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The effect of sulfur dioxide on the activity of hierarchical Pd-based catalysts in methane combustion



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ABSTRACT

SO₂ poisoning of methane oxidation over alumina-supported, $Pd@Ce_xZr_{1-x}O_2$ nanoparticle catalysts was systematically studied by means of advanced PhotoElectron Spectroscopy (PES) methods. The $Pd@Ce_xZr_{1-x}O_2$ units were synthesized and deposited on two modified-alumina supports, *i.e.* high surface area modified alumina and a model alumina prepared by Atomic Layer Deposition (ALD) of alumina on Indium Tin Oxide (ITO)/quartz slides. The model support was designed to be suitable for PES analysis and was stable to high temperature treatments (850 °C). Characterization of the high-surface-area (HSA) catalysts by X-Ray Diffraction (XRD), N₂ physisorption, CO chemisorption and Transmission Electron Microscopy (TEM) indicated formation of CeO₂–ZrO₂ (CZ) mixed-oxide crystallites that stabilize the Pd active phase against sintering. Correlation of methane-oxidation rates with PES results demonstrated two distinct mechanisms for deactivation by SO₂. Below 450 °C, the presence of SO₂ in the feed led to partial reduction of the active PdO phase and to the formation of sulfates on the Pd. Above 500 °C, poisoning by SO₂ was less severe due to spillover of the sulfates onto the oxide promoter. Pd@ZrO₂ catalysts showed the best resistance to SO₂ poisoning, outperforming analogous Pd@CZ mixed-oxide catalysts, because there was less sulfate formation and the sulfates that did form could be removed during regeneration. © 2016 Elsevier B.V. All rights reserved.

1. Introduction

Regulations for automotive emissions are becoming more stringent in order to mitigate the effects of transportation on air quality, environment and human health. Electric vehicles and hydrogenpowered, fuel-cell cars may solve these problems in the future, but hydrocarbon fuels will continue to be used in the near term. Because of fuel availability and improved efficiency, there will likely be a move towards more Natural Gas Vehicles (NGVs) and diesel-powered cars; but these both have the disadvantage that methane, a greenhouse gas, is present in significant amounts in

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http://dx.doi.org/10.1016/j.apcatb.2016.09.016 0926-3373/© 2016 Elsevier B.V. All rights reserved. their exhausts. This represents a serious challenge for complying with upcoming regulations. Low-temperature methane catalytic combustion is also important for gas turbines fueled with natural gas. Therefore, better methane-oxidation catalysts, with high activities at low temperatures and better stability, are needed.

Palladium-based catalysts are the most active materials for oxidation of methane at low temperatures, especially when they are promoted by the presence of reducible oxides such as CeO₂ [1–5]. However, they suffer from deactivation under realistic conditions, mainly because of sintering of the metal or support and because of the presence of poisons in the gas feed, including water vapor, sulfur oxides and phosphorous compounds [6–13]. SO₂, which is produced by the oxidation of sulfur compounds present in fuels and in lubricating additives [14,15], is a particularly serious poison in catalytic converters. Despite the continuing reduction of sulfur in the fuels (The Ultra-Low Sulfur Diesel (ULSD) regulation established a 15 ppm sulfur content for diesel fuel in 2006 [16]), long-term exposure to low concentrations of SO₂ is still detrimental to the catalytic activity of the exhaust after-treatment catalysts [15,17,18]. Under lean conditions and at temperatures above 200 °C, the presence of SO₂ in the exhaust leads to formation of sulfate species on both the support and the active phase [19,20]. One of the possible pathways for the formation of sulfates involves SO₂ oxidation to SO₃ by O₂ (1) and subsequent adsorption of SO₃ on the metal-oxide surface (2).

$$2SO_2 + O_2 \rightarrow 2SO_3 \tag{1}$$

$$SO_3 + M - O \rightarrow M - SO_4$$
 (2)

Alternatively, SO₂ may be oxidized by oxygen from the support or disproportionate to SO and SO₃ [21]. Formation of sulfates on supports such as alumina can also deactivate the catalyst by acting as a buffer of SO_x that prevents adsorption of SO₂ onto the active phase during exposure to gas-phase SO₂ but prolongs the poisoning effect after SO₂ removal due to a slow decomposition of the accumulated sulfates [14,15]. With a non-sulfating support, such as SiO₂, the active phase will not be protected from poisoning, leading to faster deactivation, but also to faster recovery [14].

SO₂ poisoning on reducible supports, CeO₂ and Ce_xZr_{1-x}O₂, has been widely investigated because of the importance of these oxides in catalytic converters [22–25]. It is known that ceria forms sulfates that are stable to relatively high temperatures (600 °C–700 °C) [23,26]. ZrO₂ forms fewer sulfates when exposed to the same conditions, and these are mostly confined to its surface [23]. However, conflicting observations have been made regarding the resistance of Ce-Zr (CZ) mixed oxides to SO₂ in comparison to CeO₂ and ZrO₂. Luo et al. reported that CeO₂ and CZ catalysts were affected in a similar manner by SO₂ poisoning for both the water-gas-shift (WGS) and CO-oxidation reactions [23], while Deshmukh et al. reported enhanced resistance to poisoning for the mixed oxides [25].

In this work, we investigate the effect of SO₂ poisoning on methane oxidation over hierarchical Pd@Ce_xZr_{1-x}O₂ catalysts supported on modified alumina. The synthesis of these materials relies on self-assembly of functional precursors to control the catalyst nanostructure and thereby maximize the active-phase/metaloxide promoter interfacial area [27]. This feature helps stabilize the PdO active phase at high temperature and prevents sintering during thermal annealing, which in turn leads to enhanced performance. In previous work, the hierarchical catalysts consisted of Pd nanoparticles and nanosized ceria in intimate contact, deposited on HSA Si-Al₂O₃ support (Pd@CeO₂/Si-Al₂O₃) [11,28]. These catalysts were very active for CH₄ oxidation under dry conditions [28], but deactivated in the presence of water vapor due to formation of Ce hydroxides. Replacing CeO₂ with ZrO₂ significantly improved the stability and activity in the presence of 10% H₂O vapor, with complete combustion achieved at 500 °C (rather than 600 °C for CeO₂-based catalysts) [29]. In the present work, we modified the previously reported synthesis conditions [30] in order to achieve a range of shell compositions, from pure CeO₂ to pure ZrO₂. We have also investigated the effect of SO₂ on these coreshell catalysts using X-ray photoemission spectroscopy (XPS) on model catalysts designed to closely resemble the high-surface-area materials in both surface composition and temperature treatments, while avoiding charging and low resolution problems often experienced with powdered materials [31].

