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Published in:
Journal of the American Chemical Society

DOI:
10.1021/ja00094a046

Published: 01/01/1994

Document Version
Publisher's PDF, also known as Version of Record (includes final page, issue and volume numbers)

Please check the document version of this publication:
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Link to publication

Citation for published version (APA):

Download date: 25. Nov. 2018
Atomic and Molecular Oxygen as Chemical Precursors in the Oxidation of Ammonia by Copper

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Received October 18, 1993. Revised Manuscript Received May 2, 1994.

Abstract: The role of atomic and molecular oxygen precursors in the overall catalytic cycle for ammonia dissociation is analyzed using first-principle density functional calculations. Adsorption energies for ammonia, molecular oxygen, NH₃, NO, and various intermediates and adatoms were computed from geometry optimized calculations on the model Cu(8,3) cluster of the Cu(111) surface. Reported values systematically underpredict experimental adsorption energies by 30 kJ/mol due to the finite cluster size. Attractive and repulsive lateral interactions were important in accessing accurate adsorption energies. Atomic oxygen enhances N–H bond activation; however, it also acts to poison active surface sites and inhibit ammonia dissociation kinetics. Transient molecular oxygen adsorbs weakly in both parallel (–17 kJ/mol) and perpendicular orientations (–10 kJ/mol) to the surface. Parallel adsorption appears to be a precursor for oxygen dissociation, whereas perpendicular adsorption is the precursor for ammonia dissociation. The mechanism in which hydrogen atoms are abstracted sequentially to form OOH* intermediate \([E^* (\text{apparent}) = 0 \text{ kJ/mol}]\) is favored over that in which two hydrogens are simultaneously transferred to form water directly \([E^* (\text{apparent}) = +67 \text{ kJ/mol}]\).

The transient nature of chemical precursors, however, makes it difficult to experimentally resolve their role in the mechanism. For example, oxygen has been shown to both activate strong nitrogen-, sulfur-, and carbon-hydrogen bonds on transition metal surfaces\(^\text{[13,11]}\) and also inhibit dissociation.\(^\text{[11,12,14]}\) While it is generally agreed that the formation of strong chemisorbed atomic oxygen overlayers poisons active surface sites and is responsible for inhibition, the nature of the activation process is still unclear. In this work we present a theoretical analysis of the dissociation of ammonia over copper and elucidate the role of both adsorbed atomic and molecular oxygen species. We first provide a short background of the experimental literature on ammonia oxidation before outlining the specifics of this study.

Roberts and his group have experimentally investigated the role of oxygen in the dissociation of ammonia on both Cu(110)\(^\text{[11,12]}\) and Cu(111).\(^\text{[11,14]}\) On Cu(110), they found that both the activity and the selectivity toward different NH₃ products were highly sensitive to the reaction conditions and to the order in which the reactants were introduced to the surface. For example, when oxygen was preadsorbed on the surface, a very slow exchange of oxygen and ammonia was required prior to any subsequent dissociation of ammonia to different NH₃ species. The surface amide (NH₂*) was favored at low temperatures and coverages, whereas the surface imide (NH*) and surface nitrogen (N*) were favored at higher temperatures and coverages. The activity was attributed to only a small fraction of the preadsorbed oxygen species. The remaining oxygen was thought to poison surface and inhibit the kinetics.

When oxygen and ammonia were coadsorbed (in a regime of excess ammonia), however, there was rapid dissociation of the ammonia with a high selectivity for NH* formation. The initial speculation was that a molecularly adsorbed transient oxygen precursor, similar to that which was found on Zn(001) surface,\(^\text{[10]}\) was responsible for the chemistry.

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Introduction

Both atomic and molecular oxygen surface species have been speculated as governing chemical precursors for several important transition metal catalyzed reactions. On silver, for example, weakly bound atomic oxygen which is stabilized by subsurface oxygen was cited as the dominant precursor in ethylene epoxidation,\(^\text{[1–3]}\) whereas molecular oxygen was found to control the oxidation of SO₂ and COS.\(^\text{[4]}\) In addition, there has even been speculation for both oxygen species in mediating the same catalytic reaction paths. Roberts et al., for example, have cited atomic and molecular precursors as controlling the oxidation of ammonia over Mg,\(^\text{[5]}\) Zn,\(^\text{[6]}\) and Cu.\(^\text{[7,8]}\) Clearly, the role of both atomic and molecular oxygen species as precursors is of considerable importance in understanding a significant spectrum of different selective oxidation reaction pathways.
Oxidation of Ammonia by Copper

On Cu(111), Roberts and colleagues\textsuperscript{13,14} once again found that ammonia readily dissociates in the presence of oxygen. Based on their results and the ideas proposed by Ertl for the dissociation of oxygen on platinum,\textsuperscript{15} they suggested that the mechanism for the initial dissociation might be controlled by a “hot” transient atomic oxygen species. It is speculated that these species rapidly diffuse across the surface and abstract hydrogen from the adsorbed ammonia before becoming thermally equilibrated with the surface. At high oxygen partial pressures, they found that the rate was inhibited by the presence of chemisorbed oxygen overlayers.

Clearly, the controlling chemistry in these systems and the role of oxygen is a complex function of competing elementary reaction steps. Both the overall catalytic cycle and details of the mechanism and activation barriers are, therefore, essential for a more complete and fundamental understanding of the operative chemistry. Herein, we report the results of a theoretical analysis of the mechanism of oxygen-assisted ammonia dissociation on model Cu clusters of the Cu(111) surface. We draw upon some of our previous results which were aimed at deducing (1) the electronic features responsible for the interaction of ammonia at various different adsorption sites on Cu(111) surface,\textsuperscript{16} (2) the effects of cluster size and fixed electronic core configurations on the computed ammonia adsorption energy,\textsuperscript{16} and (3) the role of chemisorbed atomic oxygen on the activation of the NH\textsubscript{2} bond.\textsuperscript{7,18}

In this work we analyze the five following paths as plausible controlling mechanistic steps in the dissociation of ammonia in the presence of oxygen.

\begin{align*}
\text{NH}_3(a) + O(a) &\rightarrow \text{NH}_2(a) + OH(a) \quad (1) \\
\text{NH}_3(a) + O(a) &\rightarrow \text{NH}(a) + H_2O(g) \quad (2) \\
\text{NH}_3(a) + O_2(a) &\rightarrow \text{NH}_2(a) + \text{OOH}(a) \quad (3) \\
\text{NH}_3(a) + O_2(a) &\rightarrow \text{NH}(a) + H_2O(g) \quad (4) \\
\text{NH}_3(a) + O_2(a) &\rightarrow \text{NH}(a) + H_2O(g) \quad (5)
\end{align*}

In the first two paths, chemisorbed atomic oxygen is proposed as being responsible for ammonia dissociation. In path 1, a single N–H bond is activated by the adsorbed oxygen, whereas in path 2 the ammonia is situated on the surface such that two N–H bonds can simultaneously be activated and water is formed directly. In paths 3 and 4, molecular oxygen is considered to be active precursor responsible for the dissociation of ammonia. In path 3, a single hydrogen is abstracted to form the OH\textsuperscript{•} intermediate, whereas in path 4 two hydrogens are simultaneously transferred to form water. The final pathway is that which was proposed by Roberts, where a “hot” transient atomic oxygen surface species (O\textsuperscript{•}) which has not been thermally equilibrated with the surface, controls NH bond activation.\textsuperscript{5,7,12,14} We used first-principle density functional calculations to analyze in detail the activation barriers and corresponding catalytic reaction cycles associated with the first four of these paths. Based on these results we speculate as to which is the dominant path and access the plausibility of the final “hot” oxygen-mediated path.

