Towards an atomic level understanding of niobia based catalysts and catalysis by combining the science of catalysis with surface science

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ABSTRACT

The science of catalysis and surface science have developed, independently, key information for understanding catalytic processes. One might argue: is there anything fundamental to be discovered through the interplay between catalysis and surface science?

Real catalysts of monometallic and bimetallic Co/Nb2O5 and Pd-Co/Nb2O5 catalysts showed interesting selectivity results on the Fischer-Tropsch synthesis (Noronha et al. 1996, Rosenir et al. 1993). The presence of a noble metal increased the C5 selectivity and decreased the methane formation depending of the reduction temperature. Model catalysts of Co-Pd supported on niobia and alumina were prepared and characterized at the atomic level, thus forming the basis for a comparison with “real” support materials. Growth, morphology and structure of both pure metal and alloy particles were studied. It is possible to support the strong metal support interaction suggested by studies on real catalysts via the investigation of model systems for niobia in comparison to alumina support in which this effect does not occur. Formation of Co2+ penetration into the niobia lattice was suggested on the basis of powder studies and can be fully supported on the basis of model studies. It is shown for both real catalysts and model systems that oxidation state of Co plays a key role in controlling the reactivity in Fischer-Tropsch reactions systems and that the addition of Pd is a determining factor for the stability of the catalyst. It is demonstrated that the interaction with unsaturated hydrocarbons depends strongly on the state of oxidation.

Key words: catalysis, model catalysts, cobalt, palladium, characterization, niobia.

INTRODUCTION

The science of catalysis and surface science have developed over the years in friendly contact, though to a large extent independently. There are, however, examples in which key information for our understanding of catalytic processes has been gained from studies in surface science, one example being ammonia synthesis (Schlögl 2008). The success of surface science in determining that nitrogen is hydrogenated after N2 dissociation on the surface was honored with the 2007 Nobel Prize for Gerhard Ertl (Ertl 2008). Even though ammonia synthesis is a special case, one might argue: does this mean that all fundamental problems in catalysis have been solved, or putting it in another way: is there anything fundamental to be discovered through this interplay between catalysis and surface science?

The answer is YES, the field is wide open, and there are many fundamental questions to be answered. They may be grouped into those connected with understand-
ing the complex materials active as catalysts and those connected with operating conditions.

Complexity often does not only lead to qualitative changes but to new phenomena that do not occur in simpler systems. They need to be understood to be able to describe catalytic phenomena reliably.

Natural gas is becoming more attractive as an energy source due to the increasing prices of oil, the huge reserves of gas and environmental issues (Noronha et al. 2000a, Nowak Ziolek 1999, Passos et al. 2000, Schmal et al. 2000, Tanabe 2003, Wachs 2005, Ziolek 2003). This has led researchers across the globe to develop more economic processes of upgrading natural gas to higher hydrocarbon fuel. The Fischer-Tropsch synthesis is an alternative route to produce liquid fuels from synthetic gas.

Monometallic cobalt catalysts are known to be effective for the synthesis of long chain hydrocarbons from hydrogen and carbon monoxide (Anderson 1984, Dry 1981). The catalytic hydrogenation (activity, selectivity and stability) of carbon monoxide on cobalt catalysts has been reported to be affected by two factors: the dispersion and the extent of reduction (Fu and Bartholomew 1985, Ho et al. 1990, Lee et al. 1988, Reuel and Bartholomew 1984). The reducibility of cobalt is closely related to the nature of the species present on the support. In particular, on supported cobalt catalysts, several cobalt species have been detected as a function of the support (Arnoldy and Moulijn 1985, Castner and Santilli 1984, Ho et al. 1992, Lapidus et al. 1991, Noronha et al. 1999, Sato et al. 1984). At least three different types of cobalt species have been reported on the alumina-supported cobalt catalysts: Co$_3$O$_4$ particles, which are more readily reduced, Co$^{2+}$ species and CoAl$_2$O$_4$, which are hardly reduced or unreduced, respectively (Castner and Santilli 1984, Chin and Hercules 1982, Stranick et al. 1987, Tung et al. 1990). Analysis revealed that catalysts with low cobalt content showed a high amount of Co$^{2+}$ species in interaction with the support. For higher cobalt loading, Co$_3$O$_4$ particles were also present, in addition to Co$^{2+}$ species (Höbel et al. 2006).

Besides the support, the addition of noble metals to a supported cobalt catalyst can also change the cobalt state and modify its activity and selectivity (Guczi et al. 1991, Kapoor et al. 1992, Niemantsverdriet et al. 1988, Schanke et al. 1995, Zsoldos et al. 1991). Generally, it has been proposed that the addition of a noble metal promotes cobalt oxide reduction (Idriss et al. 1992, Juszczyk et al. 1993, Kapoor et al. 1992, Martens et al. 1986, Sarkany et al. 1995, Van’T Blik and Prins 1986).

The addition of a noble metal to Co/Nb$_2$O$_5$ catalysts showed interesting selectivity results on the Fischer-Tropsch synthesis (Noronha et al. 1996, Rosenkir et al. 1993). The presence of a noble metal increased the C$_2^+$ selectivity and decreased the methane formation independent of the reduction temperature. However, the characterization of the reduced state was not performed, rendering a complete comprehension of the catalytic behavior difficult.

In fact, relatively few studies have been performed to study the behavior of bimetallic systems on a support able to promote the Strong Metal Support Interaction (SMSI) effect, such as Nb$_2$O$_5$ (Ahon et al. 2006, Noronha et al. 2000b). This may be explained by the difficulties in characterizing the niobium oxide supported catalysts.

