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Contraction, cation oxidation state and size effects in cerium oxide nanoparticles

Jacopo Stefano Pelli Cresi^{1,2}, Maria Chiara Spadaro^{1,2,3}, Sergio D'Addato^{1,2}, Sergio Valeri^{1,2}, Lucia Amidani⁴, Federico Boscherini^{5,6}, Giovanni Bertoni⁷, Davide Deiana³, Paola Luches^{2*}

¹ Dipartimento di Scienze Fisiche Informatiche e Matematiche, Università degli Studi di Modena e Reggio Emilia, Via G. Campi 213/a, 41125 Modena, Italy

² Istituto Nanoscienze, Consiglio Nazionale delle Ricerche, Via G. Campi 213/a, 41125 Modena, Italy

³ Interdisciplinary Centre for Electron Microscopy, Ecole Polytechnique Fédérale de Lausanne, Station 12, 1015 Lausanne, Switzerland

⁴ European Synchrotron Radiation Facility, BP 220, 38043 Grenoble, France

⁵ Department of Physics and Astronomy, University of Bologna, Viale C. Berti Pichat 6/2, 40127 Bologna, Italy

⁶ Istituto Officina dei Materiali, Consiglio Nazionale delle Ricerche, Operative Group in Grenoble, c/o ESRF, BP 220, 38043 Grenoble, France

⁷ Istituto dei Materiali per Elettronica e Magnetismo, Consiglio Nazionale delle Ricerche, Parco Area delle Scienze 37/A, 43124 Parma, Italy

* corresponding author: email: paola.luches@unimore.it

Abstract

An accurate description of the structural and chemical modifications of cerium oxide nanoparticles is mandatory for understanding their functionality in the applications. In this work we investigate the relation between local atomic structure, oxidation state, defectivity and size in cerium oxide nanoparticles with variable diameter below 10 nm, using X-ray absorption fine structure analysis in the near and extended energy range. The nanoparticles are prepared by physical methods under controlled conditions and analyzed in morphology and crystalline quality by high resolution transmission electron microscopy. We resolve here an important question on the local structure of cerium oxide nanoparticles: we demonstrate a progressive contraction in the Ce-O interatomic distance with decreasing nanoparticle diameter and we relate the observed effect to the reduced dimensionality. The contraction is not significantly modified by inducing a 4-6% higher Ce³⁺ concentration through thermal annealing in high vacuum. The consequences of the observed average cation-anion distance contraction on the properties of the nanoparticles are discussed.

Introduction

Cerium oxide based materials are attractive due to their capacity to quickly store, release and transport oxygen ions, which makes them suitable for application in different fields. Catalysts based on nanostructured cerium oxide are used for example for the abatement of toxic exhausts from car engines, and they have been shown to be active in several reactions like the water-gas shift, CO preferential oxidation or methane oxidation reaction [1]. Recent perspective applications include also energy conversion and storage devices, in which nanostructured cerium oxide is proposed as an electrode material, for example in fuel cells [2, 3] and supercapacitors [4]. Moreover, cerium oxide nanoparticles have been shown to have a great potential as antioxidant materials for therapeutic purposes [5].

The oxygen storage capacity in cerium oxide is related to the possibility for Ce ions to reversibly change their oxidation state from 4+ to 3+ under mild modifications of the ambient conditions [6]. An atomic scale understanding of the material reducibility and of the possible modifications induced by parameters like size, specific exposed surfaces, and interaction with other materials and molecules is of paramount importance in view of the possibility of designing materials with optimized functionality.

Several studies were dedicated to the investigation of the properties of cerium oxide nanoparticles (NPs) of variable size and shape obtained by different synthesis methods [6-11]. An accurate description of the electronic and atomic structure of cerium oxide as a function of NP size is indeed relevant to properly understand the size-dependent behavior of the material, which was often proved to be unexpectedly different from the bulk. The demonstrated strong dependence of the reducibility and of the oxygen storage capacity on the exposed facet derives strictly from the properties of the different surfaces [12, 13]. On the contrary, other properties arising from confinement to the nanometer range have a less trivial explanation; among these we mention as examples the predicted size-dependent oxygen vacancy formation energy, which reaches a minimum at a few nm size [14], and the peculiar delocalization of valence electrons demonstrated for sizes of approximately 2 nm [15].

