Microwave Plasma Treatment for Catalyst Preparation: Application to Alumina Supported Silver Catalysts for SCR NO\textsubscript{x} by Ethanol

Marjorie Foix\textsuperscript{1,2}, Cédric Guyon\textsuperscript{1}, Michael Tatoulian\textsuperscript{1}, Patrick Da Costa\textsuperscript{2*}

\textsuperscript{1}Chimie Paristech, LGPPTS, Plasma Chemical Engineering Laboratory, Paris, France
\textsuperscript{2}UPMC Paris 6, Sorbonne University, IJLRA, JEan Le Rond d’Alembert Institute, Saint Cyr L’Ecole, France

Email: *patrick.da_costa@upmc.fr

Received January 12, 2013; revised March 5, 2013; accepted April 12, 2013

ABSTRACT

Thermal treatment either in the presence of oxygen (calcination) or of a reducing agent (reduction) result is all the time a key issue within the preparation of a catalyst. In this work, a microwave plasma treatment was chosen as an alternative to typical calcinations, because it is a more energy efficient process. Thus, a Microwave Fluidized Bed Plasma reactor (MFBP) was employed in catalyst synthesis process under different gas compositions, such as argon and argon/oxygen mixtures over $\gamma$-alumina supported silver catalysts, which are generally used for selective reduction of NO\textsubscript{x} by ethanol. After the first catalytic tests performed in the presence of plasma treated catalyst, it can be concluded that plasma treatment process represents an interesting alternative to conventional calcination during catalyst synthesis, resulting in a more sustainable process, moreover in view of its industrial application. In order to understand the particular effect of plasma treatment, the catalysts submitted to this treatment were carefully characterized by means of thermogravimetric analysis (TGA), differential thermal analysis (DTA) and UV-VIS-NIR.

Keywords: Microwave Plasma; Fluidized Bed Reactor; Catalyst Synthesis; Silver Catalyst; NO\textsubscript{x} Reduction

1. Introduction

All around the world, the major chemical engineering processes are catalytic processes. Most the used catalysts are heterogeneous supported catalysts. An effective catalyst is generally described to have a high specific surface area in which the catalytic phase is highly dispersed. In commercial form, the final catalyst should also possess high mechanical strength and resistance to high temperatures [1].

Conventional synthesis of supported catalysts comprises four main steps, namely: 1) impregnation of the active phase; 2) maturation; 3) drying; 4) thermal activation such as calcination or reduction. Nowadays, the lack of an accurate control of some of these operations results in catalytic preparation procedures, which may be far from perfect [2,3].

In the field of catalysis, Plasma technologies have been found to be a solution in order to improve the activity or the stability of the processes [4]. Low-temperature plasmas are often used for surface processes like cleaning, activating or coating. Moreover, plasma technologies have been used within different processes as a way of activating many molecules involved, enhancing the global reaction kinetics. In example, Kiyokawa et al. [5] decomposed nitric oxide, which is one of the most notorious atmospheric pollutants, in the presence of atmospheric pressure plasma. Additionally, this method is also a promising physical tool for biological decontamination and sterilization. Deng et al. [5] described a glow discharge plasma jet and its applications in protein destruction by using a dielectric barrier discharge arrangement. Kolb et al. [6] demonstrated that an air micro plasma jet can effectively treat yeast infection on skin.

