The Influence of Hydrazine Reduction on Metal Dispersion and Support Morphology in Bimetallic Ru–Au/MgO Catalysts

A. G. SHASTRI AND J. SCHWANK

Department of Chemical Engineering, University of Michigan, Ann Arbor, Michigan 48109

Received July 9, 1985

The influence of hydrazine reduction on metal dispersion, particle size distribution, and support morphology was investigated for a series of bimetallic Ru–Au/MgO catalysts. The catalysts were prepared by impregnation of MgO with chlorine-containing precursor salt solutions. Hydrazine reduction significantly lowered the residual chlorine content of the catalysts and allowed the formation of high-surface-area MgO. This, in turn, led to much higher Ru metal dispersions as compared to that obtained after H2 reduction alone. X-Ray energy dispersive spectroscopy (EDS) proved that bimetallic Ru–Au aggregates were confined to a particle size range of less than 5 nm. Particles larger than 10 nm contained without exception monometallic Au. The hydrazine-treated catalysts did not show the suppression in catalytic activity for CO hydrogenation that was observed in H2-reduced Ru/MgO catalysts. The increased metal dispersion and restored catalytic activity of Ru in the hydrazine-treated catalysts may be linked to the absence of chlorine contamination of the support.

INTRODUCTION

In bimetallic catalysts, the nature of the support and the details of catalyst preparation and pretreatment can play a major role in controlling particle size and morphology, the extent of bimetallic cluster formation, and ultimately the catalytic behavior. Ru–Au catalysts represent an important model system for the investigation of bimetallic cluster formation between metal components which are immiscible in the bulk state. Previous work on the Ru–Au system clearly demonstrated that MgO-supported catalysts were structurally quite different compared to SiO2-supported ones (1–9). In contrast to the acidic and refractory nature of SiO2 supports, MgO is basic and can undergo massive structural changes depending on preparation and reaction conditions. Typical catalyst preparation via impregnation of simple cubic MgO with aqueous catalyst precursor solutions can lead to a transformation into hexagonal Mg(OH)2. The subsequent dehydration of Mg(OH)2 to MgO during catalyst pretreatment is accompanied by about 55% reduction in molar volume, nonuniform release of water, and a cooperative movement of Mg2+ and O2− ions (10–20). Consequently, the MgO surface created by such a dehydration process tends to be highly nonuniform and rich in defects (21). It is conceivable that these defect-rich MgO structures are quite susceptible to metal-support interactions. In the case of Au/MgO, the reaction between the basic oxide material and the precursor species AuCl4 generally results in high Au dispersions. The presence of such highly dispersed Au was found to cause structural destabilization of MgO resulting in increased rates of isotopic oxygen exchange between gas-phase and MgO oxygen (22, 23). The precursor species Ru3+, on the other hand, tends to precipitate as hydroxide, thereby causing poor Ru dispersion on MgO. Moreover, MgO seems to exert a "negative support effect" on Ru, suppressing its catalytic activity in the hydrogenolysis of ethane and in the CO hydrogenation (1). Similar activity losses were also reported for Pt/MgO after reduction in H2 at
773 K and were ascribed by the authors to the formation of a platinum–magnesia complex in which electron transfer occurs from Pt to MgO (24).

In the case of Ru/MgO, substituting hydrazine (N₂H₄) as the reducing agent in place of H₂ during catalyst preparation eliminated the suppression in activity and restored the normal catalytic behavior of Ru in CO hydrogenation (25). These observations prompted us to investigate in more detail the influence of hydrazine reduction on metal dispersions and on bimetallic Ru–Au particle formation in MgO-supported catalysts. Similar to the previously reported results on H₂-reduced Ru–Au/MgO catalysts (1, 3, 9, 25), a multifaceted characterization approach has been used to map out the size distribution and composition of metal particles in the hydrazine-reduced catalysts. Special emphasis is placed on the initial stages of metal particle formation and stability of metal dispersions as a function of temperature and reducing environment. Furthermore, the influence of Au on spill-over of atomic hydrogen from Ru sites will be discussed.