2. Experimental

2.1. Catalyst synthesis

Potassium tetrachloropalladate(II) (99.95%) was purchased from ChemPUR. Sodium borohydride (98%+) was purchased from Acros Organics. Zirconium butoxide (80% in 1-butanol), cerium ammonium nitrate (CAN, 99.99%), sodium methoxide (25% in methanol), phosphoric acid (85%), 11-mercaptoundecanoic acid (MUA, 95%), 1-dodecanethiol (DT, \geq 98%) dodecanoic acid (99%), triethoxy(octyl)silane (TEOOS, 97.5%), and all the solvents (analytical grade) were purchased from Sigma–Aldrich. Trimethyl Aluminum (TMA) was supplied by Cambridge Nanotech together with the ALD system. The Al₂O₃ support was kindly provided by SASOL (PURALOX TH 100/150). Prior to use, Al₂O₃ was stabilized by calcination at 950°C for 24 h. ITO/quartz slides were purchased from Präzisions Glas & Optik (PGS).

The hierarchical Pd@MO_x (CeO₂, ZrO₂ and Ce_xZr_{1-x}O₂) units were prepared by modifying published procedures [30]. Briefly, Pd nanoparticles protected by 11-mercaptoundecanoic acid (MUA) dispersed in THF [30], were added to a THF solution of cerium(IV) tetrakis(decyloxide) and/or zirconium butoxide, followed by the addition of a THF solution of dodecanoic acid. Typically, 10 mL of the THF solution of MUA-Pd nanoparticles (0.5 mg/mL as Pd) were slowly added to 5 mL THF solution of pre-mixed metal alkoxides, keeping the Pd:MO_x molar ratio (1:5.5) constant, followed by the addition of dodecanoic acid (1 mol vs Ce+Zr) dissolved in 10 mL of THF. Hydrolysis of the metal alkoxide in the Pd-Ce/Zr solution was carried out by slowly adding up to 1.2 mL of H₂O dissolved in 10 mL of THF over a period of 4 h (up to 120 mol vs Ce+Zr).

For preparation of HSA catalysts, the Pd@MO_x units were adsorbed in monolayer form onto modified, hydrophobic γ -Al₂O₃, prepared by reaction of calcined γ -Al₂O₃ (powdered material) with TEOOS, as discussed elsewhere [28]. The Pd(1%)@MO_x/Si-Al₂O₃ catalysts were then calcined at 850 °C in air for 5 h in order to remove the organic components and to activate the catalyst [28]. For comparison, a Pd(1%)/Si-Al₂O₃ catalyst was prepared by deposition of DT-protected Pd nanoparticles (Pd-DT NPs) on modified hydrophobic alumina and calcination at 850 °C for 5 h. The Pd-DT NPs were prepared according to a published procedure [32].

The supports for the model catalysts were prepared by ALD of Al₂O₃ overlayers of various thicknesses (2, 5 and 10 nm) on an ITO/quartz support (flat, low-surface-area material). Both TMA and water precursors were kept at room temperature, resulting in vapor pressures of about 20 Torr and 11 Torr, respectively. The deposition chamber was maintained at 250 °C and a base pressure of 0.1 Torr, with N_2 flow of 20 mL min⁻¹. The deposition steps were as follows: 1) pulse water for 0.015 s, 2) hold 5 s, 3) pulse TMA for 0.015 s, and 4) hold 5 s. The thickness of the Al₂O₃ film increased by approximately 1 Å/cycle, so that 50 cycles were used to make a 5 nm film. The Al₂O₃/ITO/quartz slides were then cut into 9×9 mm pieces with a glass cutter and treated in freshly prepared piranha solution (concentrated H₂SO₄ and 30% H₂O₂ solution in a 3:1 ratio) in order to clean the surface from any impurity and increase the hydroxyl group population. The slides were washed many times with water and acetone prior to being functionalized by reaction in a diluted solution of TEOOS (0.5 mL) in toluene (20 mL) for 2 days at room temperature. The Pd@MO_x units were then deposited from solution onto the Si-Al₂O₃/ITO/quartz slides, according to a procedure described in detail elsewhere [33]. Briefly, 0.1 mL of Pd@MO_x solution was deposited onto the support and rinsed with THF after 2 min to remove weakly adsorbed particles. The model catalysts were finally calcined at 850 °C in air for 5 h.

2.2. Catalytic measurements

Catalytic experiments on the HSA catalysts were performed in a U-shaped quartz reactor with 25 mg of catalyst sieved for grain sizes below 150 µm. Aging treatments on model catalysts were performed in a larger U-shaped quartz reactor, using an alumina tray to hold the model-catalyst slides in place. Reactions were carried out at a total gas pressure of 1 atm, with the inlet composition controlled by varying the flow rates of CH_4 , SO_2 , O_2 and Ar. The total flow rate was 83.3 mL min⁻¹ for the HSA catalysts (GHSV \approx 2 \times 10 5 mL g $^{-1}$ h^{-1}) and 21.3 mL min $^{-1}$ for the model catalysts setup (GHSV $\approx 1 \times 10^6 \text{ mLg}^{-1} \text{ h}^{-1}$, using the volume of Pd@CeO₂ added to the Al₂O₃/ITO/quartz slides). Typically, 50 ppm SO₂ was used during aging, in order to accelerate deactivation of the catalyst (SO₂ concentrations in the exhaust feed are more typically less than 5 ppm). The heating and cooling rates in all cases were fixed at 10°C min⁻¹. The composition of the effluent gases was monitored on line using a mass spectrometer. For XPS/SRPES (Synchrotron Radiation Photoelectron Spectroscopy) analysis, the catalysts were rapidly cooled to room temperature under dry conditions at the end of the aging treatment and finally stored under Ar until rapid transfer to the XPS chamber.