Other authors have proposed to use plasma technologies in the synthesis process of the catalysts [7-11]. In general, there are three main trends in catalyst preparation using plasma technologies, such as: plasma chemical synthesis of ultrafine particle catalysts [7-9], plasma assisted deposition of catalytically active compounds on various supports [7,11] and plasma enhanced preparation or plasma modification of catalysts [9]. Vissokov et al. [7]
presented a synthesis of a γ-Al₂O₃ from elementary aluminium and oxygen under plasma jet. The condensed-phase catalyst preparation can be carried out in both equilibrium and non-equilibrium plasma conditions. Rouleau et al. [9] presented a synthetic catalytic system prepared by plasma discharge spark between two metal electrodes immersed in a light hydrocarbon medium. Xia et al. [8] presented the preparation of palladium catalysts supported on structured carbon nanotubes. It was shown that the oxygen-containing groups introduced by the plasma treatment on the surface are essential to achieve a high Pd loading. There are two techniques regarding plasma enhanced preparation or plasma modification of catalyst: plasma heat treatment and plasma chemical treatment. Plasma heat treatment is carried out to replace the catalyst thermal calcination [11]. On the other hand, one of the important applications of plasma chemical treatment is the modification of zeolite acidity.

Within plasma technologies, cold plasma process particularly stands out as a considerably efficient and practical method for the selective modification of the surface of raw materials, avoiding any change in their properties. Therefore, plasma can be employed in the functionalization of several materials and activation of their surface, depending on the nature of the gas phase and experimental treatment conditions. Since, most of the time the raw materials are in powder form, Fluidized bed reactors (FBR) offer the possibility of achieving homogenous treatment as well as an accurate control of the gas-solid particles heat transfert rates [11,12]. In the literature, the plasma technology was already used after catalyst preparation in order to modify the raw material [13,14]. A recent work showed that radio frequency plasma treated catalysts led to higher efficiency in NOₓ abatement for methane SCR of NOₓ than classical calcined catalysts [15]. The aim of our work is to prepare a catalyst dedicated to NOₓ abatement in mobile sources using plasma during the synthesis process. In this context, a silver-based catalyst was chosen since the studied reaction is Ethanol SCR [16].

Upon preparation of different catalysts, the efficiency of fluidized bed plasma treatment of Ag/Al₂O₃ catalysts activation has been compared versus classical calcination in air at 500°C. Different techniques have been used for physic-chemical and activity characterization, such as temperature programmed surface reaction (TPSR), thermogravimetric analysis (TGA) and differential thermal analysis (DTA), transmission electron microscopy (TEM) coupled by energy dispersive spectroscopy (EDS), UV-Vis spectroscopy. In this study, we have also shown that the role of the microwave plasma treatment time plays a definitive role in the efficiency of alumina supported silver based materials.

2. Experimental

2.1. Catalyst Preparation and Microwave Plasma Fluidized Bed Reactor

A classical excess solution impregnation was chosen as synthesis method for catalyst preparation. Thus, g-Al₂O₃ spheres of 600 µm (170 m²·g⁻¹), chosen as a support in order to obtain a homogeneous fluidization and to avoid loss of small particles in the pumping system, were introduced in an aqueous solution containing silver nitrate. The silver loading was fixed at 2.5 wt% because this loading was found to be optimal for such catalytic application [17]. Then, the excess of solution was removed by the means of evaporation at 60°C under reduced pressure as reported elsewhere [17]. Then, the as-prepared catalysts were dried at 120°C overnight, prior to the two different treatment processes:

1) Classical calcination in oven under synthetic air (20% O₂/80% N₂) at 500°C for 2 h with a heating rate of 10°C·min⁻¹;

2) Microwave Fluidized Bed Plasma treatment in Fluidized bed reactors (FBR) reactor.

As reported in Figure 1, the Microwave Plasma Fluidized Bed Reactor (MPFBR) consists of a cylindrical glass tube of 30 mm inner diameter and 600 mm length.

After drying, impregnated alumina supported silver catalysts (r: 795 kg·m⁻³, mean-diameter of particles: 600 µm) were fluidized using the pre-treatment gases.

Moreover, we worked at the minimum fluidization conditions determined from the pressure drop versus gas flow rate [18]. All the characteristics of the reactor are described elsewhere [11].

The plasma is created as the following: two external electrodes were placed at the level of the fluidized bed and the plasma was maintained by a Microwave energy source using a Sairem generator (power range 0 - 300 W). A wattmeter was used in order to follow incident and reflected powers, its characteristics are reported elsewhere [19]. The different plasma parameters used for the final treatment of Ag/Al₂O₃ catalysts are listed in Table 1. The samples were denoted MW-X, in which X corresponds to a specific microwave treatment time or gas mixture.

