Nanoscale Advances

PAPER

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Cite this: Nanoscale Adv., 2021, 3, 1484

Received 23rd August 2020 Accepted 17th December 2020 DOI: 10.1039/d0na00700e

rsc.li/nanoscale-advances

I. Introduction

The doping process developed around 1950 has been extensively used in order to tune the electrical, magnetic, optical and structural properties of host matrices. As is the case of silicon (largely applied in the industry), n- and p-type semiconductors are obtained by introducing donor and acceptor impurities, respectively. More recently, due to its unique properties that

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Tuning the magnetic properties of $Sn_{1-x-y}Ce_{4+x}Ce_{3+y}O_2$ nanoparticles: an experimental and theoretical approach

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During the last decade, there was a substantial increase in the research on metal-doped oxide semiconductor nanoparticles due to advances in the engineering of nanomaterials and their potential application in spintronics, biomedicine and photocatalysis fields. In this regard, doping a nanomaterial is a powerful tool to tune its physicochemical properties. The aim of this work is to shine a new light on the role of the neighbouring elements on the oxidation state of the Ce-impurity, from both experimental and theoretical points of view. Herein, we present an accurate study of the mechanisms involved in the oxidation states of the Ce-ions during the doping process of SnO₂ nanoparticles (NPs) prepared by the polymeric precursor method. X-ray diffraction measurements have displayed the tetragonal rutile-type SnO₂ phase in all samples. However, the Bragg's peak (111) and (220) located at $2\theta \sim 29^{\circ}$ and $\sim 47^{\circ}$ evidence the formation of a secondary CeO₂ phase for samples with Ce content up to 10%. X-ray absorption near-edge structure (XANES) measurements, at Ce L3 edge, were performed on the NPs as a function of Ce content. The results show, on one side, the coexistence of Ce^{3+} and Ce^{4+} states in all samples; and on the other side, a clear reduction in the Ce^{3+} population driven by the increase of Ce content. It is shown that this is induced by the neighboring cation, and confirmed by magnetic measurements. The monotonic damping of the Ce^{3+}/Ce^{4+} ratio experimentally, was connected with theoretical calculations via density functional theory by simulating a variety of point defects composed of Ce impurities and surrounding oxygen vacancies. We found that the number of oxygen vacancies around the Ce-ions is the main ingredient to change the Ce oxidation state, and hence the magnetic properties of Ce-doped SnO₂ NPs. The presented results pave the way for handling the magnetic properties of oxides through the control of the oxidation state of impurities.

> exist only at the nanometer scale, a great deal of attention has been devoted to investigate transition metals and rare earth doping oxides (SnO₂, TiO₂, In₂O₃, ZnO, among others),¹⁻⁴ producing the so-called metal-doped oxide semiconductors. In these materials, among different possibilities, doping with multivalent elements results in systems with a high potential for future applications.⁵ We give special attention to the unique properties of Ce ion doping oxides. Whereas Ce4+ ions are nonmagnetic, Ce³⁺ ions act as paramagnetic centers, due to the presence of one localized unpaired valence electron. Thus, tuning the Ce oxidation state could be an effective way to handle the magnetic properties of Ce doped oxides. However, the microscopic understanding of the doping mechanism (which is the condition under which an ion prefers to occupy crystal sites in the host lattice at a magnetic oxidized state (Ce^{3+}) or nonmagnetic oxidized state (Ce^{4+}) is an open question, demanding further effort to pave the way for future technological applications of metal-doped oxides. To our knowledge, little or no attention has been paid to explain the valence state preference of these ions in doped materials. In particular, the +3



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Paper

valence state of cerium has a strong influence in the magnetic properties of the host materials and opens the possibility of the production of diluted magnetic semiconductors (DMSs) with high Curie temperature (T_c). Hence, Ce doped systems have been gathering much interest from the industrial viewpoint because of their potential to fabricate novel multifunctional semiconductor devices where the magnetic electric and optical properties are combined.

