Structure and Size of Bimetallic Palladium – Platinum Clusters in an Hydrotreatment Catalyst

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Résumé — Structure et taille de particules bimétalliques palladium – platine dans un catalyseur d’hydrotraitement — Des catalyseurs bimétalliques PdPt supportés sur alumine ont été synthétisés par imprégnation à sec de bis acénylacétonate. Les formulations Pt\textsubscript{4}Pd/Al\textsubscript{2}O\textsubscript{3} et Pt\textsubscript{6}Pd/Al\textsubscript{2}O\textsubscript{3} dédiées à l’hydrogénation des aromatiques et à la désulfuration, ont été caractérisées par spectroscopie d’absorption X, microscopie électronique à transmission, et titrage H\textsubscript{2}-O\textsubscript{2}. Il a été montré que les métaux étaient associés dans des structures de type cœur-coquille avec un enrichissement préférentiel de la surface des particules en Pd. De plus, la composition optimale pour la réaction d’hydrogénation des aromatiques en présence de soufre est dépendante de la taille des particules.

Abstract — Structure and Size of Bimetallic Palladium – Platinum Clusters in an Hydrotreatment Catalyst — Highly dispersed supported bimetallic PdPt catalysts were prepared by incipient wetness co-impregnation of alumina with platinum bis-acetylacetone Pt(Acac)\textsubscript{2} and palladium bis-acetylacetone Pd(Acac)\textsubscript{2}. The resulting Pt\textsubscript{4}Pd/Al\textsubscript{2}O\textsubscript{3} and Pt\textsubscript{6}Pd/Al\textsubscript{2}O\textsubscript{3} catalytic systems dedicated to hydro-dearomatization in presence of sulfur were characterized by X-ray absorption spectroscopy, transmission electron microscopy and volumetric H\textsubscript{2}-O\textsubscript{2} titration. Direct structural evidence is given of the presence of nanometer bimetallic particles. Moreover, the complete set of results indicates a “cherry-like” structure for the bimetallic clusters with a preferential distribution of palladium atoms at the cluster surface. Catalytic results shows that the optimum composition leading to the appropriate surface structure must be adjusted as a function of the size of the clusters.
INTRODUCTION

In both Europe and the United States of America, legislation is imposing more and more stringent reduction of the aromatic hydrocarbons contained in car diesel fuels [1]. In addition to their carcinogenic characteristic, mono- or poly-cyclic aromatics lower the diesel fuel quality by lowering the cetane index [2]. To remove these undesirable compounds, a catalytic hydroconversion, on transition metal (mainly platinum), is generally suggested.

One of the main difficulties is the presence of sulfur in the feedstock, which is still too high, even if hydro-desulphurization treatments have achieved great progress, to use a standard hydrogenation noble metal catalyst. Because, even at such a low concentration, sulphur rapidly deactivates the de-aromatizing catalyst, a thio-tolerance of the catalyst is required [3]. Various catalytic systems have already been investigated in the literature combining a thio-tolerant metallic phase with supports with different acido-basic surface properties [4-8], with addition of promoters [9-10] or alloying the Pt base catalyst with other metallic elements [11-12].

Many patents and articles claim that PdPt supported bimetallic catalysts are more thio-tolerant than pure metals [13] particularly for a selected range of Pd/Pt ratio [14-17]. Regarding academic studies, several studies have been already dedicated to the structure at the atomic scale of this bimetallic catalyst [18-22]. In some cases, contradictory results were found with no enhancement of the sulfur resistance or hydrogenation activity [23-24].

Interpretations proposed in the literature suggest that the modification of the sulphur sensitivity could be related to strong interactions between the two metals organized in core-shell structured particles. Structural and electronic characteristics depend intimately on the preparation procedure itself [25] and on the different parameters of the procedure such as the calcination temperature [26-28].

We report here a study of highly dispersed Pt-Pd catalytic systems deposited on cubic γ-alumina with different Pd/Pt atomic ratios. We have concentrated our attention on the metallic function of the catalysts. Thus, we have combined different techniques, namely X-ray absorption spectroscopy [29], TEM (Transmission Electron Microscopy) and volumetric H₂-O₂ titration. The ultimate goal of this structural and electronic investigation at the atomic scale is to give clear structural evidence of the presence (or not) of heterometallic bonds inside the catalyst. Moreover, the complete set of results allows us to propose a model regarding the distribution of the two metals inside the bimetallic particles.

