the translational energy of the product NO changes dramatically. Having a Maxwell–Boltzmann distribution at the surface temperature, it appears that we are measuring the desorption of fully accommodated molecules from a barrierless adsorption well. It is possible that the \( \text{NO}_2 + \text{H}_2 \) reaction has different dynamics, because the reaction is actually between adsorbed H and NO\(_2\), rather than the H serving to remove oxygen from the surface. However, the NO product yield and translational velocity were independent of the H\(_2\) flux, until it was too weak too remove all of the oxygen. Then the TOF spectra would begin changing, getting slower as the oxygen built up. In line with the results of the low temperature adsorption experiments on other transition metals, where the geometry of the adsorbed NO\(_2\) changed in the presence of adsorbed oxygen, a more likely explanation is that the presence of oxygen changes the NO\(_2\)–surface interaction potential. That the surface potential is appreciably different is manifested by the increase in the direct-inelastically scattered component of the incident beam.

The principal findings are that for the \( \text{NO}_2 + \text{H}_2 \) reaction, the NO product has much more translational energy than expected for equilibration at the surface temperature, but is dependent on \( T_s \). There is total energy scaling: the final energy is independent of final angle. The angular dependence of the NO intensity is nearly cosine for the higher surface temperatures, and peaked towards normal at \( T_s = 600 \) K. Also, it appears that some of the product NO decomposes to N\(_2\) and oxygen. When the H\(_2\) beam is turned off, oxygen builds up on the surface from the dissociation of NO\(_2\). Under these oxidizing conditions, NO product has an approximately Maxwell–Boltzmann velocity distribution at \( T_s \). These results add to our knowledge of NO\(_2\) reduction on rhodium. They also show how the dynamics can be radically changed by the condition of the surface, depending upon whether the oxygen from the decomposition is allowed to build up or is continuously removed.

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