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Attempts at an in-situ Raman study of Ceria / Zirconia catalysts in PM combustion

James A Sullivan,\textsuperscript{1} Petricia Dulgheru\textsuperscript{1,3}, Idriss Atribak\textsuperscript{2}, Agustín Bueno-López\textsuperscript{2}, Avelina García-García,\textsuperscript{2}

\textsuperscript{1} UCD School of Chemistry and Chemical Biology, Belfield, Dublin 4, Ireland.
\textsuperscript{2} MCMA Group, Department of Inorganic Chemistry, University of Alicante, 03690, San Vicente, Alicante, Spain.
\textsuperscript{3} Chemical Physics of Materials, Université Libre de Bruxelles, Campus de la Plaine, CP 243, B-1050 Bruxelles, Belgium

Abstract.

\(	ext{Ce}_{x}\text{Zr}_{1-x}\text{O}_2\) catalysts with various Ce / Zr contents were studied using Raman spectroscopy under different gaseous atmospheres, at different temperatures and in the presence of a model soot. Catalysts with high concentrations of Zr fluoresced at elevated temperatures making analysis of their spectra impossible. This effect became even more pronounced at higher temperatures. \(	ext{CeO}_2\) and solid solutions with relatively low concentrations of Zr showed a red shift and a decrease in intensity of the characteristic \(\text{F}_{2g}\) peak at high temperatures under different atmospheres. The magnitude of the latter effect was higher under reducing atmospheres. These changes are reversible upon cooling, showing that they relate to a lattice expansion effect rather than any major chemical change to the material. In the presence of the model soot the Raman spectra of all materials was much decreased due to the absorption of the incident and scattered radiation by the soot particles. The presence of soot does not change the relative intensities or positions of the peaks in the spectra of the solid solutions. Evidence is shown for the production of a Ce\(^{3+}\)-CO species following interaction between the soot and the surface at high temperature in an inert atmosphere.

Keywords: \(\text{Ce}_x\text{Zr}_{1-x}\text{O}_2\), PM combustion, in-situ Raman Spectroscopy, 

Introduction

Because of the detrimental effects on human health as well as the global and local environment the remediation of Particulate Matter (PM) formed in diesel and gasoline combustion chambers is an area of considerable research activity both within our group and worldwide [1-10]. The use of catalysts to promote the oxidation of PM has resulted in much research in technologies such as the diesel particulate filter (DPF) [3-4] and the diesel particulate NO\(_x\) reduction systems (DPNR) [9-10].

The use of Ceria-Zirconia solid solutions, where Zr ions are incorporated into the fluorite \(	ext{CeO}_2\) structure, in promoting this reaction has been discussed by several authors [11-15]. The rationale behind the use of these materials in this reaction is related to the oxygen storage capacity of the catalysts. \(	ext{CeO}_2\) can desorb and re-adsorb \(\text{O}_2\) through formation and consumption of a \(\text{Ce}_2\text{O}_3\) oxide
The oxygen atoms involved in this process are active for promotion of various combustion and reforming reactions [21-25] and for this reason (as well as their ability to dampen excursions to overly fuel rich or fuel lean exhaust conditions) Ce\(^{x}\)Zr\(^{(1-x)}\)O\(_2\) catalysts are used as supports in many catalytic formulations.

The characterization of Ce\(^{x}\)Zr\(^{(1-x)}\)O\(_2\) solid solutions by Raman spectroscopy has been widespread [26-30]. The structure of CeO\(_2\) is face centered cubic (fluorite type). It can be described as 8 O\(^2-\) ions linked together by the edges, with the cerium ion at the centre of the oxygen cube. One of the triply degenerate F\(_{2g}\) modes (\(\sim 465\) cm\(^{-1}\)) can be viewed as a symmetric breathing vibrational mode of the O ions around each cation [26, 27]. This mode is, therefore, very sensitive to the oxygen sub-lattice disorder resulting from exposure to reducing / oxidising atmospheres. Furthermore, Raman spectroscopy has been used to monitor the formation of a Ce\(^{3+}\) species (in the presence of CO) following reduction of a portion of CeO\(_2\) [31 – 32].