The Ce-doped SnO₂ nanostructure is a good candidate for this study. The interesting point about this type of material is that, regardless of the ions, they exhibit a ferromagnetic-like order above room temperature (at low doping content), which make them interesting materials for the next generation of spintronics, electronic, and optoelectronic devices.⁶⁻¹⁰ As is shown by Prellier et al., the coexistence of the electric and magnetic properties provides the possibility of generating a spin-valve system with enhanced magnetoresistance (MR).² Besides, several works highlight this material as promising for gas sensor applications,¹¹ such as that reported by Kumar et al. in recent publications indicating the room temperature ethanol sensing property, and the astonishing change from n-type to ptype¹² semiconductor, associated with the Ce doping level in SnO2 lattices. Also, Watts et al. showed the use of this promising material for the oxidation of organic materials and the reduction of heavy metal-ions in industrial wastewater streams.13

Using the above mentioned research, in the present work, a systematic theoretical and experimental study was carried out, focusing on the effect of cerium doping on the physical properties. X-ray diffraction measurements (XRD), transmission electron microscopy (TEM), and X-ray absorption (XANES, and EXAFS) were used for this purpose. The experimental results display the coexistence of Ce^{3+} and Ce^{4+} states in all samples, tuned by the incremental increases of Ce content. Meanwhile, our theoretical findings suggest that the change of the coordination number of the oxygen ions surrounding Ce ions provokes the oxidation state change, and this fact shows the local tuning of the magnetic properties.

II. Methods

A. Experimental approach

Ce-doped SnO₂ nanoparticles with Ce-content in the range from 0 to 30 mol% have been synthesized according to the polymeric precursor method. The undoped SnO₂ noncrystalline sample was synthesized from tin(II) citrate (Sn₂2+(C₆O₇H₄)), which was prepared from citric acid (C₆O₇H₈) and tin(II) chloride (Sn²⁺Cl₂) in agreement with the following equation:

$$C_6O_7H_8^+2Sn^{2+}Cl_2^- \rightarrow Sn_2^{2+}(C_6O_7H_4) + 4HCl$$
 (1)

Thermal annealing (TA) in the tin(π) citrate under air atmosphere was performed (see eqn (2)), promoting the oxidation from Sn²⁺ to Sn⁴⁺. The TA was made in two steps: firstly at 400 °C for 4 h; and additional calcination at 400 °C for 15 h. The latter was made in order to improve the crystallinity of the samples.

$$\operatorname{Sn_2}^{2+}(\operatorname{C_6O_7H_4}) \xrightarrow{\mathrm{TA}} \operatorname{SnO_2}$$
 (2)

Meanwhile, for the Ce-doped samples the Ce precursor is the cerium(m) nitrate $Ce^{3+}(N_3)_3$, where the amount of Ce was controlled using the Ce/(Ce + Sn) ratio, and the same thermal treatment as the undoped sample was applied. For more details on the preparation, see ref. 14. X-ray powder diffraction (XRD) patterns were obtained using a commercial Bruker diffractometer (D8 advanced) with a CuK α radiation source (=1.5406 Å). The lattice parameters, as well as the average crystallite size and residual strain were determined from the XRD data analysis: in the approach the Bragg-diffraction peak has a Lorentzian shape which is implemented in the Jana2006 program,15 the half-value breadth (β) obtained from the Rietveld refinement method was carefully corrected from the instrumental broadening (β = $\beta_{\text{sample}} + \beta_{\text{instrument}}$) using a single-crystalline Si as a standard sample. In order to separate the contribution of the size from the residual strain, the Williamson-Hall plot was used. The particle size and morphology have been determined for one of the samples by high-resolution transmission electron microscopy (HRTEM). The DC magnetic measurements were carried out using the SQUID (MPMS, Quantum Design) magnetometer, varying the temperature from 5 to 300 K, and applying magnetic fields up to ± 70 kOe. In order to study the electronic and atomic structure, X-ray absorption spectroscopy (XAS) measurements were carried out in the X-ray absorption near-edge spectroscopy (XANES) region, located in the Ce L3-edge. For these measurements, pellets were made from the powders, and the XAS spectra were carried out at room temperature in transmission and fluorescence modes, and the energy was calibrated using standard Cr and Fe foils, respectively. The XAFS2 beamline at the Brazilian Synchrotron Light Laboratory (LNLS) was used, more details of the beamline used can be obtained from ref. 16. The data were analyzed using the Athena and Artemis interfaces.17

