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Characterization of industrial catalysts by EXAFS

J. Lynch

_Institut Français du Pétrole, 1 et 4 Ave. de Bois Preau, 92500 Rueil Malmaison, France_

Abstract: Industrial catalysts are designed to present very high specific surface areas and are thus in general poorly crystallised, being often amorphous to X-ray diffraction for instance. In addition the working conditions of these solids are severe, high temperatures and pressures of reactive gases are used during both use and preparation. The penetrating power of X-rays enables _in-situ_ devices to be built so that the the processes involved in activation and deactivation of catalysts can be followed. Several families of industrial catalysts are easily amenable to X-ray absorption spectroscopy analysis. These include sulfide based catalysts, typically with over ten weight percent metal content. Problems are encountered in the study of precious metal systems, for example platinum based bimetallic catalysts supported on aluminium oxide, where metal loading is lower (less than one weight percent). Using _in-situ_ conditions enables studies of the dynamics of reactions and of the state of working catalysts. The structural modifications observed indicate that this type of study is often necessary to obtain meaningful relationships between the state of the solid and its chemical behaviour.

1. INTRODUCTION

Catalysis is a vast field of research. The present paper will only cover a small part of the field, concerned with industrial heterogeneous catalysts. For industrial processes the catalyst is deliberately "designed", usually from knowledge of the fonctionnement of already existing catalysts, to meet a specific need and is built from well controlled starting materials. Heterogeneous catalysis concerns applications in which the catalyst is in a different state (usually solid) to the reactants and products (which may be gases or liquids). This excludes for example natural biological catalysis but also homogeneous industrial catalysis using organometallic molecules. Aside from the obvious structural differences of heterogeneous catalysts, there are general differences in the conditions under which the latter are used and in their performances which make them more difficult to investigate. Heterogeneous catalysis typically takes place under conditions of high temperature and pressure. Often several functions such as selective bond breaking and cycle formation need to be performed by the same solid which will then be multi-component. Thus, although the starting materials are well characterised, the end product is hard to control.

Despite these difficulties, investigation of the environment of active species in catalysts was one of the first applications of X-ray Absorption spectroscopy (XAS)[1]. The potential of the technique was realised so early on partly because few classical methods allow access to structural and electronic information about these systems. The incentive to obtain this information comes from the industrial need to better understand how these systems work and to improve their performances. World consumption of catalysts has reached $10^{10}$ per year, about half of which is dedicated to environmental clean-up such as automobile exhaust control. The size of the market is partly due to the frequent use of precious metals, platinum is a classic example, in the formulations. On this scale, even a one per cent increase in efficiency represents a significant sum of money. The 400,000 tons of catalyst produced each year do not have an unlimited life: although they can be and are regenerated several times, on replacement they...
The sulfide particles are in general larger, partly due to the higher loadings (ten to twenty per cent) possible for this type of catalyst. With TEM and XRD one is able to define the basic structure as twisted books of MoS2 sheets a few "pages" thick. The location of the promoter atoms (inserted into the sheets or as a decoration) is not possible by these methods.
3. INFORMATION FROM XAS

It is not possible here to review XAS theory, for which there are several informative works [2]. Some comments on aspects specific to industrial catalyst studies are however relevant. XAS provides both structural information (from the extended fine structure or EXAFS) and electronic information (from the near edge structure, XANES) on specific elements in a sample. It is not a routine analysis tool however and cannot be used, as for instance electron microscopy is, for control of specific batches of industrial catalysts. The type of information we can aim at for industrial use is therefore more of a generic nature, enabling conceptual models to be built in order to understand how a class of catalysts works and how it can be improved. Although XAS can be deliberately made surface sensitive by the use of special detection systems, in the classical transmission set-up it is a bulk analysis technique. Since, however, the catalysts we are considering have been designed so that the surface is the dominant feature, even a bulk analysis will thus provide information on relevant surface properties. In addition the supported metal and sulfide catalysts may seem to have been designed for XAS analysis since the alumina support adds only a small contribution to the overall absorption. XAS does not require high vacuum, indeed the experiments may be performed under controlled atmospheres with pressures of several bars for elements with edges in the energy range over 10 keV. Several teams working on XAS studies of catalysts have constructed cells enabling the reaction of interest to be carried out during spectrum acquisition, in some cases the reaction rate can be controlled by gas chromatography so that for instance deactivation may be monitored [1,3].