The soot combustion activity and the room temperature Raman spectra of the materials we discuss here have previously been reported [33]. It has been shown that all these materials promote soot oxidation in air and NO / O\(_2\) (when reactivity profiles are compared to those of non-catalysed soot oxidations). The activity of the catalysts is inversely related to the Zr concentration. The room temperature Raman spectra of these materials has also previously been reported and it has been found that the F\(_{2g}\) peak shows a decrease in intensity and a change in position (towards higher energy) as the ZrO\(_2\) content within the solid solution is increased. This decrease is related to a change in the crystal structure which accompanies the decrease in the [Ce] (since this peak is related to a breathing mode of O\(^3-\) ions in the fluorite lattice).

In this work we describe the Raman spectra of a series of ceria zirconia mixed oxides in contact with a model Particulate Matter (Printex U [34]) under several gaseous atmospheres and following treatment at different temperatures. To our knowledge there have been no reports of a Raman spectroscopic study of the interaction between Ce\(^{x}\)Zr\(^{(1-x)}\)O\(_2\) mixed oxides and PM as a function of the gaseous atmosphere or the reaction temperature.

**Experimental**
Catalyst synthesis and Catalyst: Soot mixture preparation

CeO$_2$, ZrO$_2$ and mixed oxide (Ce$_x$Zr$_{1-x}$O$_2$) catalysts were prepared using a co-precipitation technique [33]. The catalysts were mixed in defined ratios (50 catalyst: 1 soot) with a model soot (Printex-U) to approximate a loose contact situation.

Raman Spectroscopy

Raman spectra were recorded using a Senterra III Raman Scope (Bruker) equipped with a Unilab II (Bruker) fibre optic probe. A diode laser ($\lambda = 785$ nm) operated at 25 mW was used as Raman excitation source and a germanium thermoelectrically cooled Charged Couple Device (Andorf) as detector. This probed the catalyst or catalyst soot mixture, (~200 mg) which was held in a CCR 1000 high temperature catalytic cell (Linkam). Use of the cell allowed control of the temperature and the gaseous atmosphere surrounding of the catalyst.

Thermogravimetric Analysis

TGA was carried out using a Q500 TGA (TA Instruments) thermogravimetric balance equipped with an Evolved Gas Analysis furnace. The EGA furnace was fitted with an exhaust port which allowed coupling of the TGA to a Quadrupole MS (HPR20, Hiden). This tandem TGA-MS system was able to monitor both the mass loss (as % mass loss, or derivative weight change profiles) and the composition of desorbed gases simultaneously. The catalyst soot mixtures were held in a flow of air (90 mL min$^{-1}$) and the temperature was ramped at a rate of 10 °C min$^{-1}$

Results and Discussion

Soot combustion activity and room temperature Raman spectra

The activity of these materials in promoting soot combustion (as measured using TGA rather than a fixed bed reactor) and their room temperature Raman spectra are essentially the same to those reported previously [33]. These are shown in the supporting information as Figure S1 and Figure S2.

Figure 1 shows the Raman spectra (200 – 2000 cm$^{-1}$) of the entire range of materials studied. Features in the Raman spectra of the materials between 1000 cm$^{-1}$ and 2000 cm$^{-1}$ have been ascribed to surface formats and carbonates [35-36]. These peaks become less pronounced as the ZrO$_2$ content increases.
The Raman spectra at energies around that of the $F_{2g}$ peak (between 200 cm$^{-1}$ and 650 cm$^{-1}$) also show evidence for the presence of defects once Zr is introduced into the lattice. These are shown in the inset of Figure 1 (a) where the spectra of the CeO$_2$ and Ce$_{0.8}$Zr$_{0.2}$O$_2$ materials are displayed. This has also been previously reported [26-27]. Relevant assignments of Zr-rich samples were also reported elsewhere [33].

**Effect of temperature and gaseous atmosphere on the $F_{2g}$ peak of CeO$_2$.**

Figure 2 shows the $F_{2g}$ peak of the CeO$_2$ sample recorded at room temperature, at 500 °C and again at room temperature under both a reducing H$_2$ atmosphere and an oxidizing O$_2$ atmosphere. Recall that this vibration relates to the symmetrical breathing mode of the O ions around the cation and its position and intensity are sensitive to the O concentration of the material.

