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EXAFS in situ: The effect of bromide on Pd during the catalytic direct synthesis of hydrogen peroxide

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In memory of the late Professor Benedetto Corain (July 8th 1941 - September 24th 2014), passionate chemist and teacher

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ABSTRACT

The direct synthesis of hydrogen peroxide is catalyzed by palladium catalysts supported over different solids. With the aid of a suitable plug-flow reaction cell, we carried out a preliminary X-ray absorption fine structure spectroscopy (EXAFS) characterization of a palladium catalyst supported on the commercial resin K2621 under reaction conditions (in situ). Whereas the catalyst, which in the dry catalyst presents metal Pd and PdO when fresh, is practically unaffected by the reaction medium (methanol) or by the reaction mixture ($CO_2/H_2/O_2$, 86/10/4, v/v) it undergoes an apparent reduction of part of PdO to metal Pd and some metal leaching during the reaction in the presence of bromide ions. These findings suggest the role of bromide ions as enhancers of the selective blocking of the sites responsible for the undesired formation of water, but could also entail phase modifications of the active metal.

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1. Introduction

Heterogeneous catalysts supported on commercial ionexchange resins for the direct synthesis (DS) of hydrogen peroxide from the elements, $H_2 + O_2 \rightarrow H_2O_2$, has been under investigation for several years [1–5].

This reaction is of interest to both industry [6-11] and the academia [12-24] for the small scale, on-site and on-demand production of this commodity, for which the anthraquinone autooxidation (AO) process is not convenient [12]. The key issue in this reaction is the selectivity towards H_2O_2 , because the active Pd or bimetallic Pd–Au or Pd–Pt catalysts also promote the formation of water, that is the thermodynamically favoured product.

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The addition of gold to the palladium catalysts [25] or the addition of halide ions (Cl⁻, Br⁻) to the reaction mixture, especially in combination with mineral acids [13], are the most effective ways to enhance the selectivity reported so far. While enhancing the selectivity, the halide ions decrease the reaction rate [13]. Purportedly they avoid the dissociative chemisorption of dioxygen by blocking preferentially the sites where it takes place [13,25]. This hypothesis is supported by theoretical models obtained from DFT calculations [26], but the mechanism is still unclear and sound direct evidence on the nature of their action not achieved yet.

To the best of our knowledge, no information on the role played by halide ions as DS promoters has been gathered in situ and reported so far. This kind of information is highly desirable [27] and

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in this context we have recently reported on the implementation of a cell for the XAFS analysis of solid catalysts under gas-liquid-solid conditions [28].

The XAFS (X-ray absorption fine structure spectroscopy) is a well ascribed [29,30] chemically selective technique to probe the chemical nature of the absorber (valence state) its coordination chemistry, and the local atomic structure around it. The XAFS signal is constituted by the fine oscillations that can be observed on the atomic absorption coefficient of not isolated atoms. The scattering theory describes these features as caused by the interference between the photoelectron waves outgoing from the absorber and scattered by the surrounding atomic potentials [31]. The XAFS signal is traditionally distinguished in two regions: the near edge (XANES: X-ray absorption near edge structure) and the extended (EXAFS: extended X-ray absorption fine structure) regions. In the EXAFS region a relatively simple and well ascribed formalism allows to understand the local atomic structure around the average absorber in the sample as a combination of coordination shells whose parameters (coordination numbers, coordination distances and disorder factors can be accurately determined by least square refinement procedures [31-33]. The theory required for ab-initio modelling the XANES features is generally complex because they are affected by the finest details of interatomic potentials and full multiple scattering effects take place [31]. However the XANES region is dense of structural and electronic information that can be often understood comparing the data of reference compound [30].

We report herein on the in situ XAFS investigation with this cell of a nanostructured Pd catalyst for the DS working in the presence of sodium bromide. In view of our long-standing interest into polymer-supported metal catalysts [34–38] and into their application to DS [5], this preliminary investigation was focused on a monometallic Pd catalyst supported on the commercial ion-exchange resin Lewatit K2621. The experiments performed with the in situ XAFS cell, have been carried out under atmospheric pressure. Although this condition could not be optimal for practical applications, valuable information can be anyway collected as shown by the achievements of Lunsford's and Strukul's groups in the field [19,20,23].

2. Experimental

Lewatit K2621 (sulfonated polystyrene-divinylbenzene macroreticular ion-exchange resin; exchange capacity = 1.92 mmol/g) was kindly provided by Lanxess. All the reagents and the materials were used as received from Sigma–Aldrich.