An "anomalous" lattice expansion is typically reported when the size of the investigated cerium oxide NPs is below a few tens of nm [8-10, 16-18], as opposed to the contraction reported for metal NPs since the late sixties [19-21]. As the NP diameter decreases, given the higher surface to volume ratio, the concentration of Ce^{3+} sites, possibly induced on the surface by the chemical synthesis procedures, increases. For this reason the observed lattice expansion has often been suggested to originate from the larger radius of Ce^{3+} compared to Ce^{4+} ions [8, 16-18, 22, 23]. On the contrary, theoretical studies predict that a lattice contraction is also expected in cerium oxide NPs [23, 24], and ascribe the experimentally observed expansion to the presence of oxygen vacancies [24]. However, a simple correlation between the degree of reduction and the lattice expansion with decreasing cerium oxide NP size has often been questioned, pointing out the modifications which may be induced by the synthesis or measurement conditions [25, 26]. It is clear that the outstanding question is whether CeO_2 NPs contract or expand with decreasing size, demanding for a careful study on NPs with controlled degree of reduction.

Physical methods based on gas phase synthesis have been shown to be suitable for the synthesis of NP assemblies with a good control over stoichiometry, size and architecture [27-29]. For cerium oxide NPs the oxidation state of part of the Ce ions can be reversibly modified by thermal annealing in vacuum or oxygen pressure [11, 30], in close analogy with cerium oxide epitaxial films and nanostructures .

X-ray absorption fine structure (XAFS) allows to simultaneously obtain information on the chemical state of an atomic species and on its local structural environment by analyzing an absorption edge in the near and extended energy range, respectively. In this work we apply the XAFS technique to the Ce L₃ edge in cerium oxide NPs with selected size and controlled degree of reduction, we demonstrate a contraction in the Ce-O distance compared to the bulk value and we analyze its dependence on the degree of reduction.

Experimental methods and data analysis

The NP samples were obtained using a magnetron sputtering and inert gas aggregation nanocluster source (NC200U, Oxford Applied Research), equipped with a quadrupole mass filter (QMF200, Oxford Applied Research), and installed in an ultrahigh vacuum experimental system connected to an X-ray photoelectron spectroscopy (XPS) apparatus (XR50, Specs double anode X-ray source and Phoibos 150, Specs hemispherical electron analyser) [29, 31]. For the present experiment we used a 99.9 % pure metallic cerium target and a mixture of He and Ar gases in the aggregation region. We selected NPs of variable size, below 10 nm diameter, by varying the electric power applied to the magnetron target, the Ar and He gas flux in the aggregation region, and the length of the aggregation region. The parameters used for obtaining NPs of different size are summarized in the supporting information, Table S1. The QMF was used to further select the size of the NPs before deposition. The NP deposition rate was evaluated using a quartz microbalance. In the following, the nominal film thickness is defined as the thickness of a continuous cerium oxide film with the same volume as the NP film. The NP films were deposited on a Si/SiO_x substrate for XPS and XAS measurements, while carbon coated copper grids were used as a support for transmission electron microscopy (TEM). The samples were deposited in a partial pressure of oxygen $P_{O_2} = 5 \times 10^{-6}$ mbar. In order to maximize the surface oxidation, the samples were exposed to an oxygen partial pressure $P_{\rm O_2}$ = 6 \times 10 $^{-5}$ mbar for 30 minutes after the growth. The nominal film thickness was fixed to 10 nm for the samples to be measured by XAS, while the samples for TEM had a much smaller nominal thickness, in the subnanometer range, in order to be able to identify and analyse individual particles. The TEM measurements were limited to the 4 and 6 nm NP samples, while only 4, 5 and 9 nm NP samples were measured by XAS.