The $F_{2g}$ peak of the sample held at room temperature in H$_2$ is slightly more intense than that in the equivalent spectrum recorded in O$_2$. It is clear that upon heating the material the $F_{2g}$ peak is shifted to lower energies (from $\sim$460 cm$^{-1}$ to $\sim$450 cm$^{-1}$) and its intensity is decreased. The latter effect is far more pronounced in the presence of H$_2$ than in an oxidizing atmosphere.

Upon re-cooling to room temperature the positions of the peaks returns to $\sim$460 cm$^{-1}$ and their intensity increases once more. The peak relating to the sample treated in O$_2$ is slightly more intense following the temperature cycle than it had been initially while the reverse is true of the sample which had been treated in H$_2$, *i.e.* the $F_{2g}$ peak is slightly less intense following the high temperature reduction than it had been initially.

The fact that the sample under O$_2$ at RT shows a less intense peak than the same sample under H$_2$ is puzzling. Taken at face value, this suggests that the addition of H$_2$ has increased the concentration of the cubic species responsible for the vibration, while the addition of O$_2$ has decreased this concentration. This is plainly not reasonable.
Therefore, we suggest that these relatively small intensity changes (and those noted once the materials are re-cooled following heat treatments) are within the error limits of the technique. In these experiments spectral intensity depends significantly on aspects such as the positioning of the fiber optic probe and homogeneity of the sample. Relatively small changes such as these can easily be explained by these experimental artefacts. However, the intensity changes noted upon heating (especially in the presence of \( \text{H}_2 \)) are far greater.

The individual sets of spectra showing the progression of this peak as a function of temperature in both \( \text{O}_2 \) and \( \text{H}_2 \) are shown in the supporting information (see figures S3 and S4).

The decreases in intensity at higher temperatures, coupled with the shift in the peak maxima under these conditions are important. The fact that these are, for the most part, reversible once the temperature is cooled back to room temperature suggest that they do not relate to any major chemical change taking place within the material and rather the changes are due to the thermal expansion of the fluorite lattice at 500 °C. Similar reversible effects have been noted previously with a \( \text{Ce}_{0.7}\text{Pr}_{0.3}\text{O}_2 \) sample [37] and with a range of \( \text{CeO}_{2-y} \) sub oxides [38]. They have also been noted in \( \text{V}_2\text{O}_5 \) systems [39] and have been explained by Bell and Iglesia [40]. Their explanation of these phenomena related to thermal expansion (peak position) and changes in the populations of vibrational energy levels (peak intensity / broadening) with increasing temperature.

Bell also points out that this affects the use of Raman spectroscopy for \textit{in-situ} or \textit{operando} techniques [41]. These results confirm that the position and intensity of these Raman peaks change as a function of temperature. This makes the observation of these peaks under different reaction conditions, and subsequent correlation of their behavior with any reactivity, \textit{i.e. operando} spectroscopy, fraught.

\textbf{Effect of ZrO\textsubscript{2} content on Raman spectra at higher temperature}

The presence of Zr has a further effect on the Raman spectra of the materials once the temperature is raised substantially above room temperature. In samples containing high concentrations of
Zirconia there is a large fluorescence peak seen at higher temperatures. The intensity of this fluorescence increases as a function of both [ZrO\textsubscript{2}] and T.

While it is possible that the use of a different excitation laser wavelength (from the 785 nm used in the current study) might ameliorate this effect unfortunately using our system it was not possible to determine whether this was the case. DRS spectroscopy has shown no absorbance by the materials at this wavelength. A certain amount of fluorescence (at lower T and lower [ZrO\textsubscript{2}]) can be corrected for using the Concave Rubber band treatment. Figure S5 shows this effect.

The two sets of data shown in figures 2 and discussed above (in which firstly temperature was shown to affect the position and intensity of the diagnostic F\textsubscript{2g} peak and secondly increased ZrO\textsubscript{2} content leads to fluorescence at higher temperatures) informed the future collection of spectra. Specifically, these outcomes have lead us to (a) collect spectra at room temperature following treatments in the isolated cell at higher temperatures and (b) concentrate our studies on samples containing lower ZrO\textsubscript{2} content.