2.3. Characterization techniques

The HSA catalysts were characterized by powder X-Ray Diffraction (XRD) for structural analysis, N_2 physisorption for surface area measurements, CO chemisorption for metal dispersion evaluation, Transmission Electron Microscopy (TEM)/Energy Dispersive X-Ray Spectroscopy (EDS) for morphological analysis, and X-ray Photoelectron Spectroscopy (XPS) for comparison with the model catalyst.

XRD patterns were collected on a Philips X'Pert diffractometer using a monochromatized Cu K α (λ = 0.154 nm) X-ray source in the range 10° < 2 θ < 100°. Mean crystallite sizes were calculated by applying the Scherrer equation to the principal reflection of each phase. N₂ physisorption experiments were carried out on a Micromeritics ASAP 2020C. The samples were first degassed in vacuum at 350 °C for 12 h prior to N₂ adsorption at liquid nitrogen temperature.

Pd accessibility was estimated by means of low temperature CO chemisorption measurements. Typically, 150 mg of sample were pre-reduced at 80 °C under 5.0% H₂/Ar for 30 min, degassed for 30 min at 80 °C and for 4 h at 350 °C. The samples were then cooled to -80 °C using an acetone-dry ice bath and exposed to between 2 and 20 Torr of CO. Equilibrium was considered to have been reached when the pressure change was less than 0.01% for 11 consecutive readings, taken at 30-s intervals. Adsorbed volumes were determined by extrapolation of the linear part of the adsorption isotherm to zero pressure. A 1:1 CO/Pd chemisorption stoichiometry ratio was assumed.

The samples were characterized by TEM, performed on a JEOL 2200FS microscope, equipped with a High-Angle Annular-Dark-Field (HAADF) detector, in-column energy filter and EDS.

Model catalysts were characterized by XPS and SRPES for surface chemical analysis, Ion Scattering Spectroscopy (ISS) for investigating of the chemical composition of the outmost surface layer, Atomic Force Microscopy (AFM) and Scanning Electron Microscopy (SEM) for morphological analysis, and EDS for bulk chemical analysis. XPS and SRPES experiments were carried out at the Materials Science Beamline (MSB) at the Elettra Synchrotron Light Source in Trieste, Italy. The MSB is a bending magnet beamline with a plane grating monochromator that provides light in the energy range of 21–1000 eV. The UHV end station, with a base pressure of 1×10^{-8} Pa, is equipped with an electron energy analyzer (Specs Phoibos 150) and a dual Mg/Al X-ray source. High-resolution SRPES measurements were performed on the following core levels: Pd 3d (610 eV), Zr 3d (425 eV), O 1s (610 eV), C 1s (425 eV), S 2p (425 eV), Al 2p (124.8 eV) and Si 2p (150 eV). Complementary XPS spectra of Pd 3d, Ce 3d, Zr 3d, O 1s, C 1s, S 2p, Al 2p, Si 2p, Sn 3d and In 3d core levels were recorded at the same end station, using the excitation energy of 1486.6 eV (Al K α). The apparent degree of reduction of ceria was determined from the Ce 3d spectra, using the procedure described in a previous publication [34]. Binding energies are reported after correction for charging, using adventitious C 1s as a reference.

He⁺ ions with the energy of 2 keV and the impact angle of 45° were used for the ISS analysis. AFM was performed using a Nanoscope V (Digital Instruments Metrology Group, model MMAFMLN) in tapping mode in air at room temperature, using an n-type silicon μ mash[®] SPM probe (HQ:NSC15/AL BS) with tip height of 12–18 μ m and cone angle <40° (Resonant frequency 325 kHz, force constant of ~40 N m⁻¹). SEM measurements were performed on a field-emission Scanning Electron Microscope (SEM) (Model Tescan Mira 3). High resolution was achieved using an acceleration voltage of 30 kV and a working distance of less than 3 mm. A Bruker XFlash[®] 6 | 10 Energy Dispersive X-ray Spectrometer was used to carry out elemental mapping of the sample. Characteristic X-ray radiation was excited by 20 keV primary electrons.

3. Results

3.1. High-surface-area (HSA) catalysts

The synthesis of Pd@CeO₂ and Pd@ZrO₂ particles reported in previous works [28,30] was modified in this study to allow the preparation of Pd@Ce_xZr_{1-x}O₂ particles. Ce and Zr alkoxides were mixed together before slowly adding the dispersed Pd-MUA nanoparticles to allow the reaction of both alkoxides with the carboxylic group of MUA. The Pd:(Ce+Zr) molar ratio was kept constant and the following Ce:Zr molar ratios were selected to investigate the generality of the synthetic method: (80:20), (60:40), (40:60), (20:80). Pure Pd@CeO₂ and Pd@ZrO₂ units were also prepared as references. After controlled hydrolysis, the particles were deposited on hydrophobic, silanized alumina and the materials were calcined at 850 °C for 5 h. The final Pd@MO_x/Si-Al₂O₃ catalysts were characterized by XRD to check for the formation of mixed or segregated CZ oxide phases (Figs. 1, S1 for total XRD pattern).

A complete Rietveld analysis of the CZ XRD pattern was not possible due to overlap with the signal from the Al₂O₃ support, which was more than 90 wt% of the studied materials and consisted of a mixture of metastable γ - and θ -Al₂O₃ after thermal pre-treatment [35]. Nonetheless, the partial XRD spectrum reported in Fig. 1 clearly showed the first and most intense reflection of the CZ mixed oxides, which varied between 28.5° (for CeO₂) to 30.2° (for ZrO₂). Therefore, only pseudo-cubic cell parameters were calculated in the present case, despite the fact that a transition from cubic to tetragonal crystal structure is expected in the oxide promoter with high ZrO₂ content [36]. The linear dependence of the cell parameter for the pseudo-cubic cells with the CZ composition is consistent with the formation of solid solutions between CeO_2 and ZrO_2 (inset of Fig. 1) [37]. Only in the case of CZ 40:60 a shoulder appeared at lower diffraction angles, suggesting that segregation of a small amount of a CeO₂-rich phase occurred for that sample. Phase separation has been observed previously for CZ mixed oxides with 40:60 Ce:Zr ratios, because this composition is thermodynamically unstable [36]. The decreased intensity of the main reflection with increasing ZrO₂ content is due to the lower weight loading and scattering factor of Zr.