Some questions in those contexts have been addressed and there are several review articles available. We review here a joint venture between the catalysis groups in Rio de Janeiro and the surface science groups in the Fritz Haber Institute that was triggered by the Brazilian group in trying to understand Fischer-Tropsch catalysis on cobalt based alloy systems, in which particles on a variety of oxidic supports, such as alumina and niobia, just to mention two of them, have been studied.

Based on the experimental results of the group in Rio, the following questions were posed to the surface scientists:

1. Is it possible to prepare model Pd-Co alloys systems on the supports in question?
2. Is it possible to measure properties and reactivities of those model systems that may be compared to the “real” catalysts?

We will discuss, after a short reminder of the resources and experimental conditions for such an endeavor, the understanding that has been gained through this joint
venture and try to address some of the fundamental questions that one encounters on such a journey (Carlsson et al. 2003a, b, c, Freund et al. 2003, Heemeier et al. 2002, Hill et al. 2005, Höbel et al. 2006, Mendes et al. 2006a, b, Nowitzki et al. 2007, Risse et al. 2004, Starr et al. 2005, Uhl et al. 2007).

**EXPERIMENTAL**

The real carriers were prepared from commercial materials by simple calcination in air. Niobium oxide with a specific surface area of 30 m$^2$g$^{-1}$ was obtained after calcination at 873 K of the hydrated oxide (sample AD 376 from Companhia Brasileira de Metalurgia e Mineração). Alumina was obtained from a transition alumina (sample 3996-R, area 200 m$^2$g$^{-1}$, from Harshaw) by treatment at 1473 K. The catalysts were prepared by impregnation with an aqueous solution of cobalt nitrate. The impregnated material was dried in air at 383 K for 16 h and then calcined in flowing air at 673 K for 16 h. The metal loading was always 5 wt.%. The specific surface areas of the calcined samples were 23 m$^2$g$^{-1}$ (Co/Nb$_2$O$_5$) and 9 m$^2$g$^{-1}$ (Co/Al$_2$O$_3$) (Rosenir et al. 1993).

The bimetallic catalyst was prepared on the same support by incipient wetness impregnation or co-impregnation of the support with an aqueous solution of palladium nitrate and cobalt nitrate, followed by drying at 393 K for 16 h and calcination in air at 673 K for 2 h. Table I lists the catalysts and their metal contents, measured by atomic absorption spectroscopy (Noronha et al. 2000b).

**TABLE I**

Composition of bimetallic catalysts (Noronha et al. 2000b).

| Catalyst     | Pd (wt.-%) | Co (wt.-%) |
|--------------|------------|------------|
| Pd/Nb$_2$O$_5$ | 1.3        | -          |
| Pd$_{12}$Co$_{68}$/Nb$_2$O$_5$ | 2.1        | 2.1        |
| Pd$_{12}$Co$_{68}$/Nb$_2$O$_5$ | 1.6        | 5.0        |
| Co/Nb$_2$O$_5$ | -          | 2.0        |

On the other hand, for the model catalyst, an assembly of sophisticated ultrahigh vacuum (UHV) equipment interfaced with techniques that allow application at ambient pressures have been employed. A concise description is found within the website of the Fritz Haber Institute (http://www.fhi-berlin.mpg.de/).

**RESULTS**

**Real Catalysts**

**Activation**

Figure 1 presents the Temperature Programmed Reduction (TPR) profiles for some typical catalysts. In all cases, the reduction occurs, at least, in two distinct steps whose resolution depends on the sample and/or the carrier. Pure Co$_3$O$_4$ and the mechanical mixture Co$_3$O$_4$/Nb$_2$O$_5$ are much easier to reduce than the supported catalysts. The experimental ratio H$_2$/Co, also given in Figure 1, is close to the theoretical value (1.33) for unsupported materials and for Co/Al$_2$O$_3$, but is much higher for Co/Nb$_2$O$_5$, suggesting a partial reduction of the carrier.

![Fig. 1 – TPR profiles for (a) physical mixture (H$_2$/Co = 1.34); (b) Co$_3$O$_4$ (H$_2$/Co = 1.30); (c) Co/Al$_2$O$_3$ (H$_2$/Co = 1.33) and (d) Co/Nb$_2$O$_5$ (H$_2$/CO = 1.49) (Rosenir et al. 1993).](image_url)

The results of hydrogen and oxygen chemisorption for Co/Nb$_2$O$_5$ and Co/Al$_2$O$_3$, for different reduction temperatures, are given in Table II. The oxygen uptake on the pure niobia carrier, reduced at various temperatures, and the O$_2$:H ratios are also shown.

For Co/Al$_2$O$_3$ catalysts, the hydrogen uptake is independent of the temperature of reduction, both at 573 and 773 K. For Co/Nb$_2$O$_5$ catalyst, the total hydrogen uptake decreases markedly when the temperature of reduction is increased, whereas the oxygen uptake remains
unchanged. In a second procedure, the hydrogen uptake is similar to that obtained from direct reduction, at the same temperatures. In this case, although independent of the temperature of reduction, the oxygen uptake is increased by a factor of 2. This is not yet well explained and needs further investigation. The O$_2$ : H ratio increases with the temperature of reduction, in both cases. The results of Table II also show that not only cobalt but a progressive fraction of pure niobia is reduced when the temperature of reduction is increased from 573 to 773 K.

From hydrogen chemisorption measurements, the apparent cobalt dispersion and cobalt mean particle size were calculated, with the hypothesis that the H:Co surface stoichiometry is 1. The results are given in Table III. Note that the particle size is similar for both Co/Nb$_2$O$_5$ and Co/Al$_2$O$_3$ reduced at 573 K.