All samples were characterized by *in-situ* XPS, using Al K_{α} photons. To obtain quantitative information on the oxidation state the measured Ce 3d XPS spectra were fitted using Ce⁴⁺, and Ce³⁺ components, following the procedure introduced by Skála et al [32], already used by our group for the analysis of epitaxial ceria nanostructures [33, 34].

Ex-situ scanning TEM images (STEM) in annular dark field mode (ADF) were performed to assess the size distribution and crystalline quality of the NPs. The experiments on the 6 nm sample were performed using a JEOL JEM-2200FS microscope (IMEM-CNR, Parma), equipped with a Schottky emitter working at 200 keV (point resolution 0.19 nm). This instrument was used also to acquire the selected area electron diffraction (SAED) patterns. The experiments on the 4 nm sample were performed using a double spherical corrected FEI Titan Themis cubed microscope (CIME, EPFL Lausanne) working at 200 keV, equipped with an ultra-bright Schottky (XFEG) electron source (point resolution 0.07 nm). All the ADF-STEM images have been analysed using the STEM_CELL software [35].

The XAFS measurements were performed at BM08 (LISA) beamline of the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. The measurements were acquired at the Ce L₃-edge, using a dynamically sagittal-focusing Si(311) monochromator [36]. In order to eliminate higher-order harmonics, a pair of Pt coated flat mirrors were used to set a cut-off energy of 12 keV. The XAFS spectra were acquired in the fluorescence yield mode, using a 12-element hyper-pure Ge detector. The samples were kept in nitrogen atmosphere during the transfer to the synchrotron radiation facility. A sample environment cell was used to perform the reducing thermal annealing, which consists in heating the samples to 1020 K in high vacuum (P ~ 10^{-7} mbar) for 30 min using a constant heating

and cooling rate of 7 K/min. A similar treatment was previously shown to induce a non-negligible reduction on the surface of cerium oxide ultrathin films and NP [11, 33, 37, 38]. In particular, for this work we used the same cell and the same conditions used in reference [37]. The samples were measured at room temperature (RT) before and after the high-vacuum thermal annealing (indicated as HVT). Two reference samples in powdered form were also measured in transmission mode: a CeO₂ sample, as a reference for the Ce⁴⁺ oxidation state, and a Ce(NO₃)₃·6H₂O sample, as a reference for the Ce³⁺ oxidation state.

X-ray absorption near edge spectroscopy (XANES) measurements were analysed to have information on the variation of the oxidation state in the samples. In particular, the spectra were fit using five Pearson VII distribution curves, i.e. Lorentz functions raised to a power m, to simulate the near edge features - three related to Ce^{4+} , one related to Ce^{3+} , and one related to the pre-edge - and an arctangent function to reproduce the edge jump, following reference [39]. The energy position of the centre, the FWHM and the shape of the Pearson distributions were fixed to the ones obtained from the fit of the Ce^{4+} and Ce^{3+} reference samples (see Figure S1, Supporting Information). Moreover, since the peak originating from Ce³⁺ ions occurs at a photon energy which is comparable to one of the structures corresponding to Ce⁴⁺ ions, the intensity ratio between the first two Ce⁴⁺ related features was fixed to the value obtained in the fit of the reference sample (see Figure S1, Supporting Information). Since it is not possible to resolve the Ce^{3+} and Ce^{4+} contributions to the pre-edge peak [40], its intensity was not considered in the evaluation of the Ce³⁺ concentration within the samples. We note that the reference samples for the two Ce oxidation states may contain also ions in the other oxidation state as minority components, located on some of the low coordination sites or at defect sites. This affects the accuracy of the Ce³⁺ concentration obtained from the fitting. However, all of the considerations made in this work are based on relative variations of the estimated Ce^{3+} concentration, and they are not affected by the accuracy of the numbers obtained.