EXPERIMENTAL

The catalysts were prepared by impregnating MgO (Carlo Erba, reagent grade) with a solution of RuCl₃ ⋅ H₂O (Rudi-Pont reagent grade) and HAuCl₄ ⋅ 3H₂O (Carlo Erba RPE). After drying for 4 hr at 388 K, the samples were treated under stirring in an ≈0.5 M NaOH solution of hydrazine, followed by 4 hr evacuation at 388 K. Each catalyst was assigned a code, containing the letter R for Ru, M for MgO, H for hydrazine reduction, and a three-digit number representing the atomic percentage of Ru on the basis of the total number of Ru and Au atoms in the catalyst. The hydrazine-reduced catalysts were stored in air and are referred to as “N₂H₄-only”-treated samples. Aliquots of these hydrazine-reduced samples were subsequently subjected to a second reduction step in H₂ at 673 K. These samples are labeled as “N₂H₄ + 673 K H₂” treated. The bulk composition of the samples was determined by neutron activation analysis (Table 1).

The surface area of the catalysts was determined by the single-point BET technique. N₂ at 77.3 K was used as adsorbate in a Quantachrome Monosorb surface area analyzer. The surface areas obtained via the single-point technique were generally accurate within ±5% error. Crystal structures of both the metals and the support were determined by X-ray diffraction in a Philips X-ray powder diffractometer. Average support and metal crystallite sizes were obtained from X-ray line broadening using Scherrer's equation after correcting for the instrumental contribution.

Metal particle size distributions were obtained by high-resolution transmission electron microscopy carried out in a JEOL-100CX microscope equipped with a side-entry goniometer stage. The catalyst powder was placed on holey carbon films mounted on standard copper or beryllium grids. Specimen areas protruding into the holes of the holey-carbon film were selected for microscopic examination to avoid the obscuring effect of the supporting carbon film. The microscope was equipped with an ASID-4D scanning attachment and a lithium-drifted solid-state X-ray detector for elemental analysis. Data acquisition was carried out with a multichannel analyzer connected to a ND 6620 computer. For elemental analysis of individual metal particles having a diameter of more than 4 nm, a suitable sampling area was identified and photographed in the transmission mode, and then the machine was switched over to the scanning transmission mode for elemental analysis by energy-dispersive X-ray spectroscopy. A 10-nm electron probe was focused on individual metal particles of interest and X-ray counts were acquired for 200 sec. In the JEOL microscope used, metal particles smaller than 3 nm do not provide sufficient X-ray counting statistics to derive meaningful information on individual particles. Nevertheless, some information could
be obtained by rastering the beam over a suitable region containing many small particles.

For metal dispersion measurements via gas chemisorption, a standard catalyst pretreatment routine was adopted consisting of the following steps: loading of about 0.1–0.3 g of catalyst into a static volumetric high-vacuum system and evacuation at room temperature; heating under dynamic vacuum to 673 K and further evacuation at this temperature for 4 hr; exposure to a first dose of research grade H\textsubscript{2} (33.3 kPa) at 673 K for 16 hr; 1 hr evacuation at 673 K; exposure to a second dose of research grade H\textsubscript{2} at the same temperature and pressure for 4 hr; evacuation at 673 K for 2 hr. Then, the samples were cooled to the desired chemisorption temperature under dynamic vacuum.

The dispersion of Ru and Au was measured using a recently developed stepwise chemisorption and titration procedure (8, 9). The technique is based upon differences in the chemisorption characteristics of the two metal components. H\textsubscript{2} is adsorbed only on Ru but not on Au allowing us to obtain the dispersion of Ru. O\textsubscript{2} adsorption at room temperature serves as a second independent measure of the Ru dispersion, since there is no O\textsubscript{2} adsorption on Au at room temperature. At 473 K, however, both Ru and Au chemisorb oxygen. Weakly adsorbed oxygen is removed by evacuation and then the strongly held oxygen is titrated with H\textsubscript{2} at 373 K. In our previous work we found that only oxygen adsorbed on Ru sites could be titrated with H\textsubscript{2} at 373 K while the oxygen adsorbed on Au at 373 K did not react with H\textsubscript{2} to any significant extent at this temperature. Oxygen chemisorption at 473 K and H\textsubscript{2}/O\textsubscript{2} titration at 373 K are used for obtaining the Au dispersion. A more detailed description of the technique and a critical discussion of adsorption stoichiometries and possible complications arising in this stepwise chemisorption and titration procedure have been published previously (8, 9).

RESULTS AND DISCUSSION

Support Structure and Surface Area

The BET surface areas are listed in Table 1. The "N\textsubscript{2}H\textsubscript{4}-only"-treated samples had relatively low BET surface areas. X-Ray diffraction showed that the MgO had undergone complete bulk hydration into Mg(OH)\textsubscript{2} during catalyst impregnation.

