Comprehensive Experimental and Theoretical Study of the CO + NO Reaction Catalyzed by Au/Ni Nanoparticles


INTRODUCTION

The CO + NO reaction was studied on extended single crystal surfaces, first time over 40 years ago and many investigations have been published in the succeeding decades. The reaction continues to attract attention, re-investigations have been published in the succeeding decades. In addition to its key role in the abatement of automotive emissions, it is an attractive subject for both experimental and theoretical study thanks to its apparent relative simplicity. In the case of well-defined extended single crystal surfaces, experimental results can be comparatively simple to interpret. However, with practical catalysts consisting of metal nanoparticles dispersed on metal oxide supports, additional complexities arise and comprehensive reviews devoted to the use of monometallic catalysts are available. Bimetallic systems introduce important additional flexibility into catalyst design and optimization and have therefore been extensively investigated for a wide variety of organic, petrochemical, and electrochemical reactions, as well as the CO + NO reaction.

With regard to the CO + NO reaction, the use of gold–nickel nanoparticle catalysts was recently patented by Toyota, on the basis of work conducted in both Cambridge, UK and Seville, Spain.
It was found that the methods previously used for such purposes are susceptible to coking. It is possible the nature of the active sites remains very complicated, and there is room for further investigation.

Here we report the results of a comprehensive experimental and theoretical investigation of the CO + NO reaction catalyzed by Au/Ni nanoparticles of varying composition. Experimentally, particular attention was given to the method of sample preparation and, especially, the oxidative pretreatment applied to remove extensive polymeric residues, whose presence can radically affect the nature of the working catalyst. It was found that the methods previously used for such catalysts (e.g., 300 °C in air) are inadequate. Although the catalysts underwent some sintering as a result of calcination in air, they retained their bimetallic nature and were stable against further sintering during subsequent reduction and catalytic reaction. Thorough pretreatment is an important point that is frequently overlooked or unreported in studies of metal catalysts prepared by colloidal routes, whose conclusions should therefore be regarded with circumspection.

X-ray absorption spectroscopy (XAS) results obtained under in situ reaction conditions provided key information about the structure and electronic state of the working catalysts. Specifically, these observations elucidated the nature, frequency and impact on Ni oxidation state of the Au–Ni interaction and showed that the extent of this interaction correlated well with catalytic activity. Importantly, they also showed that substantial and reversible changes in Au–Ni intermixing occurred as a function of temperature, essential information for a correct interpretation of catalytic data. This is in contrast to ex situ powder X-ray diffraction (PXRD) and X-ray photoelectron spectroscopy (XPS) methods, which obviously cannot provide structural information pertinent to reaction conditions.

Density functional theory (DFT) calculations on both smooth and stepped surfaces showed that, relative to pure Ni, the activation barrier to N + N recombination was progressively lowered by increasing amounts of Au up to ~50%, in good accord with experiment. This lowering in the activation barrier was overwhelmingly due to the destabilization of adsorbed N atoms, which bind much more strongly to pure nickel than to bimetallic sites. Moreover, this process, rather than NO dissociation, was overall rate determining. DFT calculations also showed that NiO should be catalytically inert in the CO + NO reaction. The importance of Au–Ni intermixing at elevated temperatures (~550 °C) in catalyzing the CO + NO reaction, identified here, has not been reported for other reactions catalyzed by combinations of gold and nickel, such as low temperature CO oxidation (<100 °C), where a Au-NiO core–shell structure is proposed, or methane steam reforming, where Au blocks step sites that are susceptible to coking.

Table 1. Composition and Size Information for Unsupported Nanoparticles (Top) and Supported Nanoparticle Samples (Bottom)

<table>
<thead>
<tr>
<th>catalyst and nominal metal ratio/%</th>
<th>Unsupported nanoparticles</th>
<th>Supported samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Au-Ni ICP molar ratio of unsupported nanoparticles</td>
<td>total metal loading</td>
</tr>
<tr>
<td>Au100</td>
<td>3.1 (±1.2)</td>
<td>11</td>
</tr>
<tr>
<td>Au50Ni50</td>
<td>3.7 (±1.2)</td>
<td>7</td>
</tr>
<tr>
<td>Au75Ni25</td>
<td>2.4 (±1.3)</td>
<td>15</td>
</tr>
<tr>
<td>Ni100</td>
<td>3.1 (±1.0)</td>
<td>8</td>
</tr>
<tr>
<td>Au100/α-Al2O3</td>
<td>3.1 (±1.2)</td>
<td>11</td>
</tr>
<tr>
<td>Au50Ni50/α-Al2O3</td>
<td>17 (±10)</td>
<td>7</td>
</tr>
<tr>
<td>Au75Ni25/α-Al2O3</td>
<td>24 (±7)</td>
<td>15</td>
</tr>
<tr>
<td>Ni100/α-Al2O3</td>
<td>11.0 (±3.4)</td>
<td>8</td>
</tr>
</tbody>
</table>

