XPD and XANES Studies of Ce$_{0.9}$Zr$_{0.1}$O$_2$ Nanocatalysts under Redox and Catalytic CH$_4$ Oxidation Conditions

María G. Zimicz, Fernando D. Prado, Analía L. Soldati, Diego G. Lamas, and Susana A. Larrondo

ABSTRACT: The aim of this work is to take further insight into the structural stability of cerium–zirconium catalysts under reducing, oxidizing and reaction conditions. In situ synchrotron based techniques (XANES and XPD) were used in order to determine the stability of crystal structure and the oxidation state of cerium cations in the reaction conditions prevailing in the catalytic studies. In situ XPD studies in 5 mol % H$_2$ atmosphere revealed that no structural changes occur until 870 °C. At this temperature, the sample synthesized with glycine, which presents lower crystallite agglomeration, segregated a small quantity of a reduced phase. On the contrary, the solid synthesized with lysine, with more agglomerated crystallites, does not show structural changes in all the temperature range. Reoxidation treatments in 5 mol % O$_2$ revealed that at 750 °C the segregated phase disappears and that the original cubic structure is restored. In situ XANES studies in the Ce L$_{III}$ absorption edge indicate that under catalytic reaction conditions, the degree of reduction of Ce$^{4+}$ is low, allowing the occurrence of methane oxidation. The solid is capable to deliver the oxygen of its structure when no oxygen is fed into the reactor until the 50% of cerium cations become reduced, triggering the deactivation process. Therefore, it is clear for these studies that the Ce$^{4+}$ to Ce$^{3+}$ ratio in the lattice is governing the catalytic behavior of the solid. The catalytic results collected during in situ XANES experiments are in excellent agreement with our previous catalytic studies performed with a laboratory fixed bed reactor.

1. INTRODUCTION

ZrO$_2$–CeO$_2$ mixed oxides have been widely proposed as catalysts, supports, and anode materials for intermediate-temperature solid oxide fuel cells (IT-SOFC). The elucidation of structural and chemical changes under operating conditions has become an issue of great interest. Ceria-based catalysts attracted great attention due to ceria high reducibility and oxygen storage capacity.$^{1-3}$ In a reducing environment, cerium dioxide (CeO$_2$) releases oxygen to become reduced ceria (CeO$_{2-x}$), while in oxidizing conditions this reduced ceria can reincorporate oxygen into the crystal structure recomposing back to the original cubic structure stoichiometry. This feature is usually called the oxygen storage capacity (OSC) and relays over the reversible redox process associated with the Ce$^{4+}$/Ce$^{3+}$ redox couple. However, in ceria-based materials this reduction process from Ce$^{4+}$ to Ce$^{3+}$ could lead to phase segregation, depending on the doping cation, the particle size, the specific surface, the temperature, and the surrounding atmosphere. In the case of CeO$_2$–ZrO$_2$ mixed oxides, the structural studies found in the literature are limited mostly to the region corresponding to intermediate composition (Ce$_{0.5}$Zr$_{0.5}$O$_2$). Otsuka-Yao-Matsuo et al.$^4$ and Montini et al.$^5$ reported that after reducing samples of Ce$_{0.5}$Zr$_{0.5}$O$_2$ with tetragonal structure (t$'$), a solid with the pyrochlore structure (Ce$_2$Zr$_2$O$_7$) was obtained. Vidal et al.$^6$ analyzed the influence of redox treatments on Ce$_2$Zr$_{1-x}$O$_2$ with low and high specific surface areas. They found that in Zr- and Ce-rich samples no modifications were detected, but for intermediate compositions appreciable phase segregation occurred. This phase segregation was apparently favored by the presintering of the mixed oxides. Rodriguez et al.$^7$ analyzed the physical and chemical properties of Ce$_2$Zr$_{1-x}$O$_2$ (x = 0.9, 0.66, and 0.5) oxides and reported that no phase segregation occurred when the samples were heated in 5 mol % H$_2$ in He balance from room temperature up to 1000 °C. Similar behavior was observed by Boaro et al.$^8$ who found that a sample of Ce$_{0.8}$Zr$_{0.2}$O$_2$ preserves the tetragonal

Received: June 2, 2015
Revised: July 20, 2015
DOI: 10.1021/acs.jpcc.5b05253
J. Phys. Chem. C XXXX, XXX, XXX–XXX
structure after one redox cycle, evidencing only a distortion of the \( t^* \) phase structure, probably related to the promotion of reducibility of this composition.