When the hydrazine reduction was followed by an additional reduction step in H\textsubscript{2} at 673 K, the BET surface area increased by a factor of 4 to 6, and drastic changes were observed both in the morphology of the support and in the metal particle size distribution. X-Ray diffraction gave MgO patterns with significant line broadening indicating small MgO crystallite sizes. A similar decrease in MgO particle size and increase in BET surface area upon dehydration of Mg(OH)\textsubscript{2} have been observed in our previous work (20, 21). The presence of substantial amounts (>1 wt%) of residual chlorine left behind on the MgO

<table>
<thead>
<tr>
<th>Sample code</th>
<th>Metal loading (wt%)</th>
<th>Chlorine (wt%)</th>
<th>BET surface area (m\textsuperscript{2}/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ru</td>
<td>Au</td>
<td>&quot;N\textsubscript{2}H\textsubscript{4}-only&quot;</td>
</tr>
<tr>
<td>RMH100</td>
<td>4.6</td>
<td>—</td>
<td>25.8</td>
</tr>
<tr>
<td>RMH093</td>
<td>3.28</td>
<td>0.468</td>
<td>30.4</td>
</tr>
<tr>
<td>RMH014</td>
<td>0.26</td>
<td>3.109</td>
<td>27.3</td>
</tr>
</tbody>
</table>
during catalyst preparation prevents this surface area increase during the dehydration of Mg(OH)₂; on Ru–Au/MgO catalysts which were reduced at 673 K in H₂ alone, a significant amount of chlorine contamination (1.3–2.2 wt%) was found, and consequently the BET surface areas remained low in the range of approximately 30–40 m²/g (20).

The drastic increase of the BET surface areas during dehydration of the support in the hydrazine-reduced samples indicates low residual chlorine contamination, as confirmed by the results of neutron activation analysis (Table 1). The catalyst preparation for all the MgO-supported catalysts reported here and in Ref. (20) was identical and involved the same chlorine-containing precursor salt solutions. The only difference is the H₂-versus-N₂H₄ reduction. It appears that the hydrazine treatment step is very effective in preventing the buildup of chlorine contamination on the MgO support. Similar beneficial effects of hydrazine reduction have been found for other bimetallic systems. Hydrazine reduction was vastly superior to borohydride reduction in the preparation of unsupported Pd–Au alloys, resulting in much lower Na impurity levels (30).

Wide-angle X-ray scattering (WAXS) failed to detect any Ru peaks in the samples treated with “N₂H₄-only” and with “N₂H₄ + 673 K H₂” indicating Ru particle sizes less than 4 nm. In the “N₂H₄-only”-treated samples no Au peaks were detected. However, in the “N₂H₄ + 673 K H₂”-treated samples of catalysts RMH093 and RMH014, Au peaks were found corresponding to volume average Au particle sizes of 23.4 and 20.2 nm, respectively.

**Metal Dispersions**

To obtain information about the dispersion of the individual metal components in the bimetallic catalysts, the stepwise chemisorption and H₂/O₂ titration procedure mentioned before was employed. Figure 1 shows typical adsorption isotherms obtained for catalyst RMH093. For determining the amount of strongly adsorbed hydrogen two complete isotherms were measured. The first isotherm provided information about the total gas uptake. After evacuation at the adsorption temperature (room temperature for H₂ adsorption and 373 K for H₂/O₂ titrations) a second adsorption isotherm was taken to measure the amount of weakly adsorbed hydrogen. The difference between the first and second isotherm provided the amount of strongly adsorbed hydrogen which was used for the calculation of metal dispersions. This approach takes into account the possible spill-over of atomic hydrogen to Au or support sites. To arrive at meaningful Au dispersions, the total oxygen uptake at 473 K was first measured and corrected for the weakly held amount of oxygen that could be removed by evacuation at 473 K. Lowering the temperature to 373 K generally resulted in desorption of some O₂ in addition to the weakly adsorbed amount of oxygen at 473 K (8, 9). This additional loss of O₂ was also taken into account for calculating the amount of strongly held oxygen which could not be removed by evacuation within the temperature range of 373–473 K. Finally, the Au dispersion was determined based on the difference between the corrected amount of strongly held oxygen at 473 K and the oxygen which could be titrated with H₂ at 373 K.