The reduction profiles of the bimetallic Pd-Co catalysts were very different from the profiles of Pd and Co monometallic catalysts, suggesting the existence of some interaction between the metals, (Fig. 2) (Noronha

An Acad Bras Cienc (2009) 81 (2)
NIOBIA BASED REAL CATALYSTS AND MODEL SYSTEMS

Fig. 2 – Reduction profiles of the niobia-supported catalysts. (a) Pd/Nb₂O₅; (b) Pd₃₅Co₆₅/Nb₂O₅; (c) Pd₁₅Co₈₅/Nb₂O₅; (d) Co/Nb₂O₅ (Noronha et al. 2000b).

et al. 2000b). The bimetallic catalysts showed peaks at temperatures around 400 and 700 K. The hydrogen consumption at room temperature (Table IV) was higher than the theoretical value expected for complete reduction of PdO, indicating that, besides PdO reduction, another process may occur. As PdO was completely reduced at room temperature, the peaks at 400 and 700 K can be ascribed to the reduction of cobalt oxides. However, these maxima were shifted toward lower temperatures, compared to the cobalt reduction of the monometallic Co/Nb₂O₅ catalyst, indicating that palladium catalyzes the cobalt oxide reduction.

Therefore, the palladium addition promoted not only the reduction of Co₃O₄ particles but also that of the cobalt surface phase. The magnetic measurements coupled with TPR analyses will allow us to quantify this promotional effect. Saturation of magnetization was determined from the magnetization curves for each sample reduced at increasing temperatures, and the reduction degree of cobalt was calculated (Table V).

The Pd/Nb₂O₅ catalyst showed only a diamagnetic signal. The Co/Nb₂O₅ catalyst was not completely reduced at 873 K. The reduction degree revealed only 43% of cobalt in metallic state, which was much lower than that calculated from the hydrogen uptake during the TPR. As it was discussed for the Pd/Nb₂O₅ catalyst, this difference could be attributed to a partial reduction of niobia at high temperature.

The magnetic measurements of the Pd₃₅Co₆₅/Nb₂O₅ bimetallic catalyst showed the presence of 4% of metallic cobalt even after reduction at room temperature, confirming a strong promoting effect of Pd on cobalt oxide reduction. After reduction at 473 K, the magnetic measurements indicated that 12% of cobalt was already in the metallic state (Table V). However, the hydrogen consumption measured by TPR (Table III) was higher than the amount of metallic cobalt produced, suggesting a partial transformation of Co₃O₄ to CoO in this temperature range as well. The reduction at 873 K led to a complete reduction of cobalt oxides, in opposition to the behavior observed on the Co/Nb₂O₅ catalyst. This result indicated that palladium also promoted the reduction of the Co³⁺ species since X-ray Photoelectron Spectroscopy (XPS) and TPR analysis revealed the presence of a cobalt surface phase. Thus, the magnetic measurements performed at different reduction temperatures support the TPR results, giving a further evidence of the interaction between both metals.

In the case of Pd₁₅Co₈₅/Nb₂O₅ bimetallic catalyst reduced at room temperature, the magnetic measurements did not suggest the presence of metallic cobalt. After reduction at 473 K, the amount of metallic cobalt was lower than on Pd₃₅Co₆₅/Nb₂O₅ catalyst. In other words, the promoting effect of palladium decreases with the increase of cobalt loading. However, after reduction at 873 K, all cobalt oxide was reduced to the metallic state, as observed for the Pd₃₅Co₆₅/Nb₂O₅ catalyst.

Catalytic results
The catalytic results are summarized in Table VI. The reaction rates for Co/Al₂O₃ are equal and independent of the temperature of reduction. However, when the temperature of reduction is increased from 533 to 773 K, the reaction rate with Co/Nb₂O₅ is reduced by a factor of 7. Similarly, with Co/Nb₂O₅, the Turnover Frequency (TOF) for methane formation decreases when the temperature of reduction is increased. However, the TOF
Influence of the reduction temperature (K) on activity and selectivity (Rosenir et al. 1993).
Figure 4 displays the selectivity with time on stream, for a space velocity of 6000 h⁻¹ and isoconversion (30%) for the Pd-Co catalysts, reaction temperature of 270°C and pressure of 20 bar, after reduction at 500°C. The bar diagrams of Figure 4a present product selectivities on Co/Nb₂O₅ catalyst based on moles of product carbons in the $\sum C_i$ formation with time on stream. Methane was very low (≈3.0%) and the diesel fraction (C₁₃–C₁₈) was very high (54–49%). Note that $CH_4$, $C_2$-$C_4$, $C_5$-$C_{12}$, $C_{13}$-$C_{18}$, and $C_{19+}$ correspond to saturated hydrocarbons with the carbon subscript being the number of carbon atoms in the chain. The range $C_2$-$C_4$ may also contain some $C_2$-$C_4$ olefins (ethene and propene) which were not resolved from the more pronounced saturated hydrocarbon chromatographic peaks.

Higher molecular weight olefins ($C_5$-$C_{12}$ and $C_{13}$-$C_{18}$) were detected on the Co/Nb₂O₅ catalyst.

The bar diagrams of Figure 4b clearly point out that the selectivity towards methane and $C_2$-$C_4$ hydrocarbons for the bimetallic Co-Pd/Nb₂O₅ changed with Pd addition and with time on stream. Note that the methane formation rate of this catalyst increased. Olefins, except $C_4^\text{2,4}$, were not detected before on the pure Co-containing catalysts. Light products and gasoline fractions increased, while diesel fraction decreased.

Methane was around 17–21% and the diesel fraction decreased from 53 to 37%. The addition of the second metal indicates also higher selectivity towards alkanes. Product retention and condensation are not expected to have occurred because the overall mass balance indicates that the reactor operates steady-state.

The stability of the bimetallic Pd-Co/Nb₂O₅ catalyst is shown in Figure 5 indicating good stability up to 50 h with time on stream for a reduction temperature of 500°C. It suggests that the Pd stabilizes the catalyst for a long time and from the beginning.