The reduction of particle size to the nanometric level has put in evidence its influence in the structural, electronic, and chemical properties of the solids. Colon and co-workers also studied the stability of CeO\(_2\)–ZrO\(_2\) mixed oxides during oxidation processes. They suggested that the driving force for phase segregation during these treatments is the existence of a critical particle size above which the mixed oxide tends to decompose. Crystal growth would take place until a certain critical mean crystal size is reached, when phase segregation would begin forming thermodynamically more stable phases. The authors found that for Ce\(_0.9\)Zr\(_0.1\)O\(_2\) samples, the critical size is around 15 nm, but it changes with sample composition. They suggested that for Ce-rich samples, the critical size would be higher.

In the Ce-rich region of the CeO\(_2\)–ZrO\(_2\) system, it was reported that Ce\(_0.9\)Zr\(_0.1\)O\(_2\) mixed oxide did not evidence any changes of Ce\(_0.9\)Zr\(_0.1\)O\(_2\) samples when the samples were subjected to different atmospheres (5 mol % H\(_2\)/He balance, 5 mol % O\(_2\)/He balance) was synthesized by the stoichiometric α-amino acid/nitrate combustion route, starting from ZrO(NO\(_3\))\(_2\) and Ce(NO\(_3\))\(_3\)·6H\(_2\)O. Glycine and lysine were used as fuels, and the obtained samples are referred to as GS and LS, respectively. After the synthesis process, the mixed oxides were calcined at 600 °C in air in order to eliminate any vestige of carbonaceous residues. The details of the synthesis process and the influence of the synthesis conditions on the textural and morphological properties of the samples were previously reported.

### 2. EXPERIMENTAL SECTION

#### 2.1. Synthesis of Nanopowders. Ce\(_{0.9}\)Zr\(_{0.1}\)O\(_2\) mixed oxide was synthesized by the stoichiometric α-amino acid/nitrate combustion route, starting from ZrO(NO\(_3\))\(_2\)·6H\(_2\)O and Ce(NO\(_3\))\(_3\)·6H\(_2\)O. Glycine and lysine were used as fuels, and the obtained samples were referred to as GS and LS, respectively. After the synthesis process, the mixed oxides were calcined at 600 °C in air in order to eliminate any vestige of carbonaceous residues. The details of the synthesis process and the influence of the synthesis conditions on the textural and morphological properties of the samples were previously reported.

#### 2.2. Characterization. The texture of fresh catalysts was analyzed by N\(_2\) physisorption in a Quantachrome Autosorb-1 equipment, obtaining the specific surface area and total pore volume of the solids. The size and morphology of the nanopowders were analyzed using a Philips CM 200 UT microscope equipped with an ultratwin objective lens. The electron source used was a LaB\(_6\) filament operated at 200 keV. The nominal resolution was 0.2 nm for high resolution mode. The microscope was equipped with a CCD camera for digital acquisition; contrast and illumination were adjusted linearly afterward using commercially available image treatment programs. Statistics on the particles was done with the programs ITEM and Origin 8.5. Particle size is given as the mathematical average standard deviation of the largest particle dimension. In all cases more around 200 measurements were averaged, sampling in different regions of the foil.

The crystal structure of the samples under different atmospheres (5 mol % H\(_2\)/He balance, 5 mol % O\(_2\)/He balance, and 2 mol % CH\(_4\)/4 mol % O\(_2\)/He balance) was determined by in situ X-ray powder diffraction (XPD) technique using synchrotron radiation. The time-resolved diffraction data were collected at the D10B-XPD beamline of the Brazilian Synchrotron Light Laboratory (LNLS, Campinas, Brazil) using a high-intensity and low-resolution configuration, without crystal analyzer. The energy was set at 7997.43 eV and the X-ray wavelength was 1.55031 Å. The peak profile was not significantly affected by low resolution since the peaks are intrinsically broad. The sample powders were placed on the sample-holder inside an oven with controlled temperature. Rietveld refinements were performed with FullProfuite open access software.