“Presented with respect to Au amount expected for ease of comparison. Uncalibrated values as a guide to relative composition only; values obtained are a ratio of Au M/NI L edges obtained from individual particles (see exemplar images in the Supporting Information) and the distribution given is one standard deviation. Distribution indicated based on one standard deviation of particle size count. Target loading ~0.2 mmol g⁻¹. Obtained by Scherrer analysis from the Au (311) reflection, hence Ni only sample not shown. Too small/oxidic to measure against alumina support. Japan.11 Beniya et al. used a Au/Ni(111) single crystal model system to examine the effects of surface composition on the reaction, concluding that the beneficial effect of Au incorporation was due to a reduction in the activation energy for the reaction N(a) + N(a) → N₂(g), thought to be the overall rate limiting step. Their subsequent in situ study of a single catalyst consisting of ~1:1 Au–Ni bimetallic particles supported on silica confirmed the effect of Au addition, but left open a number of key questions, including the actual condition of catalyst before and after reaction, the nature of the Au–Ni interaction, its dependence on composition, and the identity of the catalytically active sites. As the authors themselves concluded “However, the nature of the active sites remains very complicated, and there is room for further investigation.” The object of the present paper is to identify or clarify the key issues.

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EXPERIMENTAL METHODS
Catalysts were prepared on α-alumina from colloidal nanoparticles, purified by thorough washing, and by post deposition calcination to remove residual synthetic agents. A range of ex situ characterization (transmission electron microscopy [TEM], energy dispersive spectroscopy [EDS], XPS, inductively coupled plasma–optical emission spectroscopy [ICP-OES], PXRD) and in situ synchrotron XAS were used to characterize the structure of the catalyst before, during, and after reaction, and these results compared to catalyst testing conducted in a 1/4″ microreactor using both light-off and pseudo-steady-state measurements. The full details of materials preparations, characterization, and testing are provided in the Supporting Information.

DFT METHODS
Calculations were carried out using the VASP 5.4 program and full details are given in the Supporting Information. Briefly, a Ni (100) surface was modeled by a (3 × 3 × 5) supercell containing 45 atoms and a 1 Å vacuum space between the slab surfaces. This was large enough to accommodate reactants and products and to examine different Au/Ni compositions. A larger (6 × 3 × 5) supercell model was used to introduce surface steps in order to simulate the behavior of nanoparticles embodying low-coordination sites. In the geometry optimizations, all layers of the slab were allowed to relax and possible changes in cell parameters at the highest Au content were found to be negligible. Transition state calculations were done using the climbing image version of a nudged elastic band algorithm. Note, the entropic contribution of dinitrogen gas (≈0.65 eV) is omitted from energy landscapes, as it contributes a constant shift to the 0 eV reference and is therefore not significant in the calculation of energy barriers.

RESULTS AND DISCUSSION
A series of five catalysts with different Au:Ni ratios were prepared from colloidal nanoparticles, as described earlier.11,12 Electron micrographs and size distributions of the nanoparticles before deposition onto the alumina catalyst support are shown in the Supporting Information and are summarized in Table 1. Table 1 (top, columns 2 and 3) additionally shows the Au:Ni ratio obtained by ICP analysis for each composition and the Au:Ni area ratios for single particles measured by EDS, which follow the same general trend, thus confirming the presence of both Au and Ni in individual nanoparticles (further details, including exemplar images showing the EDS measurements of separate particles, are given in the Supporting Information).

