Development of high performance, Pd-based, three way catalysts

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Abstract

The use of Pd in three-way catalytic (TWC) converters of automobiles presents serious limitations which are related to sintering, poisoning and excessive selectivity for HC and CO oxidation, making NO\textsubscript{x} reduction difficult. In the present work, an advanced automotive exhaust catalyst design is applied which reduces, to a large extent, these shortcomings of Pd. This catalyst is based on dispersion of Pd on YSZ (ZrO\textsubscript{2} stabilized with 8 mol-% Y\textsubscript{2}O\textsubscript{3}) forming the internal layer of a three-layer washcoat. The external layers contain small quantities of Pt and Rh dispersed on suitable supports, such as \(\gamma\)-Al\textsubscript{2}O\textsubscript{3} and W\textsuperscript{6+}-doped TiO\textsubscript{2}, respectively. This scheme offers a high performance TWC with low light-off temperature for the conversion of all three pollutants (CO, HC and NO\textsubscript{x}), enhanced poisoning tolerance and improved thermal resistance.

Keywords: Automotive catalyst; Carbon monoxide oxidation; Hydrocarbon oxidation; Nitric oxide reduction; Palladium; YSZ

1. Introduction

Stricter automotive emission regulations will be adopted in Europe, USA and Japan, necessitating the application of various strategies to significantly reduce carbon monoxide, hydrocarbon and NO\textsubscript{x} emissions. The only available technology is the catalytic transformation of these pollutants. Thus, catalysts which are active at low temperatures for reduction of cold-start emissions and tolerant to high temperatures, together with a rational use of the rare platinum group metals (PGM) are required. Research efforts are focused on improved converter designs such as warm-up or electrically heated ones and on advanced catalytic technologies.

With respect to the latter, recent investigations have shown that palladium catalysts possess high efficiency for the oxidation of CO and HC and moderate activity for NO\textsubscript{x} reduction [1,2]. With an accurate air to fuel ratio metering system, Pd-only catalysts are efficient within a narrow operating window, significantly narrower than in the case of conventional Pt–Rh three-way catalysts. Furthermore, Pd presents some serious limitations such as rapid sintering under reducing conditions and poisoning by sulfur at low temperatures, while high Pd loadings can be too selective for HC and CO oxidation, making NO\textsubscript{x} reduction difficult. Current research efforts aim towards eliminating these disadvantages of Pd through advanced catalyst design.

In the present work, the influence of the carrier (Al\textsubscript{2}O\textsubscript{3}, SiO\textsubscript{2}, TiO\textsubscript{2}, TiO\textsubscript{2}–W\textsuperscript{6+} and YSZ) on the catalytic performance of supported Pd catalysts under CO and C\textsubscript{2}H\textsubscript{4} oxidation and
NO reduction conditions was investigated. A strong influence of the carrier on the specific activity of Pd (turnover frequency) was observed in all cases. An advanced catalyst design is applied in order to develop a high performance, Pd mainly TWC. This catalyst is based on the dispersion of Pd on YSZ, forming the internal layer of a three-layer monolith washcoat. The external layers contain small quantities of Pt and Rh dispersed on suitable supports, such as γ-Al₂O₃ and W⁶⁺-doped TiO₂, respectively. This scheme gives a high performance TWC with low light-off temperature for the conversion of all three pollutants (CO, HC and NOₓ), enhanced poisoning tolerance and improved thermal resistance.

2. Experimental

2.1. Catalyst preparation and kinetic measurements

Pd catalysts were prepared by impregnation of the supports with a PdCl₂ solution of appropriate strength so as to yield 0.5 wt.-% metal loading. Five supports were used: SiO₂, TiO₂, ZrO₂ (8 mol-% Y₂O₃), γ-Al₂O₃ and W⁶⁺-doped TiO₂. The latter support was prepared by thermal treatment of TiO₂ and WO₃ at 900°C for 5 h, a procedure which was also followed in the case of undoped TiO₂ (rutile form). The catalysts were reduced under H₂ flow at 400°C for 1 h. They were characterized with respect to exposed metallic area and metal dispersion by selective chemisorption of H₂ at room temperature. Kinetic measurements were carried out in a CSTR-type reactor [3]. Analysis of reactants and products was carried out by on line gas chromatography and IR spectroscopy. Three reactions were studied: CO and C₂H₄ oxidation and NO reduction by CO in a wide range of reactants partial pressures and temperatures (150–500°C).

In order to examine the effect of metal dispersion and metal crystallite size on turnover number, a 0.3% Pd/γ-Al₂O₃ catalyst of approximately 100% dispersion was prepared by impregnation of γ-Al₂O₃ with Pd(NO₃)₂ solution at pH 10 (NH₃). Lower dispersions, namely 63%, 38%, 24% and 4%, were obtained by calcination of this catalyst in air at 550, 650, 750 and 800°C, respectively, for 2 h.

2.2. Preparation and testing of monolithic catalysts

The Pd catalysts were prepared following the procedure which was described above. A thick aqueous suspension of the catalyst was then prepared and the catalyst particle size was reduced by wet milling for 2–3 h. A new suspension was subsequently prepared, using the catalyst of fine particles at a concentration of approximately 30%, at a pH of 3 and a temperature of approximately 40°C. A monolith specimen of cylindrical shape (16 mm diameter by 5 mm length) was immersed in this suspension for 5–10 min. When it was removed from the suspension it was blown with forced air, dried at 110°C for 1 h and calcined at 600°C for 2 h. This procedure was repeated until the desirable loading was achieved. The catalyst loading was determined gravimetrically. The monolith was placed in the reactor where the catalyst was reduced in situ, prior to initiation of the reaction.