**Methods**

Density functional quantum chemical calculations were performed to determine gas phase formation, bond dissociation, adsorption, and activation energies. Nearly all of the results reported here were derived from the Amsterdam Density Function (ADF) program of Baerends and co-worker.\textsuperscript{19–21} Some supplemental calculations of the transition barriers were determined by the DGauss program from Cray Research Inc.\textsuperscript{22} The main differences between these two methods are computational. The ADF program, for example, is based on Slater type orbitals, whereas DGauss utilizes Gaussian orbital basis sets. Adsorption energies for ammonia and other adsorbates computed from both programs were shown to be equivalent within ±0.1 kJ/mol. Both programs solve the Kohn–Sham one-electron equations and invoke the local density approximation by implementing the exchange correlation energy functional of Vosko–Wilk–Nusair.\textsuperscript{23} Nonlocal gradient corrections for the exchange and correlation terms were computed subsequent to both self-consistent-field (SCF) convergence and geometry optimization calculations and determined by the methods established by Becke\textsuperscript{24,25} and Perdew,\textsuperscript{26} respectively. All calculations reported here were performed using nonrelativistic, spin-unrestricted, frozen-core calculations. The basis sets used in both ADF and DGauss were of single-\textsuperscript{p} quality for core orthogonalization and double-\textsuperscript{p} for nitrogen oxygen and triple-\textsuperscript{p} quality for copper with polarization functions included for all valence electrons. The exponential coefficients for the basis and fit functions used in the ADF calculations are reported in the Appendix and discussed in more detail elsewhere.\textsuperscript{16,18} The basis sets and fit functions used in the DGauss program were reported by Andzelm.\textsuperscript{27} In the ADF calculations the copper basis was frozen up to and including the 3p core, whereas in nitrogen and oxygen the 1s core was frozen, whereas in the DGauss calculations all electrons were explicitly included. The analytical gradient algorithms\textsuperscript{22,28} employed in both programs enabled geometries to be optimization.

Adsorption energies were determined by optimizing the adsorbate geometry on a model Cu cluster of the Cu(111) surface (Cu–Cu bond length of 2.556 \(\text{Å}\)) and computing the energy change via eq 6

\[ \Delta E_{\text{ads}} = E_{\text{Cu cluster-adsorbate}} - E_{\text{Cu cluster}} - E_{\text{adsorbate(g)}} \] (6)

where \(E_{\text{Cu cluster-adsorbate}}\) is the energy of the adsorbate–Cu cluster complex and \(E_{\text{Cu cluster}}\) and \(E_{\text{adsorbate(g)}}\) are the energies for the free Cu cluster and the gas phase adsorbate, respectively. All adsorption energies reported herein were computed from ADF calculations, whereby the adsorption energy is determined as the energy difference of the overall complex minus its respective fragments via the Ziegler transition state method.\textsuperscript{29}

The Cu(8,3) cluster used in this work, \(I\), was chosen to mimic the essential electronic features of the Cu(111) surface while maintaining reasonable CPU expenditures. The nomenclature (8,3) refers to the arrangement of the atoms in this model; there are eight copper atoms in the surface and three atoms in the layer beneath. We have shown the effects of computed adsorption energies on cluster size and configuration for ammonia adsorption elsewhere.\textsuperscript{16} Figure 1 depicts experimental adsorption energies (Cu–Cu bond length of 2.556 \(\text{Å}\)) and computing the energy change via eq 6

\[ \Delta E_{\text{ads}} = E_{\text{Cu cluster-adsorbate}} - E_{\text{Cu cluster}} - E_{\text{adsorbate(g)}} \] (6)

where \(E_{\text{Cu cluster-adsorbate}}\) is the energy of the adsorbate–Cu cluster complex and \(E_{\text{Cu cluster}}\) and \(E_{\text{adsorbate(g)}}\) are the energies for the free Cu cluster and the gas phase adsorbate, respectively. All adsorption energies reported herein were computed from ADF calculations, whereby the adsorption energy is determined as the energy difference of the overall complex minus its respective fragments via the Ziegler transition state method.\textsuperscript{29}

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greater CPU times, which when coupled with demanding geometry optimization becomes computationally prohibitive.

Activation barriers were computed by choosing an appropriate reaction coordinate and optimizing the geometry of the adsorbates on the cluster at fixed points along the coordinate. All relevant calculations were performed using the Cu(8,3) cluster. A minimal Cu(6) surface cluster (II), however, was used as an initial scoping tool. Calculations based on the Cu(6) cluster allowed us to judiciously choose which geometry optimizations and reaction coordinate calculations to perform on the larger Cu(8,3) cluster. The central surface copper in the Cu(6) cluster is coordinatively unsaturated and as a result overestimates the nitrogen atom binding to this site. This subsequently leads to the prediction of a much weaker nitrogen-hydrogen bond. Thus, the Cu(6) cluster provides a significant lower bound on activation energies for hydrogen abstraction.

Results

The results described here are divided into four sections (1) adsorption energies, (2) lateral interactions, (3) overall catalytic energy cycles and (4) activation barriers for the four mechanistic paths outlined in eqs 1-4. A substantial portion of item 1, formation and adsorption energies, was described in some of our early work.16-18 We, therefore, only present a short review on the important features and concentrate more heavily on items 2 and 3.