2.2. Physical and Chemical Catalysts Characterization

Scanning electron microscopy (SEM) was performed on a JEOL JMS 6400 coupled with an energy dispersive spectrophotometer (EDS) analyser link EXL. For the transmission electron microscopy (TEM), the samples were deposited on a carbon film supported on a copper grid in dry conditions in order to avoid metallic silver particles [20]. Ultraviolet-visible diffuse reflectance spectra (UV-vis DRS) were collected using a spectrophotometer.
Figure 1. Experimental set-up of the fluidized bed reactor (FBR) used for the plasma surface treatment of powders.

Table 1. Experimental parameters used during the microwave plasma treatment.

<table>
<thead>
<tr>
<th>Experimental parameters</th>
<th>Catalyst code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Argon flow rate (cm³/min)</td>
<td>300</td>
</tr>
<tr>
<td>Oxygen flow rate (cm³/min)</td>
<td>0</td>
</tr>
<tr>
<td>Pressure (mbar)</td>
<td>0.70</td>
</tr>
<tr>
<td>Plasma power (Watt)</td>
<td>500</td>
</tr>
<tr>
<td>Treatment time (min)</td>
<td>10</td>
</tr>
</tbody>
</table>

(Lambda 40, Perkin-Elmer) equipped with a diffuse reflectance attachment (DRA) (Harrick). Spectra were collected at room temperature in air in the range of 190 - 500 nm with a resolution of 2 nm. As a reference we used a compressed pellet of polytetrafluoroethylene (Teflon) powder, which is superior to the conventional BaSO₄ layer with regard to both spectral smoothness and reproducibility [21]. X-ray diffraction patterns were obtained on a Seifert XRD 3000 radiation. We carried out t spectrometer using Cu Kα instrument coupled to mass spectrometry from Room Temperature (RT) up to 800°C in 80%N₂/20%O₂ mixture (100 cm³·min⁻¹, 30 mg of catalyst).

2.3. selective reduction of NOₓ by Ethanol

The catalytic runs of the NOₓ reduction by ethanol (EtOH) were performed in a fixed-bed flow reactor (10 mm inner diameter) by flowing 500 ppm NO, 2500 ppm EtOH, and 10 vol% O₂ in Ar as carrier gas with a flow rate of 184 cm³·min⁻¹ using 0.045 g of sample corresponding to a Gas Hourly Space Velocity of 190,000 h⁻¹. Temperature programmed surface reaction (TPSR) was carried out at a heating rate of 10°C·min⁻¹ up to 500°C. In order to study the reaction in steady-state conditions, isothermal steady-state reaction study was performed subsequently, from 500°C to 250°C, by steps of 50°C, to avoid adsorption/desorption processes. At each temperature to reach the steady state and thus to evaluate the NOₓ conversion into N₂, the temperature was maintained for 1 h. The temperature was increased up to 500°C. The products of the reaction were quantified by the mean of various analyzers: a micro-chromatograph Varian for N₂.
production, an infrared Ultramat Siemens detector for CO, CO₂, a chemiluminescence Ecophysics detector was used for NOₓ detection (NO and NO₂); the other compounds were followed by mass spectrometry (MKS Instrument). Only the ethanol was detected by mass spectrometry during our experimental tests, thus no oxygenated intermediates have been evidenced.

