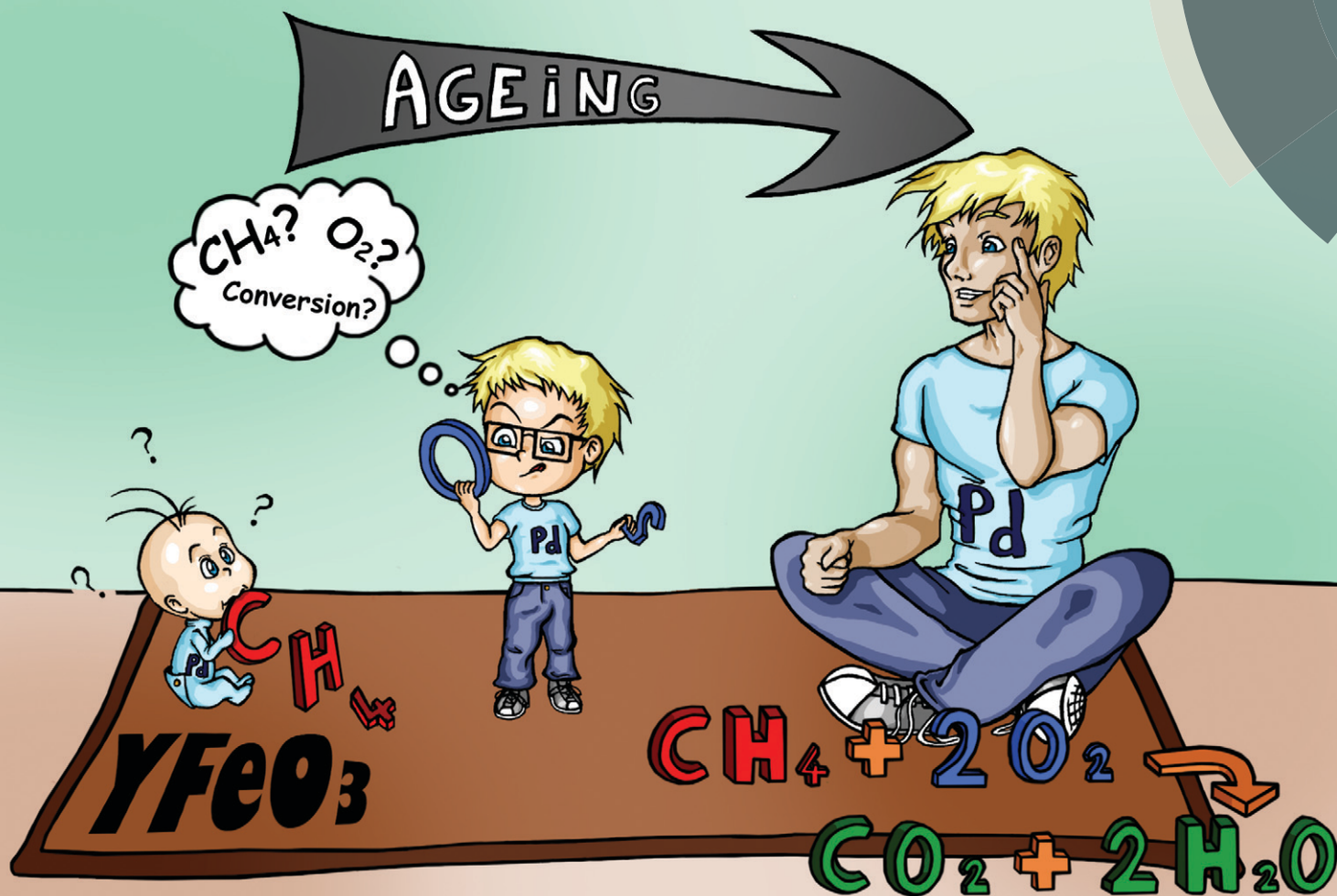


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Ageing induced improvement of methane oxidation activity of Pd/YFeO₃

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The ageing characteristics of flame-made 2 wt% Pd supported on YFeO₃ were analysed in comparison with a Pd/Al₂O₃-CeO₂-ZrO₂ three-way catalyst (TWC) with respect to structural changes and catalytic performance for methane oxidation under stoichiometric reaction conditions. Thermal treatment under lean conditions (air, 900 °C) resulted in slight decrease in the methane oxidation activity of the TWC. In marked contrast, YFeO₃-supported Pd catalysts exhibit an increase in activity after such treatment. Activity enhancement is even higher when the treatment was performed under stoichiometric conditions (air–fuel equivalence ratio, $\lambda = 1$, 900 °C). To explain this observation, in-depth characterization (BET, STEM, OSCC, XAS, and CO chemisorption) of fresh and aged catalysts was performed. Both thermal and stoichiometric ageing cause a severe sintering of the support particles and the phase transformation from hexagonal to orthorhombic YFeO₃. Despite the absence of a mixed Pd–YFeO₃ phase, the growth of Pd particles appears to be limited under the $\lambda = 1$ atmosphere. In contrast to thermally aged catalysts where large PdO particles are formed, well-defined metallic Pd nanoparticles of 10–20 nm are present after stoichiometric ageing along with higher methane oxidation activity. Although it is tempting to conclude that metallic Pd is active for methane oxidation under the given conditions, reversible and periodic partial oxidation of the large metallic particles is observed in modulation excitation high energy X-ray diffraction (HXRD) experiments designed to simulate the oscillating redox conditions experienced during operation. These results indicate that large Pd particles exhibit improved methane oxidation activity but equally confirm that activity under stoichiometric conditions is the result of a delicate equilibrium dictated by the bulk-Pd/surface-PdO pair.

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Introduction

The exhaust of modern stoichiometric engines fuelled with natural gas is controlled using Pd-based three-way catalysts (TWC).¹ Pd is considered the precious metal of choice for oxidation of methane,^{2,3} the major component of natural gas. During operation, the atmosphere around the TWC is periodically varied between slightly reducing and slightly oxidizing that allows maintenance of the emission levels within those

demanded by the existing stringent regulations.⁴ Although the TWC for natural gas vehicles is typically derived from the gasoline TWC, their chemistries can display significant differences.^{5,6} However, similarly to its gasoline counterpart, the TWC for natural gas applications suffers from deleterious ageing effects. Its activity progressively declines with mileage as a result of several intersecting phenomena such as thermal deactivation upon exposure to temperatures as high as 1000 °C and poisoning by fuel additives and impurities.^{7–14} Therefore, the growing interest in natural gas, bio-gas and synthetic natural gas as interim CO₂-poor fuel has driven efforts aiming at designing tailored TWC systems for this specific application.

PdO is considered essential for CH₄ oxidation under lean conditions.¹⁵ Selection of an appropriate oxidic support stabilizes Pd²⁺ against spontaneous reduction at high temperature.^{16–18} The decomposition of PdO to Pd is considered to cause catalyst deactivation during reaction.^{19–21} The role of metallic Pd remains an interesting and much debated issue,²² which is even more fundamental in the case of the TWC because of the oscillating redox reaction conditions. Metallic Pd is observed by X-ray absorption spectroscopy on a model TWC comprising

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Al₂O₃ and ceria–zirconia when the catalyst is exposed to high temperature (800–900 °C)⁵ and a mixed PdO–Pd state needs to be accounted for in light-off experiments under stoichiometric reaction conditions with methane as the hydrocarbon.²³ Pd reduction and sintering can be partly prevented by the formation of PdO at increasing O₂ concentration, for example during fuel cut-off.²⁴ However, methane oxidation activity under lean conditions was recently shown to increase linearly with Pd particle size ranging from 1 to 20 nm,²⁵ which is in line with previous observations.^{26,27} Ageing under reaction conditions (O₂:CH₄ = 4) also produced a more active Pd/Al₂O₃ with larger Pd particles (7–16 nm).²⁸ Restructuring of PdO crystallites upon reduction resulted in increased catalytic activity.^{15,29,30}