The mean crystallite sizes of the CZ mixed oxides were calculated to be in the range of 7–11 nm (Table S1 in the Supplementary



Fig. 1. XRD patterns of the Si-Al₂O₃ calcined at 850 °C for 5 h (grey line) and Pd@MO_x/Si-Al₂O₃ samples calcined at 850 °C for 5 h (orange: CeO₂; green: CZ 80:20; red: CZ 60:40; magenta: CZ 40:60; purple: CZ 20:80; blue: ZrO₂). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

information), which is in good agreement with TEM results (5–10 nm apparent particles size). Notably, the apparent size of CZ mixed-oxide crystallites is smaller than that of CeO_2 , as expected in view of the stabilizing effect of Zr [37]. The most intense reflection for PdO, the (101) plane, can be observed at 34°. The fitting of this line is complicated by the overlapping of other reflections, so that the mean crystallite size calculated by the Scherrer equation (15–18 nm, see Table S1 in the Supplementary information) is not very indicative of the actual size distribution of the PdO active phase in the samples, as discussed further in the section on TEM analysis.

Physisorption and chemisorption experiments on Pd@MO_x/Si-Al₂O₃ catalysts revealed that all the samples had similar surface areas, pore-size distributions, and Pd accessibility, as detailed in Table S2 in the Supplementary information. The results are comparable with those reported in previous studies on Pd@CeO_x/Si-Al₂O₃ and Pd@ZrO_x/Si-Al₂O₃ [11,29]. On the other hand, the Pd dispersion for the Pd/Si-Al₂O₃ was significantly lower due to severe sintering of Pd nanoparticles. The pore-size distributions were also consistent with previous results: Pd/Si-Al₂O₃ and the Si-Al₂O₃ support have similar pore structures, with pore sizes in the range from 10 to 50 nm in diameter, while the Pd@MO_x/Si-Al₂O₃ catalysts exhibited fewer large pores due to partial filling with the nanostructured units, along with the formation of small pores (around 10 nm in diameter) associated with the units themselves (Fig. S2 in the Supplementary information).

Representative TEM-EDS results Fig. 2, demonstrated that all Pd@MOx-based catalysts have similar morphologies: the CZ particles, identified by EDS spectra and lattice fringes, are 5-10 nm in diameter and are sometimes aggregated. For Pd@CZ 60:40, the signals of Ce and Zr observed in EDS mapping mode are always associated and the signal analysis in spot mode revealed a good agreement with the desired Ce:Zr stoichiometric ratio. The Pd signal in EDS is typically low and diffuse, though some large palladium particles were observed, similar to what was reported by Zhang et al. for similar catalysts [38]. Such a bimodal particle distribution is not detected by XRD, since the very small PdO nanoparticles will give a broad reflection that is hard to distinguish in the presence of overlapping XRD patterns from the Al₂O₃ support. The Pd/Si-Al₂O₃ catalyst showed only very large Pd particles (up to 100 nm in diameter) due to severe sintering after calcination at 850 °C for 5 h. In all cases, no apparent changes in morphology were observed after any SO₂ aging treatments, discussed in the following section (Fig. S3 in the Supplementary information).

Methane-oxidation, light-off experiments performed on each of the Pd@MO_x-based catalysts showed comparable results that were similar to those previously reported for the single-oxide catalysts (Fig. S4 in the Supplementary information) [28,30]. By comparison, the Pd/Si-Al₂O₃ catalyst showed a higher light off temperature and the usual conversion hysteresis after heating to high temperature (Fig. S4 in the Supplementary information). While light-off curves were comparable for all Pd@MO_x catalysts, sulfur tolerance was expected to vary more strongly with composition. In order to study SO₂-poisoning resistance, aging tests were performed in which the catalysts were exposed to specified concentrations of SO₂ (typically 50 ppm) at different temperatures (from 300 to $600 \,^{\circ}$ C). Introducing 50 ppm of SO₂ into the feed under dry conditions (0.5% CH₄; 2% O₂; Ar balance) caused complete and irreversible deactivation of all the studied catalysts between 300 °C and 400 °C (Fig. 3). At 450 °C, each of the hierarchical, core-shell catalysts was partially regenerated under dry conditions, recovering 50-60% of their initial conversion. The final conversion trend $Pd@ZrO_2/Al_2O_3 > Pd@Ce_{0.6}Zr_{0.4}O_2/Al_2O_3 > Pd@CeO_2/Al_2O_3.$ was Although less active, the Pd/Si-Al₂O₃ catalyst was almost completely regenerated under dry conditions. This observation suggests that the promoting effect of the metal oxides in the hierarchical catalysts is largely suppressed by SO₂ at 450 °C, while the active phase can be largely regenerated.

To look for longer-term effects, the methane conversions under dry conditions at 500 °C were monitored over each of the catalysts for 2 h during SO₂ exposure (Fig. 4A). The hierarchical catalysts deactivated sharply during the first hour of treatment, reaching a plateau at around 60% of the initial conversion. The Pd@ZrO₂ and Pd@CZ catalysts showed similar deactivation trends and were partially reactivated over time after reaching that plateau. The Pd@CeO₂ underwent a slightly slower deactivation but was not reactivated after reaching the plateau. Aging at 500 °C using different SO₂ concentrations resulted in different deactivation rates but did not significantly change the plateau conversion (Fig. S5 in the Supplementary information). The residual conversion is likely associated with the partially poisoned Pd phase, unpromoted by the metal oxide. Indeed, the Pd/Si-Al₂O₃ sample was very stable under SO₂ aging at 500 °C, indicating that the SO₂ effect on Pd is largely inhibited at this temperature. Moreover, MO_x/Si-Al₂O₃ catalysts without Pd in their formulation, prepared following a similar procedure as that used to synthesize the hierarchical catalysts, did not exhibit any methane conversion at 500 °C and conversions on these materials did not light-off until above 550 °C (Fig. S6 in the Supplementary information).