3. Results and Discussion on the Ag/Al₂O₃

3.1. A Sustainable Plasma FBR Treatment versus a Classical Thermal Treatment

The classical calcination was carried out under air at 500°C for 2 hours leading to a non-sustainable process due to use high temperatures and the consumption of industrial air. In our FBR system, the process takes place at low temperature. In the FBR system, the temperature of gas was found around 127°C ± 50°C. In order to verify this value, we used powders materials knowing their melting point precisely, these samples references, very sensitive to a change of temperature, were used to generally calibrate a bench of Kofler. Table 2 indexes the samples which were placed on sintered engine during a M.W plasma treatment to determine the temperature range. For a microwave plasma treatment (a power 500 Watt, a pressure of 0.70 mbar and an argon flow of 300 cm³ min⁻¹), the Azobetol sample melts on the contrary to the Benzil sample. Thus we can admit that the temperature of plasma M.W lies between 68°C and 95°C. This value corresponds so that we found by calculations. Thus, the temperature of plasma is definitely lower than the calcination temperature (500°C).

3.2. Influence of Plasma Treatment on the Steady-State Activity of the Catalysts in the SCR of NOₓ in the Presence of Ethanol

3.2.1. Effect of Plasma Treatment Time

The reaction of NOₓ by ethanol was carried out on all pre-treated catalyst (MW-X and calcined ones). In steady-state conditions, in isothermal range, the NOₓ reduction efficiency of alumina supported silver catalysts increased with the temperature, reaching a maximum around 400°C for all samples. Figure 2(a) compares different microwave plasma treatment with 500 W under argon as carrier gas, in which only the processing time varies. All the microwave plasma-treated catalysts show better conversion that the sample without treatment. We observe that the processing time of 30 min (M.W-2) lends on all the range of temperature to a higher NOₓ conversion compared to the calcination. Thus, at 300°C, 55% of NOₓ conversion is observed on M.W-2 vs. 43% on Calcined sample and at 400°C, 95% of NOₓ conversion is obtained with M.W-2 catalyst whereas only 80% of NOₓ conversion is achieved with the calcined sample.

We can also notice that the sample treated during 1h (M.W-3) reaches high conversions from 250°C to 370°C such as treatment M.W-2. Then from 370°C to 520°C, a decrease of NOₓ activity is observed. We can suppose that this longer treatment generated reactions at the surface and modified the active sites. Similar result has already been reported for methane SCR of NOₓ on palladium based catalysts [11] using another plasma system. The microwave plasma treatment of 10 min leads to intermediate conversions in comparison with the sample without treatment and the calcined sample.

3.2.2. Effect of Power of Plasma on the Catalytic Activity

On Figure 2(b), the effect of the power on catalyst was presented. An increase of power from 500 W and 800 W for a similar treatment time of 30 min, does not affect the NOₓ conversion on silver based catalysts. Similar effect has been found with 10 and 60 min respectively (Figure not shown). Thus, one can suppose that similar active sites remain on both samples.

3.2.3. Influence of Gas Composition of the Plasma Figure 2(c) compares plasma treatment of 500 W during 30 min under two different carrier gases: argon alone (M.W-2) and 20% oxygen/argon (M.W-4). On the two M.W. plasma treated show a better conversion than that obtained on the no-treated sample and on the calcined one. The oxygen brought in the plasma treatment does not have a particular influence on this catalytic system Ag/Al₂O₃ for 500 W. However, with higher power (800 W) (Figure 2(d)) we notice a catalyst deactivation because oxygen recombines with higher power creating unfavourable reactions [22]. All the plasma treated samples lead to higher NOₓ conversion than the untreated sample. In view of the results obtained to obtained results, the M.W-2 plasma treatment is most effective. Similar results have been obtained using another kind of plasma generator such as radio frequency generator [11].

Table 2. Range of samples reference.