Perovskite-type oxides of the general formula ABO₃ have long been recognized as oxidation catalysts^{31–34} and reported as potential NO_x storage/reduction catalysts (NSR),³⁵ TWC,^{36–38} soot combustion catalysts³⁹ and catalysts for natural gas lean burn engines.⁴⁰ In addition, they are also promising oxygen storage materials.^{37,41,42} In line with the search for metal oxides able to stabilize Pd²⁺ against reduction, they can host noble metals in an unusual oxidation state within their lattice. As a direct consequence, in terms of potential TWC components they offer the possibility of stabilizing Pd against sintering under the redox oscillations experienced during operation as has been demonstrated for use with gasoline engines.⁴³ In a very similar way, formation of a Pt-containing perovskite-type oxide (BaPtO₃ and Ba₂CePtO₆) after ageing of the Pt/Ba/CeO₂ NSR catalyst was shown to maintain Pt in a well dispersed state, with the reduced Pt state and the activity being able to be restored after mild reduction.⁴⁴ Despite the growing literature on the topic, which often assumes that the noble metal will adopt the coordination state of the B-site cation, and criticism of the self-regenerative function,⁴⁵ this property needs to be considered with care as it most likely exists within only a limited choice of phase compositions^{46–49} and is clearly temperature dependent.⁴³ Nevertheless, even in the absence of noble metal incorporation, LaFeO₃ is able to exert a strong stabilization effect that makes Pt more resistant against thermal ageing and consequently more active than when it is deposited on γ-Al₂O₃.⁵⁰ Recently, hexagonal YFeO₃ has been shown to be a promising host for Pd for CO oxidation⁵¹ and for methane oxidation under lean⁵² and stoichiometric conditions.⁵³ YFeO₃ was shown to be superior to YMn_{0.5}Fe_{0.5}O₃ with respect to the reversible incorporation of Pd within the YFeO₃ lattice.⁴⁸ YFe_{1-x}Pd_xO_{3-δ} (x = 0.1) exhibited the self-regenerative property at temperatures below 500 °C where the coordinatively unsaturated Pd reversibly segregates out of and re-incorporates within YFeO₃ in consecutive redox treatments.⁵¹ Further, the coordination of Pd with YFeO₃ changes at high temperature. The fine dispersion of Pd in the catalyst after calcination at 700 °C, where Pd is likely in the coordination state that leads to the self-regenerative CO oxidation catalyst,⁵¹ is lost after calcination at 800 °C in favour of segregated PdO, while the hexagonal structure (of the support) is still maintained.⁵² Well dispersed Pd nanoparticles on YFeO₃ grow to well defined particles of ca. 20 nm after repeated methane

oxidation cycles under stoichiometric conditions,⁵⁴ questioning the protective function of the perovskite-related material in case of high temperature treatments of YFeO₃. However, activity enhancement for methane oxidation was observed in parallel to the formation of the 20 nm Pd particles. Such an enhancement has been reported also for the La–Mn–Ce–Pd–O perovskite-type oxide where Pd is observed to segregate after ageing under simulated exhaust of a natural gas engine.⁵⁵

Given the increased methane oxidation activity of Pd/YFeO₃ after exposure to 850 °C concurrent to the growth of Pd particles that suggests possible activation after ageing, herein we address the structure and the catalytic performance of flame-made 2 wt% Pd/YFeO₃ after treatment under lean and stoichiometric conditions at 900 °C. A sample of Pd/YFeO₃ with four-fold lower Pd content was also prepared and aged under identical conditions. The methane oxidation activity under stoichiometric conditions is compared with that of a conventional model TWC.

Experimental

Catalyst synthesis

Pd/YFeO₃ (nominal 0.5 and 2 wt% Pd) was synthesized by flame spray synthesis (FSS) using an acetylene flame. Briefly, a mixture of deionized water and *N,N*-dimethylformamide (DMF, 99%, Sigma-Aldrich, 1:3) was used as solvent of the metal nitrate precursors and sprayed in an acetylene–air flame.⁵³ After synthesis, the samples were calcined in air at 700 °C for 2 h. The powder TWC was composed of Pd, alumina and ceria–zirconia (Pd/ACZ; 1.6 wt% Pd; calcined at 500 °C; SSA = 135 m² g⁻¹) and was kindly provided by Umicore.

Heat treatments, in the following designated concisely as ageing, were carried out at 900 °C for 2 h. Thermal ageing (A_T) was carried out in static air, while stoichiometric ageing (A_S) was performed using a gas mixture comprising 7000 ppm CO, 1300 ppm CH₄, 1600 ppm NO_x and 5300 ppm O₂ in He (λ = 1) at 100 mL min⁻¹ in the plug-flow reactor used for catalytic activity measurements. In the following, the samples are labelled 2Pd/YFO-x, 0.5Pd/YFO-x and Pd/ACZ-x, where x is F (fresh), A_T or A_S.

Characterization

The specific surface area was determined by the single-point BET method in 30 vol% N₂/He using a QuantaChrome ChemBET 3000 instrument. Prior to measurements, all samples were degassed in pure N₂ at 200 °C for 30 min.

The phase composition of fresh and aged catalysts was analysed by X-ray diffraction (X'Pert Pro PANalytical) using monochromatic Cu Kα₁ radiation (λ = 1.54 Å). The diffraction patterns were recorded between 20° and 80° (2θ) with an angular step interval of 0.0167°. Phase identification was performed using the ICDD reference database in the X'Pert HighScore Plus software.

The morphology of the catalysts was observed using a Jeol 2200FS transmission electron microscope equipped with a 200 kV field emission gun and a high-angle annular dark



field detector in STEM mode providing images with atomic number ($Z^{1.5-1.8}$) contrast.⁵⁶ The local composition was determined by energy dispersive X-ray spectroscopy (EDS). The samples were prepared by evaporation of an alcohol suspension on a copper grid coated with a holey carbon film. The palladium particle size distribution was obtained by measuring up to 200 particles using the software ImageJ.⁵⁷

The oxygen storage capacity complete (OSCC) defined as the total amount of desorbed oxygen was determined under static conditions using the thermogravimetric method.^{58,59} The samples were treated with 5 vol% H₂/Ar at 400 °C for 70 min using a TGA Q500 setup (TA Instruments). The measured weight loss was converted into the corresponding oxygen content assuming that the sample is fully oxidized in its initial state.

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) data were collected at a resolution of 4 cm⁻¹ using a VERTEX 70 FT-IR spectrometer (Bruker) equipped with a high temperature reaction chamber (HVC-DRP2, Harrick) and a liquid-nitrogen cooled MCT detector. Prior to CO adsorption, the samples were exposed to 20 vol% O₂/Ar (50 mL min⁻¹) at 500 °C for 30 min, followed by reduction at 300 °C for 30 min in 10 vol% H₂/Ar. After changing to Ar and cooling to room temperature, a background spectrum was obtained prior to CO adsorption in 5 vol% CO/Ar.

Extended X-ray absorption fine structure (EXAFS) spectroscopy data were obtained in the transmission mode at the Pd K-edge (24.35 keV) at the SuperXAS beamline of the Swiss Light Source (PSI, Switzerland). The spectrum of the Pd foil was simultaneously collected for energy calibration. The required sample amount was homogeneously mixed with polyethylene and pressed into pellets. The spectra were energy and background corrected and normalized with the software Athena.⁶⁰

Time-resolved high energy XRD data (86.8 keV, $\lambda = 0.143 \text{ \AA}$, $\Delta E/E \sim 1.4 \times 10^{-3}$) were collected at beamline ID15B of the ESRF (Grenoble, France) using a digital flat 2D panel detector (Pixium 4700) mounted at 900 mm from the sample position. In a concentration modulation experiment, the sample was mounted in a homemade stainless steel cell and was subjected to a sequence of 50 alternate and equally long pulses (25 s) of 1 vol% O₂/He and 1 vol% CO/He at 400 °C at 100 mL min⁻¹.⁶¹ A series of 100 time-resolved diffraction patterns were collected during a full modulation period ($T = 50 \text{ s}$). The catalyst powder (*ca.* 50 mg, 50–100 μm) was fixed between two quartz wool plugs in a homemade stainless steel reactor cell equipped with graphite windows. The temperature of the sample was recorded using a thermocouple inserted at the entrance of the catalyst bed and was constant within $\pm 1 \text{ }^\circ\text{C}$ during the isothermal pulse experiment. The outlet of the cell was connected to a mass spectrometer (OmniStar, Pfeiffer) that was used to follow signals of m/z 28 (CO), 32 (O₂), 44 (CO₂) and 4 (He) and simultaneously to read the sample temperature. The 5000 time-resolved XRD data collected in each experiment were averaged over the number of modulation periods into a single period prior to phase sensitive detection (PSD):

$$I_k^{\text{PSD}} = \frac{2}{T} \int_0^T I(t) \sin(k\omega t + \phi_k^{\text{PSD}}) dt \quad (1)$$