In parallel experiments, consecutive 50 ppm SO₂-aging treatments for periods of 30 min, 60 min or 120 min at 500 °C caused partial but irreversible deactivation of the Pd@CeO₂ and Pd@CZ samples (Fig. 4B). Conversely, the poisoning effect of SO₂ on Pd@ZrO₂ and Pd/Si-Al₂O₃ catalysts was reversible at this temperature (Fig. 4B), even after 12 h aging (data not shown). Heating the Pd@ZrO₂ catalysts that had been aged at temperatures below 500 °C in dry conditions also resulted in complete reactivation of the materials. Finally, it should be noted that the conversion trends reported in Figs. 3 and 4B were affected by chromatographic effects due to the time required for the gas to flow through the catalytic system and to reach the analyzer. Therefore, the conversion at the beginning of reactivation can only be extrapolated.

At 600 °C, under dry conditions, methane was completely converted to CO₂ over all the catalysts and there was no loss of conversion during SO₂ exposure. To increase the sensitivity of the experiment, the catalysts were diluted by addition of Al₂O₃ in a ratio of 1:3 and aged using the same conditions as those employed for the pure catalysts (Fig. 5). The diluted Pd@MO_x catalysts all achieved complete conversion at 600 °C prior to SO₂ exposure, but the diluted Pd/Si-Al₂O₃ catalyst showed only 80%



Fig. 2. Representative High-Angle Annular Dark-Field (HAADF) images (left column) and EDS mapping (right column) of samples $Pd@CeO_2/Si-Al_2O_3$ ($Pd@CeO_2$), $Pd@CeO_2/Si-Al_2O_3$ ($Pd@ZrO_2$), $Pd@ZrO_2/Si-Al_2O_3$ ($Pd@ZrO_2$) and $Pd/Si-Al_2O_3$. EDS mapping colors: blue = Al, red = Pd, yellow = Ce, green = Zr. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

conversion under these conditions. Exposure of the $Pd@CeO_2$ and Pd@CZ samples to SO_2 at this temperature caused the conversion to decrease continuously with time; however, for the $Pd@ZrO_2$ and

Pd/Si-Al₂O₃ samples, after displaying an initially sharp decline in conversion, conversions partially recovered and reached a plateau value. This behavior was reproducible and was never observed for



Fig. 3. Methane Catalytic oxidation: effect of 50 ppm SO₂ dry aging for 30 min at different temperatures on the catalytic activity of Pd@CeO₂ (orange line), Pd@ZrO₂ (blue line), Pd@Cz (red line) and Pd/Si-Al₂O₃ (black line). Conditions: 0.5% CH₄; 2% O₂; 50 ppm SO₂ (if present) Ar balance, GHSV = 200000 mL g⁻¹ h⁻¹. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 4. Methane Catalytic oxidation: (A) SO₂ dry aging at 500 °C. Pd@CeO₂ (orange line), Pd@ZrO₂ (blue line), Pd@CZ (red line) and Pd/Si-Al₂O₃ (black line); (B) Reactivation trends following consecutive SO₂ dry aging treatments at 500 °C. Conditions: 0.5% CH₄; 2% O₂; 50 ppm SO₂ (if present) Ar balance, GHSV = 200000 mL g⁻¹ h⁻¹. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 5. Methane Catalytic Oxidation: SO₂ dry aging at 600 °C and reactivation trends. Pd@CeO₂ (orange line), Pd@ZrO₂ (blue line), Pd@CZ (red line) and Pd/Si-Al₂O₃ (black line). Conditions: 0.5% CH₄; 2% O₂; 50 ppm SO₂ (if present) Ar balance, GHSV = 200000 mLg⁻¹ h⁻¹. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

CeO₂-containing catalysts. A possible explanation for the transient behavior on Pd@ZrO₂ and Pd/Si-Al₂O₃ is that SO₂ initially deactivates the PdO active phase, competing with CH₄ for oxidation. Later, SO₃ and sulfate species formed by oxidation migrate to the promoter/support. It has been previously reported that the activity of Pd/ZrO₂ catalysts can increase after SO₂ aging due to formation of a composite site between PdO and sulfates at the PdO-support interface [39]. It is noteworthy that there was also an increase in conversion with SO₂ exposure time at 500 °C for Pd@ZrO₂ and Pd@CZ catalysts (Fig. 4A), although the time scale for increasing conversion was much larger and the initial dip was not as sharp.

3.2. Model catalysts

The structure of the model catalysts used in this study is depicted graphically in Fig. 6A. A 1-mm thick quartz slide was used to support a 500-nm thick conductive layer of ITO, over which thin Al₂O₃ layers were grown by ALD. The thickness of the Al₂O₃ film was 5 nm, a value chosen so as to avoid both charging during XPS analysis and unwanted exposure of ITO after calcination (see SI for more information). To obtain a silanized, hydrophobic Si-Al₂O₃ comparable to that of the HSA Si-Al₂O₃, the slides were allowed to react with TEOOS by soaking them in TEOOS solutions, diluted with toluene, for 2 days. Finally, the slides were loaded with Pd@CeO₂, Pd@ZrO₂ or Pd@CeO₆Zr_{0.4}O₂ (Pd@CZ) particles and cal-



Fig. 6. Graphical representation of Pd@MO_x/Si-Al₂O₃/ITO/quartz model (A, not to scale); AFM 3D image and mapping of ITO/quartz (B1), ALD-Al₂O₃/ITO/quartz (B2), Pd@CZ-m calcined at 850 °C for 5 h (B3). EDS wide spectrum of calcined Pd@CZ-m (C) and SEM image of Pd@CZ-m with 1 μm view field (D1) and 10 μm view field (D2).

cined at 850 $^\circ C$ for 5 h. The catalysts are labeled Pd@CeO_2-m (m for model), Pd@ZrO_2-m and Pd@CZ-m, respectively.