In situ XANES studies were performed for the Ce L\(_{III}\)-edge in which Ce 2p electrons are excited with photons of energies above 5720 eV. These tests were carried out in the D06A-DXAS beamline of the Brazilian Synchrotron Light Laboratory. Sample preparation and method of analysis of XANES spectra were previously reported. From the fitting of the absorption profile, the evolution of the degree of reduction “\( \alpha \)” of Ce\(_{x}\)O\(_{1-x}\) cations when the samples were subjected to different atmospheres was obtained. The degree of reduction is defined as the Ce\(_{3+}\) to (Ce\(_{3+}\) + Ce\(_{4+}\)) atomic ratio.

During the in situ XPD and in situ XANES analyses, the inlet line of the reactor was connected to a gas-mixing station provided with mass flow controllers, and the outlet line was connected to a Pfeiffer QMS 422 Quadrupole Mass Spectrometer.

The catalytic activity experiments in total oxidation of methane of fresh solids were performed in a fixed bed reactor, laboratory scale, operated at atmospheric pressure. The experiments were performed in the 400–800 °C temperature range, with steps of 100 °C. The feed to the reactor was a mixture of 2 mol % CH\(_4\)/4 mol % O\(_2\) and N\(_2\) balance with a total volume flow of 333 STP cm\(^3\) min\(^{-1}\). These studies were previously reported and are presented here for comparison purposes.

### 3. RESULTS AND DISCUSSION

Results from N\(_2\) physisorption experiments are listed in Table 1. It is possible to see that the α-amino acid used in the synthesis process influences the textural characteristics of the nanopowders. Sample GS exhibits higher specific surface area and pore volume and lower degree of agglomeration (\( D_{\text{BET}}/D_{\text{XRD}} \)) than sample LS.

#### 3.1. TEM. Figure 1 shows the TEM images obtained for samples GS and LS, and the corresponding histograms for particle size distribution.

Sample GS shows a broader particle size distribution around the mean grain size of 14.7 ± 4.8 nm. This value is much higher than the determined by XPD technique. In the case of sample LS, the grain size distribution is narrower around a mean size of 7.6 ± 3.1 nm, being this value compatible with that obtained from XPD experiments. Besides, sample LS looks very agglomerated.

#### 3.2. In Situ XPD Experiments. Fresh samples exhibited cubic crystal structure corresponding to the \( Fm\overline{3}m \) space group.
The average crystallite size was calculated from the width of the (111) Bragg peak using the Scherrer equation, \( D = \frac{0.9 \lambda}{\beta \cos(\theta)} \), where \( D \) is the crystallite size, \( \lambda \) is the radiation wavelength, \( \beta \) is the peak width at half-maximum intensity, and \( \theta \) is the Bragg angle. No influence of the \( \alpha \)-amino acid used in the synthesis on the crystallite size (\( D_{XRD} \)) was observed.

The solids were heated with a heating ramp of 10 °C·min\(^{-1}\) under a reducing flow (5 mol % H\(_2\)/He balance, 50 STP cm\(^3\)·min\(^{-1}\)) from room temperature up to 900 °C with isothermal steps at 25, 300, 450, 600, 750, and 900 °C. The diffraction patterns were collected in the 2\( \theta \) = 20 – 80° angular range during the isothermal steps. In Figure 2 the cell parameters and the crystallite size evolution during the reduction process are summarized. Cell parameters for fresh samples are in agreement with values reported in the literature.\(^{15-17}\) As it was said in previous sections, mean grain size for fresh LS sample is consistent with results obtained from TEM. However, for sample GS, particle size from TEM is higher than that obtained by XPD. This discrepancy can be attributed to microstrain, which produces a greater broadening in the diffraction peak and then causing a relatively smaller estimation of the crystallite size than the observed in TEM micrographs.