where T is the length of the modulation period, ω is the stimulation frequency, k is the demodulation index, ϕ_k^{PSD} is the phase angle, and $I(t)$ is the intensity of a reflection recorded as a function of reciprocal lattice vector ($Q = 4\pi \sin(\theta/2)/\lambda$, with θ being the scattering angle and λ the wavelength of the incident radiation) and time.^{62–64} According to the theory of modulation enhanced diffraction,⁶⁵ the integral of eqn (1) can be more explicitly (and correctly) written to contain two terms, one of which depends only on the component of the system that is stimulated, whereas the second one contains both this component and the static one. PSD with $k = 2$ should provide only the response of the former component, but we did not find any difference other than intensity by using $k = 1$, which is shown here. The Simpson series was used to calculate the integral of eqn (1). The resulting new set of phase-resolved data contains only the response of species responding with the same frequency of the external stimulation. A critical point is that the changes must be reversible to the external stimulus, for example a change in concentration of the gas phase. PSD removes static signals – typically overwhelming – thus revealing the information pertaining to what are often weak signals. This method is especially useful to study real catalysts with relevant (low) Pd content, which is either silent or unlikely to be distinguished using conventional spectroscopic/diffraction methods. The resulting new set of phase-resolved data essentially contains the response of the signals (species) that are responding with the same frequency of the external stimulation (CO *vs.* O₂ concentration modulation) and on the delay with which they respond to the external stimulus.

Three-way catalytic activity

The light-off of the catalysts was measured in an exhaust mixture comprising 7000 ppm CO, 1300 ppm CH₄, 1600 ppm NO_x and 5300 ppm O₂ (He balance; 100 mL min⁻¹) between 25 °C and 850 °C. The measurements were carried out in a quartz-tube reactor ($d = 6 \text{ mm}$) using catalyst particles (150–200 μm ; 100 mg) diluted with the same volume of sea sand and firmly fixed in the middle of the reactor between two quartz wool plugs. Prior to the reaction, the catalysts were treated in 20 vol% O₂/He (50 mL min⁻¹) at 700 °C for 2 h. The gas hourly space velocity (GHSV) was approximately 60 000 h⁻¹. The outlet of the reactor was analysed using a gas chromatograph (Agilent 3000A Micro GC) equipped with two columns (PoraPLOT-Q for CH₄, CO₂, H₂O and N₂O; molecular sieve 5 \AA for CO and NO).

Results and discussion

Characterization of fresh and aged catalysts

The structural properties of fresh and aged catalysts are summarized in Table 1. The phase composition of each sample was determined by XRD. Since the same crystal phases were identified in 0.5Pd/YFO and 2Pd/YFO, only the XRD patterns of 2Pd/YFO and Pd/ACZ are presented in Fig. 1. 2Pd/YFO-F



exhibits the patterns of hexagonal YFeO_3 , while no information is obtained on the Pd-containing phase due to the overlap with the YFeO_3 phase and its good dispersion (Table 1).⁵³ Irrespective of the ageing atmosphere, hexagonal YFeO_3 is converted into the orthorhombic phase, which was reported to occur around 800 °C.^{66–68} Despite the high temperature treatment, no evidence of the presence of a Pd-containing phase is obtained. On the contrary, the inset of Fig. 1 demonstrates that metallic Pd is formed as a consequence of stoichiometric ageing. The Pd(111) and Pd(200) reflections of fcc-Pd appear at *ca.* 40 and 47°, respectively. The phase transition of YFeO_3 upon exposure to 900 °C is accompanied by a considerable loss of surface area (Table 1). Likely as a consequence of both phase transition and particle growth processes, the oxygen storage capacity complete (OSCC) is also severely diminished compared to the remarkable value measured for the fresh catalyst. The OSCC values of the aged 2Pd/YFO catalysts are *ca.* ten times higher than that of the thermally aged support suggesting that the OSCC of the aged catalysts is primarily dictated by the presence of Pd. This is also confirmed by the decreased OSCC value obtained with lower Pd loading that is still *ca.* 4–8 times higher than that of the support depending on the ageing procedure.

For comparison, the XRD patterns of fresh and aged Pd/ACZ (Fig. 1) correspond to those of $\gamma\text{-Al}_2\text{O}_3$ and cubic CZ and are very similar, confirming the thermal stability of ACZ.⁶⁹ However, the general sharpening of the diffraction peaks of CZ after both ageing procedures is indicative of some extent of sintering. This is especially evident in Pd/ACZ- A_T and is confirmed by the larger extent of surface area loss of this sample compared to Pd/ACZ- A_S . A (weak) peak corresponding to PdO becomes visible at *ca.* 55° in Pd/ACZ- A_T as well. Similarly to 2Pd/YFO- A_S , the XRD pattern of Pd/ACZ- A_S shows reflections of metallic Pd (peaks at 40 and 47°), confirming the reduction of Pd under these conditions.

HAADF-STEM was used to confirm the textural changes occurring upon ageing and to characterize the morphology of the Pd-containing phase. The STEM images of 2Pd/YFO are shown in Fig. 2d–f. Identical conclusions can be drawn for 0.5Pd/YFO (Fig. 2g–i). The sintering of YFeO_3 after exposure

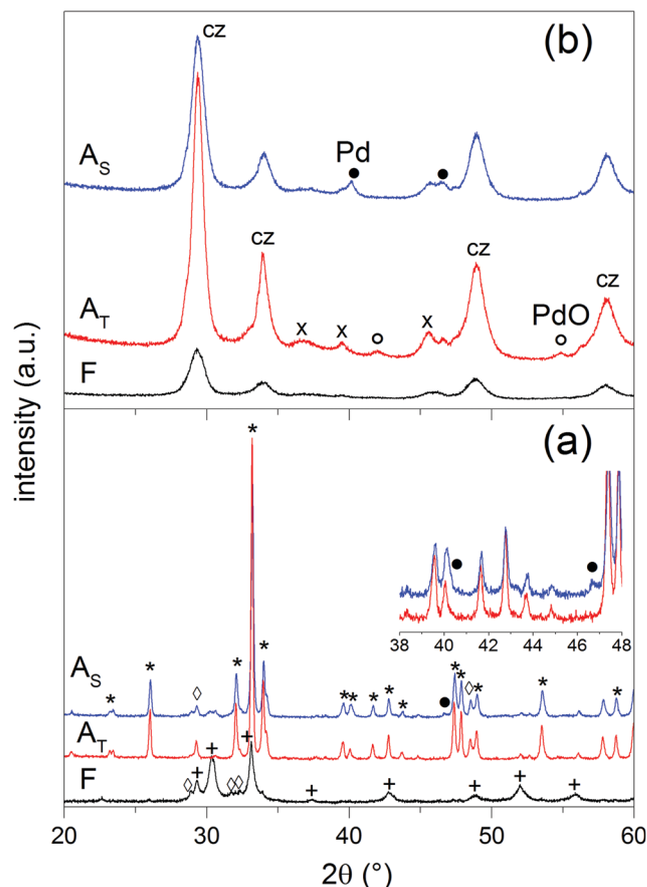


Fig. 1 XRD patterns of fresh and aged (a) 2Pd/YFO and (b) Pd/ACZ. (*) Orthorhombic YFeO_3 , (+) hexagonal YFeO_3 , (◊) Y_2O_3 , (x) $\gamma\text{-Al}_2\text{O}_3$, (cz) ceria-zirconia, (●) Pd, and (○) PdO.

to 900 °C is rather obvious from these data. Similarly, Pd particle growth is also observed to be accompanied by a change in shape that depends on the ageing procedure. Pd particles appear mostly ‘wetting’ and spreading on YFeO_3 after thermal ageing (Fig. 2e; Table 1). Plate-like Pd particles were observed to form in an oxygen containing environment at $T > 800$ °C on Al_2O_3 , while re-dispersion of Pd occurred on more coordinating metal oxides such as ZrO_2 .⁷⁰ However,

Table 1 Structural properties of fresh and aged catalysts

Entry	Condition	SSA ($\text{m}^2 \text{g}^{-1}$)	XRD ^a	OSCC ^b ($\mu\text{mol O}_2 \text{g}^{-1}$)	Particle size ^c (nm)
Pd/ACZ	F	135	ACZ	150	8 [3]
	A_T	70	ACZ + PdO	157	52 [27]
	A_S	97	ACZ + Pd	188	13 [4.4]
2Pd/YFO	F	19	h + Y_2O_3 ^d	504	3.5 [1.4]
	A_T	3	o + Y_2O_3 + h	86	91 [35]
	A_S	6	o + Y_2O_3 + h	88	19 [7.6]
0.5Pd/YFO	F	18	h + Y_2O_3 ^d	424	—
	A_T	3	o + Y_2O_3 + h	39	101 [37.5]
	A_S	6	o + Y_2O_3 + h	67	18 [5.7]
YFO	F ^e	19	h + Y_2O_3 ^d	416	—
	A_T	3	o + Y_2O_3 + h	8	—

^a h: hexagonal, o: orthorhombic. ^b Oxygen storage capacity complete, determined by TG analysis. ^c Average Pd particle size estimated by STEM. The standard deviation σ is shown in brackets. ^d Metastable monoclinic Y_2O_3 . ^e See ref. 53.