It is noteworthy, that the catalyst was stable during 50 hours and deviation of mass balance was around 20%. In addition, during the stabilization, samples after 50 h were taken and coke was analyzed by Thermogravimetry (TG) (not shown). It displays the loss of approximately 16% of carbon due to the coke deposition, which allows us to explain the deviation observed. Experiments were reproduced under similar conditions at isoconversion (around 30%).
The Anderson-Schulz-Flory equation (ASF) with these experiments displayed great deviation in the $C_{13} - C_{18}$ range (Fig. 6). As expected, the total hydrocarbon molar compositions cannot be interpreted as an ASF distribution. Although the $C_2-C_3$ anomalies and the change of the chain growth probability in the range from $C_3$ to $C_8-C_{12}$ range may be explained by $\alpha$-olefin readsorption with secondary chain propagation and the existence of two mechanisms for chain propagation, the increasing selectivity observed between $C_8-C_{12}$ for all experiments cannot be explained by these mechanisms (Ahon et al. 2006).

In conclusion, these results showed that more fundamental studies are needed to explain the behavior of a real catalyst, which was the objective to study a model catalyst, however, under UHV conditions.

MODEL STUDIES

Modeling the technical support is a very difficult task and attempts to generate knowledge at the atomic level are being launched at present. This involves research at the liquid-solid interfaces (Catalano et al. 2008, Datta et al. 1992, Kim et al. 1993, Park et al. 2006, M. Sterrer, unpublished data).

So far, model oxide supports have been prepared under UHV conditions. One uses either conductive oxide single crystals or thin films grown on top of metal single crystals. The procedures have been extensively reviewed (Bäumer and Freund 1999, Campbell 1997, Diebold 2003, Freund 1997, Freund et al. 1996, Freund and Pacchioni 2008, Freund and Goodman 2007, Henry 1998, Risse et al. 2008). The supports relevant to the present review are alumina and niobia.

Alumina has been prepared by oxidation of a NiAl(110) single crystal (Jaeger et al. 1991, Libuda et al. 1994). Its structure and properties have been the object of intense studies across the world. Finally, Scanning Tunneling Microscopy (STM) (Kulawik et al. 2006) and Atomic Force Microscopy (AFM) (Simon et al. 2008) images with atomic resolution have been published, and density functional calculations (Kresse et al. 2005) have allowed us to understand the film at the atomic level including its defect structure, which turns out to be particularly relevant when it comes to anchoring of metal nanoparticles (Schmid et al. 2006). Figure 7 shows atomic resolution of STM and AFM images together with the structural model as deduced from Density Functional Theory (DFT) calculations at areas of the film that do not contain defects. It is possible to bind metal atoms within the unit cell, even without particular defects, but this depends critically on the possibilities of charge transfer through the layer and has only been found to occur for Au. Other metals, such as Pd, do not feel this effect and we have learned to understand this (Sterrer et al. 2007).

Niobia is difficult to prepare in a well-ordered fashion. Niehus and his group came up with a strategy to prepare Niobia on a Cu$_3$Au(100) surface by predosing the surface with a “reactive” oxygen which dissolves in the alloy and then dosing Nb from a metal source which reacts with the predosed oxygen to form well-ordered niobia layers (Middeke et al. 2005). Figure 8 shows the morphology and structure of the film as imaged via STM (Starr et al. 2005). The large terraces (Fig. 8a) are separated by steps and those terraces exhibit one hexagonal atomic arrangement of oxygen atoms (Fig. 8b). The rotated domains are clearly discernable (Fig. 8c). It seems that this surface is not connected to any known surface terminations that could be deduced from known bulk structures. Understanding niobia has progressed lately but it seems that the surface structure depends on the thickness of the niobia film (Fig. 9). A thin film forms an oxygen terminated surface consisting of two hexagonal oxygen layers of 2.3 ML of Nb$^{5+}$ in between (Fig. 8d). The film exhibits a defect structure and no
Fig. 7 – STM and AFM images of alumina on NiAl(110): (a) large scale STM image showing the antiphase domain boundaries, (b) atomic resolution STM, (c) atomic resolution AFM, (d) alumina as modeled with DFT typ and side view (Kresse et al. 2005, Kulawik et al. 2006, Simon et al. 2008).

Fig. 8 – STM images of the niobia film grown on Cu$_3$Au(100). (a) Large terraces separated by steps of ~2Å in height. (b) Hexagonal structure of the niobia film with vacancy defects. The (2 × 7) superstructure cell is marked as a rectangle. (c) Two domains rotated by 90°, with each domain exhibiting characteristic surface modulation seen as stripes. Image sizes and tunneling parameters are: (a) 200 × 200 nm$^2$, VS = 1.4 V, I = 1.0 nA; (b) 15 × 15 nm$^2$, VS = 1.4 V, I = 1.0 nA; (c) 40 × 40 nm$^2$, VS = 1.4 V, I = 1.0 nA. 2(d) Cross and top views of the thin niobia film grown on Cu$_3$Au(100). The unit cell of niobia overlayer is indicated as a rhomb. The rectangle shows the (2 × 7) coincidence superstructure, which is formed between oxide and metal substrate lattices. The surface shows a hexagonal lattice with a 5.3Å periodicity (Mendes et al. 2006b, Starr et al. 2005).

Fig. 9 – HREEL-spectra measured in specular geometry, and PES spectra measured at normal electron emission for monolayer and multilayer niobia films grown on Cu$_3$Au(100) (see the text) (Uhl et al. 2007).
Nb=0 double bonds (Fig. 9). The absence of any Nb=0 double bonds is clear from High Resolution Electron Energy Loss Spectroscopy (HREELS) measurements, showing no significant intensity above ~ 900 cm\(^{-1}\) (Fig. 9a).