The catalysts were prepared from the colloidal nanoparticles by deposition onto the support, calcination and reduction. After nanoparticle deposition onto the support, temperature programmed oxidation (TPO, Supporting Information) showed that very substantial quantities of organic residue remained, arising from the nanoparticle synthesis method, despite extensive washing, as described in the experimental section in the Supporting Information. This was evidenced by an ~20% mass loss during the TPO measurements and the coincidence in temperature between the maximum in the
differential weight of the sample, and the production of both 
CO₂ and H₂O confirms that this weight loss is due to oxidation 
of hydrocarbon material. The need to remove residual polymer 
capping materials is an important and often overlooked point 
when catalysts are derived from colloidal nanoparticles, 
especially in view of the strikingly high ratio (>10:1) between 
the mass of hydrocarbon and the mass of actual metal present 
in these samples. Omission of this step could compromise both 
the validity of results and their interpretation. Because the 
TPO traces exhibited a high temperature shoulder extending to 
ca. 500 °C, all catalysts were calcined at 500 °C in an O₂/He 
flow for 1 h, to ensure complete removal of the residual 
hydrocarbon. This is especially important with respect to light-
off measurements where CO₂ production and NO con-
sumption rates were measured with catalysts derived from 
these nanoparticles. XPS of the calcined catalysts showed only 
a very small C 1s signal (see the Supporting Information), 
corresponding at most to 0.5 wt % (which likely results from 
sample handling, as is common in XPS). It is of fundamental 
importance that after exposure to reaction conditions no 
carbon was detected by XPS. The removal of capping agent by 
high temperature oxidative pretreatment was accompanied by 
a significant increase in particle size from ~3 to ~10 nm for 
the bimetallic samples, as indicated in Table 1 and by the 
micrographs in Figure 1. These materials were the uncapped, 
stable precatalysts used in this study. After reduction and 
subsequent catalytic testing, the particles maintained a stable 
size (Table 1 and Figure 1). For completeness, estimated sizes 
based on the fcc Au (311) reflection at ~78° 2θ in the PXRD 
patterns of the samples are shown in Table 1. (While the 
trends in PXRD sizes broadly agree with the numbers obtained 
from TEM, some caution should be exercised making 
comparisons as this reflection only accounts for the fcc Au-
like crystallite within the nanoparticles, and so may exclude 
some nickel-rich components, and the comparison is between a 
number and a volume distribution.) Notably, the pure gold 
sample exhibits rapid agglomeration during calcination but 
redisperses again under reaction conditions, a fact attested to 
by both microscopy (a local technique) and PXRD (a bulk 
technique). As this only occurs for the pure gold sample, it is 
not of primary importance to the system at hand. However, it 
could speculatively be suggested that this behavior in H₂ or 
CO and NO is similar to that witnessed in the presence of 
iodine during methanol carbylation reactions.16

As shown by the EDS elemental composition ratios in Table 
1, the bimetallic nanoparticles did indeed contain both Au and 
Ni; the spatial distribution of the metals and their oxidation 
states are considered below. The substantial changes in particle 
size that occurred as a result of preprocesston the nanoparticles 
to form the active catalysts (Table 1) underline the importance 
of cradle-to-grave characterization of nanoparticle catalysts, in 
contrast to the common practice of focusing attention on 
nascent materials as opposed to those that are actually used for 
and emerge from catalytic experiments.

Ex situ XPS and PXRD measurements were made at 
different stages of the catalysts’ fabrication and evolution (from 
nanoparticle precatalysts to the calcined, reduced and used 
forms). Figure 2 shows XP spectra obtained after calcination: a 
gradual shift in the Ni 2p₃/₂ signal to higher binding energies 
was apparent as the Au content was increased (full spectra 
shown in the Supporting Information). This is consistent with 
electron transfer from Ni to Au and in keeping with their 
Pauling electronegativities of 1.91 and 2.54, respectively.

Figure 2. Ni 2p XP spectra expanded in the region of Ni 2p₃/₂ as 
a function of elemental composition of the five α-Al₂O₃ supported 
samples after calcination, the nominal molar % composition is shown 
in the figure legend. The vertical dashed line is a guide to the eye, and 
it highlights the gradual shift to higher binding energies as the gold 
content in the sample increases. The spectra are calibrated to the 
Al 2p emission, which has a value of 74.0 eV for Al in the Al₂O₃ support. 
Signal intensities are as acquired and not normalized.

Smaller shifts were visible in the Au 4f and 4d spectra (Supporting Information), 
with the effect possibly being diminished due to the larger number of electrons involved. 
Attribution of the changes in Ni 2p binding energy to a core 
level shift is supported by the same trend being observed for the 
Ni LMM (3p → 2p) Auger electron kinetic energies (Supporting Information). For the pure Ni catalyst and the 
Au₅₀Ni₅₀ catalyst, XP spectra were also acquired after (i) 
reduction and (ii) catalysis using a high pressure cell attached 
to the XP spectrometer (see experimental details in the 
Supporting Information). This eliminated artifacts that would 
result from air handling of the sample. An increase of ~0.7 eV 
in the Ni 2p₃/₂ binding energy of the bimetallic Au₅₀Ni₅₀ 
catalyst relative to the pure Ni catalyst was consistently 
observed for both reduced and oxidized samples. This implies 
that Au withdraws electrons from Ni, thus inhibiting its 
oxidation. Nevertheless, for both pure Ni and the 
bimetallic 
Au₅₀Ni₅₀ catalyst the Ni 2p₃/₂ binding energy of the 
postcatalysis sample was much closer to that of the oxidized 
form than the reduced form of the catalyst, indicating 
substantial Ni oxidation in this material. More direct light is 
shed on this important issue by the XAS results obtained under 
actual working conditions (in situ), as discussed below (and 
seen in Figures 6 and 7).