B. Theoretical approach and computational details

In order to characterize the trend of Ce-ion incorporation into the SnO₂ structure, and elucidate the oxidation state, we used computational simulations at the atomistic scale to study different configurations and concentrations of the cerium doping atoms and oxygen vacancies in the SnO₂ pristine cell. Our total energy calculations are based on density functional theory (DFT),^{18,19} by employing the semilocal Perdew-Burke-Ernzerhof (PBE) exchange and correlation functional,20 using the spin-polarized generalized gradient approximation (GGA), as implemented in the Vienna ab initio simulation package -VASP,^{21,22} version 5.4.4. The Kohn-Sham equations were solved using the projector augmented wave (PAW) method^{23,24} employing the following projectors: $Sn(4d^{10}, 5s^2, 5p^2)$, $O(2s^2,$ $(2p^4)$ and Ce $(4f^1, 5d^1, 6s^2)$, where the valence states are shown in parentheses. The SnO₂ equilibrium volume (atomic forces) in the bulk-phase was reached by performing the stress-tensor minimization (total energy) calculation using a plane-wave cutoff of 829 eV (466 eV). For the Brillouin zone integration,

we used the *k*-mesh of $1 \times 1 \times 2$ for the stress-tensor and atomic force optimizations, while $2 \times 2 \times 4$ for the electronic property calculations, such as the density of states. In every calculation, Gaussian smearing of 0.01 eV was employed, and the atoms were allowed to find their equilibrium atomic position when all the forces were smaller than 0.01 eV Å⁻¹. In order to improve the limitations given by the PBE functional and to obtain a better description of the electronic band structure, a Hubbard correction within the GGA+U method proposed by Dudarev *et al.* was used,²⁵ where the Hubbard *U* parameters are 3.5 eV and 4.5 eV on Sn-d and Ce-f states, respectively.

In addition, the theoretical XANES spectra were calculated by the FEFF9 *ab initio* code.²⁶ In our simulation of Ce L3 edge XANES, we used self-consistent potentials (SCF), and full multiple scattering (FMS) for the calculation of the real-space Green's function approach. FMS calculations were performed using a Hedin–Lundqvist self-energy correction and adding a core hole in the final state rule (FSR). The potential values used for the atom species were 58, 8 and 50 for Ce, O and Sn, respectively. The cluster sizes used in the SCF and FMS routines were 4.6 Å and 6.0 Å respectively, which was enough to fulfil the required level of accuracy. Finally, the core-level broadening was reduced to match the experimental data.

III. Results and discussion

A. Experimental results

Fig. 1 provides the X-ray diffraction patterns of Ce-doped SnO₂ nanoparticles. The rutile-type tetragonal structure (JCPDS card, 41-1445), with space group, $P4_2/mnm$, has been determined for all samples. The formation of this phase is retained for each dopant content. However, for samples with Ce content above 10 mol%, a secondary phase of cerium oxide (CeO₂) was detected, which was evidenced by (111) and (220) diffraction peaks located at $2\theta \sim 29$ and $\sim 47^\circ$. This result is in agreement with Raman spectroscopy measurements which will be published elsewhere. In order to obtain further information such as the lattice constant, the average crystallite size ($\langle D \rangle_{DRX}$), average residual strain ($\langle \varepsilon \rangle$), and phase amount, the whole set of Bragg reflections were refined using the Rietveld refinement method, using Jana2006 software.²⁷

Typical Rietveld refinement analyses are shown in Fig. 1 for the 30 and 7 mol% Ce-doped samples, as can be seen, the 30 mol% XRD pattern displays the tetragonal SnO₂ and the cubic CeO_2 phases. Lattice constants *a* and *c* of undoped SnO_2 were found at 4.73 and 3.18 Å, respectively. These values are in agreement with those expected from SnO₂ NPs reported in the literature.²⁸ However, the introduction of Ce ions to the SnO₂ matrix, leads to the monotonic increase of lattice constants, and consequently to the unit cell volume (V) increase of up to 10 mol% Ce content, as shown in Table 1, and graphically in Fig. 2. This fact was associated with the substitution of Sn⁴⁺ ions by Ce^{3+} and/or Ce^{4+} ions, due to both Ce^{3+} (1.01 Å) and Ce^{4+} (0.87 Å) ions having ionic radii larger than Sn^{4+} (0.69 Å) (all for coordination VI).²⁹ This fact is in agreement with the substitutional solution of Ce to Sn ions in the rutile-type crystalline structure. On the other hand, the change of the volume rate as is



Fig. 1 (a) XRD of the Ce-doped SnO_2 nanoparticles from 0 to 30 mol%. The peaks identified by an asterisk (*) correspond to the formation of a secondary phase. Rietveld refinement of the (b) 30 and (c) 7 mol% Ce. The black points represent the experimental, the red line shows the theoretical data, and the blue bottom line the difference between them.