If one grows thicker films, one approaches the structures expected from bulk H-Nb\(_2\)O\(_5\) (Fig. 9a) surfaces. In particular, niobyl species (Nb=0) are observed and the oxidation state is represented by a Nb3d XPS spectrum at the appropriate binding energy comparable to H-Nb\(_2\)O\(_5\), while the monolayer is shifted to smaller (Fig. 9b) binding energy due to screening by the metallic substrate. Unfortunately, the multilayer exhibits some roughness. Figure 10a shows Infrared Reflection Absorption Spectroscopy (IRAS) spectra of CO adsorbed at 100 K after thermal flash to the specified temperatures and cooling back to 100 K. CO species adsorbed at 100 K after thermal flash to the specified temperatures and cooling back to 100 K. CO species adsorbed at 100 K after thermal flash to the specified temperatures and cooling back to 100 K.

As a result, the CO stretching frequency is shifted to higher wavenumbers compared to the gas phase value (2143 cm\(^{-1}\)). However, metal cations which contain partially filled d-orbitals can interact with the 2\(\pi^*\)-orbital of CO via electron back-donation from the metal to CO thus lowering the vibrational frequency (Blyholder 1964, Zecchina et al. 1996). On silica supported N\(_2\)O\(_5\) particles, Knözinger and co-workers observed the signal at 2191 cm\(^{-1}\) (i.e. blueshifted with respect to CO in the gas phase) which was assigned to CO bonded to Nb\(^{5+}\) surface atoms (Beutel et al. 1997). Therefore, the weak signal at 2189–92 cm\(^{-1}\) observed on the niobia films can be assigned to the adsorption on the Nb\(^{5+}\) cations. Concomitantly, the most prominent peak at 2141 cm\(^{-1}\) can be associated with Nb species, which are partially reduced and thus contain d-electrons through which Nb interacts more strongly with CO via back-donation effect. For example, step edges of the niobia terraces and domain boundaries can in principle expose low coordinated and partially reduced Nb cations, which may adsorb CO much stronger than regular terrace sites.

Since, for the “real” catalysts, the niobia component was dispersed on an alumina support, the modeling procedure needs one more step, namely the preparation of alumina supported niobia. Those were prepared by physical vapor deposition of Nb in an oxygen atmosphere onto the above described thin alumina film.

Figure 11 shows a series of STM images of the niobia/alumina surface prepared at room temperature at increasing niobia coverage. At 0.1Å coverage, the particles are randomly dispersed on the surface, i.e. both on and between the line defects (antiphase and reflection domain boundaries, steps) clearly visible in Figure 11a. As previously shown for other metal deposits on the alumina film, the particle spatial distribution basically reflects the degree of metal support interaction (Bäumer and Freund 1999, Heemeier et al. 2003). The absence of preferential nucleation for niobia, as shown in Figure 11, can be interpreted as a relatively strong interaction between
Fig. 10 – (a) Typical TPD spectrum of CO adsorbed on thin niobia films at 100 K. (b) IRAS spectra of CO adsorbed on thin niobia films as a function of CO exposure at 100 K. (c) IRAS spectra of CO adsorbed at 100 K and subsequently flashed to the indicated temperatures. All spectra are recorded at 100 K. The TPD and IRAS spectra for a Cu$_3$Au(100) substrate at saturation CO coverage are also shown in (b) and (c), for comparison (Mendes et al. 2006b).

Fig. 11 – STM images (size 60 × 60 nm$^2$, tunneling parameters $V_S = 3$ V, $I = 0.1$ nA) of niobia deposited on alumina films at different coverages: (a) 0.1Å; (b) 0.5Å; (c) 1.5Å; (d) 2.5Å; (e) 6.0Å; (f) apparent height of niobia particles and particle density as a function of niobia coverage (Uhl et al. 2007).
 Nb atoms and the alumina surface. Particle density increases with the coverage up to $\sim$1Å and then saturates at $\sim 2 \times 10^{13}$ cm$^{-2}$, while the average particle height is nearly constant from the onset (see Fig. 11f). Note that, at high coverage, the particle size is difficult to be determined precisely due to the tip deconvolution effect. Obviously, niobia grows in a three-dimensional mode under these conditions. Infrared spectroscopy was used to study the surface structure of the alumina supported niobia particles. The IRAS method allows one to investigate phonon vibrations at oxide surfaces as well as of adsorbates such as CO used here as a probe molecule.

The phonon region of the IRA spectra is shown in Figure 12a. The pristine alumina film exhibits a sharp peak at $\sim$ 865 cm$^{-1}$ as previously reported (Frank et al. 2001). The alumina phonon attenuates upon niobia deposition and a new band appears as a shoulder at around 900 cm$^{-1}$, ultimately resulting in a broad signal centered at 885 cm$^{-1}$ for the highest niobia coverage studied. The spectral broadening is consistent with the formation of randomly dispersed small niobia particles as revealed by STM. At the coverage of 1Å, a band centered at 986 cm$^{-1}$ emerges, which grows in intensity with the coverage. The assignment of these spectral features is facilitated by comparison with the results on a supported vanadia system previously studied in detail (Magg et al. 2003a, 2002, 2004). For convenience, the bottom curve in Figure 12a shows the spectrum obtained for the similarly prepared 3.5Å vanadia/alumina sample in the same experimental setup. Based on Raman and IR spectroscopy results, the peak at 1045 cm$^{-1}$ has been assigned to the stretching of vanadyl (V=O) groups. Meanwhile, the broad signal at $\sim$ 950 cm$^{-1}$ is associated with vibrations involving V-O-Al linkage at the vanadia/alumina interface as predicted by theoretical calculations (Magg et al. 2003b, 2002). Therefore, turning back to the niobia/alumina system, the peak at 986 cm$^{-1}$ must be assigned to the niobyl (Nb=O) stretching, and the signal at 890 cm$^{-1}$ must be assigned to Nb-O-Al interface vibrations, which should show lower frequencies as compared to vanadia/alumina, based solely on mass considerations (Nb atom is heavier than V).