3. Results and discussion

3.1. Kinetic measurements

Kinetic results of C₂H₄ oxidation over the Pd catalysts employed in the present study are shown in Fig. 1. The influence of oxygen partial pressure, at constant ethylene pressure, on turnover frequency is shown in Fig. 1a, while the influence of ethylene partial pressure at constant oxygen pressure is illustrated in Fig. 1b. In both cases, turnover frequency exhibits a maximum, illustrating the competitive adsorp-
Fig. 1. Activity (turnover frequency) of Pd catalysts supported on different carriers under C2H4 oxidation. Effect of O2 (a) and C2H4 (b) partial pressures.

Fig. 1 (continued).

The influence of the carrier on the catalytic activity of Pd under CO oxidation conditions is illustrated, in the form of an Arrhenius plot, in Fig. 2. The activation energies do not differ significantly, varying within the range of 22 to 26 kcal/mol. Fig. 3 shows the influence of the carrier on turnover frequency in the NO–CO reaction, under variable NO partial pressure. It is apparent that in all cases the specific activity of Pd is highest when it is dispersed on YSZ and it decreases in the order: YSZ > γ-Al2O3 > SiO2 > TiO2 (4% WO3) > TiO2. The high activity of Pd-YSZ catalyst may be related to the high oxygen anion mobility of YSZ which may be followed by oxygen spill-over onto the metallic sites [3]. Oxygen spill-over onto the Pd
crystallites causes an enhancement of the work function and, thus, weakening of the Pd–O bond.

The results presented in Fig. 1, Figs. 2 and 3 could be attributed to dispersion effects, if it is assumed that the three reactions are structure sensitive. The dispersion of Pd on the different carriers is different, since, their surface area varies significantly. Based on hydrogen adsorption, the dispersion of Pd was found to be: for YSZ:37%, γ-Al2O3:61%, SiO2:65%, TiO2 (4% WO3):33% and TiO2:33%.

Turnover frequencies of C2H4 and CO oxidation and NO reduction by CO are shown in Fig. 4 as a function of the average Pd crystallite size of Pd/γ-Al2O3 catalysts. The variation of the dispersion of this catalyst was achieved by sintering of the highly-dispersed formulation at different temperatures for variable period of time. It is apparent that C2H4 oxidation is a facile reaction since its turnover frequency is not at all affected by Pd dispersion, while CO oxidation and NO reduction exhibit a weak dependence on Pd crystallite size. It is important to note that the variation of specific activity with the carrier (Fig. 1, Figs. 2 and 3) can not be explained by structure sensitivity considerations and should be attributed to interactions between the metal crystallites and the carrier.

3.2. Performance of Pd-based, monolithic converter

Pd-YSZ slurries were applied to monolithic structures and then external layers of Rh/TiO2 (W6+) and Pt/γ-Al2O3 were deposited. All carriers had been promoted with 6% CeO2 and 2% La2O3. With this type of geometric configuration, Pd is protected from poisoning and alloy formation with Pt, and Rh is also protected from poisoning and undesirable interactions with Al2O3 at high temperatures. The use of YSZ as a carrier for Pd and TiO2 (W6+) for Rh, has been found [3] to improve their activity, giving a TWC of higher activity, as compared to the conventional one.

Light-off temperatures of a 3-layer catalyst containing Pt, Rh and Pd are shown in Fig. 5 for TW operation, and are compared with a conventional catalyst in which the same quantity of noble metals are uniformly distributed within the washcoat of γ-Al2O3 (both catalysts contain 0.034% Pt, 0.0086% Rh, 0.039% Pd). It is apparent that the 3-layer catalyst exhibits higher activity for all three reactions since the light-off temperature is lower in all cases. The beneficial effect of the addition of the Pd-YSZ component is also illustrated in Fig. 5 by comparison of the
performance of a 3-layer catalyst which contains Pd (0.031% Pt, 0.0085% Rh, 0.073% Pd) with that of a conventional one which does not contain Pd (0.070% Pt, 0.014% Rh). The conventional catalyst contains almost twice the amount of Pt and Rh as compared to the 3-layered one while the latter contains an amount of Pd equal twice that of Pt. It is apparent that the Pd-based catalyst exhibits significantly higher activity, even for NO reduction which is the most difficult reaction on Pd. The enhancement of activity is due to the higher activity of the Pd-YSZ combination and the inherently higher activity of the 3-layer geometric configuration.

4. Conclusions

The catalytic activity of Pd under reactions of automotive exhaust emissions treatment (CO and C_2H_4 oxidation and NO reduction) can be altered significantly using different supports. The use of Y_2O_3-stabilized ZrO_2 carrier instead of the conventional γ-Al_2O_3 improves significantly the activity of Pd. This may be due to oxygen spillover from the support to the catalyst sites. The promotional effect of the support on the activity of Pd is not related to metal dispersion or, equivalently, to metal crystallite size.

A monolithic catalyst for TW operation, based on the dispersion of Pd on YSZ, forming the internal layer of a three-layer washcoat, was developed. The external layers contain small quantities of Pt and Rh dispersed on suitable supports, γ-Al_2O_3 and W^6+-doped TiO_2, respectively. This configuration gives a high performance TWC with lower light-off temperature for the conversion of all three pollutants (CO, HC and NO_x) as compared to conventional TWC. Because Pd is protected from poisoning and alloy formation with Pt, this catalyst may also exhibit enhanced poisoning tolerance and improved thermal resistance.

References