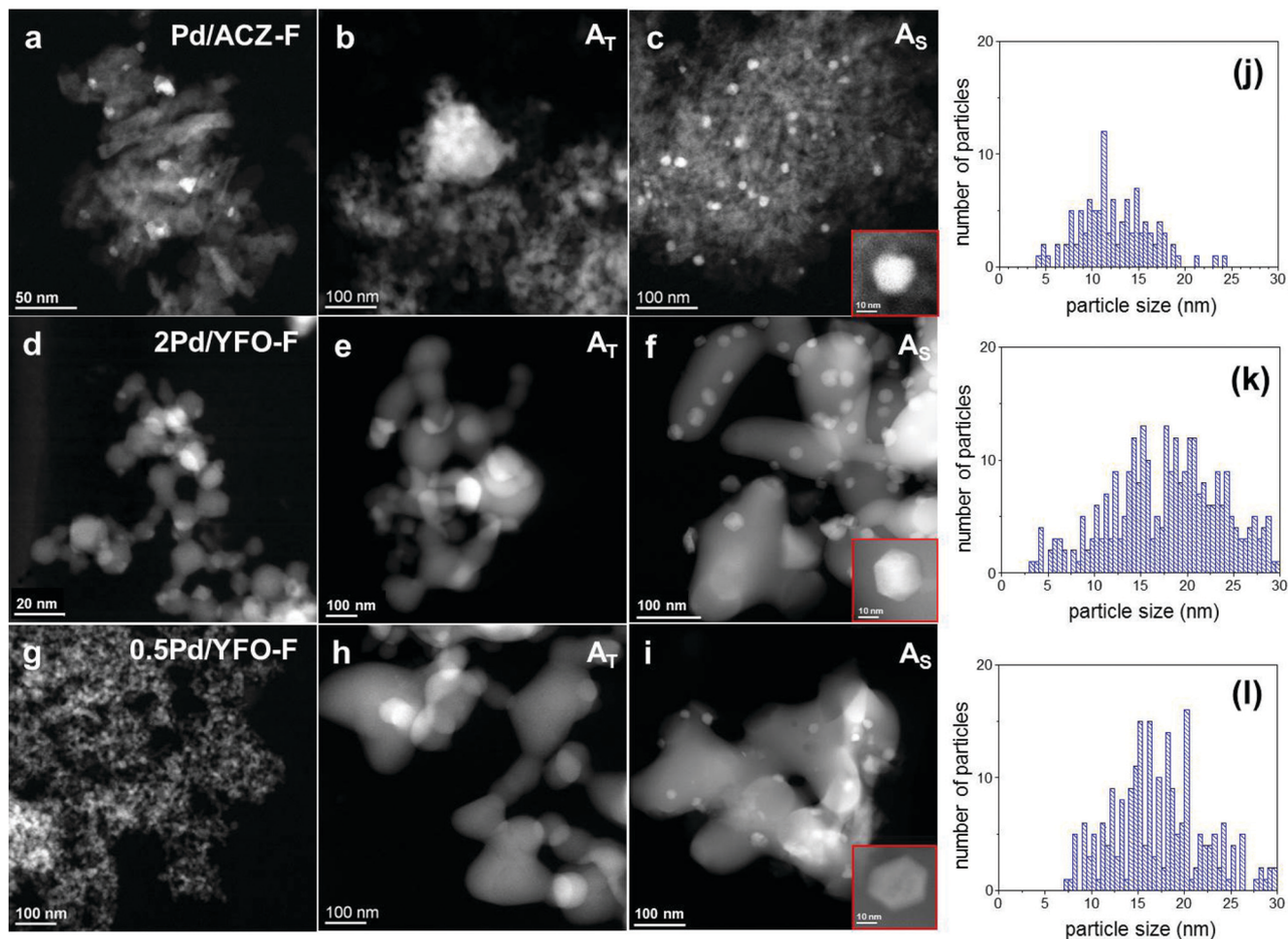


Fig. 2 HAADF-STEM images of fresh and aged (a–c) Pd/ACZ, (d–f) 2Pd/YFO, and (g–i) 0.5Pd/YFO and particle size distributions of (j) Pd/ACZ-A_S, (k) 2Pd/YFO-A_S and (l) 0.5Pd/YFO-A_S. F, fresh; A_T, thermal ageing; A_S, stoichiometric ageing.

the particles are not always easily recognizable and preclude any correct determination of particle size distribution. Assuming that the shape of Pd particles present in 2Pd/YFO-A_T is similar to that on Al₂O₃, this would indicate that the anchoring nature of YFeO₃ is lost upon ageing, in agreement with recent observations on Pd/YFeO₃ calcined at 700 °C and 800 °C.⁵² The average particle size of the finely dispersed fresh catalyst increases from *ca.* 3.5 nm to *ca.* 19 nm in 2Pd/YFO-A_S (Fig. 2f) and the particle size distribution broadens (Table 1). Additionally, Pd nanoparticles with a different morphology from those obtained upon thermal ageing are obtained. The large fraction of hexagonal projections detected in the STEM images indicates that Pd particles possess a (truncated) cubo-octahedral shape exposing (111) and (100) facets.⁷¹ The change in morphology upon stoichiometric ageing is similar to the one observed after three reaction cycles under $\lambda = 1$ conditions.⁵³ Hence, structural changes after repeated reaction cycles and stoichiometric ageing are of similar nature.

The different particle shape and the smaller particle size observed for 2Pd/YFO-A_S confirm previous results obtained on Pt/Al₂O₃ catalysts used for NO oxidation that were thermally aged at 800 °C or under a simulated lean diesel environment

(propene, CO, NO_x and O₂) at the same temperature for 5 h.⁷² Ageing under reaction conditions caused formation of truncated cubo-octahedral Pt particles with reduced size compared to the thermally aged catalyst. Hence, ageing under a reactive atmosphere produces smaller particles than thermal ageing at the same temperature.⁷³

The STEM observations on the evolution of the structure of the Pd phase on YFeO₃ are confirmed and compared with the analysis of the Pd/ACZ reference catalyst. No significant change in the size of the metal oxide is observed that could support the variations of XRD intensity. The average Pd particle size increases from 8 nm in Pd/ACZ-F to 52 nm in Pd/ACZ-A_T (Fig. 2a and b). The growth of Pd particles is again less severe under stoichiometric conditions, with the average particle diameter being 13 nm in Pd/ACZ-A_S. Both round-shaped and truncated particles can be found in this sample (Fig. 2c). The particle size distribution of Pd/ACZ by STEM is, however, greatly influenced by Pd particles deposited on CZ that are difficult to detect.