An IRAS study of adsorption of CO as a probe molecule further supports the conclusion on the oxidation state of Nb in the model systems. Interaction of CO with niobia is found to be weak and can be seen only at low temperatures (see Fig. 12b). The strongly blue-shifted frequency of CO as compared to the gas phase (2191 vs. 2143 cm$^{-1}$) implies that CO is linearly adsorbed on top of Nb$^{5+}$ sites (Beutel et al. 1997, Mendes et al. 2006b). A similar feature at 2190 cm$^{-1}$ was also obtained on the thin niobia films (not shown here), i.e. fully consistent with the above PES results. Therefore, combined STM and IRAS studies show that niobia deposited on thin alumina films in an oxygen atmosphere form small Nb$_2$O$_5$...
particles terminated by the Nb=O groups and also Nb\(^{5+}\) sites. Thermal stability of the supported niobia particles was further examined by STM and IRAS.

Figure 13 shows STM images of the 0.5Å niobia/alumina sample after a thermal flash to elevated temperatures (500–900 K) in UHV. Annealing to 700 K basically decreases the particle density partially due to sintering of the particles (some particles gain in size) and also due to niobia migration into the film, which is more clearly observed after annealing to 900 K. As previously shown for other metal particles (Heemeier et al. 2003), the interdiffusion probably goes through the line defects of the alumina film, thus resulting in their decoration (see Fig. 13d).

The 2.5Å niobia/alumina sample, shown in Figure 14a, was first flashed in UHV to 700 K, then oxidized in 10\(^{-7}\) mbar O\(_2\) at 400 K and finally at 900 K. One can see that, in contrast to annealing in UHV, ambient oxygen precludes the migration of the niobia particles, and only partial sintering occurs at 900 K.

Figure 15 shows the results of a complementary IRAS study for 2.5Å niobia/alumina sample. Initially existing Nb=O species at \(\sim 983\) cm\(^{-1}\) apparently disappear upon heating to 700 K. Concomitantly, the broad band centered at \(\sim 900\) cm\(^{-1}\) and assigned to the Nb-O-Al interface vibrations becomes sharper most probably due to a strong attenuation of the band at 865 cm\(^{-1}\) corresponding to the pristine alumina film. This effect...
can be explained by the onset of niobia migration as revealed by STM and discussed above. Mild oxidation at 400 K recovers the niobyl species without affecting the interface region. However, further oxidation at 900 K shifts the interface phonon from 915 cm\(^{-1}\) to 968 cm\(^{-1}\) (partially overlapping with the Nb=O band). This shift can be related to the structural transformations of the niobia/alumina interface since the STM results showed that overall particle morphology is not affected much by oxidation at 900 K. Note also that this finding indirectly supports our previous assignment of the ∼900 cm\(^{-1}\) band to niobia/alumina interface phonon. The effect of ambient oxygen on the thermal stability of the niobia particles can be rationalized in terms of oxygen preventing the reduction of the niobia surface and hence migration of the Nb atoms into the film. In addition, previous studies on the oxygen interaction with alumina supported particles showed that the metal particles may promote thickening of the alumina film via oxygen dissociation and subsequent diffusion of atomic oxygen to the alumina/NiAl interface (Bäumer et al. 1999, Shaikhutdinov et al. 2002). This in turn may also alter the interaction of the niobia particles and the alumina film.

Elemental metal particles have been studied extensively on a number of bulk single crystal and thin film oxide supports (Campbell 1997, Henry 1998, Freund 1997, Freund and Pacchioni 2008, Goodman 1995). For the latter case, which is the subject of the current discussion, we refer to a number of reviews. Co and/or Pd particles have been studied on aluminum (Bäumer and Freund 1999, Höbel et al. 2006), and on niobia (Bäumer and Freund 1999, Uhl et al. 2007). Co-Pd alloy particles so far exclusively on the alumina support (Heemeier et al. 2002). Co binds strongly to alumina while the metal remains in its metallic state. Figure 16a shows an STM image of 2Å of deposited Co at room temperature. There is a strong propensity to form lower coordinated metal atoms on the nanoparticle at low temperature, as may be identified by IRAS studies (Risse et al. 2003). Co particles loose those low coordinated sites and sinter upon temperature increase as revealed by STM (Carlsson et al. 2003c), ferromagnetic resonance (FMR (Hill et al. 2005, Risse et al. 2004)), and IRAS measurements (Carlsson et al. 2003c). The latter technique delivers detailed insight not only into sintering but also into the rearrangement at the surfaces of the particles, including the metal oxide interface. This is due to energetically separated surface resonances in FMR spectroscopy, which allows for separate investigation by gas adsorption at the metal vacuum and by temperature changes on the metal oxide interface (Risse et al. 2004). Pd binds less strongly to the alumina surface and forms relatively large metallic particles at room temperature. These particles are located at line defects (i.e. antiphase domains boundaries) of the film and form small, almost ideal single crystals of truncated cuboctahedral morphology. Figure 16b shows an STM image of a Pd nanoparticle distribution along with a high resolution image revealing the surface atoms in the well ordered facets. Using CO as a probe molecule, the surface sites have been characterized in detail.

Nanometer-sized Co-Pd alloy nanoparticles were generated by sequentially depositing the two constituents onto alumina. Inspired by earlier work by Henry and co-workers (Gimenez et al. 1998, Giorgio et al. 1999), different structures and compositions of the particles were obtained in a controllable way by taking advantage of the different nucleation and growth properties of the two metals.