2 EXPERIMENTAL

2.1 Catalyst Preparation

The catalysts are highly dispersed Pt, Pd monometallic and bimetallic Pt-Pd system deposited on alumina extrudates. More precisely, the support used was a cubic γ-alumina with a specific area of 280 m²/g (+/- 10 m²/g) and a pore volume of 0.7 cm³/g. Two compositions were chosen with different metal loadings corresponding to atomic Pd/Pt ratios of 4 and 6.

Pd-Pt/Al₂O₃ catalysts were prepared by incipient wetness co-impregnation of alumina with platinum bis-acetylacetonate Pt(acac)₂ and palladium bis-acetylacetonate Pd(acac)₂ according to the procedure described for example in references [30-31]. The two complexes were dissolved in toluene and the organic solution put in contact with alumina during 72 h at room temperature. After activation under airflow at 350°C during 2 h, the catalyst was reduced 2h at 450°C under nitrogen flow.

0.6% wt Pt and 0.4% wt Pd monometallic catalysts were also prepared using the same support and procedure in order to have reference catalysts.

2.2 Transmission Electron Microscopy (TEM)

TEM was carried out on a JEOL-2010 at 200 kV. After grinding in an agate mortar and sieving, the catalyst was suspended in ethanol and the suspension was then deposited on a copper grid covered with a carbon membrane. During the measurement, the sample was maintained under secondary vacuum. Histograms were constructed from the observation of around 300 particles.

2.3 Volumetric hydrogen - oxygen titration

The titration of adsorbed H₂ by O₂ has been used to determine the metallic dispersion and average particle size with a χ-sorb® apparatus. The catalysts were reduced before the titration by oxygen of the hydrogen chemisorbed on the accessible metal atoms. The reduction was performed under pure H₂ flow for 2 hours at 300°C with a intermediate stage of 1 hour at 150°C. After reduction, the catalyst was cooled under hydrogen flow and the sample cell was purged at 25°C under N₂ flow. Pulses of O₂ were repeatedly injected into the catalyst bed until no further consumption of O₂ was observed. The total volume of O₂ was then calculated by summing up the proportions of all pulses consumed. The relative accuracy of the measurement is 5%.

The titration follows equations (1) and (2):

\[ \text{Pt}_s \cdot H + \frac{3}{4} \text{O}_2 \rightarrow \text{Pt}_s \cdot \text{O} + \frac{1}{2} \text{H}_2 \text{O} \quad (1) \]

\[ \text{Pd}_s \cdot H + \frac{3}{4} \text{O}_2 \rightarrow \text{Pd}_s \cdot \text{O} + \frac{1}{2} \text{H}_2 \text{O} \quad (2) \]

The dispersions D of the monometallic and bimetallic particles are determined according to equations (3) and (4):

\[ D \% \text{wt} = \frac{V \cdot n \cdot M}{V_m} \quad (3) \]
\[
D (\%) = \frac{V \text{m} \text{ M}_{\text{Pt}} \text{ M}_{\text{Pd}}}{V_m (% \text{Pt} * \text{ M}_{\text{Pt}} + % \text{Pd} * \text{ M}_{\text{Pd}})}.
\]  

(4)

Mean particle size is obtained via the relations proposed by Van Hardeveld and Hartog [32] for regular truncated cubo-octahedron geometry:

\[
N_r = 16 m^3 - 33 m^2 + 24 m - 6
\]  

(5)

and

\[
N_r = 30 m^2 - 60 m + 32
\]  

(6)

with:

- \(m\) order of the cubo-octahedron (\(m \geq 2\)).
- \(N_r\) total number of atoms
- \(N_s\) number of surface atoms

Dispersion defined by the ratio \(D = \frac{N_s}{N_f}\) allows one to calculate successively \(m, N_r\) and the diameter \(d\) of a sphere containing \(N_f\) atoms given by:

\[
d = \frac{2(3N_rV_a)^{1/3}}{(4\pi)^{1/3}}
\]  

(7)

Metallic surface area \(S (\text{m}^2/\text{g catalyst})\) is obtained using the relation:

\[
S = \frac{V n N}{\sigma V M}
\]  

(8)

with:

- \(V_a\) atomic volume (8.9 cc/mol mean value for Pd and Pt) [33].
- \(\sigma\) number of atoms per m\(^2\) (1.27 \times 10^{19} \text{atoms/m}^2)
- \(N\) Avogadro number
- \(n\) stoichiometry metal / adsorbate
- \(V_m\) molar volume of adsorbate at T (22400 cm\(^3\)/g)
- \(M\) molecular weight (g.mol\(^{-1}\))
- \(V\) adsorbed volume (cc/g catalyst).