The oxidation state and coordination environment of Pd were investigated by X-ray absorption spectroscopy. The XANES spectra of fresh and aged 2Pd/YFO and Pd/ACZ recorded at



the Pd K-edge are shown in Fig. 3a. The spectra of a Pd foil and bulk PdO are also given for reference. No detectable metallic Pd is present in the fresh catalyst.⁵³ The white line of 2Pd/YFO-F is very close to that of Pd/ACZ-F which is typical of a supported PdO catalyst. Nevertheless, the non-phase shift corrected k^3 -weighted Fourier transform EXAFS spectrum of 2Pd/YFO-F (Fig. 3b) presents only a clear Pd–O coordination shell at 1.6 Å, while the second Pd–O–Pd shell clearly appears in Pd/ACZ-F at *ca.* 3.0 Å. The difference is due to the smaller particle size measured for the YFO-based catalyst (Table 1). The XANES and FT-EXAFS data of the thermally aged catalysts are also characteristic of PdO, but the stronger contribution of the Pd–O–Pd shell compared to the fresh catalysts confirms PdO particle growth upon ageing. After stoichiometric ageing, the presence of the Pd–Pd coordination shell at 2.5 Å at the expense of the Pd–O phase characterizes the 2Pd/YFO and Pd/ACZ catalysts. This confirms that Pd is in the metallic state, in agreement with the XRD evidence and the Pd sintering observed by STEM. Based on the presence of the Pd–Pd shell, we can observe that 2Pd/YFO-*A_T* also exhibits a contribution from metallic Pd that is evident from the small peak at *ca.* 2.5 Å. The presence of an additional scattering feature at 2.1 Å in the FT-EXAFS data of 2Pd/YFO-*A_S* could be assigned to the formation of a Pd–Fe alloy.⁴⁸

The dispersion of Pd particles was also qualitatively characterized by CO adsorption using infrared spectroscopy (Fig. 4). After reduction of the fresh catalysts at 300 °C, adsorbed CO on Pd exhibits three main signals at 2090 cm^{-1} (linearly bonded

CO on Pd⁰ corner atoms, CO_L), 2000–1950 cm^{-1} (μ_2 -bridge bonded species on Pd(100) facets or edge sites) and 1950–1800 cm^{-1} (a broad band resulting from μ_3 -hollow/multiple bonded CO on Pd(111) planes) which were assigned according to the available literature.^{71,74,75} Pd/ACZ-*A_S* is the only sample displaying a signal at 2125 cm^{-1} that could be attributed either to the presence of ionic Pd²⁺ (ref. 76) or to coordination of CO to Ce³⁺.⁷⁷ The major difference between the spectra of fresh 2Pd/YFO, 0.5Pd/YFO and Pd/ACZ lies in the overall intensity of the signals (Fig. 4), which varies in the order Pd/ACZ > 2Pd/YFO > 0.5Pd/YFO. The higher intensity of Pd/ACZ-F signals is caused by its larger specific surface area (Table 1), since more Pd can be probed by CO and by the IR radiation, whereas the larger intensity of signals of 2Pd/YFO-F over those of 0.5Pd/YFO-F is attributed to the difference in Pd loading at identical surface area values. The intensity ratio between the signals of CO_L and that of bridge-bonded CO (A_{2090}/A_{1990}) can be taken as an indicator of the contribution of CO adsorbed on Pd particle edges and corners relative to terrace sites. A larger A_{2090}/A_{1990} ratio corresponds to a smaller average particle size, which correlates well with the particle size trends measured from STEM images (Table 1). 0.5Pd/YFO-F exhibits the largest A_{2090}/A_{1990} ratio among all samples, thus explaining the difficulty in estimating its Pd particle size from STEM (Fig. 2g).

Ageing causes a severe loss of IR signal intensity in Pd/ACZ (note the scale bar in Fig. 4) and of the A_{2090}/A_{1990} intensity ratio, reflecting the loss of Pd dispersion. The intensity changes associated with the YFO-based catalysts are

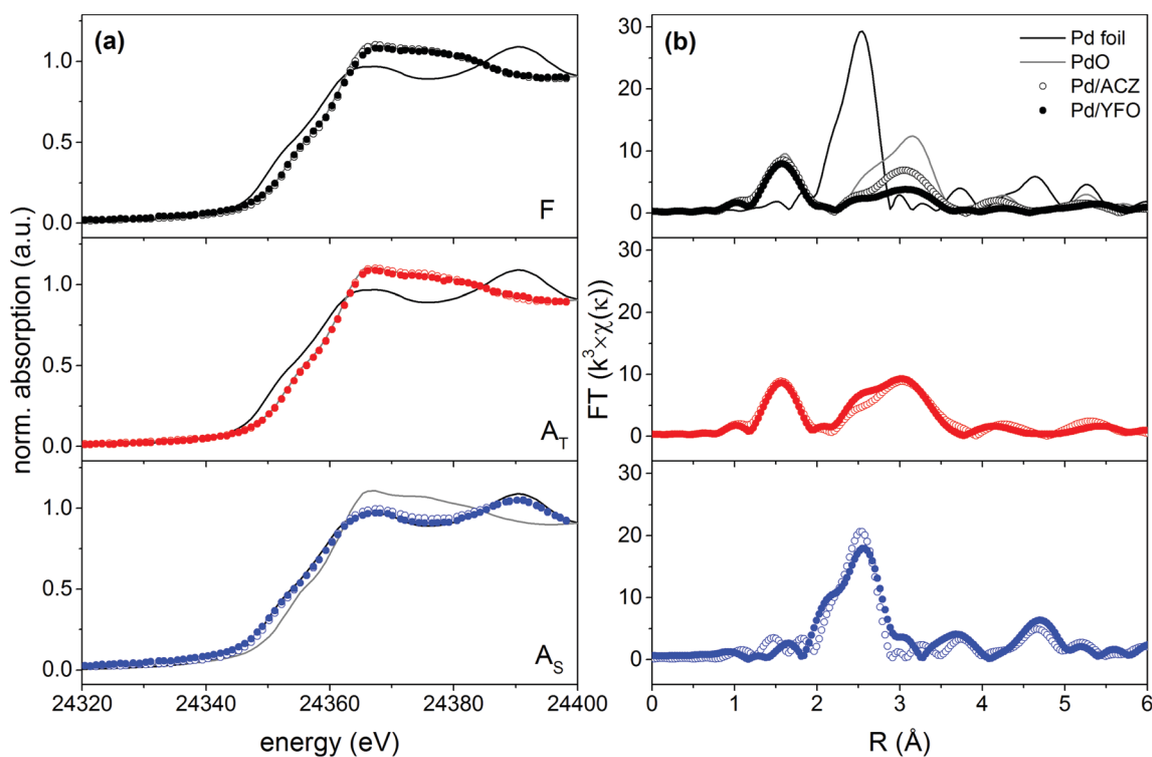


Fig. 3 (a) XANES and (b) non-phase shift corrected k^3 -weighted FT-EXAFS spectra of fresh and aged 2Pd/YFO and Pd/ACZ. F, fresh; *A_T*, thermal ageing; *A_S*, stoichiometric ageing.



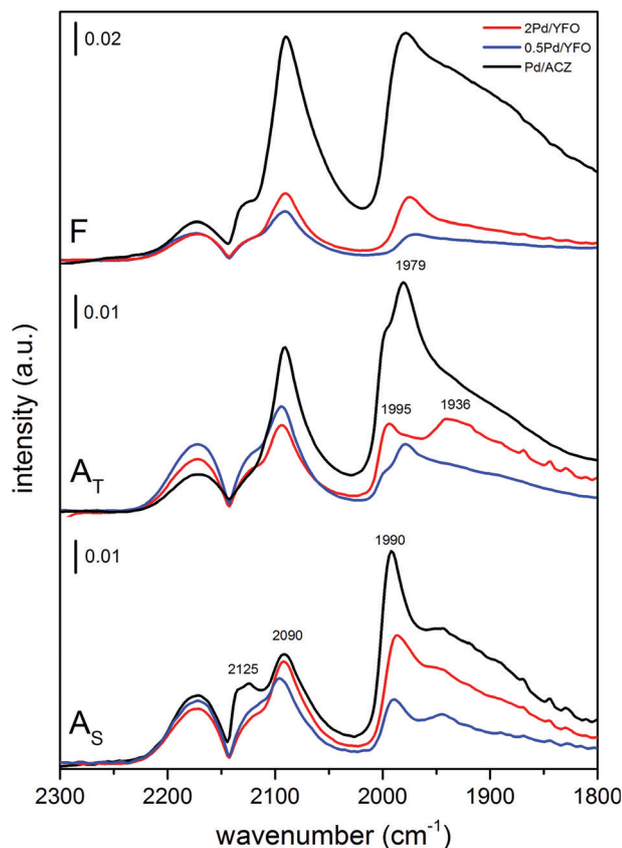


Fig. 4 DRIFTS spectra of adsorbed CO on fresh and aged 2Pd/YFO, 0.5Pd/YFO and Pd/ACZ. F, fresh; A_T, thermal ageing; A_S, stoichiometric ageing.

strikingly less evident. The position of the signals corresponding to CO_L (2090 cm⁻¹) and hollow bonded CO (1930–1800 cm⁻¹) in the aged samples is similar to those observed in fresh catalysts. The signals of the bridge-bonded CO species vary more significantly from stoichiometric ageing to thermal ageing. In the A_S catalysts, the signal is slightly shifted to 1990 cm⁻¹ compared to fresh samples (1980 cm⁻¹), but spectra of 2Pd/YFO-A_S, 0.5Pd/YFO-A_S and Pd/ACZ-A_S are very similar, likely reflecting a comparable morphology but different sizes of the Pd particles. Two to three signals corresponding to μ₂-bridge-bonded CO species are visible below 2000 cm⁻¹ in the spectra of A_T catalysts. 2Pd/YFO-A_T presents a pronounced signal at ca. 1936 cm⁻¹ that could be assigned to isolated bridge species.^{77,78} Comparison between STEM and DRIFTS data for these catalysts is also more difficult. STEM suggests the presence of large PdO particles (confirmed by EXAFS) that are challenging to clearly detect, whereas pre-reduction of the samples prior to CO adsorption may affect the aspect of the particles. Reduction of the large PdO particles of 2Pd/YFO-A_T may cause the formation of islands of metallic Pd that could account for the signal at 1936 cm⁻¹. Therefore, the difference between the spectra of A_T and A_S catalysts is attributed to the different morphologies obtained after reduction at 300 °C.