I. Adsorption Energies. The computed adsorption energies for the relevant adatoms, radical fragments, and molecular adsorbates on the Cu(8,3) cluster are summarized in Table 1. The energetics as well as the sites and optimized Cu-adsorbate bond distances are reported. Ziegler10 has recently reported that density functional methods typically predict structural bond lengths to within 0.01 Å and organometallic bond energies within about 20 kJ/mol. In general, and as expected, the heats of chemisorption reported in Table 1 demonstrate the relative ordering:

molecular adsorption < radical fragment adsorption < adatoms

Molecular adsorbates have a weak electrostatic surface interaction and tend to prefer onefold coordination sites, whereby Pauli repulsion is minimized.16,18 Adatoms and hydrogen deficient radical fragments such as NH and OH, however, bind much more strongly and prefer the higher threefold coordination sites. The overlap between the free adatom p_x and p_y orbitals with the metal surface s orbitals provides additional stabilization at the higher coordination sites which helps to increase the binding energy. Intermediate radical fragments, such as NH_2, prefer twofold coordination. Interestingly, a number of these adsorption values correspond with values reported by Shustorovich13 based on bond order conservation calculations through a linear relationship, such as the one which was illustrated in Figure 1.

The adsorption of molecular oxygen is an important step in the overall cycle so we gave considerable attention to optimizing the binding of O_2 to the surface. Both perpendicular and symmetric parallel adsorption over threefold coordination sites were found to be favorable steps and within 10 kJ/mol of one another. In the parallel arrangement, the oxygen-oxygen bond is stretched from 1.21 to 1.43 Å, and each oxygen appears to take on a considerable negative charge. This geometry is very similar to the experimental results reported by Madix for O_2 on Ag(111) where the oxygen-oxygen bond was found to bind parallel to the surface with an O-O distance of 1.41 Å.32 The bond length follows the value reported for the gas phase O_2 (1.41 Å)33 which

Table 1. ADF Computed Adsorption Energies for Molecules, Radicals, and Atoms on the Model Cu(8,3) Cluster

<table>
<thead>
<tr>
<th>adsorbate</th>
<th>site</th>
<th>atoms defining site</th>
<th>distance to nearest Cu atom (Å)</th>
<th>adsorption energy (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH_3 on Cu(8,3)</td>
<td>1-fold</td>
<td>1</td>
<td>2.41</td>
<td>-16</td>
</tr>
<tr>
<td>O_3 on Cu(8,3)</td>
<td>1</td>
<td>2-fold</td>
<td>-42</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>3-fold</td>
<td>-17</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>4-fold</td>
<td>-10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H_2O on Cu(8,3)</td>
<td>1-fold</td>
<td>1</td>
<td>2</td>
<td>+1</td>
</tr>
<tr>
<td>NO on Cu(8,3)</td>
<td>3-fold</td>
<td>2-3-8</td>
<td>2.18</td>
<td>-83</td>
</tr>
<tr>
<td>NH_2 on Cu(8,3)</td>
<td>1-fold</td>
<td>1</td>
<td>1.99</td>
<td>-121</td>
</tr>
<tr>
<td>NH on Cu(8,3)</td>
<td>1-fold</td>
<td>1-2-3</td>
<td>2.25</td>
<td>-164</td>
</tr>
<tr>
<td>OH on Cu(8,3)</td>
<td>1-fold</td>
<td>1-5-6</td>
<td>1.84</td>
<td>-258</td>
</tr>
<tr>
<td>OOH on Cu(8,3)</td>
<td>3-fold</td>
<td>2-3-8</td>
<td>2.14</td>
<td>-105</td>
</tr>
<tr>
<td>N on Cu(8,3)</td>
<td>3-fold</td>
<td>1-5-6</td>
<td>1.90</td>
<td>-339</td>
</tr>
<tr>
<td>O on Cu(8,3)</td>
<td>3-fold</td>
<td>2-3-8</td>
<td>1.91</td>
<td>-401</td>
</tr>
<tr>
<td>H on Cu(8,3)</td>
<td>1-fold</td>
<td>1</td>
<td>1.57</td>
<td>-133</td>
</tr>
<tr>
<td>2-fold</td>
<td>2-3</td>
<td>1.74</td>
<td>-129</td>
<td></td>
</tr>
<tr>
<td>3-fold</td>
<td>1-2-3</td>
<td>1.79</td>
<td>-160</td>
<td></td>
</tr>
<tr>
<td>3-fold</td>
<td>1-2-3</td>
<td>1.68</td>
<td>-175</td>
<td></td>
</tr>
</tbody>
</table>

suggests that each oxygen takes on a considerable negative charge. The computed Mulliken charges are given in Table 2. Due to arbitrariness of the Mulliken population analysis to apportion the density in a chemical bond to its representative atoms, the results in Table 2 provide only an approximate qualitative measure of charge and charge transfer. The surface O2, species picks up a significant charge (−0.62) from its neighboring copper atoms (4, 5, and 8 in Table 2), which is in good agreement with the results reported by Fischer and Whitten34 (−0.76 for O2 bond tilted on Cu(001)). This is somewhat lower than value for chemisorbed atomic oxygen (−0.75). The combination of the long O−O bond and the enhanced surface charge transfer to this bond suggests that the parallel arrangement is a likely precursor state for the O2 dissociation.

Interestingly, while both the O2 surface-species and chemisorbed atomic oxygen (Table 2) pick up significant charge from the Cu surface, the charge-transfer is not as high as two electrons. The value for atomic oxygen on Cu(8,3), −0.75, is in good agreement with that reported by Illas et al.,35−0.96 and −0.62, and Mattisson et al.,36−0.85, for O on Cu(4,1). Through a detailed analysis of (1) the projected O orbitals in the Cu50 wave function, (2) the dipole moment curves for O and various point charges interactions with Cu5 cluster, and (3) the participation of the Cu 3d orbitals in the Cu−O bond, Illas and Bagus37 were able to extend approximate Mulliken results to find a better estimate of the charge on the oxygen, −1.5 electrons. The good agreement between the Mulliken charges reported here and those determined by Illas and Bagus suggest that the actual charge on the oxygen on the Cu(8,3) cluster is closer to −1.5. This also agrees with the experimental results of Clendening et al.,38 who find small increases in the work function induced by oxygen and hence a charge transfer from the metal to the oxygen less than two electrons.

For O2 adsorbed perpendicular to the surface, the optimized oxygen−oxygen distance (1.30 Å) was found to be significantly shorter than that for O2−parallel case. The 1.30 Å distance is close to the reported 1.36 Å for O2 (g). In addition, there is also a decrease in the overall charge transfer, −0.33 on O2. While the parallel mode of O2 adsorption is slightly favored on the clean Cu surface, the perpendicular mode, as we shall see, becomes favored in the presence of coadsorbed or preadsorbed ammonia species due to a reduction in the repulsive interactions.