AFM analysis of the model catalysts was performed at different stages in the preparation procedure (Fig. 6B). The bare ITO surface shows characteristic worm-like structures (Fig. 6B1) which become slightly more defined in shape after Al₂O₃ deposition (Fig. 6B2). The final structure obtained after depositing Pd@MO_x, followed by calcination at 850 °C, shows a similar morphology, but with some aggregates on the surface (In Fig. 6B3, see the white spots in the 2D image, blue in the 3D image). Similar to the results reported in a recent paper by Zhang et al. [38], SEM/EDS analysis revealed aggregates composed of Pd with CeO2, ZrO2 or CZ on top of small features that covered the Al₂O₃ layer (Fig. 6D). A representative EDS spectrum of Pd@CZ-m is reported in Fig. 6C. The spectra obtained from different spots on the model catalysts revealed a uniform composition with the expected Pd:(Ce + Zr) and Ce:Zr molar ratio (1:5.5 and 6:4 respectively). CeO₂- and ZrO₂-based catalysts also showed the expected Pd:MOx molar ratio. No differences in surface morphology were observed for CeO₂, ZrO₂ or CZ-containing samples.

Because the SRPES signals for Pd 3d and Zr 3p overlap $(3p_{1/2}: 343 \text{ eV}, 3p_{3/2}: 330 \text{ eV})$, only Pd 3d spectra of Pd@CeO₂-m samples are reported here in Fig. 7. Calcination temperature had a significant effect on the Pd 3d region. Reference Pd@CeO₂-m sam-

ples calcined at 500 °C showed the Pd spin-orbit split doublet (Pd $3d_{5/2}$ and Pd $3d_{3/2}$) at 337.2 eV and 342.2 eV (Fig. 7A1), which can be assigned to PdO, based on previous XPS and X-ray Absorption Near Edge Structure (XANES) analysis of similar materials [40]. A shift of the PdO signal to higher BE (almost 1 eV) with respect to a bulk value is likely due to a size effect [41–47]. Calcination of the Pd@CeO₂-m samples to 850 °C causes partial agglomeration of some of the nanostructures to form Pd and CeO₂ particles in the range of 50–100 nm (Fig. 7B2) [38]; and this agglomeration gives rise to an additional contribution in the Pd 3d spectra, more typical for bulk PdO (366.5 eV) (Fig. 7B2).

Because the relative contribution of the larger agglomerates in photoemission is expected to increase with excitation energy, we examined the surface:bulk signal contribution by tuning the SRPES excitation energy. In agreement with this, the signal at higher BE, tentatively assigned to PdO NPs, was relatively more intense at lower excitation energies, implying that this feature is associated with a considerably smaller structures. An alternative explanation is that the two contributions to the SRPES signal could be due to a Pd-Ce mixed oxide and a bulk PdO. For example, the Pd 3d signal of Pd–Ce mixed oxides has been reported to have BE of 338 eV BE [48]. Since the mixed oxides would likely segregate at higher temperatures, resulting in bulk PdO formation, this could also explain the



Fig. 7. Left: Pd 3d SRPES spectra at different excitation energies for Pd@CeO₂-m calcined at 500 °C (A1) and 850 °C (B1). 500 eV: red line, 640 eV: green line, 880 eV: blue line. Grey lines are guidelines to the eye. Right: representative SEM images of Pd@CeO₂-m samples calcined at 500 °C (A2) and 850 °C (B2); bars: 200 nm. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 8. XPS spectra of Pd 3d region of Pd@CeO₂-m, fresh (black line) and $350 \degree C SO_2$ aged (red line). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

two features in the spectra [49]. However, we regard this explanation as less probable since the synthesis of the core-shell materials starts with pre-formed Pd nanoparticles and complete disruption of the PdO and incorporation into CeO_2 seems unlikely to occur at 500 °C. Also, this latter explanation does not agree with the observation of a PdO (101) reflection in the HSA catalysts calcined at 850 °C (Fig. 1) or with previous EXAFS studies on similar systems [11,40]. Due to the lower resolution, only one contribution averaging the signals associated to PdO NPs and aggregates was observed in XPS (Fig. 8).

In order to study the effect of SO₂ aging at different temperatures, the model catalysts were treated under similar conditions to those used with the HSA catalysts (except for higher GHSV) and then transferred under inert atmosphere to the analysis chamber for analysis by XPS/SRPES. First, the catalysts were aged for 2 h under dry methane-oxidation (0.5% CH₄, 2.0% O₂) conditions with 50 ppm SO₂ at 350 °C, 500 °C or 600 °C. Then, after XPS analysis, the aged catalysts were regenerated under dry methane-oxidation conditions, without SO₂, for 2 h at the same temperature at which they were aged. After SO₂ aging at 350 °C, the XPS spectrum showed a peak at 335.2 eV, which can be assigned to metallic Pd (Fig. 8) [20,50,51]. Partial reduction of PdO could result from SO₂ oxidation to SO₃ by PdO lattice oxygen. The complete re-oxidation of the then formed metallic Pd would not be possible under the conditions of this study at 350 °C, as previous XANES results on HSA Pd@CeO₂/Si-Al₂O₃ suggest, because of the low temperature and oxygen pressure involved [11]. A similar effect has been reported by Venezia et al. after aging Pd-based catalysts on TiO₂ or SiO₂ at the same temperature overnight [20,50] and by Liotta et al. for Pd/CeO₂ catalysts operated under lean conditions and 10 ppm of SO₂ [51].