**Effect of PM on Raman spectrum**

Figure 3 shows the Raman spectrum of Printex-U. Two broad vibrations can be observed: a very strong Raman vibration in the 1190 cm\textsuperscript{-1} – 1472 cm\textsuperscript{-1} region centred at 1330 cm\textsuperscript{-1}, and another broad vibration mode in the 1600 cm\textsuperscript{-1} – 1963 cm\textsuperscript{-1} region centred at 1695 cm\textsuperscript{-1}. The spectrum also shows a small peak in the 1472 cm\textsuperscript{-1} – 1600 cm\textsuperscript{-1} region with a maximum appearing at 1550 cm\textsuperscript{-1}.

Sadezky et al. [42] have recorded the Raman spectra of pure graphite and observed a very strong band at 1550 cm\textsuperscript{-1} (the G band) and have attributed it to a pure graphite lattice vibration with E\textsubscript{2g} symmetry. A less intense band at 1360 cm\textsuperscript{-1} (referred to as the D1 band) has been ascribed to defects in the graphite lattice. Another weak band (the D2 band) has been observed in pure graphite at 1620 cm\textsuperscript{-1} (and was also ascribed to defects in the graphite lattice).

Other workers [43-47] have studied different types of synthetic carbonaceous PM and graphitic materials and concluded that the Raman spectra of carbonaceous particulate matter can be interpreted in terms of highly disordered graphitic structures. According to these researchers a band at approximately 1360 cm\textsuperscript{-1} is very specific for disordered carbon samples (the transition is forbidden in pure graphite) and a vibration appearing at approximately 1580 cm\textsuperscript{-1} can be attributed to an ideal graphite lattice vibration with E\textsubscript{2g} symmetry.
Printex-U displays all these vibration modes in the 1000 cm$^{-1}$ – 1800 cm$^{-1}$ region making the deconvolution between surface carbonate species, which are always present on these oxide surfaces (see Figure 1) in the presence of CO$_2$, and Printex-U very difficult (see above).

Figure 4 compares the Raman spectra of Printex-U with those of CeO$_2$ and a CeO$_2$ / Printex-U mixture. The mixed sample was black, in contrast with the light yellow CeO$_2$. In Figure 4, in order to scale the spectra onto one plot, the spectrum for CeO$_2$ was reduced by a factor of 25 while that for Printex-U was reduced by a factor of 5. This scaling is required because the intensity of the Raman signal is very much decreased in the presence of PM. The addition of relatively small (2% of the mixture by mass) amounts of soot has a dramatic effect on the intensities of the spectra collected, i.e. the Raman Effect is much decreased in presence of Printex-U.

According to Li et al.[48], when the excitation laser and scattered light are strongly absorbed by the sample, smaller amounts of scattered light escapes and therefore the observed Raman signal is weakened. This effect can be further seen in Figure 6 where the Raman spectra of two of the samples in the presence and absence of PM are shown. The Raman spectrum in the presence of PM is considerably weaker than when the pure sample is analysed.

The spectra of the oxide – soot samples also lack features relating to the G and D bands of the PM (see above). It should be noted that these samples are mixtures in which the PM concentration is diluted by a factor of 50 compared to the pure Printex U sample shown in Figure 3.

The fact that clear bands relating to Printex-U are not observable at these low concentrations, coupled with the fact that increasing the Printex-U concentration in the mixture further quenches the Raman spectra of the oxides of interest adds a further difficulty to attempts to observe soot combustion in-situ using Raman spectroscopy. However, the removal of soot through oxidation can be followed indirectly using this effect (see below).

Apart from the aforementioned decrease in Raman intensity associated with the absorption of laser and scattered light by the PM, almost no differences can be observed when comparing the Raman spectra shown in Figure 5. The overall decrease in intensity applies to the entire spectrum. This makes less intense peaks appear significantly smaller and the smaller peaks seem to disappear. These weaker peaks are removed but the position and relative intensities of the major carbonate related peaks and the F$_{2g}$ peak remains the same in the presence and absence of soot – suggesting there is no redox reaction interaction between the two components which leads to an observable (using Raman spectroscopy) removal of oxygen from the fluorite phase.
Raman Spectra confirming PM oxidation.