Time-resolved high energy XRD

The surface dynamics of the fresh and aged catalysts was addressed using high energy X-ray diffraction combined with the modulation excitation approach.^{61,79} High energy XRD is a suitable method because it provides information on the whole material in a snapshot and in a time-resolved manner. It has been shown that subtle structural changes occurring during reaction that are not visible in XRD can be enhanced if the experimental data are collected under pulsed conditions.⁶¹

In this work, the modulation experiment consists of alternate and repeated CO vs. O₂ pulses that should simulate the protocol typically used to determine the dynamic OSC.⁸⁰ We restrict the discussion to fresh and aged 2Pd/YFO catalysts whose time-resolved XRD patterns recorded at 400 °C are dominated by the hexagonal and orthorhombic YFeO₃ structures, respectively (Fig. 5a). Similarly to the XRD of Fig. 1, and despite the exposure to high temperature during ageing and the resulting structural changes mentioned above, there remains still no clear indication about the Pd-containing phase because of the systematic overlap of Pd and PdO reflections with those of the support oxide. A completely different picture is obtained from the phase-resolved data of Fig. 5b, where reversible structural variations during the CO–O₂ pulses become clearly visible. By comparison with ICSD diffraction data (Fig. 5b, top), the reflections that became visible at 2.78 Å⁻¹ (corresponding to 2θ = 40° for CuKα radiation) and 3.22 Å⁻¹ (46.5°) are assigned to Pd metal, whereas the broader features at 2.36 Å⁻¹ (33.6°) and 3.74 Å⁻¹ (54.6°) are assigned to PdO reflections. The assignment is also corroborated by the opposite sign of the signals of Pd and PdO. The reflections of the Pd-containing phases experience only an amplitude change in response to the modulation. On the contrary, the weaker changes associated with the dominant reflections of YFeO₃ involve both amplitude and *d* spacing variations. The signals exhibit a complex differential profile that is given by the overlap of various close reflections. In agreement with similar experiments conducted on Pd/ACZ,⁶¹ the *d* spacing variation during the modulation experiment is associated with the OSC characteristics of the material, therefore oxygen release in CO (decrease of *Q*) and oxygen uptake in O₂ (increase of *Q*).

The major difference between the catalysts is the amplitude of the changes emphasized by PSD. The overall intensity of the changes observed in 2Pd/YFO-F is small compared to that of the aged catalysts. Only the oscillations of hexagonal YFeO₃(111) at 2.32 Å⁻¹ (2θ = 33°) and Pd(111) at 2.78 Å⁻¹ are obvious. This is due to lack of long range crystal order in the fresh catalyst because of the nano-size of the support and Pd. The reflection of Pd(111) in the fresh catalyst appears very broad for the same reason. The difference in the YFeO₃ crystal structure (hexagonal and orthorhombic) may also influence the extent of the observed changes related to the support oxide, because the surface oxygen content of the material is structure dependent.⁶⁷ The changes around the Pd and PdO reflections are most pronounced in 2Pd/YFO-A_S, whereas those associated with orthorhombic YFeO₃ are evident only in 2Pd/YFO-A_T



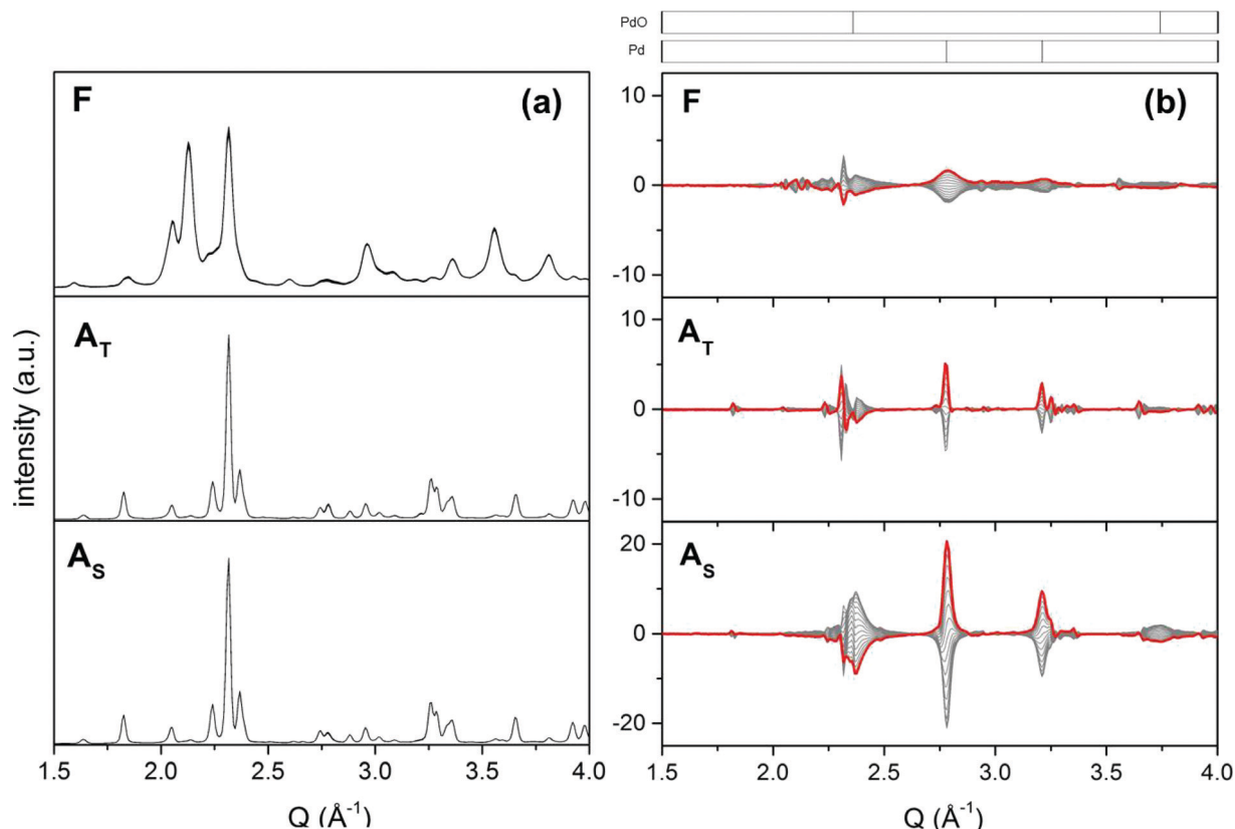


Fig. 5 (a) Time-resolved high energy XRD data of fresh and aged 2Pd/YFO obtained in a CO vs. O₂ modulation experiment at 400 °C. (b) Corresponding phase-resolved data. The red profile is the in-phase pattern. F, fresh; A_T, thermal ageing; A_S, stoichiometric ageing.

as a result of the lesser contribution from PdO. Thus, the oxygen exchange properties of the aged catalysts rely mainly on the oxidation and reduction of Pd particles in agreement with the above observations on the low OSCC values.⁸¹ We cannot establish quantitatively how relevant is the structural variation observed for YFeO₃.

The temporal evolution of the Pd(200) reflection (3.22 \AA^{-1}) for the three catalysts is reported in Fig. 6a. No kinetic data for the two YFeO₃ phases are shown because of the high level of noise and the very small changes induced by the modulation experiment that allow only for a qualitative evaluation of the data. Pd(200) is selected because the strongest Pd(111) reflection (2.78 \AA^{-1}) interferes with those of the YFeO₃ phases, especially in 2Pd/YFO-A_T. The data also clearly evidence the difference in amplitude discussed above. It is clear from the kinetic data that the Pd and PdO reflections show opposite behavior in the phase-resolved data of Fig. 5b because Pd species reduce and re-oxidize in the CO–O₂ pulses. Besides the marked difference in amplitude of the changes, the delay at which changes occur in the CO pulse is also different. Whereas PdO of 2Pd/YFO-A_T reduces immediately in the presence of CO (Fig. 6a), the development of the Pd(200) reflection sets in only after *ca.* 10 s in 2Pd/YFO-F and 2Pd/YFO-A_S and is completed within a few seconds.