As temperature is increased, increments in cell parameter and crystallite size are observed, in agreement with results reported for CeO\(_2\).\(^{18}\) The increment in the cell parameter is related to the expansion of the unit cell induced by the partial reduction of Ce\(^{4+}\) to Ce\(^{3+}\), which according to Shannon and Prewitt\(^{19}\) have cationic radii, for the 8-fold coordination, of 0.97 Å (Ce\(^{4+}\)) and 1.14 Å (Ce\(^{3+}\)), respectively. The initial average crystallite sizes are in the 5–6 nm range. Around 750 °C the size starts to increase reaching a value in the 6–7 nm range at 900 °C, with an increment of nearly 25% for sample GS and above 50% for sample LS. This different behavior is a consequence of the different agglomeration ratio of the two powders. Sample LS has a much higher agglomeration ratio that favors the development of sintering necks and crystallite growth. As it was previously reported, the agglomeration ratio is strongly influenced by the \( \alpha \)-amino acid used in synthesis, which governs the speed of the gel autoignition.\(^{11}\) It is important to address that the size growth of the crystallites is much lower than observed by other authors, who reported increments reaching 300% at 900 °C for the same composition but different synthesis methods, pointing out the higher thermal stability of samples synthesized by gel-combustion method.\(^{20}\)

The Journal of Physical Chemistry C

Figure 1. TEM images with two magnifications and particle size distribution for fresh samples (a) GS and (b) LS.

Figure 2. Variation of structural parameters with temperature in reducing atmosphere: (a) variation of \( D_{XRD} \) and (b) variation of cell parameter “a”.

DOI: 10.1021/acs.jpcc.5b05253
J. Phys. Chem. C XXXX, XXX, XXX--XXX
From the diffraction patterns collected on heating in reducing atmosphere, it is clear that sample LS retains the cubic structure in all the temperature range (Figure 3).

However, sample GS evidences phase segregation at 900 °C. At this temperature, a new "reduced" phase coexists with the original cubic structure. Perrichon et al. reported that when CeO₂ is reduced at temperatures above 800 °C, an expanded Ce₂O₃₋ₓ phase and cubic and hexagonal Ce₂O₃ phases are formed. Bernal et al. observed segregation of Ce₂O₃ from CeO₂ during reduction at 1000 °C, addressed similar results. In our case, due to the low intensity and resolution of the peaks assigned to the "reduced" phase, it was not possible to determine its structure and composition with certainty, but they likely correspond to a reduced CeO₂₋ₓ phase. Even when well-crystallized phases are present, it is difficult to establish the structure with certainty. Boaro et al. investigated the structural properties of Ce₀.₅Zr₀.₅O₂ mixed oxide, before and after redox treatments. By XRD and Raman analysis they found that the original symmetry of the t₄ phase is lost upon reduction at 1273 K and a new phase is being formed. They suggested that there is a coexistence of tetragonal and pseudopyrochlore phases. The pyrochlore structure in ceria–zirconia mixed oxides is usually reported to form at high temperature under reducing conditions.

In order to determine the temperature of segregation of the new phase during reduction treatment, shorter diffraction patterns were collected for sample GS in the 2θ = 26–36° region, where the main peak of the segregated phase appeared. The data collection was fast enough to ensure a temperature change below 10 °C during each pattern collection. The obtained results are plotted in Figure 4. It is clear that the reduced phase appears in the 885–895 °C range.