Along with the reduction of the Pd, there was a slight increase of the XPS signal at higher BE following SO₂ aging at 350 °C. The shift of the signal is less than that expected for Pd in the form of PdSO₄ and we suggest that this might be indicative of some interaction with sulfates groups, in agreement with the literature [6,14]. After 2 h under dry methane-oxidation conditions at 350 °C, the XPS spectra did not change appreciably from those of the SO₂ aged catalyst. An irreversible poisoning of the PdO active phase would explain the similar behavior observed for HSA catalysts, regardless of the presence and composition of the promoter (Fig. 3). At higher temperature (500 °C to 600 °C) SRPES spectra of the Pd 3d region did not change after SO₂ aging or regeneration, suggesting that the interaction of PdO with sulfates is weaker and reversible. Accordingly, the



Fig. 9. S 2p region of SRPES spectra of SO₂ aged and regenerated samples at (A) 500 °C (and (B) 600 °C (orange: Pd@CeO₂-m; red: Pd@CZ-m; blue: Pd@ZrO₂-m; thick line: aged; thin line: regenerated); (C) S 2p region of XPS spectra of SO₂ aged samples at 500 °C (red line) and 600 °C (green line) (top: Pd@CeO₂-m; middle: Pd@CZ-m; bottom: Pd@ZrO₂-m). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Pd/Si-Al₂O₃ catalyst was very stable during dry SO₂ aging at 500 °C (Fig. 4A) and a sharp transient deactivation at 600 °C could only be observed on the diluted catalyst (Fig. 5). These observations are in accordance with thermodynamic calculations on PdSO₄ formation from mixtures of O₂ and SO₃, predicting PdSO₄ decomposition below 400 °C [52].

Independent of the SO₂ aging temperature, the S 2p region of the SRPES/XPS spectra showed only a signal corresponding to sulfate species (168–172 eV BE). The use of SRPES was crucial in order to observe an intense and resolved S 2p signal, thanks to the much higher photoionization cross-section (0.83 Mbarn at 425 eV; 0.022 Mbarn at 1486.6 eV) [53] and shorter information length, which favors signal from surface species. Indeed, after 500 °C aging, sulfates were only observed in SRPES spectra and not in XPS spectra (Fig. 9A,C), which also showed an overlapping Zr 3d signal due to Al K α_3 (1476.8 eV) and K α_4 (1474.8 eV) lines. After 2 h of regeneration at 500 °C, sulfates were partially removed from Pd@ZrO₂-m, while the sulfates remained on the Pd@CeO₂-m and Pd@CZ-m samples, a finding consistent with the regeneration trends observed with the HSA catalysts (Fig. 4).

During regeneration of the catalyst, only a small fraction of sulfur species desorbed, suggesting that a rearrangement of sulfates on the surface could lead to reactivation and that TPD studies may give misleading results for evaluation of SO₂ poisoning resistance. At 600 °C, more sulfates are formed (even if a decrease in conversion was observed only in diluted samples) and these could be observed both in SRPES and XPS spectra (Fig. 9B,C). Based on the cross section corrected photoemission signal of S and Ce (and/or Zr) in XPS, we estimate a MO_x:S molar ratio of 5:1 for CeO₂ and CZ and 7:1 for ZrO₂. These ratios are consistent with about 1 monolayer (ML) of sulfate being formed on the surface of the oxide nanoparticles, considering that the XPS information depth is around 2 nm, which corresponds to about 6 ML of CeO₂ in the close-packed (111) direction. Regeneration under dry conditions at 600 °C resulted in desorption of sulfates from the surface/subsurface of all the samples (Fig. 9B), consistent with similar trends observed in the recovery of conversion (Fig. 5).

Previous FTIR [26], TPD [23] and XPS [54] studies reported sulfate desorption from Ce₁Zr_{1-x}O₂ in the range of 600 °C to 700 °C. The desorption of sulfate species from pure ZrO₂-based materials is much less investigated, with TPD results showing higher desorption temperatures than for CeO₂ sulfates (800 °C) [23]. We suggest that the surface sensitivity of SRPES/XPS analysis allowed us to observe a partial desorption of sulfates from ZrO₂, occurring at lower temperature, which could not be detected by TPD. The difference of

observed sulfate stability could also be due to the atmosphere in which desorption was performed, which can affect decomposition of S-species [17]. Indeed, Colussi et al. observed different TPD profiles on SO₂-aged Pd/CeO₂/Al₂O₃ samples when the desorption was carried out in reaction conditions or inert atmosphere [17], suggesting a role of *in-situ* produced water in the SO₂ desorption mechanism.

Interestingly, the BE for the S 2p features in Fig. 9 was different for the Pd@CeO₂-m and Pd@ZrO₂-m samples, implying that sulfates bonded to Zr or Ce cations are distinguishable. Since the spectra of Pd@CZ-m are very similar to that of Pd@CeO₂ both in shape and formation/desorption trends, it appears that primarily Ce sulfates are formed on Pd@CZ-m samples and that these are similar to sulfates formed on CeO₂. In agreement with this, the Ce 3d and Zr 3d XPS spectral regions show modifications after SO₂ aging that depend on the MO_x composition (Fig. 10). In particular, Ce 3d spectra of Pd@CeO₂-m (Fig. 10A) and Pd@CZ-m (Fig. 10C1) aged at 600 °C are similar and exhibit more intense Ce(III) contributions than what is observed on the fresh catalyst, probably due to reduction of CeO₂ by SO₂ with consequent formation of cerium sulfate. By contrast, Zr 3d spectra for Pd@ZrO₂-m (Fig. 10B) are different from spectra for Pd@CZ-m (Fig. 10C2), revealing sulfate formation on Pd@ZrO₂-m after aging at 500 °C and 600 °C, while the Pd@CZ-m spectra do not change. This is consistent with the formation of sulfates only on Ce cations in CZ materials, which leaves the chemical environment of Zr cations unaffected.

Fig. 11 shows the O 1 s spectra of the fresh samples, exhibiting peaks typical of bulk oxygen (529 eV for CeO₂ and 531 eV for ZrO₂) and hydroxyl species (532 eV)[55]. SO₂ aging leads to an increase in the intensity of the higher BE contribution (532 eV) associated with OH groups (Fig. 11). These results, taken together with the trends observed in S 2p spectra, suggest that the variation of O 1 s signals is mostly due to oxygen from sulfate species.