<table>
<thead>
<tr>
<th>Material name</th>
<th>Melting point (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azobenzol</td>
<td>68.0</td>
</tr>
<tr>
<td>Benzil</td>
<td>95.0</td>
</tr>
<tr>
<td>Acetanilide</td>
<td>114.5</td>
</tr>
<tr>
<td>Phenacetin</td>
<td>134.5</td>
</tr>
<tr>
<td>Benzanilide</td>
<td>163.0</td>
</tr>
<tr>
<td>Salophen</td>
<td>191.0</td>
</tr>
<tr>
<td>Dicyandiamide</td>
<td>210.0</td>
</tr>
<tr>
<td>Saccharin</td>
<td>228.0</td>
</tr>
</tbody>
</table>
Figure 2. NO\textsubscript{x} conversion in N\textsubscript{2} as function of temperature during the steady-state reaction of NO\textsubscript{x} reduction by EtOH over Ag/Al\textsubscript{2}O\textsubscript{3} catalyst; (a) As a function of different microwave plasma treatments at various time of treatment (♂) M.W-1, (□) M.W-2, (△) M.W-3. And (★) calcination, (●) without treatment; (b) As a function of different microwave plasma treatments at various powers: (□) M.W-2, (■) M.W-5. And (★) calcination, (●) without treatment; (c) As a function of different microwave plasma treatments at various carrier gas at 500 W: (□) M.W-2, (○) M.W-4. And (★) calcination, (●) without treatment; (d) As a function of different microwave plasma treatments at various carrier gas at 800 W: (■) M.W-5, (▲) M.W-6. And (★) calcination, (●) without treatment. (Reaction conditions: 500 ppm NO, 2500 ppm EtOH, and 10 vol% O\textsubscript{2} in Ar at a flow rate of 184 cm\textsuperscript{3}·min\textsuperscript{−1} over 0.045 g alumina spheres catalyst (GHSV = Ag/190,000 h\textsuperscript{−1})).

3.3. Characterization of the Plasma Treated Catalysts by Means of TPSR of NO\textsubscript{x} in the Presence of Ethanol

Temperature Programmed Surface Reactions (TPSR) were performed on the following samples: 1) untreated and 2) calcined samples and 3) all microwave plasma treated catalyst. Figure 3(a) shows the influence of the treatment time of microwave plasma. We can notice that the TPSR results are in agreement with the results obtained in steady-state conditions. The plasma-treated catalyst-named M.W-2 (500 W-Ar-30 min) allows a better adsorption of species NO\textsubscript{x}, with 400°C this treatment reaches 88.8% NO\textsubscript{x} conversion. This conversion is very close to that obtained by the calcined sample with around 81% NO\textsubscript{x} conversion at 365°C. Whereas a M.W-1 plasma treatment (500 W-Ar-10 min) only leads to 57.2% of conversion and on M.W-3 (500 W-Ar-60 min) only 80.6% of conversion is obtained. We can see that its conversion decreases to reach at the end of the rise of temperature the M.W-1 plasma treatment. If the treatment plasma M.W is prolonged then the catalyst will lose its effectiveness. On Figure 3(b), we compare the influence of the power of the plasma used. We notice that the all plasma treatments allow a good activation of catalyst since the NO\textsubscript{x} concentration are always lower for such catalysts compared to the sample without treatment. A NO desorption higher than 500 ppm is observed for all the treatments in temperature. Whereas for the untreated sample, we distinguish that a NO desorption around 250°C indicating that the sites of adsorption are not all free sites. Figure 3(c) shows the influence of the nature of the carrier gas. The obtained results are comparable with those obtained in steady-state conditions (Figure 2). The M.W-4 plasma treatment (500 W-20% oxygen/argon-30 min) leads to lower conversion of NO\textsubscript{x} than the M.W-2 plasma treatment under Argon. This effect of the carrier gas 20%O\textsubscript{2}/Ar is also reproduced with microwave plasma with 800W. On Figure 3(d), one can see that a sample treated with a plasma under argon (M.W-5) converts more NO\textsubscript{x} at high temperature than a sample treated with a microwave plasma under 20% oxygen/argon (M.W-6).

3.4. Influence of Microwave Plasma Treatment on TGA-DTA

The effect of the plasma activation was followed by means of DTG-DTA, analyses were performed on: 1) untreated freshcatalyst; and 2) microwave treated catalysts; and 3) calcined catalysts.