To determine whether the process was reversible or not, a reoxidation treatment was performed afterward. The sample GS was cooled down to room temperature in the former reducing atmosphere, and then was heated up to 900 °C with the same temperature program used in the reduction process, in an oxidizing flow of 5 mol % O₂/He balance (50 STP cm⁻³·min⁻¹). The diffraction patterns were collected at the intermediate isothermal steps. In Figure S it is possible to observe that the reduced phase persists at temperatures below 750 °C, while at this temperature the reduced phase disappeared and the original cubic phase was recovered. In a previous in situ XANES study of reduction and reoxidation processes of sample GS, we found that the complete reoxidation of Ce³⁺ cations is reached at 750 °C.13 This result is pointing out that the segregated phase disappears when cerium cations are totally reoxidized, confirming that the segregated phase is a reduced ceria phase. In our previous paper we have also reported that even though samples GS and LS have the same composition and crystal structure, the reduction and reoxidation processes of these samples are very different, suggesting the influence of crystallite size and agglomeration grade on them. At 800 °C, sample GS reached a 34% of reduction, while the sample LS suffered a reduction of 39%. From XPD studies it is clear that no phase segregation takes place in both samples until 800 °C. However, structural stability of sample LS seems to be greater since there is no segregation of secondary phases up to 900 °C. The different behavior observed between the two samples may be due to the higher degree of agglomeration of the crystallites in the sample LS, allowing a faster rate of crystallite growth (Figure 1) and an easier accommodation within the structure of the Ce³⁺ cations generated in the reduction process.

Finally, in situ XPD and XANES studies were also performed in operating conditions (gas flow, feed composition, temperature, etc.) similar to those used in previous catalytic experiments performed in our laboratory with a fixed-bed reactor. In this case, the samples were heated at 10 °C/min from room temperature up to 750 °C, with isothermal steps at 25, 450, 600, and 750 °C, in a mixture of 2 mol % CH₄ and 4 mol % O₂ (He balance). Diffraction patterns were collected during the isothermal periods. No structural changes were detected in sample LS. Sample GS also retained the cubic structure, but an additional peak centered at 26.5° appeared during the isothermal period.

Carbon formation can occur both by decomposition of methane (eq 1) and by the Boudouard reaction (eq 2). It is important to remark that eq 1 is highly endothermic and then favored at high temperature. On the contrary, Boudouard reaction is exothermic and carbon formation is favored at temperatures below 550 °C.

\[ \text{CH}_4 \leftrightarrow \text{C} + 2\text{H}_2 \]  

(1)
From mass spectrometry data collected during the in situ XPD studies (Figure 7), it can be observed that oxidation products are detected from 400 °C, approximately. The signal corresponding to carbon monoxide starts to appear at 450 °C, and its signal increases with temperature increment. Therefore, the carbon deposits could be possibly produced through eq 2.

3.3. In Situ XANES Experiments. In situ XANES experiments were performed at the Ce LIII-edge (see Figure 8). A temperature ramp of 10 °C/min and a feed consistent of 2 mol % CH₄ and 4 mol % O₂ diluted in He were used. XANES spectra were collected from room temperature up to 800 °C. After 30 min at 800 °C, oxygen was removed from the feed to the reactor, and for 30 min XANES spectra were collected in this atmosphere of 10 mol % CH₄/He. XANES spectra were fitted as a linear combination of standard spectra, as it was previously reported. 13,18 XANES experiments in methane oxidation conditions were conducted only for sample GS, due to the poor catalytic activity of sample LS previously observed in conventional catalytic tests.14 In Figure 9a it is shown the degree of reduction versus time on stream corresponding to these experiments, and in Figure 9b, mass spectrometry data collected during the whole experiment are plotted.

During reaction in the feed containing CH₄ and O₂ the solid showed a very low degree of reduction reaching a steady state with a percentage of reduction, “α”, of 5%. In this situation, a dynamic equilibrium between reduction and reoxidation of
Ce⁴⁺ cations was achieved, which allows the occurrence of methane total oxidation. Spectrometry data (Figure 9b) showed the continuous decrease of CH₄ and O₂ signals, evidencing the catalytic activity of the solid for methane oxidation. Total oxidation products (CO₂ and H₂O) are prevalent in the low temperature range. At approximately 650 °C a small quantity of CO is detected, indicating that methane partial oxidation is a secondary reaction.

Our previous catalytic results (Figure 10) obtained in our laboratory with a fixed bed reactor have shown an excellent performance, reaching 80% of methane conversion at 700 °C in a feed composition consisting of 2 mol % CH₄ and 4 mol % O₂ (N₂ balance), being the methane total oxidation the prevalent reaction in all the temperature range.