4. Discussion

First, the observations here demonstrate that the previously reported synthesis of $Pd@CeO_2$ and $Pd@ZrO_2$ particles by selfassembly can be extended to form materials with mixed-oxide shells, $Pd@Ce_xZr_{1-x}O_2$. The pre-formed core-shell particles can be deposited on both high-surface-area (HSA), functionalized alumina or on model Si-Al₂O₃ surfaces to achieve a similar, controlled morphology. In all of the materials, the oxide shell was found to promote catalytic activity for methane oxidation. The series of hierarchically structured catalysts supported on Si-Al₂O₃ also had



Fig. 10. XPS spectra of pristine samples (black line) and SO₂ aged samples at 500 °C (red line) and 600 °C (green line). (A) Ce 3d region of Pd@CeO₂-m; (B) Zr 3d region of Pd@ZrO₂-m; (C1) Ce 3d region and (C2) Zr 3d region of Pd@CZ-m. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 11. O 1 s region of SRPES spectra of SO₂ aged and regenerated samples at (A) 500 °C (and (B) 600 °C (orange: Pd@CeO₂-m; red: Pd@CZ-m; blue: Pd@ZTO₂-m; thick line: aged; thin line: regenerated); (C) O 1 s region of XPS spectra of SO₂ aged samples at 500 °C (red line) and 600 °C (green line) (top: Pd@CeO₂-m; middle: Pd@CZ-m; bottom: Pd@ZTO₂-m). SRPES and XPS spectra of fresh samples are reported for reference (black lines). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

comparable surface areas and active-phase accessibilities. These properties make them good candidates to study the effect of the oxide promoter composition on SO_2 poisoning.

The methods used in preparing the model catalysts in this study are also of general interest for investigating the effects of temperature and operating conditions. The approach is based on the ability of ALD to form uniform, thin films having precise thicknesses and a wide range of possible final compositions. Indeed, model supports could be prepared from almost any metal oxide by varying the ALD precursors [56]. This allows support composition to be varied systematically in order to study the effect of metal-support interactions on the catalyst performance, morphology (by SEM) and chemical state (by XPS/XANES). Also, model catalysts prepared in this way can be tested and analyzed repeatedly, making it possible to study the effect of aging and regeneration treatments together with properties for adsorption, desorption or redistribution of surface species. The lower surface area of model catalysts, that affects the range of achievable GHSV, is the only drawback of this powerful method of investigation.

SO₂ poisoning of Pd-based catalysts in lean catalytic combustion of methane proceeds via two distinct mechanisms, depending on the aging temperature. At lower temperature (<450 °C), SO₂ irreversibly poisons PdO, causing a partial reduction to Pd and the formation of sulfates in close proximity to the active phase. The temperature is not high enough to allow the spillover of sulfate species from the active phase to the support/promoter and to promote a complete re-oxidation of the poisoned catalyst [11,57]. At higher temperatures, PdO is more resistant to poisoning, thanks to the spillover of sulfates from the active phase and/or the direct reaction of SO₂ with the promoter. However, since the promoting effect of the metal oxide is turned off by sulfate poisoning, the catalysts deactivate until a plateau is reached, corresponding to the active phase residual activity.

The effect of the oxide promoter composition on SO₂ poisoning is also temperature dependent. At 500 °C, Pd@ZrO₂ catalysts showed the best sulfur resistance among the samples, being only reversibly deactivated thanks to partial sulfate desorption. The performance of Pd@CZ catalysts is intermediate between that of the two pure oxides, suggesting that no cooperative effect of Ce and Zr takes place. On the other hand, at 600 °C, Pd@CeO₂ and Pd@CZ catalyst against transient deactivation, acting as a sink for SO₂, while Pd@ZrO₂ deactivates in a similar way as Pd/Si-Al₂O₃. However, the poisoning effect of SO₂ is less relevant than at 500 °C, since the deactivation is observed only for very high GHSV. These results make Pd@ZrO₂ catalysts a more suitable candidate for real application, taking into account the better stability in the presence of H₂O with respect to Pd@CeO₂ catalysts.

The chemical-state and surface sensitivity of photoelectron spectroscopy also allowed for the observation of preferential sulfation of Ce over Zr in CZ mixed oxides. The chemical environment of Zr cations only changed after sulfation of ZrO₂, while in the case of CZ no evident differences were observed. These results strongly suggest that sulfate species are associated with individual metal cations, producing a first evidence of what was tentatively proposed by Luo et al. based on TPD and pulse-reactor experiments [23].

5. Conclusions

The self-assembly methodology described previously [32] was modified in order to synthesize nanostructured Pd@Ce_xZr_{1-x}O₂ $(Pd@MO_x)$ units in the whole compositional range (0 < x < 1). The synthesis of dispersed Pd@MO_x allowed the preparation of a series of high-surface-area Si-Al₂O₃ supported catalysts and model catalysts having similar nanostructure and surface chemistry. Comparison of results on the two types of catalysts allowed the SO₂ poisoning of methane oxidation on Pd-based catalysts to be systematically studied to elucidate the role of the MO_x promoter and the aging conditions. At lower temperatures (<450 °C), the PdO active phase is irreversibly poisoned by SO₂ due to interaction with sulfates which are not able to spillover to the support/promoter. At higher temperatures (>500 °C), poisoning is slowed by formation of sulfate species on the oxide promoter. Due to partial decomposition of sulfates at 500 °C, Pd@ZrO2-based catalysts showed the best sulfur-poisoning resistance, attaining complete regeneration even after prolonged aging, and thus they are the best candidates for real application. Pd@Ce_{0.6}Zr_{0.4}O₂ catalysts showed intermediate sulfur tolerance compared to Pd@CeO₂ and Pd@ZrO₂, in agreement with previously reported results [23]. The high chemical sensitivity of PES techniques provided direct evidence for previously suggested formation of sulfate species on individual metal cations in $Ce_x Zr_{1-x}O_2$ mixed oxides [23]. Finally, the modelcatalyst approaches developed here should allow the study of metal-support interactions in other catalytically relevant systems by simply varying the ALD-deposited thin film composition.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.apcatb.2016. 09.016.

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