The extended X-ray absorption fine structure (EXAFS) was acquired up to k = 9 Å⁻¹. The available *k*-range was limited by a small Cr K adsorption edge signal, possibly originating from diffused light scattering from the walls of the sample environment cell. The EXAFS spectra were processed using the Athena software package [41]. Starting from the fluorite structure of cerium oxide, the theoretical scattering functions were simulated *ab-initio* using the FEFF program [42]. Quantitative data analysis was performed by fitting the EXAFS spectra with the simulated scattering functions, including only the Ce-O first shell single scattering path, by means of the Artemis code [43]. The presence of multiple excitations and the mixed valence nature of the excited state [44, 45] has been neglected, given the expected weak intensity of the resulting features and the limited energy range available for the analysis.

Results and Discussion

A series of cerium oxide NP samples with different nominal diameter were grown with the procedures described in the experimental section. The surface stoichiometry of the NPs after deposition and exposure to oxygen was evaluated by analyzing Ce 3d XPS spectra acquired *in-situ* (Figure 1). The dominant features in the spectra of the samples originate from three doublets corresponding to emission from Ce⁴⁺ ions with different final states, indicated as v - u, v''- u'', and v''' - u''' in Figure 1, following Romeo et al. [46]. As the NP diameter decreases the spectra show a progressive increase of intensity in the binding energy region between v and v'' and between u and u'', corresponding to the most intense v' - u' doublet from Ce³⁺ ions (Figure 1). The Ce³⁺ surface concentration in each sample was evaluated by fitting the spectra with Ce⁴⁺ and Ce³⁺ related doublets [32], and the results

are reported in Table 1. The Ce³⁺ surface concentration in the 4 nm NPs is above 20 % and it decreases with increasing NP diameter, in agreement with the presence of a higher concentration of low-coordinated surface sites in NP with smaller diameter. It has to be noted that the accuracy obtained by this method is limited to approximately \pm 10%. Moreover, the measured values may well be systematically overestimated due to a demonstrated photon beam-induced surface reduction of the NP surface in ultra-high vacuum [11].



Figure 1: Ce 3d XPS spectra (black curve) of cerium oxide NPs with different average diameter. The fitting using Ce^{3+} and Ce^{4+} related components are also shown (red curve). The binding energy positions of the Ce^{4+} and of the Ce^{3+} -related doublets are indicated in the top and bottom of the figure, respectively.

ø (nm)	Ce ³⁺ (%)
4	23 ± 2
5	18 ± 2
6	14 ± 2
9	10 ± 2

Table 1: Ce^{3+} surface concentration in the cerium oxide NP samples of different diameter, evaluated from the fitting of Ce 3d XPS spectra. The uncertainties represent the precision of the Ce³⁺ concentration values, rather than their accuracy.

The morphology, crystal quality and size distribution of the cerium oxide NPs were investigated by high resolution STEM measurements. A representative image for NPs with selected diameter of 6 nm is shown in Figure 2 a. The corresponding high resolution image, shown in the figure inset, shows that most of the NPs have a single-crystalline fluorite structure (Fm $\overline{3}$ m space group) with evident (111) crystal planes fringes.



Fig.2: ADF-STEM images of cerium oxide NPs of 6 nm (a) and 4 nm (c) average diameter; insets: higher magnification of selected NPs; b) and d) size distribution extracted from the STEM images in a) and c), respectively, and fit with a lognormal distribution.

The shape of the NPs is polyhedral and the (111) and (100) surfaces are preferentially exposed. Most of the NPs are single crystalline, while few of them are polycrystalline, as already observed in previous work on NP grown in the same way [30]. The size distribution of the 6 nm NP sample (Figure 2 b) is rather narrow; a fit with a log-normal distribution has the maximum at 6.2 nm and a full width at half maximum (FWHM) of 1.1 nm. The STEM images corresponding to NPs with selected diameter of 4 nm, shown in Figure 2 c, confirm that even the NPs with smallest size analyzed in this work, maintain their individuality and the high degree of crystal quality. In this case the size distribution (Figure 2 d) has the maximum at 4.3 nm and it is slightly broader (FWHM=1.7 nm) than the distribution adopted for the QMF in the two samples (U/V = 0.100 for the 6 nm sample, U/V = 0.069 for the 4 nm sample). This choice results from a compromise between an appreciable NP deposition rate and a small dispersion of the NP size. The TEM-SAED patterns are compatible with the fluorite structure (see figure S3). However, we could not extract precise information on the long range lattice parameters from the positions of the rings, which appeared weak and broad, given the high dilution of the nanoparticles and their small size.