Note that this semi-empirical formula gives mean diameter values very close to those calculated with more realistic models as proposed in reference [23].

2.4 Electron Micro Probe Analysis

For EMPA (Electron Micro Probe Analysis) characterization, sample extrudates were embedded in a metacrylate resin, polished with SiC paper and coated with carbon black. Measurements were performed with a JEOL JXA 8800R microprobe fitted with four WDS spectrometers. Electron beam conditions were 20 kV accelerating voltage, 40 nA beam current and a focused probe. The Al K\(\alpha\), Pt M\(\alpha\) and Pd L\(\alpha\) lines were acquired using respectively TAP, PETH and PETH crystals. Counting times on both peak and background were respectively 10, 20 and 20 s. Elemental quantification was performed using a ZAF procedure, oxygen being calculated by stoichiometry (\(\text{Al}_2\text{O}_3\)). About 24 analysis points were acquired every 50 \(\mu m\) along the diameter on five different sample extrudates.

2.5 Extended X-ray Absorption Fine Structure (EXAFS)

X-ray absorption experiments (with both \textit{in situ} and \textit{ex situ} reduced catalysts) were performed at the LURE laboratory using DCI, a storage ring operated with an electron energy of 1.85 GeV and a current between 260 and 360 mA. Data were collected on the EXAFS IV station, a beam line placed after a bending magnet which is equipped with a double crystal monochromator, Si(111) for the Pt L\(\text{III}\)-edge (11564 eV) and Si(311) for the Pd K-edge (24352 eV), and a double borosilicate mirror for the rejection of harmonics. Measurements were done either in transmission mode (using ionisation chambers filled with air to absorb around 20% of the X-ray beam in the first ion chamber and the remainder of the X-ray beam in the second ion chamber) or in fluorescence mode using a multi-detector element. More precisely, transmission mode was used to perform experiments at the Pd K-edge whereas fluorescence mode was used at the Pt L\(\text{III}\)-edge. Only the Pd\(_{60}\)Pt catalyst was studied by EXAFS spectroscopy.

At the Pd K-edge, the spectrum was recorded twice with a step of 4 eV (from 24150 to 25350 eV) and 2 s per point. At the Pt L\(\text{III}\)-edge, each spectrum was recorded twice with a step of 2 eV (from 11450 to 12450 eV) and 5 s per point. At the Pd K-edge, as well as the Pt L\(\text{III}\)-edge, the estimated energy resolution of the monochromator was approximately 3 eV. The energy/angle calibration was performed using either a Pd foil (thickness 7.5 \(\mu m\)) or a Pt foil (thickness 7.5 \(\mu m\)) as references. After sieving, particles in the 100-200 \(\mu m\) range were either loaded into a sample holder sealed by self-adhesive tape made of Captor\texttrademark for experiments in transmission mode (Pd K-edge) or agglomerated and loaded into a sample holder (Pt L\(\text{III}\)-edge).

The Pd K-edge was recorded at room temperature under atmospheric pressure. Regarding the experiments in fluorescence mode, the sample was placed in a special furnace for \textit{in situ} treatment with entrance and exit windows made of boron nitride that is transparent to the X-ray beam [34-35].

The \textit{in situ} reduction of the catalyst was performed at 350°C, under pure hydrogen, during 1 hour. The Pt L\(\text{III}\)-edge was recorded at room temperature under 1 bar pressure of pure nitrogen. Note that \textit{in situ} reduction of the catalyst was followed by XANES (X-ray Absorption Near Edge Structure) measurements. Normalized EXAFS spectra were isolated from the experimental data using a standard procedure [36-37].