From Figures 4(a) and (b), presenting the thermogravimetric and differential thermal analysis results, the initial weight loss with a maximum close to 100°C corresponds to the loss of physisorbed water [23]. The TGA curve indicates a major weight loss up to 200°C. Hence,
this loss can be attributed to NO$_3$ ligands. In Figure 4(b), the TGA plot shows exothermic peaks in the temperature range 200°C - 550°C. Furthermore, the loss of NO$_3$ ligands from the surface corresponds to the weight loss observed between 200°C and 550°C.

TGA results indicate that the calcination at 500°C has totally eliminated these nitrates.

The TGA of the calcined samples at various temperatures were presented in Figure 5. This study will allow us to follow the nitrates elimination on the silver based catalysts. First of all, no effect is observed on the decomposition of nitrates when a calcination at 100°C is performed. Then, with a treatment in air at 200°C, we notice a first change. Thus a 200°C treatment leads to a decrease of the ligand-support interaction. Finally, the 400°C treatment reduces the range of temperature clearly related to the nitrate species which goes from [200°C - 550°C] to [350°C - 550°C].

Finally, we analyzed the various samples treated by plasma M.W-1, M.W-2 and M.W-3. In the Figure 6(a), we can follow the effect of the processing time plasma on the decomposition of nitrates of [200°C with 550°C]. We can notice that the more the processing time increases, the less there are nitrate species, which will desorb. The processing time microwave plasma has a positive influence on the decomposition of nitrates.

The effect of power used in plasma reactor was also studied (Figure 6(b)). We can notice that when the power increases to 800 Watt, it remains a low concentration of nitrate on the surface of catalyst. Thus, we can conclude that an increase of the microwave plasma power leads to a positive influence on the decomposition of nitrates.

On Figures 6(c) and (d), we follow the influence of oxygen in the mixture of the carrier gas on the decomposition of nitrates, for M.W. plasma with 500 Watt and 800 Watt respectively. We can notice that when the power is higher than 500 W the presence of oxygen in the plasma treatment shifted the temperature range of decomposition of nitrates. The M.W-4 plasma treatment leads to weak the nitrate-support bonds. Nevertheless, using the M.W-6 treatment under 800 Watt under 20% O$_2$/Ar, it remains some nitrate species on the surface of catalyst. As conclusion, we can note that the effect of the microwave plasma power is more important than the nature of the gas in plasma itself on the decomposition of nitrates. These characterizations confirm the results obtained in catalytic activity (Figures 2 and 3). We can also conclude that the M.W. plasma treatments in FBR are not as efficient as a typical calcination for overall nitrate species removal. However, a plasma treatment for 30 min in Ar led to a higher NO$_x$ removal in SCR of NO$_x$ by ethanol reaction.

Thus, we can suppose that these nitrates species that are left on the surface during the plasma treatment process can play an important role in SCR of NO$_x$ by ethanol.
Figure 4. TGA analysis. (i) of 2.5% silver ong-Al₂O₃ spheres: (a) untreated sample, (b) calcination; (ii) of 2.5% silver ong-Al₂O₃ spheres: (a) untreated sample, (b) calcination.

Figure 5. DTG analysis of 2.5% silver ong-Al₂O₃ spheres after various temperature of calcination: (×) 500°C—2 h, (◇) 400°C—2 h, (△) 300°C—2 h, (□) 200°C—2 h, (◇) 100°C—2 h, (●) without treatment. Heating rate: 10°C/min, carrier gas: air, final temperature: 800°C.
reaction. Further in-situ investigations will be performed on those samples in order to draw a SCR mechanism.

3.5. Identification of Active Silver Species

It is known that during the calcination process, AgO or AgO2 phases are then obtained from the decomposition of silver nitrate as it was described by Musi et al. [17]. The specific surface area (BET) is about 170 m²·g⁻¹ for all studied samples. No silver particles (AgO, Ag0) are detected by TEM coupled by EDS even EDS measurements show that silver is well dispersed on the support as Ag/Al ratio is always constant (Figure not shown).