The adsorption values reported in Table 1 were all determined using the same 11 atom asymmetric Cu(8,3) cluster, as is depicted

### Table 2. Computed Mulliken Charges for Atomic Oxygen, Molecular-Oxygen Bound Parallel, and Molecular Oxygen Bound Perpendicular to the Cu (8.3) Cluster Surface

<table>
<thead>
<tr>
<th>no.</th>
<th>atom</th>
<th>Cu(8.3)</th>
<th>Cu(8.3)−O</th>
<th>Cu(8.3)−O2 (parallel)</th>
<th>Cu(8.3)−O2 (perpendicular)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cu</td>
<td>0.1900</td>
<td>0.193</td>
<td>0.188</td>
<td>−0.100</td>
</tr>
<tr>
<td>2</td>
<td>Cu</td>
<td>0.066</td>
<td>0.016</td>
<td>0.005</td>
<td>0.036</td>
</tr>
<tr>
<td>3</td>
<td>Cu</td>
<td>0.006</td>
<td>0.077</td>
<td>−0.0003</td>
<td>0.083</td>
</tr>
<tr>
<td>4</td>
<td>Cu</td>
<td>−0.029</td>
<td>0.240</td>
<td>0.1481</td>
<td>0.033</td>
</tr>
<tr>
<td>5</td>
<td>Cu</td>
<td>−0.029</td>
<td>0.240</td>
<td>0.1481</td>
<td>0.033</td>
</tr>
<tr>
<td>6</td>
<td>Cu</td>
<td>0.006</td>
<td>0.077</td>
<td>−0.0003</td>
<td>0.083</td>
</tr>
<tr>
<td>7</td>
<td>Cu</td>
<td>−0.066</td>
<td>0.016</td>
<td>0.005</td>
<td>0.036</td>
</tr>
<tr>
<td>8</td>
<td>Cu</td>
<td>−0.098</td>
<td>0.235</td>
<td>0.117</td>
<td>0.061</td>
</tr>
<tr>
<td>9</td>
<td>Cu</td>
<td>0.038</td>
<td>0.062</td>
<td>−0.004</td>
<td>−0.055</td>
</tr>
<tr>
<td>10</td>
<td>Cu</td>
<td>0.023</td>
<td>0.010</td>
<td>0.006</td>
<td>−0.0004</td>
</tr>
<tr>
<td>11</td>
<td>Cu</td>
<td>0.023</td>
<td>0.010</td>
<td>0.006</td>
<td>−0.0004</td>
</tr>
<tr>
<td>12</td>
<td>Cu</td>
<td>0.023</td>
<td>0.010</td>
<td>0.006</td>
<td>−0.0004</td>
</tr>
<tr>
<td>13</td>
<td>O</td>
<td>−0.745</td>
<td>−0.341</td>
<td>−0.674</td>
<td>−0.255</td>
</tr>
<tr>
<td>14</td>
<td>O</td>
<td>−0.277</td>
<td>−0.255</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


by I above. One can question how cluster size and cluster symmetry (or lack thereof) will affect the binding energetics on this Cu(8,3) cluster. The result of adding an additional Cu atom to the second layer to form Cu(8,4) changed the binding energy of both atomic nitrogen and NH3 at the center threefold site by at least as 4 kJ/mol. The local environment about a particular site was found to be much more important. Threefold sites situated directly atop of a Cu atom in the second layer were found to be energetically less favorable for adsorption by 40–50 kJ/mol for both atomic nitrogen and NH3 than for those threefold sites directly above a second layer threefold hollow site.

In an effort to prove the latter question on symmetry, the binding of different adsorbates on the two symmetric copper clusters, Cu(7,3) III and Cu(10,3) IV, was examined. The adsorption energy of ammonia on Cu(7,3) was within 1 kJ/mol of the value reported on Cu(8,3) suggesting that cluster symmetry has little effect. The adsorption energies of atomic oxygen and atomic nitrogen, on the other hand, were reduced from 401 to 362 and from 339 to 288 kJ/mol, respectively, in going from Cu(8,3) to Cu(10,3). While cluster symmetry may be responsible for the small but measurable lowering of the adsorption energies, the more likely explanation is due to changes in the cluster Fermi level.

### II. Lateral Surface Interactions

Our previous results on the adsorption of ammonia on Cu(111) and the electronic effect of coadsorbed atomic oxygen demonstrated the significance of lateral surface interactions on measured adsorption energies.16,17 Lateral interaction energies are defined here as the enhanced or diminished adsorption energy of an adsorbate on a surface due to the presence
of additional surface species. The controlling enhancement or reduction should be governed predominantly by changes in the electronic surface structure, i.e., through-metal interactions rather than through-space interactions. A schematic of the adsorbate arrangement on the surface for the above example is depicted in case I of Table 3. Ammonia, which prefers the onefold adsorption site (Cu1 in I or A1 from Table 3), is stabilized by 32 kJ/mol via a through-metal attractive interaction with the neighboring threefold oxygen adsorbate (A2). A detailed electronic analysis of this type of interaction was presented elsewhere.17,18,38 The attractive interaction can be rationalized along the lines of bond order conservation39,40 and least metal atom sharing principles.38

The presence of the oxygen adsorbate weakens the bonds between nearest-neighbor (Cu2 and Cu3) and next-nearest-neighbor metal atoms (Cu1–Cu4–Cu7–Cu9) which subsequently strengthens the binding at next-nearest-neighbor adsorption sites, such as the atop position for ammonia A1 (Cu1). The adsorption of ammonia in the presence of O2*: and the adsorption of NH2 in the presence of OH*: are two similar examples of the enhanced binding effects for case I type adsorption. Ammonia binds 17 kJ/mol more strongly, and NH2 binds 5 kJ/mol more strongly in the presence of O2*: and OH*: respectively.

The case II and case III adsorption schemes depicted in Table 3 present situations where both adsorbates prefer higher coordination and sit at adjacent threefold sites. In the case II situation, the adsorbates (A1 and A2) share two metal atoms (i and ii) in forming adsorbate–metal bonds. This “sharing” considerably weakens the metal–adsorbate bonds and gives rise to a large repulsive interaction between the two adsorbates. For example, the adsorption of NH on the Cu(8,3)–O cluster, where the oxygen and NH share two metal atom centers, was found to be 178 kJ/mol less favorable than for the adsorption of NH on the clean Cu(8,3) surface. According to the principle of least metal atom sharing,38 one should expect a large repulsive interaction. The strength of the repulsive interaction, however, can be considerably reduced by moving to a scenario where the adsorbates share only a single metal atom center (i), case III. This explains the results in Table 3, where the repulsive interactions are now reduced to 25–30 kJ/mol. These values are now in line with general experimental adsorbate–adsorbate repulsive interaction energies on transition metal surfaces. In general the results suggest that only case I and case III situations are likely on the Cu(111) surface.