Fourier transforms (FT) of the \(k^3\) weighted EXAFS functions were obtained using a Kaiser type window ranging between 3.35 and 13.00 \(\AA^{-1}\) beyond the Pd K-edge and...
between 2.04 and 8.80 Å⁻¹ beyond the Pt L₃-edge (the spectra are noisy above those limits). In this work, all FT are calculated and presented without phase correction. The inverse FT were obtained in the range between 0.169 and 0.318 nm for palladium and between 0.120 and 0.380 nm for platinum. Palladium and platinum metallic foils were employed as references for the Pt-Pt and Pd-Pd bonds [38], while a metallic alloy PtPd was used for Pd-Pt bonds [39].

Regarding the metal-oxygen bonds, the reference powder PdO was used for the Pd-O bond and the reference powder PtO₂ was used for the Pt-O bond. Structural parameters (coordination numbers and interatomic distance) of the reference compounds used in this study are summarized in Table 1. A two-shell least-squares fitting procedure (in k-space) using a single scattering EXAFS formulation was used to extract the coordination number (N), distance (R) and Debye-Waller factor (\(\Delta\sigma\)). Figure 1a and 1b show the quality of agreement obtained in k and R space between filtered experimental data and numerical simulation for the Pd K-edge EXAFS spectrum of the Pd-Pt catalyst. Note that the numerical simulation corresponds to a typical residual of the order of 10⁻².

### Table 1

<table>
<thead>
<tr>
<th></th>
<th>Pd foil</th>
<th>Pt foil</th>
<th>Pd-Pt alloy</th>
<th>PdO</th>
<th>PtO₂</th>
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<tbody>
<tr>
<td>N</td>
<td>12</td>
<td>12</td>
<td>12</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>R (Å)</td>
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<td>2.76</td>
<td>2.73</td>
<td>2.00</td>
<td>2.05</td>
</tr>
</tbody>
</table>

2.6 Catalytic Tests: Hydrogenation of Aromatic Feed with Sulfur Addition

Hydrogenation runs were carried out in a bench scale plant test containing an isothermal fixed bed reactor operating in up-flow mode. Feedstock was a deeply desulfurized Light Cycle Oil (DLCO) which has been prepared by hydrotreating a pure Light Cycle Oil (LCO). LCO is a middle distillate refinery product generated in a fluid catalytic cracking (FCC) unit. The main properties of DLCO are summarized in Table 2. The DLCO has a very low sulfur and a high aromatic content. The sulfur content of this feed has been increased artificially up to 200 ppm S by adding dibenzothiophene (DBT). This procedure allows evaluating the effect of the amount of sulfur on the hydrogenating properties of catalysts for a same aromatic composition of the feed.

Hydrogenation of DLCO feed was carried out under the following conditions: 25 cm³ of catalysts in 1.2 mm extruded shape diluted in 25 cm³ of SiC (in the 1-3 mm particle size range), a total pressure of 6 Mpa, reaction temperature of 300°C, liquid hourly space velocity (LHSV) of 5 h⁻¹, hydrogen-to-feed oil ratio (without recycle) of 450 NL/L.

The catalyst was dried in situ under pure H₂ flow for 4 h at 150°C before being reduced at 300°C for 2 h. After reduction, the catalyst was cooled to 150°C. The initial DLCO feedstock was introduced and the test conditions were adjusted. The change of feedstock containing different sulfur contents was realized after stabilization of performances. In practice, about 300 hours are necessary to stabilize the catalysts. During this first period, the catalysts are passivated by...
sulfur. After that passivation, about 100 hours are necessary to stabilize the catalysts when DBT is added to the feedstock.

The percentage and type of aromatics (total aromatics, mono-aromatics and poly-aromatics) in the feed and the products were measured by near infrared (NIR) spectroscopy correlated to the Fitzgerald method based on mass spectrometry (MS) measurements. The cetane number was also determined by NIR spectroscopy correlated to ASTM D613 (cetane motor). The NIR spectra were recorded on nitrogen purged Bomem MB160 FT-NIR spectrometers equipped with a DTGS (deuterated tri glycide sulfide) detector, operating in transmission mode with a resolution of 4 cm⁻¹. Total amount of sulfur in the products was measured by X-ray fluorescence.