In the UV-Vis spectroscopy was used in order to investigate the silver species on the different used catalysts. On calcined samples (Figure 7(a)), peaks at 212, 260, 290, 330, and 437 nm are observed. The peaks at 212 and 260 nm are generally attributed to the 4d¹⁰ to 4d’s transition of highly dispersed Ag⁺ cations [24-26]. From the literature, the peak found at ca. 290 nm is linked to Ag₉⁺ clusters [25]. However, Martinez-Arias et al. proposed that this peak could be due to electronic transitions of reduced Ag species [27]. This peak could also correspond to the shifted peak of silver aluminate, usually observed at 230 nm [25]. The peak at 330 nm is generally assigned to Ag₈⁺ clusters [24-27]. The last one at 437 nm corresponds to metallic silver particles [24-27]. There are more oxidized silver species with the plasma treatment. Furthermore, there is the same band for the cluster species between the plasma treatment and calcination.

Moreover, there are even more metallic species with the plasma treatment. Thus, metallic silver Ag0 promotes the reaction of NOx reduction at the low temperatures and the reaction of oxidation of the reducing agent [17, 25, 28, 29]. Figure 7(a) indicates that the samples M.W-1, M.W-2 and M.W-3 present the same species than those present on untreated samples. Figure 7(c) shows that the plasma treatment M.W-4 leads to the presence of Ag₈⁺ clusters (at 290 nm). These cationic cluster species are present in lower concentration on M.W-4 plasma treated samples. This observation can explain the low catalytic activity of these latter samples. In Figures 7(b) and (d), we can conclude that the species present on M.W-5 (800 Watt) samples are similar than those present on the calcined sample.

4. Conclusions

Microwave Fluidized bed plasma reactor treated alumina silver based catalysts lead to higher NOx reduction efficiency in ethanol SCR than calcined catalyst. Despite a lower nitrate desorption, as observed by DTG studies, it has been clearly shown that microwave plasma process leads to the formation of different and more reactive silver species. Further in-situ investigations are in course in order to explain these differences. In this study, we used a microwave source generator. Thus microwave plasmas
~ 210nm $\rightarrow$ Ag$^+$
~ 230nm $\rightarrow$ AgAlO$_2$

~ 290nm and ~ 350nm $\rightarrow$ Ag$^{2+}$
~ 390nm $\rightarrow$ Ag$^0$

(a)
present the advantage to possess higher density of energy in comparison to radio frequency plasmas. Furthermore, positive effects could be expected regarding the elimination of ligands and the use lower plasma treatment times. All the plasma treated samples lead to higher conversion of NO\(_x\) than a sample without treatment. The influence of the parameters such as the process time, the power and the nature of the carrier gas have an effect on the decomposition of nitrates and thus on the reduction of NO\(_x\). The increase of the microwave plasma power has a positive influence on the decomposition of nitrates. The plasma treatment with 500 Watt can be associated with a calcination of 400°C. We can thus conclude that a classical calcination can be easily replaced by a plasma treatment such as microwave plasma treatment on alumina silver based catalysts, which are potential catalysts for SCR NO\(_x\) by Ethanol. For 800 Watt treatment, we can conclude that it presents the same effect than calcination at 500°C. In order to draw a mechanism for SCR of NO\(_x\) on activated microwave plasma alumina supported catalysts, further in-situ investigations will be performed. The aim will be to determine the role of the nitrates species that are left on the surface during the plasma treatment process versus a classical calcination process.

Moreover, other supports will be studied to show the benefits of plasma activation process in FBR. Thus, zeolites and mesoporous materials (SBA-15) are actually in use. The first results confirm those observed on alumina-supported catalysts. Finally, Optical Emission Spectroscopy (OES) is in course to have a better understanding of the plasma itself and its interaction with the surface of catalysts [30,31].

REFERENCES


344-353. doi:10.1016/j.jcat.2005.01.034