While XPS substantiates the occurrence of Ni → Au 
electronic transfer, it provides no indication of the range of this 
effect, nor of the nanoparticle structure. PXRD was therefore 
used to investigate the possibility of alloy formation within 
nanoparticles. The PXRD patterns (Supporting Information) 
were dominated by features attributable to the α-Al₂O₃ support, 
but a number of reflections attributable to fcc Au 
were apparent. As expected, no Ni or NiO reflections were 
visible against the Al₂O₃ background, due to the much smaller 
atomic number of Ni versus Au. Given the different atomic 
radii of the two metals, alloy structures should show substantial 
shifts in 2θ positions for the metallic reflections (Vegard’s law).

Figure 3 summarizes the PXRD data for the different samples 
before and after calcination, and also after catalytic testing; also
shown are the expected position of the reflections for each composition. Clearly, at every stage, the samples showed no evidence of significant alloy crystal phases when examined ex situ: only a very small incorporation of Ni into Au was evident in line with the relative concentrations in each sample. However, as we shall see, XAS measurements carried out under in situ conditions unambiguously demonstrated that extensive alloy formation occurred at 550 °C in reaction gas, and that it was reversible with temperature. Clearly, reliance on ex situ characterization techniques can be seriously misleading in the interpretation of catalytic data.

The five catalysts (based on two pure metal and three bimetallic samples) were tested for activity in the CO + NO reaction by means of light-off measurements, followed by steady state measurements recorded after stable performance was reached at 550 °C. The latter procedure was important in confirming catalyst stability and the establishment of a steady state, i.e., the observed reactivity was truly catalytic and did not result from reaction of NO with residual synthetic reagents (e.g., polymer capping agent). Note that this protocol was not followed in previous studies of this system.12,13,17 Table 2 summarizes the resulting steady state reaction data. All catalysts exhibited low NO formation at 550 °C, a temperature that falls within the typical operating window for such catalysts; very conspicuously, the Au2Ni3 and Au50Ni50 compositions delivered much higher NO conversions than the others. The similar CO and NO conversions (Supporting Information) confirm that it was indeed the intended reaction that was being studied, unperturbed by artifacts due to consumption of residual synthetic agents initially present on the nanoparticles. Light-off profiles are shown in more detail in Figure 4, which reveals some important points. The profiles refer to NO consumption, but, as shown by the full profiles for each catalytic reaction (Supporting Information), the overall process was highly selective and essentially stoichiometric with respect to CO/NO (small quantities of N2O were produced only at low temperature). Initially, all catalysts were fully reduced. The Ni catalyst showed pronounced low temperature activity, quickly deactivated, and only slowly regained activity with subsequently increasing temperature. Pure gold was almost inactive throughout. Au75Ni25 was slightly less active than pure nickel, but it did not display the low temperature activity shown by pure Ni. The two most active catalyst compositions (Au33Ni67 and Au50Ni50) lit-off at a similar temperature and exhibited similar kinetics, with a steep rise in activity at ~400 °C. With increasing temperature, despite similar masses, loadings, and reactor geometry (i.e., the same mass transfer characteristics), the Au50Ni50 composition was more active at the highest temperatures, reaching saturation before Au33Ni67. These trends are explained well by the in situ XAS experiments described below (and seen in Figures 6 and 7).

It has been proposed that the beneficial effect of Au in enhancing the activity of Au/Ni catalysts for the CO + NO reaction was due to inhibition of Ni oxidation, a conclusion based on experiments with a Au50Ni50 nanoparticle system similar to that used here.12,17 Subsequently, based on single crystal model studies, the same group proposed a different explanation; namely, that Au/Ni electronic interaction decreased the activation barrier to N + N recombination, thought to be the rate limiting step.12 Fundamental under-

Table 2. Steady State Conversion and Selectivity for Five Catalysts Tested under a NO–CO Feed

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>NO Conversion (%)</th>
<th>CO Conversion (%)</th>
<th>N2O Selectivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni100/α-Al2O3</td>
<td>25</td>
<td>25</td>
<td>0.1</td>
</tr>
<tr>
<td>Au33Ni67/α-Al2O3</td>
<td>100</td>
<td>98</td>
<td>0.0</td>
</tr>
<tr>
<td>Au50Ni50/α-Al2O3</td>
<td>100</td>
<td>100</td>
<td>0.0</td>
</tr>
<tr>
<td>Au2Ni3/α-Al2O3</td>
<td>20</td>
<td>19</td>
<td>0.8</td>
</tr>
<tr>
<td>Au100/α-Al2O3</td>
<td>6</td>
<td>6</td>
<td>0.3</td>
</tr>
</tbody>
</table>