The decrease in the total aromatics content in DLCO was assumed to be pseudo-first-order. The rate constant was defined by the following equation:

\[ k_a (h^{-1}) = \ln \left( \frac{C_{\text{feed}}}{C_{\text{product}}} \right) \times \text{LHSV} \]

with:
\( C_{\text{feed}} \) concentration of aromatic in feedstock and
\( C_{\text{product}} \) concentration of aromatics in product.

### 3 RESULTS

#### 3.1 Main Characteristics

Table 3 summarizes the main characteristics of the catalysts. Nominal compositions are Pd₆Pt and Pd₄Pt. Both elements are very homogeneously distributed along the radial dimension of the extrudates as shown by the R coefficient, very close to 1 (Fig. 2). The symmetry of the two profiles is consistent with a close association of the two metals at least at the resolution of the Castaing microprobe analysis (about 1 µm). The dispersions and the mean diameters determined from H₂-O₂ titration measurements are similar for the two catalysts.

#### 3.2 Transmission Electron Microscopy (TEM)

Regarding the support, electron microscopy shows that the specimen is dominated by a single morphology. More precisely, the different TEM photographic pictures show small plates of cubic γ-alumina extrudates, of 1-10 nm in length. A TEM photograph and the particle size distribution of the Pd₆Pt catalyst, obtained by a statistical analysis of several micrographs, are presented on Figure 3. Most of the particles have a size around 1 nm and the average size by number is 1.8 nm. Larger particles up to 6 nm are also present but to a much lower extent. The probe size is too large compared to the size of the particles to determine the distribution of the different metal elements of the individual catalyst particles.

#### 3.3 Atomic Structure: EXAFS Measurements

The XANES part of the absorption spectrum for Pd₆Pt catalyst is plotted in Figure 4. After in situ treatment under H₂, the platinum atoms are reduced to metallic state as indicated by the similarity in the intensity of the white line of the Pt foil to that of the catalyst [40-41]. Moreover, a comparison of the EXAFS part of the absorption spectra for Pd₆Pt catalyst (Fig. 5) clearly indicated that the local environment of Pt atoms essentially consists of metal atoms. This is confirmed by the shape and the position of the FT modulus (Fig. 6) as well as by the coordination numbers given by the numerical simulation of the EXAFS oscillations (Table 4). More precisely, quantitative analysis gives 3.4 for the average number of platinum atoms and 6.7 for average number of palladium atoms around platinum atoms.

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>Pt wt%</th>
<th>Pd wt%</th>
<th>Pd/Pt molar ratio</th>
<th>R (Pt)</th>
<th>R (Pd)</th>
<th>Support specific surface area (m²/g)</th>
<th>O₂ volume (cc/gcatalyst)</th>
<th>Dispersion %</th>
<th>Mean Diameter (nm)²</th>
<th>Metallic surface area (m²/gcatalyst)³</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd₄Pt</td>
<td>0.6</td>
<td>0.18</td>
<td>4.3</td>
<td>0.98</td>
<td>0.96</td>
<td>265</td>
<td>0.47</td>
<td>84</td>
<td>1.3</td>
<td>124</td>
</tr>
<tr>
<td>Pd₆Pt</td>
<td>0.42</td>
<td>0.67</td>
<td>6.1</td>
<td>0.96</td>
<td>0.96</td>
<td>265</td>
<td>0.59</td>
<td>85</td>
<td>1.3</td>
<td>151</td>
</tr>
</tbody>
</table>

1 calculated with equation (3) or (4).
2 calculated with equation (7).
3 calculated with equation (8) and corrected by the % weight of metal.
Diameter (µm)

Concentration

0 500 1000

Concentration

0 500 1000

Figure 2
Metallic repartitions of platinum and palladium for both Pd₆Pt/Al₂O₃ and Pd₆Pt/Al₂O₃ catalysts by EPMA.

Figure 3
TEM micrographs (with large metallic particles) and histogram of the Pd₆Pt catalyst.

Figure 4
XANES spectra collected at the Pt L₃ edge for the Pd₆Pt catalyst (A) compared to the alloy PtPd metallic foil (B) and the Pt metallic foil (C).