The structural changes are reversible and Pd re-oxidation occurs in all catalysts. However, it is a slower process than

reduction and encompasses the whole O₂ pulse. The corresponding MS data (Fig. 6b) show that CO₂ evolution takes place over the whole CO pulse in the case of the fresh catalyst. In agreement with the temporal evolution of the Pd(200) reflection, the only little gas phase CO that is observed appears after *ca.* 10 s from admittance of CO and in correspondence of the decrease of CO₂ evolution. Therefore, 2Pd/YFO-F is the most active to oxidize CO, which is facilitated by the rather high SSA and high OSCC value compared to the aged catalysts (Table 1). Ageing causes an earlier CO breakthrough in the gas phase and a corresponding earlier breakthrough of gas phase O₂ in the oxidation pulse. However, 2Pd/YFO-A_S is able to consume CO up to about half the total duration of the CO pulse, a fact that is reflected in the broader CO₂ profile than that of 2Pd/YFO-A_T and which agrees well with the retarded Pd reduction observed in the high energy XRD data. Hence, in this type of experiment 2Pd/YFO-A_S demonstrates better oxygen storage capacity than 2Pd/YFO-A_T. The catalyst aged under stoichiometric conditions possesses highly reactive surface oxygen species giving high CO₂ evolution under oscillating conditions, whereas the thermally aged catalyst is characterized by poorer oxygen exchange properties. Given the similar structure of the support, this difference is associated with the redox properties of the metallic Pd particles of 2Pd/YFO-A_S. The discrepancy with the OSCC values is explained by the milder reduction conditions and the repeated redox pulses



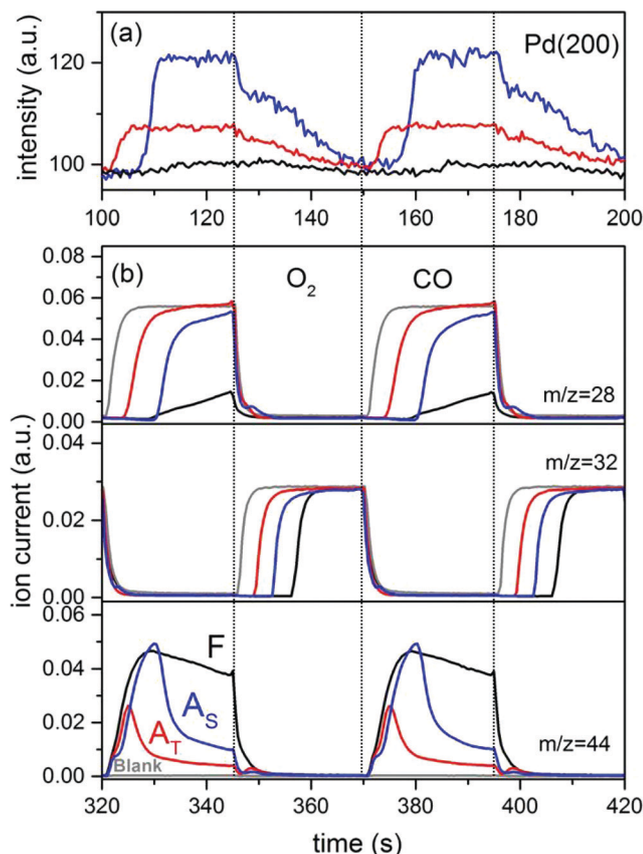


Fig. 6 (a) Temporal evolution of the Pd(200) reflection obtained from the time-resolved high energy XRD data of the CO vs. O₂ modulation experiment at 400 °C. (b) Corresponding online MS data. F, fresh; A_T, thermal ageing; A_S, stoichiometric ageing.

applied during the pulse experiment compared to the thermogravimetric experiment. Therefore, the metallic Pd particles observed on 2Pd/YFO-A_S appear crucial for maintaining the oxygen exchange properties of the catalyst.

Based on the Scherrer equation using the Pd(111) reflection of the in-phase XRD pattern of each catalyst (Fig. 5b, red), the estimation of the Pd crystallite size provides 2.6, 13.5 and 7.6 nm for 2Pd/YFO-F, 2Pd/YFO-A_T and 2Pd/YFO-A_S, respectively. The estimation is strongly affected by the assumption that all catalysts possess spherical (and unstrained) particles, which does not correspond to the STEM images especially in the case of 2Pd/YFO-A_T. Finally, the estimation is also affected by some overlap of the selected reflection with those of YFeO₃. The values obtained considering these issues are lower than the particle size values determined by STEM (Table 1). The kinetic data of Fig. 6a indicate a slow re-oxidation that is possibly not able to involve the full particle volume. Hence, the estimation of the crystallite size based on the net variation of the Pd(200) reflection indicates that a large fraction of Pd but likely not all Pd responds to the CO–O₂ modulation, *i.e.* is repeatedly oxidized and reduced.

To conclude this section, high energy XRD data provide a qualitative measure of the facility of Pd to undergo a redox cycle. The large metallic Pd particles observed in 2Pd/YFO-A_S

(*ca.* 20 nm) are subject to re-oxidation under periodic redox conditions at 400 °C suggesting that this could well be the situation also under stoichiometric reaction conditions.

Three-way catalytic activity

Fig. 7 shows the light-off curves of CH₄ obtained for fresh and aged catalysts under stoichiometric conditions. The T_{50} for CH₄ and T_{50} for CO ($T_{50}^{\text{CH}_4}$ and T_{50}^{CO}) are summarized in

Table 2. A systematic improvement in $T_{50}^{\text{CH}_4}$ upon ageing is observed in the perovskite-type catalysts. Despite the structural changes observed after ageing involving both Pd and support particle growth, $T_{50}^{\text{CH}_4}$ of the 2Pd/YFO-A_S catalyst is shifted by more than 100 °C to lower temperature after stoichiometric ageing, attaining those of Pd/ACZ-F and Pd/ACZ-A_S. Similar activity improvement by nearly 80 °C is observed also for 0.5Pd/YFO-A_S. The effect of thermal ageing is less pronounced, but $T_{50}^{\text{CH}_4}$ is shifted by 50 °C to lower temperature in both YFO-based catalysts. For comparison, the activity of Pd/ACZ is stable upon ageing. Only thermal ageing increases $T_{50}^{\text{CH}_4}$ by 40 °C contrary to the behavior of Pd/YFO catalysts.

This deactivation is attributed to the more significant sintering of PdO particles (Table 1). In contrast to Pd/ACZ, the aged YFO catalysts do not reach full CH₄ conversion at 800 °C, which is attributed to the low surface area of both the Pd and the YFeO₃ support and to the limited oxygen storage capacity of the aged catalysts.

Contrary to CH₄, CO oxidation activity of 2Pd/YFO, 0.5Pd/YFO and Pd/ACZ decreases upon ageing (Table 2) and the structural changes produced by thermal ageing appear more severe likely as a result of Pd particle growth.^{82,83} This is in agreement with observations on Pd-based catalysts aged on a dynamometer.⁸⁴ Besides, the YFeO₃ support alone exhibited deactivation upon consecutive reaction runs.⁵³ Therefore, the activity enhancement is peculiar to CH₄ chemistry on Pd. The influence of water on the feedstock still remains to be assessed to ascertain whether a true activity enhancement would also be observed since water has been reported to decrease the catalytic activity of Pd.^{85,86}

It is clear from the characterization and the activity data that activation is coincident with the formation of well-defined metallic Pd particles. Because of the similar structural changes, the activity improvement appears to be of the same nature as that observed for 2Pd/YFO-F during consecutive reaction cycles under stoichiometric conditions.⁵³ The formation of metallic Pd particles is accompanied by the loss of surface area of the catalyst and by the phase transition from hexagonal to orthorhombic YFeO₃. However, because the OSCC values after ageing seem to be associated with Pd rather than with the presence of YFeO₃, the support does not appear to be responsible for the improved catalytic activity after ageing. This indicates that the degree of coordination of Pd with YFeO₃ after prolonged exposure to 900 °C is low and likely not