Figure 6 shows the effect on the intensity of the Raman spectra of treating a Ce$_{0.8}$Zr$_{0.2}$O$_2$ / Printex-U mixture in an oxidizing environment (NO / O$_2$) at elevated temperatures. Because of the fluorescence problems at higher temperatures (see Figure S5) these spectra are recorded at room temperature before and after the high temperature treatment. It is clear that, while the relative intensities and positions of the individual peaks in each spectrum are unchanged following the treatment, the overall intensity of the spectra is significantly increased following the excursion to higher temperatures in NO / O$_2$. It should be noted that the final intensity is still approximately 8 times lower than it had been in the absence of PM (Figure 5). We ascribe the increase in intensity to the oxidation of PM at the temperatures of the NO / O$_2$ treatment and the subsequent lessening of the quenching effects seen above, i.e. there is less PM in the system to absorb the incident and scattered radiation.

Formation of Ce$^{3+}$ – CO species from reaction between Ce$_x$Zr$_{1-x}$O$_2$ and PM

Figure 7(a) shows the TGA profile of a sample of Ce$_{0.8}$Zr$_{0.2}$O$_2$ – PM mixture which was held in a He flow at 500 °C for 250 min. The mass spectrometer profiles for O$_2$ (m / e = 32) and CO$_2$ (m / e = 44) evolved during the experiment are also shown. During the 250 min of the experiment approximately 1.6% of the mass of the mixture was lost. This mass loss was accompanied by the removal of CO$_2$ and O$_2$ from the system. The CO$_2$ profile shows maximum production of CO$_2$ at the beginning of the experiment followed by an exponential decrease. This was mirrored by the production of O$_2$ which began at a relatively low level and subsequently increased throughout the course of the experiment.

These observations can be explained as being due to desorption of O$_2$ from the ceria containing materials (and the concomitant formation of Ce$^{3+}$) [16-23]. Initially this O$_2$ (rather than leaving the system) reacted with the PM and formed CO$_2$. As the concentration of proximate PM in the system decreased the volumes of CO$_2$ produced decreased and the amount of O$_2$ leaving the system un-reacted increased.

Figure 7(b) shows a series of normalized Raman spectra (where they were normalized using the intensity of the F$_{2g}$ peak (see Figure S6)) showing essentially the same experiment. The spectra show a Ce$_{0.8}$Zr$_{0.2}$O$_2$ catalyst admixed with soot, the same sample following a 4 h treatment in He and again following a 30 h treatment in He. In Figure 7(b) the development of a peak at approximately 2100 cm$^{-1}$ is clearly visible. This peak is not present in oxidized Ce$_{0.8}$Zr$_{0.2}$O$_2$. Peaks in this spectral region have previously been noted by several authors [31-32, 49-50] and are specific to reduced CeO$_2$ samples. Swanson and co-workers have assigned the species as a Ce$^{3+}$ – CO vibration. This suggests that either atomic oxygen on the surface of the Ce$_{0.8}$Zr$_{0.2}$O$_2$ catalyst reacted with PM, or that a desorbed O$_2$ reacted with PM to form CO or CO$_2$ which subsequently adsorbed on the formed Ce$^{3+}$ sites. The re-adsorption of carbon-containing gaseous products on the catalyst leads to the exchange
of oxygen between the adsorbed gases and the catalyst, as previously demonstrated [51]. The Ce\(^{3+}\)-CO species observed by Raman spectroscopy may be reaction intermediates of these oxygen exchange processes.