*1000 ppm of NO, 1000 ppm of CO, He balance. Each catalyst was generated by pretreating immobilized nanoparticle samples by oxidation in synthetic air (O2 21%, He balance) at 300 °C (heating rate = 10 °C min−1, hold time = 1 h) before cooling and switching gas feed to 5% H2, He balance for reduction. Reduction was conducted in situ by heating at 10 °C min−1 to 550 °C for 1 h and cooling under 5% H2 in He before admitting the reactive gases. The amount of 60 mg of sample was loaded into the reactor in each case. Total gas flow rate = 100 sccm, GSV = 1.6 L min−1 gcat−1.
standing of the origin and mechanism of activity promotion by Au in this important reaction is essential for rational catalyst development. Theoretical calculations described below establish, among other things, that reduced Ni sites are required and that the Ni → Au electronic interaction does indeed promote N + N recombination. The significance of both factors will be discussed. Importantly, in situ XAS measurements enabled conclusions to be drawn about both the Ni oxidation state, which does not correlate with activity, and the frequency of Au–Ni interactions present under reaction conditions.

X-ray absorption near edge structure (XANES) spectroscopy was used to probe the Ni oxidation state in the samples following calcination, reduction and after being used as catalysts. Figure 5 shows the Ni K-edge XANES region for all the Ni-containing samples, which were loaded into the XAS cell in their calcined form (measured in He at 25 °C, black spectrum), reduced at 550 °C in H2, cooled and then exposed to a CO + NO feed; spectra were acquired at different temperatures during this sequence, as shown. Typically, three or more spectra were acquired under each set of conditions to confirm establishment of a stable state and then averaged. It is immediately apparent that the so-called white line (peak height at the edge), whose intensity is commonly taken as indicating oxidation state, changed abruptly between calcination (black), where its high intensity signifies extensive Ni oxidation, and subsequent high temperature reduction in hydrogen (red), where its strong attenuation indicates a mostly reduced form of Ni. Exposure to reaction conditions resulted in rapid oxidation of the pure Ni sample while in contrast the Au50Ni50 sample remained substantially reduced; both Au25Ni75, which was relatively active, and Au75Ni25, which was no more active than pure Ni, became significantly oxidized. However, for a number of metals it has been shown that the shift in edge position provides a more reliable gauge of oxidation state.19 The observed edge positions under reaction conditions (relative to Ni foil) for a range of temperatures are illustrated in Figure 6. These indicate that nickel in Au50Ni50 was the most reduced (∼10.3 eV versus Ni foil), followed by Au75Ni25 and Au25Ni75 (∼11.3 and ∼11.7 eV, respectively), and finally pure Ni (∼12.5 eV). While this analysis shows that the presence of Au definitely did contribute to moderating the oxidation state of Ni, there is no significant correlation with the catalytic activity data, where Au50Ni50 and Au25Ni75 were both substantially better than the essentially inactive Au75Ni25 and Ni100 samples.

In situ extended X-ray absorption fine structure (EXAFS) spectroscopy was very revealing, allowing an estimation of the coordination number for different types of metal–metal interactions under reaction conditions. In particular, it enabled distinguishing between Ni–Ni and Ni–Au (using Ni K-edge data) and between Au–Au and Au–Ni (using Au L-edge data). Fitted data from both edges shows the same general trends in coordination numbers, while Ni–O interactions shown by Ni K-edge fits (see the Supporting Information) corroborate the Ni oxidation state trends seen in the XANES

![Figure 5](image.png)

Figure 5. In situ Ni K-edge XANES spectra of the four Ni-containing samples as a function of gas conditions and indicated temperatures. Energy scale corrected to Ni foil edge at 8333 eV.
data. However, the clearest picture is provided by the coordination numbers derived from the Au L-edge (Figure 7); these were least prone to error, particularly for Au$_{25}$Ni$_{75}$

![Figure 6](image6.png)

**Figure 6.** In situ Ni K-edge positions as a function of gas conditions and temperature. Energy scale corrected to Ni foil edge at 8333 eV whose position is indicated by the dashed red line at the bottom of the graph; even under hydrogen the Ni was not quite fully reduced. The region corresponding to steady state reaction conditions is highlighted.