Let us now assume that Pd is deposited on a surface already covered by Co particles. Owing to the higher mobility of Pd on the surface, it will be trapped at Co particles before reaching the line defects. The STM image presented in Figure 17 essentially corroborates this expectation. The arrangement of particles found for this sequence strongly resembles that for pure Co, thus suggesting particles with a Co core and a Pd shell. If, on the other hand, Pd is deposited first, the less mobile Co atoms should partly cover the Pd crystallites and partly nucleate between them. The STM image indeed shows triangular and hexagonal crystallites as well as the large number of new small clusters between them.

To verify the surface composition of the alloy particles, we performed both temperature programmed desorption and IRAS experiments using CO as a probe molecule (Carlsson et al. 2003a). In this context it is important to know that CO prefers different adsorption sites on
Fig. 16 – Sequential deposition of Co and Pd and vice-versa onto a thin alumina film at 300 K. The ball models schematically show the structure of the systems investigated: gray: NiAl substrate; blue: Al₂O₃ film; light blue: defects of the alumina film; yellow: Pd atoms; orange: Co atoms (Hansen et al. 1999, Heemeier et al. 2002).

Fig. 17 – STM images (100 nm × 100 nm) taken after depositing 2 Å Pd and 2 Å Co alone. The ball models schematically show the structure of the systems investigated: gray: NiAl substrate; blue: Al₂O₃ film; light blue: defects of the alumina film; yellow: Pd atoms; orange: Co atoms (Heemeier et al. 2002).

The interaction of Co with niobia is completely different. Even though we have to stress that so far, only the thin niobia film has been studied with respect to Co deposition, what is revealed as likely to be representative for thicker films also.

Figure 19 shows STM images of Co deposited at 300 K on the niobia films at two Co coverages, 0.5 Å (a and c) and 2 Å (c and d), respectively. It is clear that small particles are homogeneously dispersed on the surface and coalesce at higher coverage. However, the majority of these particles are only 1.5–2 Å in height. This
implies that Co essentially wets the surface and forms two-dimensional, monolayer islands at 300 K. Since the formation of these islands can be kinetically limited, we have examined the surface after annealing to 500 K, which resulted in large well-shaped monolayer islands at low Co coverage (Fig. 19c). At high Co coverage, the ill-defined and relatively rough surface, with a corrugation amplitude of ca. 1Å, is observed in Fig. 19d. Note that this morphology is very different from Co deposited on thin alumina films, in which three-dimensional particles up to 2 nm in height are formed under these conditions (Hill et al. 1998).

The electronic structure of the Co species on the niobia films was studied by Photoelectron Spectroscopy (PES) using synchrotron radiation. Figure 20 shows PES spectra before and after Co deposition at 300 K. The clean film is characterized by the binding energies (BE) of the Nb 3d$^{5/2}$ level at 206.4 eV and of the O 1s level at 529.8 eV. Note that the BE of the Nb level is $\sim$0.6 eV lower than for Nb$^{5+}$ in Nb$_2$O$_5$ powders ($\sim$207 eV (Morris et al. 2000)) due to the well-documented screening effect of the metal substrate underneath oxide films. For Co deposited at 300 K the PE-spectra show two species characterized by BE of Co $2p_{3/2}$ at 778 and 781 eV. These values are typical for metallic (Co$^0$) and oxidized (Co$^{3+}$, like in Co$_3$O$_4$ or CoO) states, respectively [XPS and AES Database, Thermo Electr. Corp. http://www.lasurface.com/database/element.php]. The Co$^{3+}$:Co$^0$ integral ratio increases from 0.1 to 0.3 when the spectrum is measured at grazing incidence. This indicates that Co$^{3+}$ species are on the surface, while metallic Co is mostly located in the subsurface region. In principle, this could be explained by the formation of the Co particles with oxidized Co$^{3+}$ species on the surface modified by Nb$^{5+}$ as revealed by CO IRAS. However, the STM images presented in Figure 19 did not show the three-dimensional particles but single layer islands, which are typical for the morphology of metal-on-metal systems. Therefore, it seems likely that Co migrates through the film and forms a metallic layer directly bonded to the Cu$_3$Au(100) substrate. In this model, the oxidized Co$^{3+}$ species are formed on or incorporated into the niobia film, thus resulting in a partial reduction of the niobium cations. The formation of two Nb species is clearly seen in the Nb3d spectrum of the Co/niobia sample, which splits into two components, centered at 207.1 and 206.2 eV for the Nb 3d$^{5/2}$ level, (after deconvolution not shown here). The ratio between these two signals does not depend on the detection angle as shown in Figure 20, which suggests that all Nb species lie in a single layer as in the original film (see Fig. 8d). The component at 207.1 eV deserves a comment since the binding energy is higher than in the original niobia film, where Nb is already in its highest oxidation state, 5$^+$. We tentatively attribute this effect to the presence of cobalt ions and reduced niobia ions in the film, which probably pin the Fermi level at a different position. Although the precise structure of this mixed Co-Nb oxide phase needs further studies, the PES results clearly show that Co deposited on the niobia film can be readily oxidized at room temperature, which is in contrast to Co deposits on the alumina films (Carlsson et al. 2003a, Hill et al. 1998).