The only carbonated product detected in the effluent gas was carbon dioxide (CO₂). Probably carbon monoxide (CO) was not detected due to the lower sensitivity of the gas-chromatography technique compared with mass spectrometry. Besides, carbon balance between feed and effluent was in the 100% ± 5% range with no observation of traces of carbon deposits in the scanning electron microscopy micrographs of the spent catalysts.

As it was mentioned above, mass spectrometry data collected during in situ XPD studies reveals that methane total oxidation starts at 400 °C, and methane partial oxidation at 450 °C, approximately. These temperatures are lower than those observed during in situ XANES analyses due to differences in sample preparation. In XPD studies, the samples are in powder form, with a good contact between gas-phase and active sites leading to a greater contact time. In XANES analyses the samples are prepared by mixing the powders with boron nitride and pressed into self-supporting discs, so the sample is diluted leading to a lower contact time, and active sites are less accessible than in XPD experiments.

After stabilizing the temperature at 800 °C, the oxygen was removed from the feed, and XANES spectra were collected...
delivered oxygen from its structure, suffering a consequent reduction. This behavior is maintained until the solid reaches a maximum level of reduction and it deactivates. From mass spectrometry data, it can be observed that in the first minutes without oxygen, the oxygen delivered from the solid is enough to produce the total oxidation of methane. As sample reduction continues, the capability of delivering oxygen reduces, and the signal of total oxidation products (CO₂ and H₂O) starts to decrease, while carbon monoxide signal becomes predominant. The other important conclusion is that this capability remains until 50% of Ce⁴⁺ reduction is reached, which means that 25% of the oxygen atoms are released. It is worth mentioning that when the solid reaches a maximum reduction percentage (∼50%), an abrupt decrease of all the reaction products and an increase in the CH₄ signal is detected, indicating the deactivation of the catalyst. The degree of reduction reached at the end of the experiment is consistent with the previously reported maximum reduction reached in 5 mol % H₂—He balance atmosphere observed by XANES experiments.¹³

These results confirm the capability of the ceria—zirconia catalyst to activate the methane molecule and exchange the lattice oxygen with methane leading to water and carbon oxides. The other important conclusion is that this capability remains until 50% of Ce⁴⁺ reduction is reached, which means that 25% of the oxygen atoms are released. It is worth mentioning that the reduction of two Ce⁴⁺ cations to Ce³⁺ releases one oxygen atom. Depending on the oxygen content in the mixed oxide, we can observe the prevalence of products corresponding to total or partial oxidation of methane. This is an indication that Ce⁴⁺ to Ce³⁺ ratio governs the activity of the material.

4. CONCLUSIONS

From in situ XPD experiments, we observed a higher growth of average crystallite size in sample synthesized with lysine as α-amino acid (LS) than in that synthesized with glycine (GS) due to the higher degree of agglomeration, which facilitates the sintering process. As it was previously reported, the agglomeration ratio is strongly influenced by the α-amino acid used in the synthesis process.

From the diffraction patterns collected on heating in reducing atmosphere, it is clear that sample LS retains the cubic structure in all the temperature range, while sample GS evidences the segregation of a reduced ceria-based phase at temperature in the 885—895 °C range. We confirmed that segregation is reversed by reoxidation. We believe that the agglomeration of the crystallites plays an important role in phase segregation. The more agglomerated solid (LS) shows a faster rate of crystallite growth that facilitates the accommodation of Ce³⁺ cations within the structure.

In situ XANES studies confirmed our previous catalytic results obtained by conventional catalytic tests in a fixed-bed reactor indicating that sample GS is an excellent catalyst for total oxidation of methane. In situ XANES studies in Ce-LIII absorption edge pointed out that the cerium cations were reduced up to a 5 mol % during the oxidation of methane, reaching a stable condition. Furthermore, it was possible to confirm that the solid is capable to deliver the oxygen of its structure when no oxygen is fed into the reactor. In this last case, the catalyst maintains its catalytic activity for a limited period of time, while cerium cations are reduced. This process continues until 50% of cerium cations become reduced, triggering the deactivation process. Therefore, it is clear for these studies that the Ce⁴⁺ to Ce³⁺ ratio in the lattice is governing the catalytic behavior of the solid, which operates according to the Mars—Van Krevelen model, exchanging the oxygen of the lattice with the methane molecule.