![Figure 7](image7.png)

**Figure 7.** EXAFS derived Au–Ni average first nearest neighbor coordination numbers derived from in situ Au L-edge measurements as a function of pretreatment and reaction conditions for the three bimetallic systems studied (full details of the fitting and fitting parameters, and corresponding Ni K-edge fits are given in the Supporting Information). In a true random alloy of fcc AuNi, each gold atom is expected to have 3, 6, or 9 Ni nearest neighbors for the Au$_{25}$Ni$_{75}$, Au$_{50}$Ni$_{50}$ and Au$_{75}$Ni$_{25}$ compositions, respectively. The error bars shown indicate the uncertainty in coordination number for the specific fit from which the data shown were derived.

data, which could not be fitted reliably at the Ni edge due to the low Ni concentration (Supporting Information). As shown in Figure 7, there is a good correlation between the most reactive catalyst and the Au–Ni coordination number. Recall that these measurements are from the perspective of the gold component. Although it may seem surprising that the Au$_{50}$Ni$_{50}$ sample has a higher Au–Ni coordination number than the Au$_{25}$Ni$_{75}$ (where there are 50% more Ni atoms to potentially interact with when measured from the gold atoms’ perspective), this observation accords with the catalytic behavior: the higher the Au–Ni coordination number, the better the catalyst.

The initially high activity of Au$_{25}$Ni$_{75}$ may reflect the different time scales of the light-off and spectroscopic measurements. Reduction in hydrogen followed by CO + NO at 150 °C resulted in an increase in Au–Ni coordination number followed by a decrease at higher temperatures. During XAS measurements, each set of conditions required ~15 min acquisition time and ~15 min for changing temperature and stabilizing the sample. This corresponds to an elapsed time of ~2 h in going from hydrogen to CO + NO at 400 °C, where reactivity differences between Au$_{50}$Ni$_{50}$ and the Au$_{75}$Ni$_{25}$ samples were apparent. In contrast, light-off measurement taken at 5 °C min$^{-1}$ required only 50 min. It therefore seems very likely that the higher activities in the low temperature region of the light-off for the Au$_{25}$Ni$_{75}$ catalyst, comparable to that of Au$_{50}$Ni$_{50}$, reflected an initially higher Au–Ni interaction frequency similar to that of Au$_{50}$Ni$_{50}$ at low temperature. Subsequently, at higher temperatures and in CO + NO, reactivity decreased due to partial Au/Ni segregation.

A final important observation was the clear decrease in Au–Ni coordination that occurred when cooling the Au$_{50}$Ni$_{50}$ sample after use as a catalyst under reaction gases; by the time this sample had cooled to 150 °C, the Au–Ni coordination number had already decreased very substantially. Indeed, the temperature-dependent increases and decreases in Au/Ni intermixing evident in Figure 7 underline the importance of in situ measurements in the acquisition of catalytically relevant information. They also explain why evidence of increased mixing was not seen in the ex situ PXRD and XPS measurements described above, which were recorded after cooling and removal of the reactive gases.

The crucial role of temperature and particle size in determining when intermixing of Au and Ni occurs is nicely illustrated by the work by Sopoušek et al., who showed how the phase diagram for miscibility of these metals changes in going from the bulk to nanometric length scales. For 13 nm particles, they calculated the depression in temperature for formation of a mixed (Au + Ni) fcc single phase versus segregation into the separate metals. Figure 8 shows their results, which are directly relevant to an understanding of our work. Also indicated in the figure are the compositions of our nanoparticles and the reaction temperatures that we used.

![Figure 8](image8.png)

**Figure 8.** Phase diagram for Au–Ni alloying versus separate fcc phases as a function of composition (data taken from ref 17). The gray dashed line is the thermodynamic phase boundary in the bulk, and the solid black line is for a 13 nm nanoparticle. The horizontal dashed red line highlights the steady state reaction temperature of 550 °C, and the vertical lines indicate the bimetallic compositions used in this study (their nominal compositions are indicated, and the actual weight percent plotted is based on the ICP measured composition of the supported nanoparticles).
Figure 8 provides a clear indication that at 550 °C extensive intermixing of Au and Ni can be expected in all cases, and that cooling below that temperature should lead to major changes in nanoparticle constitution, in accord with our catalytic results. The coordination numbers actually identified by EXAFS are of course smaller (on average) than a true alloy (highest are 3–5), but this still points conclusively to the fact that in the working material at temperature the intermixing to gold nickel is occurring. Of course, the presence of reactive gases could alter the picture to some degree, but the essential message remains: in situ characterization of bimetallic catalysts under operating conditions is an essential prerequisite for acquisition of meaningful data that enable reliable interpretation of catalytic measurements.