This approach is justified in view of our previous observation that oxygen chemisorbed on monometallic Au/SiO₂ and Au/MgO cannot be removed by evacuation at 473 K (8, 9). Recent experiments on Au single crystals following our proposed chemisorption procedure at 473 K failed to detect any dissociative chemisorption of molecular oxygen (26, 27). However, it is well known that atomic oxygen will chemisorb on Au surfaces and desorb at temperatures around 600–680 K as O₂ gas (27, 28). Our supported gold catalysts show strong chemisorption of oxygen at 473 K. The adsorbed oxygen can easily be removed by
evacuation at 623 K, a behavior which is typical for desorption of atomic oxygen from Au. This indicates that there must be some source of atomic oxygen present in our supported catalysts. One possible source could be the oxide support itself. In previous work we discovered that impregnation of oxides with highly dispersed Au can dramatically increase the isotopic oxygen exchange activity of MgO and SiO₂ (22, 23). The isotopic oxygen exchange process is likely to involve some breakage of molecular oxygen bonds either on the oxide surface or on the gold/oxide interface. Some atomic oxygen could also be generated on trace impurity sites. Our neutron activation results, however, indicate that if there are any metal impurities in our catalysts they must be of the order of parts per million levels. Of course, in the case of bimetallic Ru–Au catalysts, Ru might be the source of atomic oxygen for chemisorption on Au sites via spillover.

The oxygen species adsorbed on Au do not react to any measurable extent with molecular H₂ at 373 K, a temperature at which the reactivity of H₂ with oxygen adsorbed on Ru is very high. This offers the opportunity to employ H₂/O₂ titration at 373 K to discern between Ru and Au surface sites. It is very crucial to keep the titration temperature at 373 K, since on monometallic Au catalysts the reaction between H₂ and adsorbed oxygen reaches a measurable rate at temperatures exceeding 423 K (see Table 2).

The results of the stepwise chemisorption and titration procedure are summarized in Table 3. From the gas uptake values, the number of both Ru as well as Au surface atoms can be determined, based on adsorption stoichiometries of Ru/H = 1,
Ru/O₂ = 1, and Au/O₂ = 4. If Ru and Au was present in monometallic particles only without formation of any bimetallic particles, one could translate the metal dispersion directly into surface average particle sizes \( d(\text{Ru}) \) and \( d(\text{Au}) \). These estimated particle sizes are also listed in Table 3. One interesting observation made in this context is that the relative amount of weakly adsorbed hydrogen that can be removed by evacuation at 373 K decreases as the gold content increases. This phenomenon is not restricted to the hydrazine-reduced Ru–Au/MgO catalysts discussed in this paper, but is also seen in the results reported previously on H₂-reduced Ru–Au/MgO and Ru–Au/SiO₂ catalysts (8, 9). One explanation could be that weakly adsorbed atomic hydrogen is highly mobile and can migrate around on the surface of metal particles and on the support. Because of the low Au–H bond strength, atomic hydrogen species arriving at gold sites tend to recombine to molecular H₂ and desorb into the gas phase. However, atomic hydrogen spilled over onto the support has a higher residence time than on Au sites and requires evacuation for removal. This interpretation is in agreement with the observation that on bimetallic Ru–Au no increased hydrogen uptake takes place as compared to monometallic Ru with similar Ru metal loadings (8, 9). Thus our technique of subtracting the weakly adsorbed gases corrects for spill-over of atomic hydrogen if it should be taking place under our experimental conditions.

**Particle Morphology and Composition**

In order to compare the influence of “N₂H₄-only”-vs-“N₂H₄ + 673 K H₂” reduction on support morphology and metal particle size distribution, transmission electron microscopy (TEM) and energy dispersive X-ray spectroscopy (EDS) were carried out. The massive changes in support morphology and the significantly increased metal particle sizes after “N₂H₄ + 673K H₂” reduction can readily be seen in the representative micrographs for each sample (Fig. 2). The corresponding changes in metal particle size distributions are shown in the histograms of the N₂H₄-reduced and (N₂H₄ + H₂)-reduced catalysts (Fig. 3). The main feature of the histograms is that the number of very small particles (<1 nm) is not significantly affected by the H₂ reduction at 673 K, while the particles with an initial size of more than 1 nm agglomerate readily into larger particles, thus shifting the size distribution toward higher values.