seen in Fig. 2(a) suggests a change in the Ce^{3+}/Ce^{4+} rate, therefore, a shift of the oxidized state of cerium ions from Ce^{3+} to Ce^{4+} , and for higher concentrations up to 10 mol%, the formation of the CeO_2 secondary phase, indicating the occurrence of a doping saturation limit, tuned by the change of valence state from Ce^{3+} to Ce^{4+} . This fact will be corroborated later by magnetic and XAS measurements. The average

Table 1	Parameters ob	tained from the	Rietveld refinemen	t of the X-ray	diffractogram of	Ce-doped SnO ₂	NPs
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Ce (mol%)	<i>a</i> (Å)	<i>c</i> (Å)	$V(\text{\AA}^3)$	$\langle D \rangle$ (nm)	$\langle \varepsilon \rangle$ (%)	CeO ₂ (%)
0	4.7344	3.1844	71.38	5.6	-0.385	0
1	4.7287	3.1831	71.17	4.1	0.023	0
2	4.7397	3.1887	71.64	3.9	0.380	0
5	4.7416	3.1918	71.76	3.7	1.376	0
7	4.7493	3.2036	72.26	3.7	1.191	0
10	4.7525	3.2017	72.31	3.3	0.989	0
20	4.7628	3.2110	72.84	2.8	1.864	${\sim}3$
25	4.7522	3.2216	72.75	2.4	1.784	${\sim}4$
30	4.7495	3.2151	72.52	2.2	1.565	${\sim}5$



Fig. 2 Evolution of the lattice volume (a), mean particle size $\langle D \rangle_{\text{DRX}}$ (b), and residual strain $\langle \epsilon \rangle$ for the Ce-doped SnO₂ nanoparticles (c).

crystallite size and residual strain were determined by the Williamson–Hall plot method implemented within the Jana2006 software.²⁷ The $\langle D \rangle_{\text{DRX}}$ has expected decreasing behavior with the Ce content change. This effect is correlated with surface segregation of the dopant ions on the particle surface.^{30,31} Due to the aliovalent nature of cerium (Ce³⁺ and Ce⁴⁺), the doping of the SnO₂ matrix can produce an undoubtedly different type of substitution process. However, when analyzing the literature, for example, in Ti⁴⁺-doped SnO₂ (ref. 32) and Al³⁺-doped SnO₂, both have continuous decreasing crystallite size with increasing doping concentration.³³ This suggests that the type of doping element has no dependence on the final particle size. This indicates that the monotonic

crystallite size is correlated with the surface segregation of the dopant ions on the particle surface,^{30,31} leading to an increase in the nanocrystals' surface energy, and reducing the average crystallite size with increasing dopant concentration. This migration process continued until it reached the nucleation regime where a second phase emerges.³⁴ Our samples change around 10%, and afterward, a tendency to be constant is expected as shown in Fig. 2(b). The average crystallite size reduction is accompanied by an increase of the residual strain (see Table 1) associated with the cerium ions located in sites incorporated into SnO₂ as a solid solution, provoking local distortions that lead to the breaking of the symmetry of the SnO₆ octahedral structure.

In order to assess the physical crystallite size, and compare it with the crystallite size obtained by XRD results, transmission electron microscopy images were obtained. Fig. 3 shows one of the TEM images obtained for the 10% Ce-doped SnO₂ NPs. The particle sizes were obtained from the TEM images, and subsequently, a particle size histogram was made using the Sturges method, for more details see ref. 35. The bottom-right inset in Fig. 3 provides the obtained histogram. As observed, this histogram is well modeled by a log-normal distribution: $(D) = 1/(\sqrt{2\pi} \sigma D) \exp(-(\ln^2(D/D_0))(2\sigma^2))$, the red line shows where D_0 is the median value and σ is the polydispersity index. The median particle size value was assessed using $D_{\text{TEM}} =$ $D_0 \exp(\sigma^2/2)$, and the value was estimated at 4.1 \pm 0.3 nm, which is slightly higher than that obtained by XRD suggesting a surface amorphous phase, which could be associated with Ceion surface segregation. Furthermore, an interplanar distance of ~ 0.33 nm was obtained (see left-top inset in Fig. 3) and is associated with the interplanar spacing of (110) planes.