The Ce L₃-edge XANES spectra of the cerium oxide NP samples before and after the thermal annealing are shown in Figure 3. For all samples the XANES line shape shows dominant features which originate from Ce⁴⁺ ions, namely a peak at 5732.6 eV (peak B in Figure 3) ascribed to a screened transition from 2p levels to the 5d band, and one at 5739.6 eV (peak A in Figure 3) ascribed to unscreened transitions between the same levels [47]. In addition, the spectra show a low energy shoulder at 5728.2 eV (peak C in Figure 3) ascribed to crystal-field splitting of the 5d band [48] and a poorly resolved pre-edge feature (peak D in Figure 3) ascribed to dipole-forbidden 2p-4f transitions [49].



Figure 3: Ce L₃-edge XANES spectra of the cerium oxide NP samples before (blue curves) and after (red curves) the reducing high vacuum thermal annealing (HVT). The fits of the spectra, using the Ce^{4+} and Ce^{3+} related components shown at the bottom, are also reported (dashed line).

After the thermal annealing at 1020 K in vacuum the edge is slightly shifted towards lower photon energies and the shape of the first feature changes, in agreement with a non-negligible degree of reduction of the samples. The Ce^{4+} and Ce^{3+} concentrations in the samples were quantified by fitting the spectra with Ce^{4+} and Ce^{3+} components and the results are shown in Table 2. We note that also in this case all the considerations made are based on the variation of the degree of reduction in the different samples, rather than on their absolute values, which can be affected by a non-negligible inaccuracy, and possibly systematically overestimated due to eventual beam-induced surface reduction.

	Ce^{3+} (%)	Ce ³⁺ (%)	
ø (nm)	as prepared	after HVT	
4	9 ± 1	13 ± 1	
5	13 ± 1	19 ± 1	

9	13 ± 1	17 ± 1

Table 2: Ce^{3+} concentration in the cerium oxide NP samples of different diameter, evaluated from the fitting of Ce L₃edge XANES spectra, before and after the reducing high vacuum thermal treatment (HVT) at 1020 K. The uncertainties represent the precision of the Ce³⁺ concentration values, rather than their accuracy.

In spite of this, the relatively low degree of reduction detected by XANES measurements on the cerium oxide NPs before the reducing thermal treatment indicates that the bulk of the NPs is well oxidized, although the Ce^{3+} species represent a non-negligible minority component. The comparison with the systematically higher values of Ce^{3+} concentration detected by XPS, which is a surface sensitive technique, indicates that the Ce^{3+} species are mainly localized at the surface. The slightly higher Ce^{3+} concentration in the samples with larger diameter possibly reflects a slightly higher bulk defectivity in the larger samples. The reducing thermal annealing induces a moderate increase of the Ce^{3+} concentration in all samples. The difference between the Ce^{3+} concentration before and after the treatment is comparable in the three samples within the error, which indicates that the process induces a comparable fraction of Ce^{3+} ions independent on their size and on the initial defectivity. A similar thermal annealing treatment in ultra-high vacuum has been previously shown to induce a relevant reduction of the NP surface [11, 30]. The mild reduction observed by XANES, which is sensitive to the whole NP volume, is an indication that the reduction process involves mainly the NP surface.

To have information on the local atomic structure of the NPs with controlled oxidation state, the EXAFS energy region of the spectra was acquired right after the XANES for each sample. The magnitude of the Fourier transforms of the EXAFS spectra of the NP samples before and after the high-vacuum thermal annealing are shown in Figure 4.



Figure 4: Magnitude of Fourier transforms of Ce L_3 -edge EXAFS spectra before (blue curves) and after (red curves) the reducing high vacuum thermal treatment (HVT) for three different cerium oxide NP samples. The position of the peak maximum is marked with blue and red ticks. The fits are also reported as dashed lines.