**TABLE 2**

H₂/O₂ Titration on 3.46 wt% Au/MgO as a Function of Temperature after Preadsorption of 3.46 μmol of O₂/g of Catalyst at 473 K

<table>
<thead>
<tr>
<th>H₂/O₂ titration temperature (K)</th>
<th>H₂ uptake (μmol/g catalyst)</th>
<th>mole of H₂ consumed/mole of adsorbed O₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>373</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>423</td>
<td>1.25</td>
<td>0.36</td>
</tr>
<tr>
<td>473</td>
<td>1.87</td>
<td>0.54</td>
</tr>
</tbody>
</table>

**TABLE 3**

Summary of Stepwise Chemisorption and H₂/O₂ Titration Results on “N₂H₄ + 673 K H₂”-Treated Catalysts

<table>
<thead>
<tr>
<th>Sample code</th>
<th>H₂ (293 K)</th>
<th>O₂ (293 K)</th>
<th>O₂ (473 K)</th>
<th>H₂/O₂ (373 K)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H₂ uptake (cm³/g)</td>
<td>H₂/Ru d(Ru) (nm)</td>
<td>O₂ uptake (cm³/g)</td>
<td>O₂/Ru d(Ru) (nm)</td>
</tr>
<tr>
<td>RMH100</td>
<td>1.70</td>
<td>0.333</td>
<td>2.8</td>
<td>—</td>
</tr>
<tr>
<td>RMH093</td>
<td>0.65</td>
<td>0.179</td>
<td>5.1</td>
<td>0.83</td>
</tr>
<tr>
<td>RMH014</td>
<td>0.013</td>
<td>0.045</td>
<td>20.3</td>
<td>0.024</td>
</tr>
</tbody>
</table>
Preliminary electron $\mu-\mu$ diffraction studies indicate that the alignment of metal particles with respect to the oxide substrate changes as a function of metal particle size. We suspect that these differences in particle alignment might influence the agglomeration characteristics.

Table 4 summarizes the number average particle sizes obtained from electron microscopy ($d_n = \Sigma n_i d_i / \Sigma n_i$) along with the computed values of surface average particle sizes ($d_s = \Sigma n_i d_i^2 / \Sigma n_i$) and volume average particle sizes ($d_v = \Sigma n_i d_i^3 / \Sigma n_i$) to allow a direct comparison with chemisorption and WAXS results. $n_i$ represents the number of particles of size $d_i$. The good agreement in the Ru particle size results obtained by chemisorption (Table 3) and by electron microscopy (Table 4) indicates that there is no significant problem associated with hydrogen spillover, in contrast to recent observations on the Ru–Cu system (29).

To obtain information concerning the composition of individual metal particles, EDS was carried out using the scanning transmission mode. In view of the small signal-to-noise ratio and the low X-ray count obtained on such small metal particles, only a qualitative analysis can be attempted. Following a previously proposed strategy (25), we grouped the EDS spectra into four categories, namely, spectra showing signals of Ru only, Ru with a trace of Au, Au with a trace of Ru, and Au only. Figure 4 shows examples of typical EDS spectra obtained on individual particles. Table 4 summarizes the EDS results obtained on the "$\text{N}_2\text{H}_4 + 673\text{K H}_2$"-treated catalysts RMH093 and RMH014. For each catalyst, 50 metal particles 2–11 nm in diameter were analyzed. Table 4 shows the percentage of these metal particles belonging to one of the four above-mentioned composition categories. There was no significant difference in the EDS spectra of the "$\text{N}_2\text{H}_4$-only"- and the "$\text{N}_2\text{H}_4 + 673\text{K H}_2$"-reduced catalysts. The monometallic gold particles listed in Table 4 as "Au only" were without exception larger than 9 nm.

**Catalyst Microstructure**

Interesting details concerning the microstructure of the catalysts are provided utilizing the rich data base accumulated through the various characterization techniques. We will now discuss each of the three samples in turn. First, the monometallic Ru sample RMH100 has a very high metal dispersion corroborated by TEM, chemisorption, and WAXS results which are in excellent agreement. The Ru-rich sample RMH093 shows good agreement between microscopy, chemisorption, and WAXS as far as Ru is concerned. However, there is an important discrepancy between the average Au particle size as derived from chemisorption and WAXS. The WAXS particle size is by more than a factor of 3 larger than the chemisorption particle size. In contrast to chemisorption, WAXS requires a certain degree of long-range order and coherence length, and is not sensitive to metal in a state of very high disper-
FIG. 2. Representative transmission electron micrographs of catalysts. (a) Sample RMH100, "N₂H₄-only" treated. (b) Sample RMH100, "N₂H₄ + 673 K H₂" treated. (c) Sample RMH093, "N₂H₄-only" treated. (d) Sample RMH093, "N₂H₄ + 673 K H₂" treated. (e) Sample RMH014, "N₂H₄-only" treated.