Fig. 4(a) displays the magnetization (*M*) as a function of magnetic field (*H*) at room temperature for undoped and Cedoped SnO₂ NPs, after the subtraction between the diamagnetic background signal (holder sample) and the paramagnetic susceptibility (obtained from the fit at high magnetic field values, as seen in the inset of Fig. 4(a)), the room temperature ferromagnetism (RTF) contribution was obtained. With respect to pure SnO₂, an S-shape curve in the central evidence of a ferromagnetic contribution, with a magnetization saturation of $M_{\rm S} \sim 0.07 \times 10^{-3}$ emu g⁻¹, which is associated with the exchange coupling of the spins of electrons trapped in oxygen vacancies (F-center), mainly located on the particle surface, as



Fig. 3 High-resolution TEM image of the 10.0 mol% Ce-doped sample with the particle size histogram, where the interplanar distance of 0.33 nm corresponds to the (110) plane of the SnO₂ rutile-type structure.



Fig. 4 (a) Magnetization curves for the undoped and Ce-doped SnO₂ NPs (the inset shows the magnetization signal before subtracting the paramagnetic and diamagnetic contributions in the 10% Ce-doped NPs). (b) Saturation magnetization of the ferromagnetic contribution as a function of the Ce content. (c) Percent of Ce³⁺-ions as a function of the Ce content.

was discussed in a previous publication.³⁶ As can be seen, from Fig. 4(b), we have a substantial increase of the $M_{\rm S}$ (~129%) from ~0.07 × 10⁻³ to ~0.16 × 10⁻³ emu g⁻¹, mapped from 0 to 1 mol% Ce content. A plausible source of the strengthening of the ferromagnetism, can be associated with the exchange interaction of cerium ions with non-zero magnetic moment (4f¹) *via* the F-centers. The latter, are electrons trapped in oxygen vacancies, likewise, the strengthening of the

ferromagnetism is linked to the amount of V_0 which is in agreement with the PL results associated to V_0 in the same samples.³⁷ Furthermore, above the 1 mol% concentration the magnetic saturation shows a monotonic decrease of up to 10 mol%, which is associated with the reduction of the Fcenters due to an enrichment of the surface segregation and the change of oxidized state from Ce³⁺ to Ce⁴⁺ after increasing the doping content. The latter is produced during the synthesis process, where the amount of Ce³⁺ (*x*) and Ce⁴⁺ (*y*) in the SnO₂ host matrix, and Ce⁴⁺ (*z*) in the CeO₂ secondary phase depends on the Ce content according to the following equation,

$$Sn_{2}^{2+}(C_{6}O_{7}H_{4}) + (x + y + z)Ce(NO_{3})_{3} \xrightarrow{IA} SnCe_{x}^{4+}Ce_{y}^{3+}O_{2} + zCeO_{2}$$
(3)

However, the maximum value of the $M_{\rm S}$ was obtained for 10% Ce content, which is interesting in itself. The origin of the room temperature ferromagnetism is still unclear, but surface defects seem to be crucial for its appearance due to small CeO₂ NPs (which has been evidenced by Raman measurements) with a high concentration of oxygen vacancies (V_O), which are the main type of defects in ceria NPs.³⁸ The oxygen vacancies provoke the rise of Ce³⁺ ions to form Ce⁴⁺-V_O-Ce³⁺ complexes. These ingredients could be the key for the RTF observed in this sample.^{39,40} On the other hand, from the paramagnetic susceptibility ($\chi_{\rm P} = M/H$) and using eqn (1), the concentration of Ce³⁺, x(Ce³⁺), was determined and is displayed in Fig. 4(c).

$$\mu_{\rm eff}^2 x ({\rm Ce}^{3+}) N_{\rm A} \mu_{\rm B} = 3A k_{\rm B} T \chi_{\rm P} \tag{4}$$

Here, μ_{eff} is the effective magnetic moment (2.4 μ_{B} for the Ce³⁺ions), N_{A} the Avogadro's constant, k_{B} the Boltzmann's constant, *A* the molecular weight, and *T* the temperature. The result suggests a high concentration of Ce³⁺ (~45%) in the 1% cerium



Fig. 5 DC susceptibility (χ_{DC}) as a function of temperature (T) recorded at H = 1 kOe. Symbols represent experimental data, whereas solid lines show fitting using the Curie-Weiss law.