A peak around 1.8 Å, marked with a blue tick in Figure 4, corresponding to single scattering from first shell oxygen atoms, is the dominant feature in the spectra of the as prepared samples. After the thermal annealing the peak, marked with a red tick in Figure 4, is systematically shifted towards larger Ce-O distances and its amplitude is slightly increased. A first-shell fitting allowed to have quantitative information on the local structure of the different samples and on the evolution with the high-vacuum thermal annealing. In the fitting the many body amplitude reduction factor was fixed to the value obtained from the fit of the reference CeO₂ sample spectrum (see Figure S2, supporting information). The free fitting parameters were the energy origin shift, the coordination number, the variation of the Ce-O interatomic distance, and the Debye-Waller factor. The fitting was performed in the ranges k = 2.5 - 7.0 Å⁻¹ and R=1-3 Å on k²-weighted spectra. The results obtained are summarized in Table 3, which reports also the values obtained on the reference CeO₂ sample, using the same k and R range.

ø (nm)	treatment	Ν	σ (Å ²)	E ₀ (eV)	ΔR(Ce-O) (Å)
9	as prepared	7(2)	0.007(7)	3(2)	-0.06(3)
9	after HVT	7.5(1.5)	0.006(4)	2.4(1.7)	-0.08(2)
5	as prepared	7.6(1.2)	0.011(4)	2.1(1.3)	-0.084(19)
5	after HVT	6.1(5)	0.005(2)	3.0(7)	-0.061(9)
4	as prepared	7(2)	0.006(6)	2(2)	-0.08(3)

4	after HVT	7.8(9)	0.008(3)	2.8(1.0)	-0.076(14)
CeO ₂ reference		8	0.006(3)	4.7(1.1)	-0.046(16)

Table 3: Parameters obtained by the fitting of the Ce L₃-edge EXAFS spectra of the three different NP samples before and after the high-vacuum thermal annealing (HVT): coordination numbers (N), Debye-Waller factors (σ), energy origin shift (E₀), and variation of the Ce-O interatomic distance (Δ R(Ce-O)). The values obtained for the CeO₂ reference sample are also reported. The numbers in parentheses indicate the standard deviation on the last or last two digits.

The correlation between the coordination numbers and the Debye-Waller factors in the fitting procedure prevents from drawing reliable conclusions on the evolution of the two parameters with the NP diameter and high-vacuum thermal annealing. We note that the obtained values are not very different from the ones in the reference CeO_2 sample, within the error bars, confirming the dominant fluorite bulk-like coordination of Ce ions in the NPs. On the contrary, the observed values for the Ce-O distance deserve some considerations. The NP samples show a small, but systematic, compression of the Ce-O interatomic distance with decreasing NP size. The measured contraction as a function of the reciprocal average diameter of the NPs is reported in Figure 5.



Figure 5: Contraction of the Ce-O interatomic distance evaluated from the fitting of the Ce L_3 edge EXAFS spectra of the NPs of different diameter and of the CeO₂ reference sample as function of the reciprocal average diameter. The solid line represents a linear fit of the data. The error bars reported in graded color indicate the standard deviations on the Ce-O distance obtained from the fitting.

The majority of previous works on the structure of cerium oxide NPs report a progressive lattice expansion with decreasing particle diameter between 2 and 10 nm. The expansion has been frequently ascribed to a higher concentration of Ce³⁺ ions at smaller NP size, and to the larger ionic radius of Ce³⁺ with respect to Ce⁴⁺. A recent work by Paun et al. proved that the modifications of lattice parameter are not fully correlated with the NP stoichiometry, since also well oxidized NPs showed a non-negligible lattice expansion, ascribed to the negative Madelung pressure at reduced dimensionality [50]. Assuming the thermodynamic model developed by Vermaak [19], which relates the relative volume variation $\Delta V/V$ to the reciprocal radius 1/r with a linear dependence involving the surface stress f, and the bulk modulus B $\left(\frac{\Delta V}{V} = -\frac{2f}{B}\frac{1}{r}\right)$, other works ascribe the observed expansion to a negative surface stress, in analogy with metallic NPs, although in the latter case the surface stress is positive and it leads to a lattice contraction. The strong dependence of the lattice parameter on the synthesis methods and on the measurement conditions has been discussed in detail . In a theoretical study Loschen et al. predicted a contraction of the average interatomic distance in cerium oxide clusters with size of approximately 1-2 nm without oxygen vacancies, and they ascribed the interatomic distance modification to dimensional effects, in close analogy to the behavior observed for metallic NPs. At present, only two works reported an experimental evidence for interatomic contraction in cerium oxide NPs. Nachimuthu et al. observed a contraction in NPs with