Pd on the niobia has also been studied. A combined TDS and Sum Frequency Generation (SFG) study has been performed. Again CO has been used as a probe.
Fig. 20 – Co2p, Nb3d and O1s regions of photoelectron spectra measured at two different geometries as depicted in the inset. The spectra for Nb and O core levels are shown before and after 2Å of Co have been deposited at 300 K (Mendes et al. 2006b).

molecule. The adsorption of CO on Pd nanoparticles supported by Nb2O5/Cu3Au(100) was investigated by combined SFG and TPD studies from 100 K to 500 K and from UHV to 100 mbar. Annealing Pd-Nb2O5/Cu3Au(100) higher than 300 K lead to irreversible structural changes, indicated by a ~50% loss of the CO adsorption capacity and pronounced changes of the adsorption states. Changes in the vibrational frequencies and phase in CO-SFG spectra suggest alterations in the electronic structure of the model catalyst by metal-support interaction, i.e. formation of “mixed Pd-NbOx compounds”, while simple structural changes or sintering of the Pd particles can be excluded. Furthermore, the new surface composition is different from pure Nb2O5, i.e. migration of Nb2O5 over Pd particles cannot explain the current findings. These effects were observed both under UHV and ambient pressure and may contribute to the catalytic properties of Nb2O5 supported metal nanoparticles.

After having studied the morphological, structural, electronic and adsorption properties of the materials in question, it is very important to move on to their reactivity which in turn allows us the most detailed comparison to the real catalyst. So far, reactivity studies on model systems have only been performed on Co and Co based alloy particles on alumina (Carlsson et al. 2003b, c). Those reactivity studies are governed by the fact that the target reactions are of Fischer-Tropsch type, i.e.

\[(2n + 1)H_2 + nCO \rightarrow C_2H_{2n+2} + nH_2O\]

Co is a well known Fischer-Tropsch catalyst which, however, deactivates upon oxidation. It has been proposed that the addition of Pd promotes the reduction of Co oxide by activating hydrogen. In order to shed light on this question we have studied adsorption and reaction of molecules relevant for a Fischer-Tropsch scenario, i.e. oxygen, hydrogen, carbonmonooxide, and ethene. The point to note is the influence of oxygen on the state of oxidation of the particles and the concomitant influence on the reaction behavior. A rather detailed study on oxygen adsorption and sample annealing employing a combination of XPS, IRAS, TPD, and FMR shows that oxidation involves diffusion of oxygen into the model support and, after stabilization, a combination of surface and subsurface oxygen is formed, leaving the XPS detectable Co in a metallic state but suppressing CO adsorption at and above room temperature. Pd deposition onto the Co particles leads to the formation of a Pd shell.
Two limiting cases have been studied: i) incomplete Pd shell, ii) thick Pd shell. While the incomplete shell leads to facilitated oxidation, the full Pd shell suppresses it entirely.

Hydrogen and CO adsorption is facile on the non-oxidized particles, but is severely inhibited on the oxidized bimetallic and pure Co particles; Pd particles become covered with a chemisorbed oxygen overlayer which still allows a small amount of H$_2$ adsorption. This suggests that oxidized Co or Co-Pd particles should be less active than unoxidized ones in the Fischer-Tropsch conversion of CO+H$_2$ to higher hydrocarbons because CO and H$_2$ cannot adsorb. The interaction of ethylene with the bimetallic Co-Pd particles is similar to the interaction of ethylene with Pd or Co particles, in which it is $\pi$-bonded below 230 K and di-$\sigma$ bonded from $\sim$ 230 to 270 K. At 270 K it begins to desorb or decompose to form ethylidyne, liberating hydrogen. In contrast, on the oxidized Co-containing particles, ethylene is only able to adsorb in the $\pi$-bonded state and to form the di-$\sigma$ bonded state. The formation of ethylidyne, is inhibited as well. To slightly different degrees, the bimetallic or monometallic particles are active for ethylene hydrogenation; the oxidized particles are essentially non-active for ethylene hydrogenation. These reactive studies indicate that the reduction of CoO to metal could be of prime importance for the Fischer-Tropsch reaction.

SYNOPSIS
What have we learned?
• It is possible to prepare and characterize different model support materials such as alumina, niobia and alumina supported niobia and characterize them at the atomic level, thus forming the basis for a comparison with “real” support materials. Model studies have provided detailed information of the support surfaces which are difficult to deduce for the real catalyst support. Clearly, the full complexity of the real material has not been modeled yet, and further studies are needed. However, the strategy to build up models starting with the simple systems and procedures bottom up clearly is the strategy of choice.
• Growth, morphology and structure of both pure metal and alloy particles may be studied within model systems. In particular, alloy formation may directly be followed providing useful information for studies on real catalysts in which such a proof is difficult to deliver.
• It is possible to follow the strong metal support interaction suggested by studies on real catalysts on the model systems for niobia supports and not for alumina supports. The suggested formation of Co$^{2+}$ penetration into the niobia lattice can be fully supported.
• Due to the well defined morphology of model systems, it is possible to get deduce the distribution of the oxide and the metal in the particles. The same is true for the distribution of the metal composition upon alloy formation.
• It is shown for both real catalysts and model systems that Co oxidation plays a key role in controlling the reactivity of the system and that the addition of Pd is a determining factor with respect to the interaction with hydrogen and oxidation.
• It is demonstrated that the interaction with unsaturated hydrocarbons depends strongly on the state of oxidation of the particles.

So we have learned a lot but certainly not enough to say we have a full understanding of the phenomena in this particular case. Work has to be done on both sides: Models have to be made more stable to endure higher temperatures for treatment. Also, more frequently techniques have to be applied, such as sum frequency generation that allows us to go to ambient conditions. More sophisticated techniques have to be applied to real catalysts, such as aberration corrected transmission electron microscopy, which would allow us to look at those systems closer to the atomic level and compare them with scanning probe data gained on the models.

We believe that such a joint effect between groups working on catalysis with an applied touch and groups trying to improve a more basic understanding of the phenomena is very beneficial for both sides and the scientific community as a whole because, by working collaboratively, one develops not only scientific understanding but also trust in the work of others. The latter is crucial when it comes to employ the power of applied and basic research in combination to benefit our societies in the 21st century.

An Acad Bras Cienc (2009) 81 (2)
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An Acad Bras Cienc (2009) 81 (2)
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