In either case this appears to be evidence for the reaction between the Ce\(_{0.8}\)Zr\(_{0.2}\)O\(_2\) catalyst and the PM under inert gas conditions. This conclusion is consistent with previous studies of the ceria-catalysed soot combustion mechanism carried out using isotopically labeled oxygen in a TAP reactor [52]. It was concluded that the driving force for the pure ceria-catalysed soot combustion is O\(_2\) uptake by ceria, which destabilizes surface oxygen entities delivering active oxygen species from the ceria to soot. On the contrary, in La\(^{3+}\)-doped ceria catalysts (which have improved redox properties relative to ceria), the oxygen in the lattice itself oxidizes soot and subsequently gas phase oxygen fills the lattice vacancies.

On the other hand changes in the shape and position of the F\(_{2g}\) peak following these treatments are not seen (see figure S6). If significant proportions of the Ce\(^{4+}\) content of the material had been reduced to Ce\(^{3+}\) (through loss of O\(_2\) from the fluorite lattice) we might have expected this peak to shift (as was the case in Figure S2).

**Conclusions**

Collecting Raman spectra from these mixed oxides in the presence of PM under reaction conditions is complicated by several effects.

Firstly, under conditions where the samples are irradiated with a 785 nm laser, the Zr-containing materials fluoresce at high temperatures. The extent of this fluorescence is correlated with the concentration of ZrO\(_2\) within the materials. Secondly, in the absence of soot at high temperature there is also a change in the position and intensity of the characteristic F\(_{2g}\) band. This change involves a movement of the peak to lower energies (due to thermal expansion of the absorbing particle) and a decrease in the peak intensity (related to a broadening of the peak due to the occupancy of significant numbers of vibrational sub-levels at these temperatures). These changes were more pronounced when the gaseous atmosphere was reducing (H\(_2\)) but were also present when the atmosphere was oxidizing (O\(_2\)). These peak shifts were reversible upon cooling the samples in the reactive atmospheres.

Thirdly, the Raman Effect is much decreased in the presence of soot due to the absorption of both the incident laser light and the scattered signal. This results in far less intense spectra in the presence of soot and also in a changing spectral intensity during reaction as soot was removed from the in-situ cell.
Finally, the production of surface carbonates as a result of soot combustion is masked by the bands due to the soot itself and it is not possible to de-convolute these signals.

These observations led us to collect all subsequent spectra at room temperature following whatever treatments (temperature / gaseous environment) the catalysts (or catalyst soot mixtures) were subjected to.

A Ce$_{0.8}$Zr$_{0.2}$O$_2$ catalyst in contact with Printex-U developed Raman bands relating to a Ce$^{3+}$–CO species following an extended treatment in inert gas at 500 °C. This suggests that the catalyst material was able to transfer an O atom from the catalyst surface to the soot, forming CO or CO$_2$, portions of which subsequently adsorbed onto the reduced catalyst.

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**References**


Figure 1. The 200-2000 cm\(^{-1}\) region of the Raman spectra of Ce\(_x\)Zr\(_{(1-x)}\)O\(_2\). Plot (a) compares the spectra of CeO\(_2\) (lower) and Ce\(_{0.8}\)Zr\(_{0.2}\)O\(_2\) (upper), plot (b) compares the spectra of Ce\(_{0.6}\)Zr\(_{0.4}\)O\(_2\) (lower) and Ce\(_{0.4}\)Zr\(_{0.6}\)O\(_2\) (upper) and plot (c) compares the spectra of Ce\(_{0.2}\)Zr\(_{0.8}\)O\(_2\) (lower) and ZrO\(_2\) (upper). The inset of plot (a) shows the region around the F\(_{2g}\) peak.)
Figure 2. $F_{2g}$ peak of CeO$_2$ recorded at room temperature (□, ■), at 500 °C (△, ▲) and again at room temperature (○, ●) in a H$_2$ atmosphere (filled symbols) and an O$_2$ atmosphere (empty symbols). Spectra have been corrected for fluorescence using the Concave Rubber Band function.
Figure 3. Raman spectrum of Printex U, a model soot.
Figure 4. The 1000 cm\(^{-1}\) – 2000 cm\(^{-1}\) region of the Raman spectra of CeO\(_2\) (\(\bigcirc\)), CeO\(_2\)-U (\(\star\)) and Printex U (\(\square\)). Inset shows the 400 – 500 cm\(^{-1}\) region (F\(_{2g}\) peak) from the same spectra. For clarity, profiles were displaced and the CeO\(_2\) and Printex U spectra were reduced by factors of 25 and 5 respectively.
Figure 5. Raman Spectra of CeO$_2$ (a) and Ce$_{0.8}$Zr$_{0.2}$O$_2$ (b) in the presence (lower plots) and absence (upper plots) of Printex U. The spectra in the presence of soot have been increased by factors of 5 and 20 respectively.
Figure 6. Raman spectra of a Ce$_{0.8}$Zr$_{0.2}$O$_2$ catalyst soot mixture before (lower profile) and after (upper profile) a treatment in NO / O$_2$ at 500 °C. Spectra were recorded at room temperature.
Figure 7 (a) TGA and evolved O$_2$ and CO$_2$ profiles recorded during an experiment where a Ce$_{0.8}$Zr$_{0.2}$O$_2$ – PM mixture was subjected to a flow of He at 500 °C.