diameter below 15 nm and they related it to the decrease in coordination of Ce ions [39]. In another work, only NPs with diameter above 15 nm showed an increasing lattice contraction with decreasing diameter, while smaller NP exhibited a reverse trend, due to a dominant lattice expansion induced by oxygen vacancies . Moreover, an almost isotropic contraction of the Ce-O distance was observed on epitaxial cerium oxide nanostructures of a few tens of nm lateral size and ~ 1 nm thickness [51]. In this case the in-plane contraction was ascribed to epitaxial effects, while the out-of-plane contraction, unexpected considering the bulk elastic theory, was hypothesized to originate from reduced dimensionality effects. In the present study, we could measure simultaneously the cerium local structural environment and its oxidation state. We detected a non-negligible Ce-O contraction even in NPs with a small concentration of Ce³⁺ sites. Figure 5 shows that the Ce-O bond length has a linear dependence on the surface to volume ratio S/V, in turn proportional to the reciprocal diameter 1/d. This is a clear indication that the contraction is a dimensionality related effect, in close analogy with metallic NPs and in agreement with theoretical predictions [23, 24]. Our data demonstrate that in cerium oxide NPs the dimensionality-induced bulk contraction dominates over the expected local expansion around Ce³⁺ sites, resulting in an overall decrease of the average Ce-O distance. We believe that the clean conditions used in the preparation and measurement of the NPs and the low concentration of defects allowed to clearly identify the dimensionality-induced contraction. Note that diffraction techniques, often employed to determine the lattice variation in previous studies of cerium oxide NP, are only sensitive to well-ordered crystalline phases, while the XAFS technique used in the present study detects the average Ce-O distance in all the Ce-containing phases present in the sample, including possible less ordered phases. The assumption of a dimensionality-related origin of the observed contraction is strengthened by the fact that defective low-ordered zones, eventually present in the samples, are expected to have an expanded Ce-O distance with respect to the bulk value. The high-vacuum thermal annealing, observed to induce a moderate increase of Ce³⁺ concentration, does not significantly modify the observed contraction within the uncertainties in the measurements (Table 4), confirming that dimensionality is largely dominant over stoichiometry-related effects.

The contraction in cation-anion distance is indeed expected to modify some of the properties of the NPs, since it originates from a stronger average bonding. In general, a different band structure is expected, with consequences e.g. on the optical properties [52], or on the mechanical properties [53]. A possibly lower mobility of oxygen ions and electrons may also be induced. The lower electron mobility may be an advantage for some applications, like for example fuel cell electrodes in which a low electric conductivity may be required. On the other hand, a lower oxygen ion mobility will certainly limit the oxygen storage capacity, which is typically required to be maximized in most applications. In spite of this, we do not foresee a reduced surface reactivity in the NPs here investigated, due to the low coordination of the cerium ions on the surface, and the non-negligible degree of surface reduction measured in vacuum, which is expected to favor the reversible uptake and release of oxygen ions.

Conclusions

Cerium oxide NPs with diameter below 10 nm have a dominant single crystalline fluorite structure and they show a contraction of the Ce-O distance compared to the bulk value, in contrast with the frequently observed lattice expansion with decreasing NP size. The contraction has a linear dependence on the surface-to-volume ratio and it is ascribed to a compressive strain arising from reduced dimensionality. The size-induced strain dominates over the local expansion expected at the Ce^{3+} sites, due to the low concentration of reduced ions in the investigated samples. The results here

described can help to understand the relationship between structure and functionality in systems based on cerium oxide NPs.

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