AUTHOR INFORMATION

Corresponding Author
*Tel: +54.11.4709-8158. E-mail: slarrondo@citedef.gob.ar or slarrondo@unsam.edu.ar.

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS
This work was supported by the Brazilian Synchrotron Light Laboratory (LNLS) under proposals D06A-DXAS-9949 and XRDI-14413, Agencia Nacional de Promoción Científica y Tecnológica (Argentina, PICT 2010 No. 0322, PICT 2013 No. 1493, PICT 2013 No. 1587, PICT 2013 No. 1032) and Ministerio de Defensa (PIDDEF No. 011/11). A.S. acknowledges the valuable support of Dr. H. Troiani (CONICET, Metals Division, Bariloche Atomic Centre) in the TEM measurements.

REFERENCES
(1) Terribile, D.; Trovarelli, A.; de Leitenburg, C.; Primavera, A.; Dolcetti, G. Catalytic Combustion of Hydrocarbons with Mn and Cu-Doped Ceria—Zirconia Solid Solutions. Catal. Today 1999, 47, 133—140.
(2) Bozo, C.; Guilhaume, N.; Garbowski, E.; Primet, M. Combustion of Methane on CeO₂—ZrO₂ Based Catalysts. Catal. Today 2000, 59, 33—45.
(3) Pengpanich, S.; Meeyoo, V.; Rirksonboon, T. Catalytic Oxidation of Methane Over CeO₂—ZrO₂ Mixed Oxide Solid Solution Catalysts Prepared Via Urea Hydrolysis. Appl. Catal., A 2002, 234, 221—233.
(4) Otsuka-Yao-Matsuo, S.; Omata, T.; Izu, N.; Kishimoto, H. Oxygen Release Behavior of CeZrO₄ Powders and Appearance of New Compounds κ and τα. J. Solid State Chem. 1998, 138, 47—54.
(5) Montini, T.; Bañares, M. A.; Hickey, N.; Di Monte, R.; Fornasiero, P.; Kaspar, J.; Graziani, M. Promotion of Reduction in
Ce0.9Zr0.1O2 Mixed Oxides: The Pyrochlore Structure as Effect Rather Than Cause. Phys. Chem. Chem. Phys. 2004, 6, 1–3.

(6) Boaro, M.; Desinan, S.; Abate, C.; Ferluga, M.; de Leitenburg, C.; Trovarelli, A. Study on Redox, Structural and Electrical Properties of Ce0.9Zr0.1O2 for Applications in SOFC Anodes. J. Electrochem. Soc. 2011, 158 (2), P22–P29.

(7) Vidal, H.; Kașpar, J.; Pijolat, M.; Colon, G.; Bernal, S.; Cordón, A.; Perrichon, V.; Fally, F. Redox Behavior of CeO2–ZrO2 Mixed Oxides: II. Influence of Redox Treatments on Low Surface Area Catalysts. Appl. Catal., B 2001, 30, 75–85.

(8) Rodriguez, J. A.; Wang, X.; Liu, G.; Hanson, J. C.; Hrbej, J.; Peden, C. H. F.; Iglesias-Juez, A.; Fernández-Garcia, M. Physical and Chemical Properties of Ce1–xZrO2 Nanoparticles and Ce1–xZr2O5 (1 1 1) Surfaces: Synchrotron-Based Studies. J. Mol. Catal. A: Chem. 2005, 228, 11–19.

(9) Colon, G.; Valdivieso, A.; Pijolat, M.; Baker, R. T.; Calvino, J. J.; Bernal, S. Textural and Phase Stability of Ce0.7Zr0.3O2 Mixed Oxides Under High Temperature Oxidising Conditions. Catal. Today 1999, 50, 271–284.

(10) Zhang, F.; Hanson, J. C.; Robinson, R. D.; Herman, P.; Siu-Wai, C. Phases in Ceria–Zirconia Binary OXide (1–X)CeO2–XZrO2 Nanoparticles: The Effect of Particle Size. J. Am. Ceram. Soc. 2006, 89 (3), 1028–1036.