One major handicap to XAS analysis however is that one sees an average over the whole specimen. Close control of the range of particle sizes is necessary in order to provide interpretable data. In addition, as the temperature is increased, the thermal disorder becomes important. Recently molecular dynamics simulations, taking into account the increased freedom of movement of atoms at the surfaces of small particles, have shown that standard EXAFS analysis underestimates coordination number and therefore particle size, with the effect being more pronounced as the temperature increases [4]. It is interesting to note that this work was able to be verified by simultaneous XRD using a combined in-situ set-up. The possibility of combining XAS with classic or anomalous XRD to investigate both short range and long range order opens up new fields of investigation. EXAFS at high temperatures on catalysts is often limited to the analysis of the first coordination shell. Information on the particle shape, or more generally on the three dimensional arrangement of atoms, cannot then be extracted. The type of structural model which can be built from EXAFS data is in these cases fairly qualitative, often a proposed model is only one of several which may fit the data. Structural constraints on the model imposed by XRD, as for those imposed on the chemical composition by X-ray fluorescence and on the size by TEM enable a closer approach to the reality. Thus XAS cannot hope to replace classic laboratory techniques but must work in harmony with them. The major role of XAS stems from the fact that the information provided, even that which must be interpreted semi-quantitatively, is often not available from any other technique.

4. EXAMPLES OF APPLICATIONS TO CATALYSTS

As for XAS theory, it is not possible to cover the whole field of applications of the technique to catalysts. Several reviews exist [5] and the major EXAFS conferences contain special sections on catalyst applications. The two examples treated here demonstrate how even a qualitative overview of XAS data can be of precious help in constructing a model of a catalyst.

4.2. Sulphides

As mentioned above, XAS analysis of these catalysts is facilitated by the relatively high metal loading. The active phase is achieved by sulfuration of oxides under a $\text{H}_2 + \text{H}_2\text{S}$ mixture. The particles are highly reactive and although sealed sample holders may be loaded in a glove box, analysis is best carried out in-situ after sulfuration. This means in general a dedicated apparatus since sulfur is likely to adsorb onto the walls of the cell and is a poison for other catalysts such as metals. The type of information provided by EXAFS is illustrated in figure 2 by a study at the tungsten L\text{III} edge of model tungsten on alumina based catalysts, one of which is promoted by nickel. Although tungsten loading and sulfuration conditions are the same for the two catalysts, simple visual inspection and comparison with WS\text{2} and WO\text{3} references reveals that the unpromoted tungsten is not completely sulfided.

An estimation of the degree of sulfidation may be obtained from the ratio of W-O to W-S bonds. In the case of the promoted catalyst one also observes a lower co-ordination to sulfur for the first shell (5 instead of 6 for WS\text{2}), but especially for the higher shells due to the small particle sizes. Although particle sizes may easily be obtained by TEM for these systems, the incomplete first shell co-ordination of the major active metal is not detectable by other means and suggests an explanation for the catalytic activity in desulfuration. The activation of the catalyst may be followed by in-situ as shown by figure 3 from a study at the Mo K edge of a Ni-Mo/alumina catalyst under flowing $\text{H}_2 + \text{H}_2\text{S}$ mixture. The decrease in Mo-O coordination accompanied by the formation of Mo-S bonds can be followed as a function of temperature.
Figure 2 Comparison with reference oxide and sulfide compounds of unpromoted and Ni promoted sulfided tungsten on alumina catalysts (x axis in Å).

Figure 3 Evolution of a Mo-Ni/alumina catalyst during sulfidation at (a) 20°C, (b) 200°C, (c) 400°C (x axis in Å).

The promoter environnement has been studied at the Ni K edge for different nickel loadings expressed as $\alpha = \text{Ni}/(\text{Ni}+\text{Mo})$ from 0.05 to 0.95. The results shown in table 1 are coherent with the formation of nickel sulfides for high loadings (nickel in NiS$_2$ has a sulfur coordination of 6 at 0.237 nm). This is indeed confirmed by XRD.

At lower loadings where no Ni containing structure is observer by DRX, the EXAFS results suggest the approach to an analogy to the Müller complexes [6] based on the structural element Mo$_2$NiS$_6$ with a Ni-S distance of 0.219 nm and a coordintaion number of four. Returning to the reactivity of these solids, it is noteworthy that on contact with air the return to an oxidised, sulfate like, environnement is visible first in the Ni EXAFS, the change in the Mo coordination being slower. This strongly supports the idea of a decoration of the MoS$_2$ sheets by surface nickel.
Table 1 Coordination numbers N and distances R for nickel at various loadings in sulfided Ni-Mo on 
alumina catalysts.