Table 2 Magnetic parameters calculated from modified Curie–Weiss law fittings

Ce (mol%)	$C \text{ (emu K g}^{-1} \text{ Oe}^{-1}\text{)}$	$\theta_{\rm cw}$	$\chi_0 \left(emu \; g^{-1} \; Oe^{-1}\right)$	$\mu_{ m eff}$
1	0.0 10=6	15.05	15.0 × 10 ⁻⁸	1.00
1	9.8×10^{-1}	-15.35	15.2×10^{-1}	1.09
2	8.3×10^{-6}	-9.93	$6.9 imes10^{-8}$	0.71
5	$9.8 imes10^{-6}$	-7.64	$7.7 imes10^{-8}$	0.49
7	16.6×10^{-6}	-15.26	8.0×10^{-8}	0.54
10	34.7×10^{-6}	-50.48	$31.2 imes10^{-8}$	0.65
20	5.5×10^{-6}	-5.07	15.1×10^{-8}	0.18
25	11.1×10^{-6}	-4.31	$24.3 imes10^{-8}$	0.24
30	$5.7 imes10^{-6}$	-7.83	22.6×10^{-8}	0.16

content, decreasing with the cerium content. On the other hand, Fig. 5 shows the susceptibility as a function of the temperature for all samples. As we can be seen in Fig. 5 a positive susceptibility for the undoped SnO₂ sample is seen, suggesting Pauli paramagnetic behavior. Meanwhile, all doped samples present typically paramagnetic or paramagnetic-like behavior. In this regard, the thermal dependence of the susceptibility was modeled by the Curie-Weiss law. It means that we have the coexistence of paramagnetism with a ferromagnetic-like contribution in samples with low Fe content. The data of all samples were fitted to the modified Curie-Weiss law ($\chi = \chi_0 + C/(T - \theta_{cw})$), where C is the Curie constant, θ_{cw} is the Curie-Weiss temperature, and χ_0 is the temperatureindependent susceptibility. The obtained parameters are listed in Table 2.

The high negative values of θ_{cw} (see Table 2) suggest the occurrence of antiferromagnetic correlations among the Ceions dispersed in the SnO2 matrix. The effective magnetic moments (μ_{eff}) calculated from the Curie constant are in the range 0.71 to 0.65 $\mu_{\rm B}$ (see Table 2). Due to the expected magnetic moment values for the Ce³⁺ ($\mu_{eff} = 2.54 \mu_B$) and Ce⁴⁺ ($\mu_{eff} = 0 \mu_B$) the low and decreasing tendency of the effective magnetic moment suggests a monotonic reduction of the Ce³⁺-ions as the cerium content increases.



Fig. 6 Room-temperature Ce L3-edge XANES spectra obtained for cerium-doped SnO₂ NPs.

Fig. 6(a) shows the X-ray absorption near-edge spectra (XANES) at Ce L3-edge, for the Ce-doped SnO₂ NPs. In order to study the valence state of the cerium ions all spectra were compared with the spectra obtained for the standard



Fig. 7 (a) Fit of the XANES spectrum of the 7 mol% cerium-doped SnO₂ NPs using three components. (b) Percent of Ce³⁺-ions as function of the Ce content obtained from X-ray absorption.

compounds $Ce(C_2H_3O_2)_3\cdot H_2O$ and $CeO_2,$ for more details see the previous work.^{41}

A visual inspection of the spectra's shape suggests the coexistence of Ce^{3+} and Ce^{4+} in all Ce-doped samples, with a clear reduction in the amount of Ce^{3+} -ions as the Ce content increases. However, *via* deconvolution of the recorded spectra, the quantitative Ce^{3+} population can be assessed, as already reported in the literature.^{42,43} In this regard, the spectra have been fitted using three peaks, one for the Ce^{3+} -ions (located at 5727 eV) and two for the Ce^{4+} -ions (located at 5731 eV and 5738 eV). A typical fitting of the XANES spectra for the 7 mol% cerium-doped SnO₂ NPs is shown in Fig. 7(a).