(11) Zimicz, M. G.; Fábregas, I. O.; Lamas, D. G.; Larrondo, S. A. Effect of Synthesis Conditions on The Nanopowder Properties of Ce0.6Zr0.4O2. Mater. Res. Bull. 2011, 46 (6), 850–857.

(12) Rodríguez-Carvajal, J. Fullprof Suite, version 2007.

(13) Zimicz, M. G.; Larrondo, S. A.; Prado, R. J.; Lamas, D. G. Time-resolved in Situ XANES Study of The Redox Properties of Ce0.9Zr0.1O2 Mixed Oxides. Int. J. Hydrogen Energy 2012, 37 (19), 14881–14886.

(14) Zimicz, M. G.; Lamas, D. G.; Larrondo, S. A. Ce0.9Zr0.1O2 Nanocatalyst: Influence of Synthesis Conditions in The Reducibility and Catalytic Activity. Catal. Commun. 2011, 15, 68–73.

(15) Acuña, L. M.; Fuentes, R. O.; Fantini, M. C. A.; Lamas, D. G. Relation Between Distortions in The Oxygen Sublattice and The Local Order of Zr in Nanostructured ZrO2–CeO2 Mixed Oxides. J. Phys. Chem. C 2014, 118, 11445–11453.

(16) Kaneko, H.; Taku, S.; Tamaura, Y. Reduction Reactivity of CeO2–ZrO2 Oxide Under High O2 Partial Pressure in Two-Step Water Splitting Process. Sol. Energy 2011, 85, 2321–2330.

(17) Fornasiero, P.; Di Monte, R.; Rao, G. R.; Kaspar, J.; Meriani, S.; Trovarelli, A.; Graziani, M. Rh-Loaded CeO2-ZrO2 Solid-Solutions as Highly Efficient Oxygen Exchangers: Dependence of The Reduction Behavior and The Oxygen Storage Capacity on The Structural-Properties. J. Catal. 1995, 151, 168–177.

(18) Alayoglu, S.; An, K.; Melaet, G.; Chen, S.; Bernardi, F.; Wang, L.; Lindeman, A. E.; Musselwhite, N.; Guo, J.; Liu, Z.; Marcus, M. A.; Somorjai, G. A. Pt-Mediated Reversible Reduction and Expansion of CeO2 in Pt Nanoparticles/Mesoporous CeO2 Catalyst: In Situ X-Ray Spectroscopy and Diffraction Studies Under Redox (H2 and O2) Atmospheres. J. Phys. Chem. C 2013, 117, 26608–26616.

(19) Shannon, R. D.; Prewitt, C. T. Effective Ionic Radii in Oxides and Fluorides. Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem. 1969, 25, 925–1048.

(20) Rodriguez, J. A.; Hansob, J. C.; Kim, J. Y.; Liu, G.; Iglesias-Juez, A.; Fernández-Garcia, M. Properties of CeO2 and Ce1.3Zr0.7O2 Nanoparticles: X-Ray Absorption Near-Edge Spectroscopy, Density Functional, and Time-Resolved X-Ray Diffraction Studies. J. Phys. Chem. B 2003, 107 (15), 3535–3543.

(21) Perrichon, V.; Laachir, A.; Bergeret, G.; Frety, R.; Toumayan, L.; Touret, O. Reduction of Cerias with Different Textures by Hydrogen and Their Reoxidation by Oxygen. J. Chem. Soc. Faraday Trans. 1994, 90, 773–781.

(22) Bernal, S.; Blanco, G.; Cauqui, M. A.; Cifredo, G. A.; Pintado, J. M.; Rodríguez-Izquierdo, J. M. Influence of Reduction Treatment on The Structural and Redox Behaviour of Ceria, La/Ce and Y/Ce Mixed Oxides. Catal. Lett. 1998, 53 (1–2), 51–57.

(23) Boaro, M.; Pappacena, A.; Abate, C.; Ferluga, M.; Llorca, J.; Trovarelli, A. Effect of Redox Treatments on Ce0.5Zr0.5O2 Based Solid Oxide Fuel Cell Anodes. J. Power Sources 2014, 270, 79–91.