The chemisorption results give a strong indication that in catalyst RMH093 some Au must be present in a state of high dispersion below the detection limit of WAXS. At this point let us recall the EDS results indicating that some of the small (<4 nm) particles contained Ru with a trace of Au. Thus, it seems reasonable to model...
these small particles as either monometallic Ru particles or Ru particles having some Au adsorbed on their surface.

On the Au-rich catalyst RMH014, TEM showed after "N$_2$H$_4$-only" treatment faint scattering centers attributable to Ru together with large monometallic Au particles. In the "N$_2$H$_4$ + 673 K H$_2$"-treated
sample, the small Ru scattering centers were not seen (Figs. 2f,g). Interestingly, EDS analysis and WAXS failed to detect the presence of any large Ru particles even though the calculations based on chemisorption indicated an average Ru particle size of 20.3 nm. Similarly, a much larger Au particle size is derived from the chemisorption data than seen in WAXS or TEM. Therefore, the model previously proposed for catalyst RMH093, where gold adsorbs on the surface of Ru and consequently blocks part of the Ru surface from gas adsorption, is not applicable in the case of RMH014. Both Ru and Au seem to be partially inaccessible to gas chemisorption. If
one can account for the inaccessibility of gases to a part of the metal sites. A careful examination of the micrographs (Figs. 2f,g) shows that indeed some support platelets are superimposed one on top of the other. It is conceivable that small metal particles get sandwiched between two MgO platelets in the course of the restructuring events of the support which occur during the dehydration of Mg(OH)$_2$. Metal particles covered by thin MgO overlayers would be detectable in both WAXS and TEM experiments but unable to chemisorb gases.

Table 5 presents the average metal particle sizes of the "N$_2$H$_4$ + 673K Hz"-treated catalysts (RMHXXX series) as estimated from chemisorption and measured by WAXS. For comparison, previously reported data on Hz-only-reduced Ru-Au/SiO$_2$ (RSXXX series) (8) and Ru–Au/MgO (RMXXX series) catalysts (9) are also included. Reduction of MgO-supported catalysts in N$_2$H$_4$ can lead to substantially higher Ru and Au dispersions compared to the conventional H$_2$ reduction. It should be

one envisions a scenario where some Ru and Au particles are buried under MgO structures during the dehydration process accompanying the H$_2$ reduction at 673 K, then one can account for the inaccessibility of gases to a part of the metal sites. A careful examination of the micrographs (Figs. 2f,g) shows that indeed some support platelets are superimposed one on top of the other. It is conceivable that small metal particles get sandwiched between two MgO platelets in the course of the restructuring events of the support which occur during the dehydration of Mg(OH)$_2$. Metal particles covered by thin MgO overlayers would be detectable in both WAXS and TEM experiments but unable to chemisorb gases.

Table 5 presents the average metal particle sizes of the "N$_2$H$_4$ + 673K Hz"-treated catalysts (RMHXXX series) as estimated from chemisorption and measured by WAXS. For comparison, previously reported data on Hz-only-reduced Ru–Au/SiO$_2$ (RSXXX series) (8) and Ru–Au/MgO (RMXXX series) catalysts (9) are also included. Reduction of MgO-supported catalysts in N$_2$H$_4$ can lead to substantially higher Ru and Au dispersions compared to the conventional H$_2$ reduction. It should be

one envisions a scenario where some Ru and Au particles are buried under MgO structures during the dehydration process accompanying the H$_2$ reduction at 673 K, then one can account for the inaccessibility of gases to a part of the metal sites. A careful examination of the micrographs (Figs. 2f,g) shows that indeed some support platelets are superimposed one on top of the other. It is conceivable that small metal particles get sandwiched between two MgO platelets in the course of the restructuring events of the support which occur during the dehydration of Mg(OH)$_2$. Metal particles covered by thin MgO overlayers would be detectable in both WAXS and TEM experiments but unable to chemisorb gases.