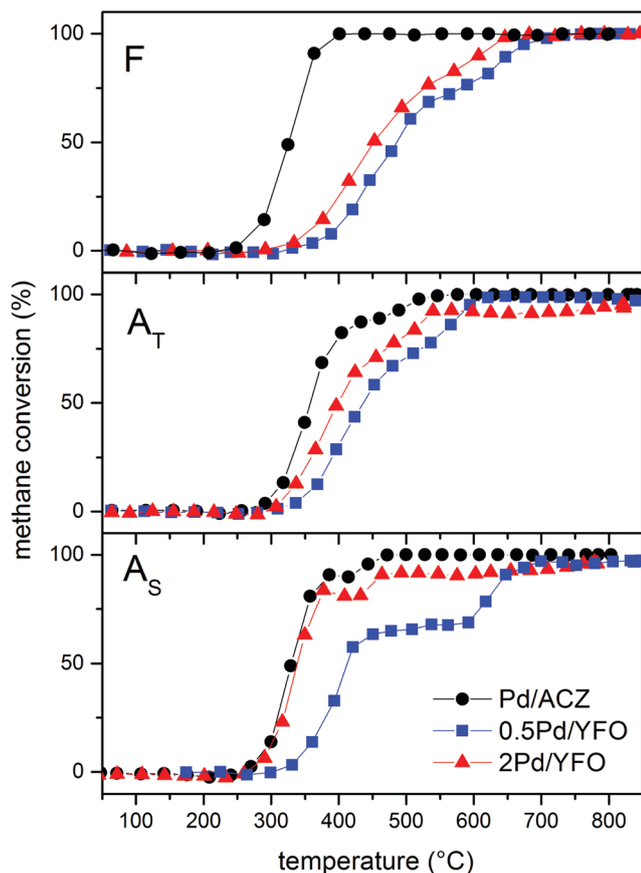


Fig. 7 CH_4 conversion curves of fresh and aged 2Pd/YFO, 0.5Pd/YFO and Pd/ACZ. F, fresh; A_T , thermal ageing; A_S , stoichiometric ageing.

Table 2 Catalytic activity of the fresh and aged catalysts

Entry	$T_{50}^{\text{CH}_4}$ (°C) ^a			T_{50}^{CO} (°C) ^b		
	F	A_T	A_S	F	A_T	A_S
Pd/ACZ	320	358	329	116	206	189
2Pd/YFO	452	399	338	164	228	204
0.5Pd/YFO	487	435	413	171	174	236

^a Temperature of 50% CH_4 conversion. ^b Temperature of 50% CO conversion.

of the same ‘self-regenerative’ nature as LaFeO_3 . The Pd– YFeO_3 interaction was shown to change already from a sample calcined at 700 °C to one calcined at 800 °C, the latter exposing PdO-like particles on hexagonal YFeO_3 .⁵² However, no phase transition was observed, indicating that the hexagonal structure may have been stabilized by the interaction with Pd. In contrast, $\text{YFe}_{1-x}\text{Pd}_x\text{O}_{3-\delta}$ ($x = 0.1$) was shown to be a self-regenerative catalyst for CO oxidation,⁶⁶ but no high temperature treatment was performed in previous studies. Our interpretation of the low temperature self-regeneration *versus* the high temperature segregation of Pd is in terms of geometric factors; however, this needs further confirmation. The periodic segregation–incorporation of Pd in YFeO_3 is favored

at low temperature due to the fact that Pd can adopt a coordination similar to that of tetrahedral Fe ions of hexagonal YFeO_3 .⁵¹ In the flame-made catalyst, Pd is deposited on the YFeO_3 oxide already in the fresh state. Some extent of interaction is observed, but it is likely not sufficient to avoid the phase transition to orthorhombic YFeO_3 .⁵³ In this phase, the segregation of Pd is favored because Y^{3+} (1.18 Å)⁸⁷ does not help in stabilizing a Pd^{3+} -like cation in the octahedral Fe sites, contrary to the larger La^{3+} (1.36 Å)⁸⁸ in LaFeO_3 . The strong tendency of Pd^{3+} to disproportionate to Pd^{2+} and Pd^{4+} , when it is in an oxide lattice, can be overcome by the relatively small octahedral tilting due to the largest rare earth ion being in the A-site.⁸⁹

It is tempting to associate the improved catalytic activity with metallic Pd based on the experimental observations. The particle size dependence (structure sensitivity) and the role of metallic Pd *vs.* PdO are still debated issues in methane oxidation.^{4,19,90–95} This uncertainty is predominantly associated with the different experimental conditions used for the determination of catalytic activity and the difference in nature of the various catalysts. Moreover, the debate concerns primarily methane oxidation under lean conditions, where it is accepted that PdO is formed during reaction. Few studies have been reported for oxidation under low partial pressures of oxygen as it is the case for a TWC. The characterization data, together with the activity data of Fig. 7, demonstrate that discrete metal particles are favorable for methane oxidation, which is in agreement with various observations by others.^{25,26,90} This is not only the case for 2Pd/YFO but also for the reference catalyst, a conventional TWC, where particle growth occurs despite the presence of ceria–zirconia, and the activity of Pd/ACZ- A_S is unchanged compared to that of Pd/ACZ-F. An important piece of information is derived from the observation that both the large Pd particles and the possibly large PdO particles undergo a reversible reduction–oxidation under redox conditions. Therefore, under the stoichiometric conditions used in this work, oxygen is activated on the large Pd particles of 2Pd/YFO- A_S . On the large PdO particles formed in the thermally aged Pd/YFO- A_T catalysts where an activity improvement is induced to a lesser extent, the activation phenomenon could be explained with the partial reduction of PdO during reaction, thus forming metallic Pd on the PdO surface.³⁰ EXAFS and DRIFTS data confirm this possibility. Therefore, a mixed PdO_x/Pd phase is present during reaction in both aged catalysts.

Finally, the light-off profiles of Pd/ACZ- A_S and 2Pd/YFO- A_S exhibit a clear loss of CH_4 conversion by 10–20% around 350–450 °C (Fig. 7). This is more pronounced in 0.5Pd/YFO- A_S (400–650 °C), likely as a consequence of the lower Pd content. This behaviour is characteristic of supported PdO catalysts under lean conditions and is due to spontaneous PdO reduction.^{19,20,93} The activity loss is slightly more pronounced in the samples aged under stoichiometric conditions because of the presence of large metal particles, thus diminishing the amount of available Pd atoms. These particles can be partially oxidized as demonstrated above, but reduction during



reaction occurs at lower temperature because of the low oxygen content in the feed under $\lambda = 1$ conditions.²³ Further study under reaction conditions should be performed to confirm this behavior and the low temperature reduction of PdO.

Conclusions

The catalytic performance of Pd/YFO with 0.5 and 2 wt% Pd content was determined after high temperature treatment. Two ageing protocols were chosen, thermal ageing at 900 °C in air and stoichiometric ageing at 900 °C under reaction conditions comprising CH₄, CO, NO and oxygen. The reference TWC was aged under identical conditions for comparison. A gradual activity improvement was observed from the performance of the fresh Pd/YFO catalyst to the thermally aged one and to the sample aged under stoichiometric conditions. Pd loading affected only the extent of improvement, with the highest loading providing a better activity. After stoichiometric ageing, 2Pd/YFO was as active for methane oxidation as Pd/ACZ. In contrast to Pd/YFO catalysts, Pd/ACZ was less active after thermal ageing. For comparison, CO oxidation activity systematically decreased upon ageing indicating that the observed improved oxidation activity is peculiar to CH₄.

Structural characterization of the aged catalysts revealed that the activity improvement of Pd/YFO after stoichiometric ageing is accompanied by the formation of well-defined metallic Pd particles of ca. 20 nm. In the case of thermal ageing that produces less activity enhancement, Pd remains predominantly oxidized. The interaction of Pd with YFeO₃ is poorer after ageing probably as a result of the phase changes observed in the support that prevent coordination of Pd within the YFeO₃ lattice. Therefore, the observed activity improvement is mostly attributed to the changes related to the shape and oxidation state of Pd particles. However, it has been demonstrated that both Pd and YFeO₃ remain highly dynamic with respect to exposure to periodic redox conditions. Large metallic and oxidic Pd particles periodically reduce and oxidize. Therefore, this study provides evidence that a mixed PdO_x/Pd pair is at work during reaction and that large metal particles are not detrimental to methane oxidation under stoichiometric conditions.

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