DFT Results. In the case of Rh catalysts, CO + NO reaction has been studied many times and it is universally accepted that NO dissociation is the reaction-initiating step, although there is no consensus in regard to other catalytic systems, including those studied here. Another key process, namely, N + N recombination has been proposed as the overall rate-limiting process. As we shall demonstrate in this section, for Ni-based catalysts, detailed calculations provide clear confirmation that NO dissociation is not rate-controlling, whereas the associative N + N reaction is rate-controlling. We also show how addition of Au to Ni switches the rate-limiting process for N + N recombination from the association of adjacent N atoms to the surface diffusion of N. The energetics of both processes have been examined in detail, with particular reference to (i) the effects of adding Au to Ni and (ii) the effect of surface steps. Examination of the latter aspect is relevant to rationalization of nanoparticle behavior where surface properties may differ significantly from those of extended facet planes due to the presence of steps, edges, and other low coordination sites. In addition, given the experimental results, which showed substantial oxidation of Ni in certain circumstances, we investigated whether NiO could contribute to the observed catalytic behavior. All calculations were based on (111)-oriented slabs of Ni, modified by introduction of Au atoms (see the Supporting Information for details), step sites, or both. (In the interest of brevity, we...
shall refer to the N + N associative reaction as “recombination”, even though this designation is not strictly accurate.

**NO Dissociation on Ni and Au/Ni.** NO adsorption and dissociation on smooth pure Ni (111) is exothermic by 3.5 eV with an activation barrier of 1.38 eV. This is in good agreement with earlier work (1.31 eV).\(^ {22}\) The effect of introducing Au atoms was investigated by examining three different initial adsorption sites and eight different final configurations for the resulting O and N adatoms. The most favorable pathway corresponded to a somewhat reduced exothermicity (\(\sim\) 1 eV), while the activation barrier remained essentially unchanged and too high to account for the experimentally observed facile dissociation of NO on Ni on both single crystals\(^ {23, 24} \) and supported catalysts\(^ {25} \) at around room temperature. We therefore examined NO on a stepped Ni (111) surface. This was found to strongly adsorb NO and greatly reduce the barrier for dissociation to 0.24 eV. Given the much higher activation barriers found for various modes of the N + N reaction, we may conclude that the latter reaction, and not NO dissociation, was always rate-limiting.

**NO Dissociation on NiO.** As noted above, our experimental results showed that oxidized Ni was present under certain reaction conditions. We therefore tested the possibility that the reaction initiating step, NO dissociation, could occur on NiO. The (100) NiO surface was represented by a (4 \(\times\) 4 \(\times\) 4) supercell that took account of the magnetic properties of NiO, with Ni atoms ordered in alternate high and low spin layers along the [111] direction. The adsorption geometry of NO was vertically atop a Ni atom with the N atom above the surface. All other NO geometries did not produce a bound state. Four different geometries for the N+O products were tested, the most stable corresponding to the O atom above a surface Ni with the N atom in a bridge site between Ni and O atoms. This best-case process is highly endothermic (\(\sim 4.3\) eV), from which we may conclude that NiO is not a catalytically active phase for the CO + NO reaction.

\(\text{N} + \text{N} \rightarrow \text{N}_2\) **“Recombination” Reaction.** The flat Ni (111) surface was modeled by a (3 \(\times\) 3) slab containing five layers of metal and a vacuum space of 15 Å in the c direction. This generates a supercell of \(\sim 7.5\) Å in the a and b directions, similar to models used in the literature, and sufficient to obtain converged adsorption energies in systems similar to those studied here.\(^ {22, 26} \) Upon adsorption, N atoms acquire a substantial negative charge (Bader formal charge \(\sim\) N\(^{-1}\)) which gives rise to the 0.62 eV diffusion barrier that must be overcome for two well separated N atoms to arrive at adjacent sites (Figure 9a). However, subsequent reaction to form a N\(_2\) adsorbed intermediate involves crossing a large 1.69 eV barrier (Figure 9a), and it is this chemical reaction step that becomes the rate-determining step on the pure Ni surface. Desorption of the N\(_{2ad}\) final state is endothermic by \(\sim 0.76\) eV (Figure 9a), in accord with direct experimental determination of this quantity,\(^ {27, 28} \) which adds confidence to the quantitative validity of the calculations. Introduction of a monatomic step increases the diffusion barrier to 0.8 eV (Figure 9b), which nevertheless remains smaller than the corresponding chemical barrier (1.06 eV, Figure 9b), so that even at step sites on pure Ni the chemical reaction remains rate-controlling (Figure 9b).