Figure 5
EXAFS data collected at the Pt L₃ edge for the Pd₆Pt catalyst after in situ H₂ reduction (A) compared to the alloy PtPd metallic foil (B) and the Pt metallic foil (C).
A similar approach was carried out at the Pd K-edge. The XANES part of the absorption spectra collected at the Pd K-edge after in situ reduction of the Pd₆Pt/Al₂O₃ catalyst (Fig. 7) indicates clearly that the palladium atoms are reduced to metallic state [42]. Also, a comparison of the different EXAFS parts for Pd₆Pt catalyst above the Pd K-edge shows that the local environment of Pd atoms essentially consists of metal atoms (Fig. 8). Finally, FT modules as well as the values of the different coordination numbers, confirmed these observations (Fig. 9). Regarding the metallic particles present inside the bimetallic catalyst, the analysis of the EXAFS modulations present beyond the Pd K-edge shows for the local environment of palladium 6.9 Pd atoms at 0.273 nm and 0.9 Pt atoms at 0.277 nm.

As previously underlined [43], we have in the case of bimetallic clusters, the following relationship:

\[ R_{\text{PtPd}} = R_{\text{PdPt}} \]
\[ \sigma_{\text{PtPd}} = \sigma_{\text{PdPt}} \]
\[ C_{\text{Pd}} N_{\text{Pd}} = C_{\text{Pt}} N_{\text{Pt}} \]

where \( C_A \) and \( C_B \) corresponds to the atomic concentration of the two metals.

The different measured bond lengths in the samples gave results in good agreement with those calculated in the experimental references. Pd-Pt bond lengths measured at Pd K-edge and Pt L₃-edge are not exactly the same but are coherent with the estimated error of the measurement. Pd-Pt bond lengths measured at Pd K-edge and Pt L₃-edge are not exactly the same but are coherent with the estimated error of the measurement.

\[ R_{\text{PtPd}} = R_{\text{PdPt}} \]
\[ \sigma_{\text{PtPd}} = \sigma_{\text{PdPt}} \]
\[ C_{\text{Pd}} N_{\text{Pd}} = C_{\text{Pt}} N_{\text{Pt}} \]
length in the catalyst is typical of metallic material. Pt-Pt bond length is a little short but remain acceptable. The equality regarding the Debye-Waller factor is also respected. Finally, the relationship between the coordination numbers $N_{\text{PdPd}}$ (average number of palladium atoms around platinum) and $N_{\text{PdPt}}$ (average number of platinum atoms around palladium) is respected, the atomic ratio between platinum and palladium being equal to 6.

3.4 Catalytic Results: Hydrogenation of Aromatic Feed with Sulfur Addition

The Figure 10 shows that the activity of both bimetallic catalysts decreases when the sulfur content of the feed increases. Nevertheless, for each sulfur content in the feed, the Pd$_6$Pt catalyst is found more active than the Pd$_4$Pt catalyst in terms of rate constants. The Figure 11 presents the aromatic contents and the cetane number measured for the products using the Pd$_6$Pt and Pd$_4$Pt catalysts. The higher cetane numbers obtained with the Pd$_6$Pt catalyst are in accordance with its higher efficiency in hydrogenation in presence of sulfur. This appears essentially by a better mono-aromatics saturation, the gain in poly-aromatics saturation being less important. On the other hand, at 20 ppm of sulfur in the feed, the Pd$_6$Pt catalyst allow to reach a PAH (Poly Aromatic Hydrocarbons) target which is inferior to 2 wt % (in the conditions used) contrary to the Pd$_4$Pt catalyst for which the PAH content equals 4 wt %.

Besides the aromatic reduction, hydrodesulfurization is also performed. The sulfur content of products versus sulfur content of the feed is presented in Figure 12. We note that the Pd$_6$Pt catalyst is more efficient in desulfurizing the feed than the Pd$_4$Pt catalyst when the sulfur content of the feed increases (between 20 and 50 ppm of sulfur in the feed, the conversion differences are negligible, whilst they are significant at higher sulfur content).