<table>
<thead>
<tr>
<th>α = Ni/(Ni+Mo)</th>
<th>N (±1)</th>
<th>R (nm ±0.005)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>5</td>
<td>0.220</td>
</tr>
<tr>
<td>0.2</td>
<td>4</td>
<td>0.223</td>
</tr>
<tr>
<td>0.3</td>
<td>5</td>
<td>0.225</td>
</tr>
<tr>
<td>0.6</td>
<td>6</td>
<td>0.235</td>
</tr>
<tr>
<td>0.9</td>
<td>5</td>
<td>0.237</td>
</tr>
</tbody>
</table>

4.1. Supported metals

The activation of these catalysts provides another example of the possibilities of in-situ studies. The 
intensity of the white line XANES feature on the L_III edge has been linked to the density of empty d type 
electron states [1]. By the use of reference compounds, for which the d band electron density has been 
calculated, it is possible to calibrate the intensity in terms of empty states. It is thus possible to follow the 
approach to the metallic state of a calcined sample of Pt/alumina for which the EXAFS shows the local 
coordination of the platinum to be six oxygens.

Figure 4 The number of empty d band states as a function of temperature for a Pt/alumina catalyst under 
flowing hydrogen. The horizontal lines represent data from bulk reference compounds.

In figure 4 one observes a slow reduction of the platinum under hydrogen as the temperature is raised to 
350°C. At this temperature the empty d band state density is equivalent to 0.9 electrons par atom, 
comparable to the theoretical value of one in the 5d^9 configuration of an isolated platinum atom. There 
follows a sharp fall to the value of bulk platinum (0.33 e^-/atom) by 400°C. This simple example can be 
compared directly with temperature programmed reduction (TPR) of the catalyst to confirm the 
findings. In the case shown here one does indeed observe the major peak of hydrogen consumption in 
the range of 300°C to 400°C. In the case of bimetallic catalysts, where it is not possible to attribute TPR 
peaks directly to one or other of the metals, the element specificity of XAS is highly valuable.

For the platinum phase formed by high temperature reduction, the simplest model that fits the EXAFS 
data is of a particle containing 10 to 20 atoms, based on the bulk fcc structure. No firm evidence for 
bonding to the support oxygens is provided: the metal - support interaction may be weak or may be 
affected by a large statistical distribution of distances. These considerations are once again only semi 
quantitative but help to eliminate a large number of possible models incompatible with the EXAFS 
results.

In the study of bimetallic catalysts a major question to be addressed is the formation or not of an alloy 
phase. The preparation procedures normally used, involving successive impregnation or mixtures of salts 
of the two metals, do not guaranty the formation of an alloy and modifications to the catalytic activity 
may be interpreted as due to reactions occuring first on one monometallic phase then on another, close
by on the support but not in intimate contact. One example in which the formation of an alloy is evident from the EXAFS results is a model Pt-Zr catalyst. Here the particles are fairly large, around 10 nm in diameter, leading to high signal to noise ratios in the data. Figure 5 shows the pseudo-radial distribution functions around platinum of a bimetallic Pt-Zr and a monometallic Pt/alumina catalyst prepared so as to have the same particle size.

Figure 5. Pt LIII edge data for monometallic Pt and bimetallic Pt-Zr catalysts (x axis in Å).

The large difference in the distributions comes from the phase difference between platinum and zirconium neighbors leading to a beat in the EXAFS spectrum and a splitting of the first neighbor peak. The fit to the data in the Pt-Zr case, with four Zr and eight Pt neighbors at 0.285 nm, is in close accord with the hypothesis of formation of a Pt3Zr alloy. Again no interaction with the support is detected by EXAFS, although for large particles this is not surprising as any metal atoms at the interface will be in a minority.

The above case, with large particles, is for EXAFS practically equivalent to the analysis of bulk samples. The acquisition times used were of the order of ten minutes. As the particle size decreases the number of neighbors decreases and for bimetallic systems they may be of two types when an alloy is formed. For realistic systems, with subnanometer particles, one is thus faced with the need to extract more information from data where the signal to noise ratio is intrinsically lower. Acquisition times are thus much longer, of the order of hours.