The effects on the reaction of adding gold to the surface were investigated by stepwise addition of Au atoms, allowing the system to relax at each stage. Increasing amounts progressively reduced the activation barrier for the chemical reaction, while increasing the diffusion barrier (see Supporting Information). The reduction of the chemical barrier is a consequence of the partial negative charge on Au atoms that arises from Ni \(\rightarrow\) Au charge transfer (see the Supporting Information). Their repulsive interaction with the initial state consisting of two adjacent N\(^{-}\) species destabilizes the latter, resulting in a decreased activation barrier for the subsequent chemical reaction step. Note that, although the N\(_{ad}\) state is higher in energy than the reference state, the interaction of the N adatom with the surface is strong (we calculate this to be 4.7 eV on this surface) so a significant concentration of N adatoms at the surface resulting from NO molecule dissociation is to be expected. At a composition of Au\(_{11}\)Ni\(_{34}\), the chemical reaction barrier was very markedly reduced by \(\sim 1\) to 0.68 eV (Figure 9c), making it significantly lower than the diffusion barrier of 1.1 eV (Figure 9c). The composition Au\(_{11}\)Ni\(_{34}\) in the bulk corresponds stoichiometrically to the highly active Au\(_{25}\)Ni\(_{75}\) catalyst so the quantity of gold is in the right ball-park. However, it must be cautioned that we cannot know the working surface composition: this stoichiometry and choice of structure was made simply to permit theoretical exploration of the effect of “no gold” vs “gold intermixing” scenarios to enable us to interpret the experimental observations. As a result of the inclusion of gold, the identity of the rate-determining step switches from the chemical reaction step to that of the diffusion of N atoms to occupy adjacent sites, as shown in Figure 9c.

Finally, addition of Au at step sites increases both the diffusion barrier and the chemical reaction barrier. This can be rationalized through the gold atoms at the step edge being negatively charged, Au\(^{−}\). Since the N adatoms are also negatively charged, the resulting repulsive interaction disfavors their diffusion from the terrace to these sites, thus inhibiting reaction. For example, at the composition Au\(_{30}\)Ni\(_{61}\), these quantities are 2.08 and 2.17 eV, respectively, much more activated than they are at pure Ni steps (0.80 and 1.06 eV, respectively), so that bimetallic step sites should be inactive and have no effect on the overall reaction rate.

### CONCLUSIONS

Incorporation of Au greatly increased the activity of Ni toward the CO + NO reaction due to electronic interaction between the two metals, rather than solely by maintaining the presence of metallic Ni sites, which are important for inducing the dissociation of NO.

Postreaction XPS showed that, with both pure nickel and bimetallic catalysts, the Ni underwent substantial oxidation. In situ EXAFS spectroscopy at reaction temperature and in the presence of reaction gas showed extensive and reversible intermixing of Ni and Au in the two most active catalysts, which does not occur at room temperature. This underlines the critical importance of such measurements to the correct interpretation of catalytic performance, compared to more conventional and common methodologies based on ex situ postmortem observations carried out at room temperature.

Although the presence of Au did moderate nickel oxidation as a result of Ni \(\rightarrow\) Au charge transfer, the oxidation state of Ni did not correlate with activity across the range of compositions studied. This is in disagreement with an earlier proposal based on the study of only a single catalyst composition. The 50/50 catalyst was the most active. In situ XAS unambiguously demonstrated that extensive Au–Ni alloy formation occurred at 550 °C in reaction gas, that it was reversible with...
temperature, and that there was a good correlation between the most reactive catalyst and the Au–Ni coordination number at reaction temperature.

Oxidative pretreatment to remove hydrocarbonaceous residues inevitably introduced during nanoparticle synthesis is of critical importance to making valid measurements of subsequent performance. This step, which preserved the bimetallic nature of the nanoparticles, albeit at the expense of an increase in particle size, if omitted left large quantities of hydrocarbonaceous residues on the catalysts, thus compromising interpretation of results. XPS using a high-pressure cell that prevented air exposure revealed that the catalyst surfaces were entirely free of carbon after exposure to reaction conditions, confirming the validity of our procedures and results.

The experimental findings are fully consistent and very well aligned with the results of DFT calculations which indicated that (i) N + N recombination rather than NO dissociation is rate-controlling and (ii) addition of gold and consequent electronic interaction reduces the overall barrier for the rate-controlling and (ii) addition of gold and consequent electronic interaction reduces the overall barrier for the rate-controlling

reaction. Theory also shows that NiO, which may be present under reaction conditions, is catalytically inert.

The insights provided by this work should assist the design of lower cost bimetallic catalysts intended for the replacement of materials based on platinum group metals. They also emphasize the importance of correct materials pretreatment procedures and appropriate experimental methodologies required for the reliable acquisition and interpretation of results.

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Notes
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