Figure 7(b) Normalised Raman spectra of a fresh sample of CeZ20 - U fresh (○), and exposed to an inert atmosphere at 500 °C for 4 hours (☆) for 30 hours (△).
Figure S1

Figure S1 Activity of a selection of the different solid solutions in promoting soot combustion in Air (lower plots) and NO/O₂ (upper plots) as measured using TGA. Activity in each case is proportional to [Ce] in the catalyst. The non-catalysed combustion is shown in each case as a dotted line.
Figure S2. Room temperature Raman spectra showing the variation in the F$_{2g}$ peak as [Zr] in the solid solution changes. CeO$_2$ (□), Ce$_{0.8}$Zr$_{0.2}$O$_2$ (○), Ce$_{0.6}$Zr$_{0.4}$O$_2$ (◇), Ce$_{0.4}$Zr$_{0.6}$O$_2$ (●), Ce$_{0.2}$Zr$_{0.8}$O$_2$ (△) and ZrO$_2$ (■)
Figure S3. Showing 400 - 500 cm\(^{-1}\) region of the Raman spectra of CeO\(_2\) recorded at: RT (□), 100 °C (○), 200 °C (▲), 300 °C (△) 400 °C (◇), and 500 °C (■) while being exposed to 10 mL min\(^{-1}\) H\(_2\).
Figure S4. Showing 400 - 500 cm\(^{-1}\) region of the Raman spectra of CeO\(_2\) recorded at: RT (□), 100 °C (○), 200 °C (★), 300 °C (△) 400 °C (◇), and 500 °C (■) while being exposed to 10 mL min\(^{-1}\) O\(_2\).
Figure S5

Under the spectral collection conditions used on our work samples containing higher concentrations of Zr fluoresce (see figure S5 (a) and 3(c)). A certain amount of this fluorescence can be corrected using concave rubber band correction (see figure S5(b)). However, at the highest loadings of Zr within the lattices concave rubber band correction is no longer effective (see figure S5(d)).

Figure S5. Raman Spectra recorded during exposure of Ce$_{0.4}$Zr$_{0.6}$O$_2$–PM mixtures (plots (a) and (b)) and Ce$_{0.2}$Zr$_{0.8}$O$_2$–PM mixtures (plots (c) and (d)) to NO/O$_2$ between RT and 500 °C. RT ($\square$), 100 °C ($\bigcirc$), 200 °C ($\star$), 300 °C ($\triangle$), 400 °C ($\Diamond$), 500 °C ($\bullet$). Plots (a) and (c) show the raw spectra while plots (b) and (d) show the effect of concave rubber band correction on the spectra.
Figure S6. Normalised $F_{2g}$ region of the Raman spectra of a Ce$_{0.8}$Zr$_{0.2}$O$_2$ sample following the treatments shown in Figure 7. Fresh Ce$_{0.8}$Zr$_{0.2}$O$_2$ - PM (○), Ce$_{0.8}$Zr$_{0.2}$O$_2$ - PM exposed to an inert atmosphere for 4 hours at 500 °C (☆) and Ce$_{0.8}$Zr$_{0.2}$O$_2$ - PM exposed to an inert atmosphere for 30 hours at 500 °C (△).