4 DISCUSSION

Regarding the size of the metallic particles, the complete set of data, coming from HRTEM, volumetric $\text{H}_2$-$\text{O}_2$ titration...
and X-ray absorption spectroscopy, indicate the presence of nanometer scale metallic particles for the Pd₆Pt catalyst. The results given by each technique are quite coherent if we take into account the different advantages and limitations of each characterization technique. Since the limit of detection of TEM is about 6 to 8 Å, very small metallic particles (less than 6 Å in diameter) may exist on the catalyst, which were not detected by TEM. However, whereas the TEM technique can distinguish several populations of metallic particles with different sizes, X-ray absorption spectroscopy and volumetric H₂/O₂ titration can only provide an average particle size. The key point of the EXAFS data is to point out the existence of heteroatomic bonds. At both Pd K-edge and Pt LIII-edge, bonds between palladium and platinum were characterized. Several studies [44-47] have used the synchrotron radiation specific X-ray absorption spectroscopy to give direct structural evidence of the presence or not of bimetallic clusters. The fact that our experiments are performed in situ allows us to build a structural model through the structural characteristics at the atomic level obtained with the different techniques. But what is the distribution of the two metals inside the bimetallic particle?

From a theoretical point of view [48-49] we have now the possibility to describe in a realistic way the electronic structure of nanometer scale metallic clusters and thus to understand both their atomic and chemical rearrangements [50-51]. Moreover, the element with the lowest surface energy is indeed found enriched at the surface: Pd in Pt–Pd and Pt in Pt–Mo, Pt–Rh and Pt–Ru.

From an experimental point of view, the segregation of Pd has been observed in several studies [52]. More recently, nanometer sized particles of platinum and palladium have been prepared through a reaction of platinum and palladium aqueous salts in water-in-oil microemulsions. M. Yashima [53] noticed that the Pt particles are crystalline while the significantly smaller palladium particles are mostly amorphous and often adsorbed on the surface of platinum particles. The presence of palladium at the surface of the nanoparticles has been also pointed out by J. Solla-Gullon [54].

Finally, we have to underline that such distribution between the two metals is not systematic. For example, L. D’Souza [55] found through XPS and CO adsorption studies that the structure of the particles consists of a palladium core and a platinum shell.
In the case of an homogeneous system for which the core of the cluster is composed of \(N(\text{Pt})\) atoms Pt and surface is made of \(N(\text{Pd})\) atoms Pd, we have the relationship:

\[
[N_{\text{PtPt}} + N_{\text{PtPd}}]_{\text{Pt edge}} = 12 > [N_{\text{PdPt}} + N_{\text{PdPd}}]_{\text{Pd edge}}.
\]

Where:
- \(N_{\text{PtPt}}\) is the average number of platinum atoms around platinum (Pt edge),
- \(N_{\text{PtPd}}\) is the average number of palladium atoms around platinum (Pt edge),
- \(N_{\text{PdPt}}\) is the average number of platinum atoms around palladium (Pd edge),
- \(N_{\text{PdPd}}\) is the average number of palladium atoms around palladium (Pd edge).

This hypothesis is based on the absence of monometallic clusters. In the present study, the total number of bonds engaged by the metallic element is more important for platinum i.e. \(N_{\text{PtPt}} (= 3.4) + N_{\text{PtPd}} (= 6.7) = 10.1\) than for palladium i.e. \(N_{\text{PdPt}} (= 0.9) + N_{\text{PdPd}} (= 6.9) = 7.8\). This suggests that the particles have a “cherry” structure: a shell of palladium surrounds a core of platinum atoms.

This approach is consistent with TEM analysis where it appeared that the specimen is dominated by a single morphology of particles. Thus, simple calculations of coordination numbers corresponding of spherical bimetallic particles of different sizes have been considered. In Figure 13, we have selected the case where a shell of palladium atoms surrounds a core of platinum atoms, the core size being considered as a parameter. A simple observation of this figure shows that an agreement can be found between the experimental results and the set of these simple calculations. In order to evaluate the uncertainty for the composition of the surface of the metallic cluster, we have chosen another family of clusters. In Figure 14, the core of platinum atoms contains 13 atoms and the number of shells of palladium atoms is considered as a parameter. Due to the size distribution, it is quite clear that the composition of the surface cannot be precisely determined in terms of the number of surface shells of palladium atoms. But it remains that, in the majority, the particles were found bimetallic, with a “cherry” structure with a preferential distribution of palladium atoms at the cluster surface.