Table 5 presents the average metal particle sizes of the "N$_2$H$_4$ + 673K Hz"-treated catalysts (RMHXXX series) as estimated from chemisorption and measured by WAXS. For comparison, previously reported data on Hz-only-reduced Ru–Au/SiO$_2$ (RSXXX series) (8) and Ru–Au/MgO (RMXXX series) catalysts (9) are also included. Reduction of MgO-supported catalysts in N$_2$H$_4$ can lead to substantially higher Ru and Au dispersions compared to the conventional H$_2$ reduction. It should be
noted that the Ru dispersion achieved in the monometallic Ru/MgO catalyst RMH100 is comparable to that of the Ru/SiO₂ catalyst RS100. This result is quite remarkable for conventional wet impregnation of an MgO support with RuCl₃ precursor salt solution followed by H₂ reduction at 673 K. N₂H₄ solutions in aqueous Na(OH) obviously exert a neutralizing influence on the acidic RuCl₃ precursor solution leading to different metal deposition and nucleation characteristics. The metal dispersions after N₂H₄ reduction are very high. Subsequent H₂ reduction at 673 K leads to considerable metal particle growth and agglomeration probably due to the high metal mobility during the phase transformation and restructuring of the support. Nevertheless, the final result is a much higher ruthenium dispersion than expected for MgO-supported Ru–Au catalysts prepared by wet impregnation and H₂ reduction. Once the N₂H₄-reduced catalysts have been treated in H₂ at 673 K, the particle size distribution appears to be rather stable even under prolonged exposure to typical catalytic reaction environments. The activity-suppressing “negative support effect” of MgO was found to be absent in the N₂H₄-reduced Ru/MgO catalyst RMH100, and the catalytic activity for CO hydrogenation was comparable to that of metallic Ru (25).

CONCLUSIONS

It is generally difficult to achieve high Ru metal dispersions by wet impregnation of MgO with chlorine-containing precursors such as RuCl₃ · H₂O. The basic nature of MgO causes the precipitation of Ru in the form of large Ru(OH)₃ particles. Prereduction of the catalysts in hydrazine prior to reduction in H₂ at 673 K leads to much higher Ru metal dispersions. The differences in Ru metal dispersion appear to be linked to the morphology and the chlorine content of the support. Chlorine tends to prevent the formation of high-surface-area MgO. Hydrazine treatment leads to almost complete removal of residual chlorine contamination from the MgO support. In the absence of chlorine contamination, the BET surface area of MgO can increase significantly upon dehydration, and consequently high metal dispersions are facilitated. Numerous small metal particles are seen, some of them apparently in the form of small, two-dimensional rafts completely wetting the support. It should be noted that the chlorine-free, hydrazine-treated Ru/MgO catalyst RMH100 exhibits a CO hydrogenation activity similar to that of Ru metal or Ru/SiO₂, while conventionally prepared Ru/MgO catalysts containing relatively high amounts of residual chlorine have severely suppressed catalytic activity. The suppression of the activity of MgO-supported Ru could be either a direct consequence of the presence of chlorine in the vicinity of Ru or an indirect effect of chlorine via a modification of the MgO morphology or the Ru/MgO interface. MgO morphologies are known to be highly sensitive to the preparation route, impurity content, and thermal history. These factors strongly influence the adsorption characteristics of MgO. It is quite likely that these variations in adsorption characteristics carry to some extent over into the deposition of metal precursor ions on the support. This could change the nucleation and growth of metal particles and result in different manifestations of metal–support interactions.

The formation of bimetallic Ru–Au particles does not seem to be significantly influenced by hydrazine treatment. As in the previously investigated H₂-reduced Ru–Au/MgO catalyst series, bimetallic Ru–Au particles are restricted to particle sizes of less than 5 nm. Evidence for encapsulation of some metal particles by platelets of MgO was discovered. The encapsulation does not seem to be a direct consequence of hydrazine reduction, but may be linked to the structural changes of the Mg(OH)₂/MgO system during dehydration.
ACKNOWLEDGMENTS

The authors thank Dr. S. Galvagno and Dr. G. R. Tauszik for their contribution to catalyst preparation and characterization, and Dr. A. K. Datye and Dr. J. Y. Lee for their helpful suggestions. Skillful experimental assistance in part of the work was provided by Mr. M. Ballard. Financial support by the National Science Foundation and the Army Research Office is gratefully acknowledged.

REFERENCES