The preparation of the catalyst Pd/K2621f is described in ref. [4]. The detailed description of the catalytic cell for in situ XAFS measurements is reported in ref. [28]. In a typical experiment, an aqueous solutions of [Pd(NH₃)₄]SO₄ (0.0256 g, 0.188 mmol, corresponding to 1% Pd (w/w), in the final catalysts) was added to a suspension of 1.0104 g of K2621 in 10 ml of distilled water, after swelling for 2 h. The suspension was let to react overnight on a swirling plate. The solid was recovered by vacuum filtration, carefully washed with distilled water $(3 \times 10 \text{ ml})$ on the gooch filter and dried overnight at 110°C. The filtrate was combined with the washing liquor and analyzed with ICP-MS for the quantitative determination of the unreacted metal. After the ion-exchange, the residual amount of Pd in the mother liquor was less than 0.1% of its starting amounts. Consequently, the uptake of palladium was complete and the experimental metal loading practically corresponds to the nominal one for each sample.

The beige (very light brown-yellow) solid was suspended in 50 ml of 37% aqueous formaldehyde solution and heated up to reflux condition with an oil bath for 3 h. The black product was recovered by vacuum filtration and carefully washed with distilled water (3 \times 10 ml). The material was dried in oven at 110 $^\circ\text{C}$ overnight.

XAFS measurements at Pd K-edge (24,350 eV) in transmission geometry have been collected at ELETTRA-XAFS [39] and ESRF-BM08 [40] beamlines. The structural EXAFS (extended-XAFS) signal has been extracted and quantitatively analyzed using ESTRA and Fitexa programs, respectively [41].

The catalyst was characterized by EXAFS at room temperature in the dry state (polyvinylpirrolydone pellet), in the methanolswollen state, with the alcohol flowing through the catalyst placed inside the measurement cell after swelling with methanol, and under reaction conditions, respectively with or without 10 ppm of NaBr dissolved in methanol. For the latter experiment a mixture of methanol (or a 10 ppm NaBr solution in methanol), pre-saturated with CO₂, H₂ and O₂, of H₂ and of O₂ were continuously let to flow through the catalyst bed inside the measurement cell. As described in Ref. [28] each gas was separately fed with a dedicated syringe at atmospheric pressure at the desired flow rates. Flow rates (CO₂: 1.72 ml/min, O₂: 0.20 ml/min, H₂: 0.08 ml/min; CO₂/H₂/O₂ = 86/10/4, v/v) were set as to keep the gas mixture out of the explosive range [42]. The liquid was fed using a peristaltic pump.

The same procedure has been employed also for the off-line catalytic runs for the quantitative analysis of H_2O_2 and the concentration of H_2O_2 has been measured by iodometric titration.

3. Results and discussion

In this work we investigated with EXAFS in situ a palladium catalyst obtained during a previous investigation of ours [4] under halogen-free conditions to prevent the accidental retention of halide ions. For this purpose $[Pd(NH_3)_4]^{2+}$ was used as the metal precursor and ion-exchanged for H⁺ in the resin Lewatit K2621. The immobilized metal complex was the reduced with 37% aqueous formaldehyde at reflux.

Since the goal of this investigation was to simply check the reliability of the apparatus for the characterization in situ of the catalysts for the DS reaction, we did not attempt any quantitative estimate of the production of H₂O₂ and of the selectivity towards it during the XAFS measurements. The production of H₂O₂ was only qualitatively shown upon the addition of an acidic solution of $TiOSO_4$ [43] to samples of the liquid phase collected at the cell outlet. However, representative off-line experiments with the same experimental set-up, carried out in our laboratory, showed that under these conditions the concentration of H₂O₂ is around 0.06 mM and 0.02 mM in the absence and in presence of bromide ions, respectively. These values are quite low, but it should be borne in mind that the reactor was not optimized for catalytic performance. Nevertheless, these outcomes are in agreement with the literature, showing that bromide ions as the selectivity enhancers produce a significant decrease of the catalytic activity [12,13].

The results of the analysis of the EXAFS measurements collected under different experimental conditions at room temperature are summarized in Table 1. In Fig. 1, EXAFS signals from the as prepared catalysts (a) and after the treatment with 1.0 ml/min methanol for 2 h (b) are reported with their best fit curves, as an example. The contributions to the total EXAFS from the metal and oxidized (PdO) Pd phases are also shown (lower part of each panel).

In the as-prepared catalyst we found both metal Pd and PdO. After some preliminary tests, the metal phase was assumed to have the fcc structure. The interatomic distances of metal Pd are constrained to the fcc structure, leaving only the fcc edge (a_{Pd}) as free parameter in the refinement [44]. The a_{Pd} = 3.88(1)Å is in good agreement with the literature value $(a_{Pd}$ = 3.89Å) for bulk Pd. The oxide phase (PdO) was included with signals from two

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