In contrast to the monometallic and the large particle cases for which no particle-support interface can be defined from EXAFS, for bimetallic Pt-Sn catalysts recent work [7] has shown the existence of Pt-O bonds as well as Pt-Sn and Pt-Pt bonds in the reduced solid. The XANES of platinum confirms that the element is in a reduced state. Table 2 resumes the structural parameters deduced for monometallic and bimetallic catalysts after calcination (the results are identical for the two catalysts) and after reduction. The EXAFS results provide not only proof of intimate contact between the two metals but also evidence for an improved particle-support contact, which is sufficiently well defined to be detected.

Table 2. Platinum LIII EXAFS results for calcined and reduced supported metal catalysts (estimated precision is 15% for the coordination number N, 0.002 nm for the distance R).

<table>
<thead>
<tr>
<th>nearest neighbor</th>
<th>calcined Pt or Pt-Sn</th>
<th>reduced Pt/alumina monometallic</th>
<th>reduced Pt-Sn / alumina bimetallic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>R (nm)</td>
<td>N</td>
</tr>
<tr>
<td>O</td>
<td>6</td>
<td>0.201</td>
<td>0.0</td>
</tr>
<tr>
<td>Pt</td>
<td>0</td>
<td>-</td>
<td>6.4</td>
</tr>
<tr>
<td>Sn</td>
<td>0</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

The clear increase in strength, or at least in coherence, of the metal support interaction, attributed to the presence of tin in the coordination sphere, is of considerable importance in understanding the resistance to sintering of platinum-tin catalysts, which has itself been the subject of XAS and TEM investigation. Under repeated cycles of reduction and oxidation the monometallic catalysts lose their activity due to the formation of large particles, clearly visible in TEM reaching sizes of several tens of nanometers. The bimetallic catalysts on the other hand retain both their activity and the high dispersion of the metal up to six cycles.

The use of in-situ cells has allowed the study of catalysts in contact with hydrocarbons at high temperatures (300°C and 460°C). In a recent study [8] the behaviour of monometallic Pt/alumina and
bimetallic Pt-Re/alumina samples was compared. The reforming of n-heptane was chosen as a test reaction. When the monometallic catalyst is working (as monitored by gas chromatography) the platinum L₂3 EXAFS is modified with the appearance of new bonding due to low atomic number neighbors at short distances. Analysis showed these to be carbon atoms. Although it is notoriously difficult to distinguish atoms of close atomic number, the possibility of Pt-O bonds was eliminated based on the phase behaviour of the EXAFS oscillations but also on chemical reasoning: the formation of Pt-C bonds is to be expected in a working catalyst. No other modifications to the metal particle structure were observed. In the case of monometallic catalysts Pt-C bonds were detected (with a coordination number of 1.2 at 0.194 nm) at both temperatures studied. For the bimetallic catalyst only at the lower temperature were Pt-C bonds observed.

The fact that the Pt-Re catalyst was still functioning at high temperature may mean that contact of the hydrocarbons is occurring only with the rhenium atoms, but the platinum-carbon interaction itself may also be strongly modified. Indeed at lower temperatures, where carbon bonds are seen, both the coordination number (2.0) and more significantly the distance (0.209 nm) are different. A modification of such magnitude in the Pt-C bond length seems to indicate a change in the type of bonding in the bimetallic system compared to the monometallic case. There is indeed a significant improvement for Pt-Re catalysts in resistance to the formation of coke, one of the major causes of deactivation in industrial applications.

5 PERSPECTIVES FOR XAS

The examples shown above demonstrate that significant information can be obtained from EXAFS at a qualitative level on model systems close to industrial catalysts. This information is extremely valuable in defining a general view of the structure of catalysts and how they work. The possibility to use EXAFS in-situ has enabled modifications to the structure with temperature and in contact with reactants to be followed. No attempt has been made here to develop atomic scale models in three dimensions of the catalyst particles or of their interactions with promoters and hydrocarbon molecules as this is far from being a straightforward process. Indeed, for the greater part of the systems studied, it is impossible using EXAFS alone to build such a definitive model. Two technological developments are likely to improve the situation: access to new high intensity sources and coupling of EXAFS to other techniques. In particular for the supported metal catalysts studied, where the loading is 1% or lower by weight, the signal to noise ratio of the data prohibits extraction of information from outer shells of coordination which would be useful in construction of a three dimensional model. The coupling of EXAFS to diffraction has already been mentioned as being of great importance in studying particle sizes. In other examples shown here, such as the activation of molybdenum sulfide catalysts, simultaneous diffraction data would also be invaluable in identifying three dimensional phase formation.

The data presented here were acquired at the LURE facility in Orsay in collaboration with H. Dexpert and D. Bazin to whom many thanks are due.

References