Before proceeding to the calculation of overall reaction cycles, we first discuss the model used for the relative positioning of adsorbates on the cluster. We attempt to minimize artifacts of cluster size effects and establish an appropriate model of the real surface. Ammonia which binds weakly to the cluster will be much more sensitive to location (interior vs exterior) than the strongly chemisorbed atomic oxygen. Therefore the more realistic arrangement of NH3 and O adsorbates is to have ammonia sit atop the central copper site with atomic oxygen closer to the edge. The inverse situation, where NH3 is placed at an edge site and oxygen located toward the center of the cluster, will be less representative of an extended surface because of the enhanced stability of NH3 at the unsaturated edge site. Here we analyze the analogous situation of NH2 and O on Cu(8,3). The binding energy was 140 kJ/mol stronger for the case where NH2 was placed at an edge twofold site (with oxygen positioned in the center of the cluster) than the case in which NH2 was bound at a central edge site (with oxygen situated at an exterior hollow site). This increased binding energy (for NH2 at the edge of the cluster) is directly due to the excessive stabilization of NH2 at unsaturated metal sites. The more likely model of the extended surface is therefore the one in which atomic oxygen binds at the exterior with NH3 and NH2 occupying more central adsorption sites.

### III. Overall Catalytic Cycles

The adsorption energies, gas phase formation energies, and lateral interaction energies provided the basis for computing reaction enthalpies for elementary adsorption, surface reaction, and desorption steps. Overall catalytic cycles were fashioned from these steps and are delineated
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Two mechanistic paths exist for the cycles controlled by atomic oxygen. In the first, atomic oxygen abstracts a single hydrogen from neighboring ammonia to form surface amide and hydroxyl groups (eq 1). In the second, two of the ammonia hydrogens are simultaneously transferred from the adsorbed ammonia to the surface oxygen to yield gaseous water and NH\(^*\) directly. This is followed by a final H- abstraction to form N\(^*\). The first two steps in both cycles refer to the adsorption of NH\(_3\) and atomic oxygen. The intermediate steps SR1-SR4 are the hydrogen surface reaction steps depicted in Table 4. The final terms, N\(_2\) (g), NO\(^*\), NO(g), and H\(_2\)O removal, refer to recombinative desorption of nitrogen adatoms to N\(_2\)(g), surface reaction of N\(^*\) and O\(^*\) to form NO\(^*\), recombinative desorption of NO, and recombinative of surface hydroxyl groups to form H\(_2\)O(g), steps, respectively.

in terms of the controlling mechanistic paths defined in eqs 1-4 above. Each of these are described here in turn.

**Atomic Oxygen Controlled.** Two mechanistic paths exist for the cycles controlled by atomic oxygen. In the first, atomic oxygen abstracts a single hydrogen from neighboring ammonia to form surface amide and hydroxyl groups (eq 1). In the second, two of the ammonia hydrogens are simultaneously transferred from the adsorbed ammonia to the surface oxygen to yield gaseous water directly and a surface imide (eq 2). The elementary adsorption, surface reaction, and desorption steps required for the two overall cycles which follow these two different paths are outlined in Table 4A,B. The four different surface reactions are numbered in order. The corresponding reaction enthalpies for each of these steps are assembled into the overall potential energy profile for each of the two cycles, and the results are shown in Figure 2A,B.

By way of example, the adsorbate-cluster structure for each of the basic steps for the first scheme from Figure 2a are depicted along with corresponding stoichiometric coefficients in the cycle presented in Figure 3. In the first step, ammonia is adsorbed on the surface in the presence of oxygen at +48 kJ/mol. The negative sign simply indicates that the step is exothermic. Additional oxygen, which is required for the overall cycle, is dissociatively adsorbed leading to a strongly bound chemisorbed atomic oxygen (−148 kJ/mol). The high exothermicity of this step indicates that any ad-oxygen that is formed on the surface will be very difficult to remove and thus acts to poison active surface sites. The initial ammonia dissociation step, SR1, was found to be 128 kJ/mol more likely in the presence of oxygen than on the clean Cu(111) surface; however, at +48 kJ/mol some form of thermal energy is still required for this endothermic step. Subsequent dissociation steps, SR2 (+6 kJ/mol) and SR3 (−48 kJ/mol), to form NH\(^*\) and N\(^*\) help to drive the energetics of the overall cycle. Thermodynamics indicate that the final nitrogen surface species favor recombination with additional N\(^*\) and desorption as N\(_2\) (−214 kJ/mol) over recombination with surface oxygen and desorption as NO (+46 kJ/mol). However under steady state reaction conditions, chemisorbed oxygen, which is strongly bound to the surface, will be the most abundant surface intermediate (MARI). The number of N-O collisions will, therefore, be far greater than the number of N-N collisions, thus implying that NO formation will dominate over N\(_2\) formation. Lastly, the recombinative of surface hydroxyl groups to form water and chemisorbed atomic oxygen was found to be highly endothermic and is speculated as a plausible rate controlling step. The overall reaction energy for the complete catalytic cycle was computed to be −625 kJ/mol which is in very good agreement with the estimate of −611 kJ/mol from thermochemical data.

Figure 2 (part B) represents the overall potential energy diagram for the atomic oxygen mediated path in which two hydrogens are simultaneously transferred in the surface dissociation step and water is formed directly. This cycle is identical to that in Figure 2A with two exceptions: (1) less atomic oxygen is required to carry out the overall reaction and (2) the number of surface reactions is reduced from two to three due to the simultaneous transfer of two hydrogens in the initial dissociation step. The most striking feature of the cycle is the considerable energy required to dissociate the two hydrogens from ammonia. At +92 kJ/mol, this step is now +44 kJ/mol more endothermic than that for the abstraction of a single hydrogen (Figure 2A).

**Molecular Oxygen Controlled.** The overall cycles for the molecular paths, eq 3 and 4, are reported in Table 4B and 3, and their corresponding potential energy profiles depicted in Figure 4. The path mediated by a single hydrogen transfer to form the OOH\(^*\) intermediate is shown in part a. In this scheme, ammonia and molecular oxygen are coadsorbed and react to form OOH\(^*\) and NH\(_2\)\(^*\). A series of ammonia adsorption calculations at various O-O surface angles indicated that the coadsorbed molecular oxygen prefers to sit perpendicular to the surface. This angle decreases somewhat as the hydrogen abstraction reaction proceeds to help shorten the distance that the hydrogen must travel. Subsequent dissociation steps lead to O\(^*\), OH\(^*\), and NH\(_2\)\(^*\) products. To complete the series of abstraction steps, 1 mol of dissociated O\(_2\) was required for the final step: NH\(_2\)\(^*\) + O\(^*\) → N\(^*\) + OH\(^*\). The initial surface dissociation step for this scheme, SR5, is now exothermic and appears more likely than the considerable endothermic steps reported for dissociation on the clean Cu(111) surface and the atomic-oxygen-assisted dissociation. The desorption steps are the same as those reported in Table 4A and Figure 2A.