TEM or EXAFS experiments were not performed on the \(\text{Pd}_4\text{Pt}\) catalyst. But since we used the same preparation techniques for the two catalysts and since the results of \(\text{H}_2-\text{O}_2\) volumetric titration were very comparable, the particle size for the two catalysts will be assumed to be the same.
Figure 15
Theoretical predictions [23, 57] for the Pd surface segregation versus Pd bulk concentration of Pd₄Pt and Pd₆Pt catalysts. Surface composition of particles is obtained by the intersect of the curves 100 or 111 orientation and the straight lines.

Figure 16
Pd surface compositions versus particle size calculated from the model described in references [23, 57].

The activity (7.2 h⁻¹) found for the Pd rich catalyst (Pd₄Pt) is slightly higher than one could expect considering the simple increase of exposed metallic surface area keeping the same particle size (6.6 h⁻¹).

One can assume that going from the Pd₄Pt to the Pd₆Pt, not only the number of active sites but also their intrinsic activity are increased. The Pd₆Pt catalyst has therefore an overall intrinsic activity in presence of sulfur higher than the Pd₄Pt catalyst.

Moreover, considering that more Pt atoms are accessible at the surface of the particles in the case of Pd₄Pt, and that the intrinsic activity of Pt is much higher than Pd (by a factor of 1.7), Pd₄Pt should be more active than Pd₆Pt without synergetic effect. Neither can a simple co-operative effect between Pd (hydrogen supplier) and Pt (aromatic hydrogenation site) as proposed in reference [23] explain our catalytic results.

Surface composition is clearly the key point for the understanding of the catalytic properties of bimetallic catalysts. Theoretical calculations as described in references [23, 56] allow the surface composition of small aggregates to be predicted. Application of this model to our case is presented on Figure 15. One can see that for cuboctahedron of order 2 (diameter of about 1 nm), the atomic surface composition of the Pd₄Pt is 90% palladium, the complement being platinum (10%), whereas in the case of Pt₄Pd these values are respectively 95% and 5%. On the basis of this theoretical calculation, we can say that more Pd atoms are accessible at the surface of the particles of the Pd₄Pt compared to the Pd₆Pt catalyst.

On Figure 16, are plotted, for the two bulk compositions, the evolution of Pd surface composition versus the particle size for a truncated cuboctahedron structure (order 2, 3, 4, 5 and 6). For particles smaller than 2 nm, the surface concentration decreases more quickly for the low Pd concentration Pd₄Pt than for Pd₆Pt. The bulk concentration in Pd₄Pt is not high enough to provide sufficient Pd atoms to have a full coverage of the surface of all the metallic particles. This point highlights the fact that the optimum bulk composition is size dependent and the Pd/Pt molar ratio of 4 often reported in literature as the best composition is appropriate in the case of large particles.

Keeping in mind that the real particle size distribution includes particles smaller than 1 nm, one can suppose that, in the case of the Pd₄Pt catalyst, a significant fraction of the overall metallic surface area corresponds to a Pd surface composition much lower than 100% with a significant fraction of exposed Pt atoms. The available amount of Pd, which tends to segregate, is lower than the number of surface sites and full coverage is not achieved. As was observed in a previous study [3], the optimum hydrogenation activity in presence of sulfur is obtained for a precise structure of the active site: a layer of Pd covering a Pt core with adsorbed sulfur coming from the testing conditions. This is not the case for the surface structure of Pt₄Pd and the better results for the Pd rich catalyst (Pd₄Pt) can be associated with this difference in terms of Pd surface composition.

The synergy between palladium and platinum can also be shown if we compare rate constants, determined for the starting feed containing 20 ppm sulfur, of pure Pd and Pt monometallics as shown on Table 5. Because rate constants of the bimetallic catalysts are both higher than those of pure monometallic catalysts, no additivity effect, whatever the size of the catalytic site, can account for this experimental result.
In addition, the sulphur/metal ratios measured on the used catalysts after hot washing with toluene and taking into account the sulphur adsorbed on the alumina carrier, are significantly lower for the bimetallic catalysts (0.1 and 0.2) than for monometallic Pt (0.5). This difference of amount of adsorbed sulphur between mono and bimetallic catalysts has also been observed in a previous study [3] and related to the core-shell structure. The Pd surrounding the Pt atoms inhibits the sulfidation of the Pt atoms, as shown by EXAFS measurements, where no Pt-S bonds in the bimetallic catalysts were observed.

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