The final path analyzed, eq 4, was that for molecular-mediated oxygen dissociation and the simultaneous abstraction of two hydrogens to form gaseous water. The steps required for this scheme are outlined in Table 4D and depicted in Figure 4B. The overall potential energy profile is the most appealing of all those studied. The cycle moves to lower energies at each subsequent elementary step until the final endothermic recombinative desorption steps. In addition, this path completely eliminates the highly exothermic oxygen dissociation step which leads to strong chemisorbed atomic oxygen that poison active surface sites.

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**Figure 2.** The overall energetics for the catalytic reaction cycle 2NH\(_3\)(g) + O\(_2\)(g) → N\(_2\)(g) + H\(_2\)O(g) for the mechanisms controlled by chemisorbed atomic oxygen. The two cycles shown follow two different reaction paths: (a) sequential hydrogen transfer steps to form NH\(_2\)\(^*\), NH\(^*\), N\(^*\), and OH\(^*\) and (b) simultaneous transfer of two hydrogens to form gaseous water and NH\(^*\) directly. This is followed by a final H- abstraction to form N\(^*\). The first two steps in both cycles refer to the adsorption of NH\(_3\) and atomic oxygen. The intermediate steps SR1-SR4 are the hydroxyl groups to form gaseous water and NH\(^\bullet\) directly. This is followed by a final H- abstraction to form N\(^\bullet\). The first two steps in both cycles refer to the adsorption of NH\(_3\) and atomic oxygen. The intermediate steps SR1-SR4 are the hydroxyl groups to form gaseous water and NH\(^\bullet\) directly. This is followed by a final H- abstraction to form N\(^\bullet\). The first two steps in both cycles refer to the adsorption of NH\(_3\) and atomic oxygen.
The initial surface dissociation step

$$\text{NH}_3(a) + \text{O}_2(a) \rightarrow \text{NH}(a) + \text{O}(a) + \text{H}_2\text{O}(g)$$

(4)

is now thermodynamically quite favorable at $-184 \text{ kJ/mol}$.

**Activation Energies and Reaction Mechanism.** In general, the overall reaction energies for each step suggest that recombinative hydroxyl removal as water and the initial hydrogen abstraction step are the most likely candidates for rate-controlling. While the former process is the more endothermic, entropy is an important factor. At higher temperatures, recombinative desorption is considerably enhanced due to the entropic contributions, and the surface reaction steps become more predominant in controlling the mechanism. We, therefore, analyzed the activation barriers for each of the initial surface reaction steps shown in eqs 1-4.

**Atomic Oxygen.** We reported the activation barrier for ammonia dissociation in the presence of atomic oxygen in a previous communication. The dissociation was modeled as a two-step process: N–H bond activation with the nitrogen fixed...
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Figure 4. The overall energetics for the catalytic reaction cycle 2NH₃(g) + O₂(g) → N₂(g) + H₂O(g) for the mechanisms controlled by molecular oxygen. The two cycles shown follow two different reaction paths: (a) sequential hydrogen transfer steps to form NH₂*, NH*, N*, and OOH* and (b) simultaneous transfer of two hydrogens to form gaseous water and NH₂ directly. This is followed by a final H-abstraction to form N*. The first two steps in both cycles refer to the adsorption of NH₃ and molecular oxygen. The intermediate steps SR2-SR6 are the hydrogen surface reaction steps depicted in Table 4. The final terms, N₂(g), NO*, NO(g), OOH* Diss., and H₂O removal, refer to recombinative desorption of nitrogen adatoms to N₂(g), surface reaction of N* and O* to form NO*, recombinative desorption of surface OOH* to form O* and OH*, and recombination of surface hydroxyl groups to form H₂O(g), steps, respectively.

at the onefold site and surface diffusion of the NH₂* product to the stable twofold binding site. The initial N–H activation step was found to be controlling with an activation barrier of +132 kJ/mol. The surface reactants, transition complex, and surface products for this reaction along with the associated activation energy and overall reaction energy are depicted in Figure 5.

The detailed reaction coordinate for the simultaneous activation of two NH bonds by atomic oxygen was analyzed here on the Cu(6) cluster. The N–H bond stretch was chosen as the reaction coordinate. All other adsorbate-surface variables were optimized at each position along this coordinate. The results are shown in Figure 6. As the two N–H bonds stretch, the resulting NH* fragment rotates toward the threefold site which is adjacent to the H₂O species that is being formed. This is illustrated by the decreasing N–H surface angle along the reaction coordinate in Figure 6. All three oxygen–copper bonds elongate as gaseous water is formed. The two O–Cu bonds closest to the nitrogen-bound adsorbate, however, increase somewhat more substantially. At the N–H stretch of 1.5 Å, for example the oxygen–copper distances are 2.10 Å for O–Cu2 and O–Cu3 and 1.874 Å for O–Cu6. This increase in the bond distance is attributed to the repulsive interaction between the oxygen which still maintains some interaction with the surface and the neighboring NH fragment which is forming. The release of water to the gas phase essentially removes this repulsive interaction and thus allows the surface inside to be formed toward the front of the cluster. Had the water remained adsorbed, the NH* species would have more likely moved toward the back of the cluster which is in accord with the principle of least metal atom sharing. The transition state occurs as the N–H bond is stretched 0.32 Å from its original equilibrium distance (1.028 Å). This corresponds to an activation barrier of 204 kJ/mol. However, as was discussed earlier, this simple Cu(6) cluster underpredicts the NH bond strength and therefore provides only a lower bound on the energy barrier. The true activation energy is therefore >204 kJ/mol, which makes this an unlikely reaction path.

Molecular Oxygen. The N–H bond stretch was also chosen as the reaction coordinate for the abstraction of a single hydrogen by molecular oxygen to form OOH*. Reaction coordinate calculations were performed on both the Cu(6) and Cu(8,3) clusters, and the results are shown in Figure 7 and 8, for the two respective clusters. As discussed earlier, O₂ initially prefers the near-perpendicular adsorption geometry in the presence of NH₃. The O–O surface angle decreases very slightly as the reaction proceeds to shorten the distance and hydrogen must travel to the neighboring oxygen. However, once the hydrogen begin binding with the oxygen, the O–O surface angle returns to the perpendicular state. In general, the results in Figure 7 and 8 demonstrate...
that as the N–H bond is elongated the Cu–N distance decreases, the O–Cu distance decreases and the O–O distance increases. This corresponds with the development of a stronger Cu–N, O–Cu bonds and a weaker O–O bond and follows along the lines of the BOC principle. The low activation energy (+29 kJ/mol) and the shorter N–Cu bonds on the Cu(6) cluster are clear evidence of the Cu(6) model's overprediction for Cu–N binding and hence underprediction for N–H bond breaking. This helps to establish the role of the Cu(6) for providing lower limits on N–H dissociation barrier. The Cu(8,3) system yields a much more accurate representation and predicts an activation energy of +67 kJ/mol. The results presented in Figure 8 are from DGAuss calculations. The activation energy result was repeated using the ADF program and found to be within 5 kJ/mol (+62 kJ/mol).

This reaction is quite late in the N–H stretch. It is not until the surface amide diffuses to the more favorable twofold site and rotates about the surface-nitrogen bond to a more favorable bonding arrangement where the remaining N–H bonds are perpendicular to the Cu–Cu bond (as is shown in the final two structures in Figure 8) that the reaction energy becomes favorable (ΔE < 0). From Figure 6 it is clear that the NH₂ prefers the twofold site which is one copper atom removed from the binding of the OOH* species. This arrangement is similar to that shown in case I above, whereby the through-metal interaction stabilizes the binding of both adsorbates. One last point worth noting is the difference between the three Cu–O bond lengths. In path 2 above, where atomic oxygen abstracted two hydrogens, the water and NH* species were in close proximity, and therefore repulsive interactions lead the Mu2 and Mu3 to be somewhat longer. In this path (eq 3), however, the through-metal interaction increases the binding and hence decreases the O–Cu2 and O–Cu3 bond lengths.

In the final mechanistic path, eq 4, molecular oxygen acts to simultaneously abstract two hydrogens from the coadsorbed ammonia to form water directly. The N–H bond stretch was once again chosen as the reaction coordinate. The results are depicted in Figure 9 on the Cu(6) cluster and suggest that the activation energy for this path is greater than +172 kJ/mol. The activation barrier for this path is considerably higher than that for the abstraction of a single hydrogen. This is attributed to the longer initial N–H distances (1.66 vs 2.41 Å for path 4) and the necessity to activate two bonds rather than one. In fact, if one compares the N–H bond length for the initial adsorbates in path 3 (1.098 Å on Cu(6) and 1.07 Å on Cu(8,3)) to those for path 4 (1.021 Å on Cu(6) and 1.03 Å on Cu(8,3)), it appears that the NH bond for path 3 is initially activated as a result of the short initial O–H distance. There is no initial activation, however, for path 4.
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Figure 8. Reaction coordinate calculations for the path 3 dissociation of ammonia over Cu(8,3). Molecular oxygen abstracts a single hydrogen from the coadsorbed ammonia to form NH2* and OOH* intermediates. The N–H stretch is taken as the reaction coordinate. ΔE is the change in energy with respect to the initial adsorbed state and is reported in kJ/mol. Bond lengths are reported in Å.

An energetically more favorable reaction coordinate for this path was found to be the stretch of the weak oxygen–oxygen bond. A summary of the reaction coordinate results for this step are provided in Figure 10. The activation energy for this path is +134 kJ/mol, and the transition state appears to be early in the N–H stretch (+0.032 Å) but late with respect to the O–O stretch (+1.02 Å). As the O–O bond is stretched, the Cu–O bond becomes shorter, and the two hydrogens are separated from the chemisorbed ammonia to form water in the gas phase and a surface imide which sits threefold. The results from path 2 (Figure 6) demonstrated that the NH* preferred to move forward on the cluster toward the releasing water. However, in path 4, as is shown here in Figure 10, the surface imide actually prefers to move toward the threefold site at the back of the cluster, away from the threefold chemisorbed oxygen. Had the NH* moved to the threefold site directly in front of it both the NH* and O* would have two metal atom neighbors, a case II scenario, which was found to be considerably repulsive.

Nitric Oxide Formation. While recombination of nitrogen and oxygen adatoms and desorption as NO is thermodynamically less favorable than recombinative desorption of N2, the substantially higher oxygen surface coverages make the NO path much more likely. To further investigate this we computed the detailed potential energy profile for the adsorption and dissociation of NO over the Cu(8,4) cluster. Figure 11 depicts the important structural results and a summary of the corresponding energetics, while Figure 12 provides a more complete potential energy profile. NO adsorbs molecularly at the threefold adsorption site (N–Cu and N–O bond distances of 2.18 and 1.15 Å, respectively) and releases 83 kJ/mol. This adsorption energy agrees quite well with the experimental value of 105 kJ/mol reported by Conrad and Ertl for NO on Ni(111). Nickel is expected to be more active than copper due to vacancies in its valence electron band, and thus the adsorption energy on Ni(111) should be somewhat greater. The computed adsorption energy also follows the ab initio MCSCF results reported by Bagus et al. for NO atop Cu(1,4) which indicate adsorption energies of 80–100 kJ/mol.

A linear transient between the optimized NO reactant and the dissociated product state was chosen as the reaction coordinate for NO dissociation reaction. The activation barrier was
energies (in atomic units) at particular points along the coordinate, and state. Bond lengths are reported in Å.

The various reaction paths for the dissociation of ammonia studied herein are summarized in Table 5. While the Cu(111) surface helps to lower both the endothermicity and the activation barrier for ammonia dissociation considerably, the substantial activation barrier and reaction enthalpy indicate that little dissociation will occur over the clean Cu(111) surface. The presence of atomic oxygen decreases the activation barrier and the endothermicity even further. The activation energy is now low enough that moderate dissociation is likely at high temperatures and very low surface coverages. The high exothermicity which accompanies the dissociative adsorption of oxygen, however, indicates that chemisorbed oxygen will poison the surface and impede ammonia dissociation kinetics. All of these results are consistent with the experimental findings presented by Roberts and colleagues on Cu(110)8,11,12 and Cu(111)14 where at low coverages of preadsorbed oxygen, ammonia dissociation occurs but at a rather slow rate. At higher coverages the ammonia dissociation path shuts down, and chemisorbed oxygen overlayers poison the surface.

Of the two candidates of atomic-oxygen-assisted dissociation, we find that the sequential hydrogen dissociation path (eq 1) is more likely. In this path, the ammonia hydrogen and surface O* are significantly closer, only one N-H bond needs to be cleaved, and the strong Cu-O bond does not have to be broken. The activation barrier is over 72 kJ/mol lower in energy, and the overall reaction energy 44 kJ/mol is more exothermic for the mechanism based on single hydrogen transfer steps over the mechanism for the simultaneous transfer of two hydrogens.

The results presented above and summarized in Table 5 clearly indicate that molecular oxygen acts as chemical precursor in the dissociation of ammonia. Both the sequential and simultaneous hydrogen abstraction paths (paths 3 and 4) are largely exothermic. Molecular oxygen is weakly bound perpendicular to the Cu surface with an adsorption energy of ~10 kJ/mol. Electron back donation from the surface to the π* orbital of O2 weakens the O-O bond which is reflected by the longer O-O distance (O2*) = 1.36 Å, O2(g) = 1.21 Å). The weakly bound molecular oxygen can then readily abstract hydrogen from the coadsorbed ammonia. The initial distance between the ammonia hydrogen and the oxygen atom that is transferred to is now only 1.66 Å for path 3, whereas for the atomic oxygen abstraction the distance was 2.45 Å. While the overall thermodynamics favor the simultaneous hydrogen abstraction to form water directly, the activation barriers indicate that the sequential path which is 67 kJ/mol less costly predominates. The relatively close initial O-H distance and the need to break only one NH bond once again are likely explanations why path 3 is favored over path 4. In addition, these explanations help to clarify the change in the reaction coordinate in going from path 3 (controlled by the N-H stretch) to path 4 (controlled by O-O stretch). In path 3, the N-H stretch is activated by the close proximity N-H hydrogen and the neighboring oxygen. In path 4, however, the oxygen-hydrogen distance is significantly longer thus reducing N-H bond activation. NH bond scission is therefore much more costly than stretching of the weak O-O bond. In the end, the energy required for the significant O-O stretch in path 4 is greater than the energy needed to cleave the activation NH bond in path 3.

Under the typical UHV conditions of Roberts's11,14 and at low oxygen partial pressures, the apparent activation energy for NH3 dissociation is really a measure of both the intrinsic NH bond scission and equilibrium adsorption constants for NH3 and O2. For path 4 the apparent activation barrier is +47 kJ/mol, whereas for the more likely path 3 the apparent activation barrier is 0 kJ/mol. Path 3 is essentially a nonactivated route for molecular oxygen dissociation of ammonia under these conditions. Molecular oxygen dissociation may be in direct competition with this path for the consumption of O2. Boostma et al.,13 however, determined that the apparent activation for O2 dissociation over...
Figure 10. Reaction coordinate calculations for the path 4 dissociation of ammonia over Cu(8,3). Molecular oxygen acts to abstract a two hydrogen from the coadsorbed ammonia to form NH*, O*, and gaseous water. The O–O stretch is taken as the reaction coordinate. ∆E is the change in energy with respect to the initial adsorbed state and is reported in kJ/mol. Bond lengths are in Å.

Cu(111) is an activated process with a barrier of 8–17 kJ/mol. Molecular oxygen–ammonia dissociation is therefore the favored route. All of these results directly correspond to the experimental evidence from Roberts who demonstrated that ammonia readily dissociates in the presence of coadsorbed oxygen and confirm the role of a transient oxygen precursor. At higher partial pressures of oxygen the surface essential becomes poisoned by chemisorbed atomic oxygen species and the rate severely inhibited. This is also consistent with Roberts findings.

Finally, the “hot” atomic oxygen mediated mechanism proposed by Roberts (path 5) which cannot be completely ruled out is somewhat inconsistent with our findings. The scission of the NH bond by reaction with hot atomic oxygen requires that energy from the transient oxygen be transferred to the NH bond. The timescale for this step, however, is comparable with that required for thermal equilibration. Thus, the reaction via the molecular precursor path appears more likely.

Conclusions

Ammonia adsorsbs on a model of the Cu(111) surface at −32 kJ/mol. In the absence of coadsorbed oxygen species, the dissociation of ammonia over copper is energetically an unfavorable process with an activation barrier of +344 kJ/mol and an overall reaction enthalpy of +176 kJ/mol. In the presence of coadsorbed oxygen species, lateral interactions are important in determining both the adsorption and the activation energies. Atomic oxygen promotes N–H bond activation, by lowering the activation energy to +132 kJ/mol. The strong atomic oxygen surface bond, however, leads to poisoning of surface sites which inhibits the overall dissociation kinetics. The atomic pathway in which hydrogen is transferred sequentially to form OH* is favored over the simultaneous path leading to the direct formation of water.

Molecular oxygen adsorsbs favorably to Cu(111) both in parallel and perpendicular orientations to the surface. The parallel mode is a precursor state for O₂ dissociation where a significant charge transfer from the surface to O₂ π* orbital weakens the oxygen–oxygen bond as is demonstrated by the long O₂ bond (1.43 Å). The perpendicular mode, on the other hand, is the precursor state for ammonia dissociation. Coadsorbed molecular oxygen enhances the adsorption of ammonia by 17 kJ/mol and acts to lower the barrier for N–H bond activation. The sequential hydrogen abstraction pathway, which leads to the OOH* formation,
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Surface Reactants

\[ \Delta E_{\text{ads}} = -83 \text{ kJ/mol} \]

Transition State

\[ E^* = +88 \text{ kJ/mol} \]

Surface Products

\[ \Delta E = -46 \text{ kJ/mol} \]

Figure 11. Summary of reaction coordinate calculations for NO dissociation. The surface reactant state refers to the adsorption of NO on Cu(8,4). The transition state energy and the overall reaction energy are measured with respect to the gas phase, i.e., NO(g) and clean Cu-(8,4). Bond lengths are reported in Å.

intermediate, while 100 kJ/mol less exothermic than the path for the simultaneous hydrogen transfer, is kinetically favored by 67 kJ/mol.

The sequential-hydrogen-transfer-transient-molecular-oxygen path is the most likely of all those studied. Its apparent activation energy is zero, i.e., a nonactivated process. All paths controlled by transient atomic oxygen, including the not transient oxygen case proposed by Roberts, require at least an initial activation barrier of +17 kJ/mol. At low to moderate temperatures the hot atomic oxygen path will be kinetically insignificant in comparison with the molecular path. As the temperature increases, however, these other pathways also become feasible. Chemisorbed atomic oxygen is formed, which in turn inhibits ammonia dissociation kinetics. At high enough temperatures or coverages the dissociation reaction eventually shuts down due to the formation of strong chemisorbed oxygen overlayers.

Finally, based solely on overall enthalpy, the recombination of nitrogen adatoms and desorption of N\textsubscript{2} was found to be favored over the recombinative removal of NO. However, under reaction conditions both the entropy and the selective collision frequency favor NO formation.

Acknowledgment. We thank Professor E. J. Baerends and his colleagues at the Free University of Amsterdam for their invaluable technical input and the use of the ADF program and G. J. C. S. van de Kerkhof and Dr. P. R. Davies for their initial efforts on this project. This work was supported by the Dutch Science Foundation (Stimulus project) and Eindhoven University of Technology. We kindly acknowledge the computational resources allocated from the National Computing Facilities (NCF) Foundation under project SC-183.

Supplementary Material Available: Tables of exponential coefficients for ADF basis and fit functions (1 page). This material is contained in many libraries on microfiche, immediately follows this article in the microfilm version of the journal, and can be ordered from the ACS; see any current masthead page for ordering information.