In Fig. 7(a), the green peak in the pre-edge region is related to the final states of 2p5d with delocalized d character at the bottom of the conduction band.⁴² The blue peaks located at 5731 and 5738 eV are associated with Ce^{4+} and correspond to $2p^4f^15d^1$ and $2p^4f^05d^1$ final states, respectively.⁴³ The orange peak located at 5727 eV has its origin in the specific white-line spectra of Ce^{3+} ions and has been used in the estimation of their concentration.^{42,43}

Estimation of the Ce³⁺-content for all doped samples has been obtained using the spectral area ratio of the peaks Ce³⁺/ (Ce³⁺ + Ce⁴⁺). Our results show a higher concentration of Ce³⁺ for the 1 mol% (~80%). This value is larger when compared to the value obtained from the magnetic measurements (~45%). This fact could be closely linked to the occurrence of antiferromagnetic interactions, leading to antiparallel alignment $\uparrow\downarrow\cdots\uparrow\downarrow$ of neighboring Ce-ions. This antiparallel alignment drives an apparent reduction in the effective magnetic moment of the Ce-ions in the paramagnetic phase, leading to low concentration values of the Ce³⁺. Moreover, calculated XANES

spectra for Ce-doped SnO2 at Ce L3-edge using ab initio FEFF code are also shown in Fig. 8. The input files for the FEFF code were initially generated using a crystallographic model according to the XRD measurements where Sn4+ ions are replaced by Ce⁴⁺ ions. The cluster used in this calculation radius presents a radius of 6.0 Å with 68 atoms around the absorber Ce atom, which is also shown in Fig. 8. According to this crystallographic model, Ce atoms in Sn sites are surrounded by 6 O atoms in the first shell and 2 Sn atoms in the second shell. As can be seen in this figure, the calculated XANES spectra with 6 O atoms in the first shell shows a higher intensity for the feature around 5733 eV, which is, as stated above, directly related to Ce^{4+} ions. The calculated XANES spectra with 4 O atoms in the first shell (2 O vacancies) and 2 Sn atoms in the second shell, and the calculated XANES spectra with 4 O atoms in the first shell (2 O vacancies) and 1 Ce and 1 Sn atom in the second shell, are also shown in Fig. 8. As can be seen in this figure, a decrease in the intensity for the feature around 5738 eV is observed.

Hence, for two O vacancies we have, in the former case, one Ce^{3+} ion whereas in the latter we have two Ce^{3+} ions. It means that each Ce atom must be coordinated with two O vacancies to give a Ce^{3+} ion as is shown in the DFT *ab initio* simulation. Further, the calculated XANES spectra reproduce satisfactorily the experimental spectra for concentrations lower than 10% Ce-doped samples. In other words, O vacancies are associated with the decreasing oxidation state from +4 to +3, demonstrating a majority of Ce^{3+} ions in these samples. Thus, the reduced tendency of the amount of Ce^{3+} -ions as a function of Ce content is in agreement with magnetic and X-ray absorption measurements.

B. *Ab initio* investigation of the oxidation state of the dopant Ce ions

We performed *ab initio* simulations of Ce-doped SnO_2 to provide an atomistic understanding of the ingredients that determine the oxidation state of the Ce-ions. In our



Fig. 9 The more likely defect structures after optimization by DFT calculations of Ce-doped SnO_2 : (a) 1Ce_{Sn} , (b) $1\text{Ce}_{\text{Sn}}1\text{V}_{\text{O}}$, (c) $1\text{Ce}_{\text{Sn}}2\text{V}_{\text{O}}$, (d) $2\text{Ce}_{\text{Sn}}1\text{V}_{\text{O}}$, (e) $2\text{Ce}_{\text{Sn}}2\text{V}_{\text{O}}$ and (f) $3\text{Ce}_{\text{Sn}}2\text{V}_{\text{O}}$. The tin, oxygen and cerium atoms are represented by gray, red and green colors. The dashed circle is a quide for the eye of the surrounding vacancy.

respectively.



Fig. 8 Theoretical XANES spectra for Ce-doped SnO₂ samples,

considering three clusters: 6 O atoms in the first shell and 2 Sn atoms

in the second shell (black solid line), 4 O atoms in the first shell and 2 Sn

atoms in the second shell (purple solid line), and 4 O atoms and 1 Sn and 1 Ce atom in the second shell (magenta solid line), around Ce

absorption. The cluster used in each calculation is also illustrated,

where green, red, gray and ivory spheres represent the Ce absorber

atom, O atoms, Sn atoms and Ce atom in the second shell,

Table 3 Point defect structures as exhibited in Fig. 9, defect energy formation (E_t), oxidation state of Ce atom (OS), magnetization energy (ME), total spin angular moment (SOM) at the level of the PBE/GGA scheme

Structures	$E_{\rm f}$ (eV)	OS	ME (meV)	SOM
1Ce _{Sp}	7.237	+4	_	0
1Ce _{Sn} 1V _O	1.651	+4	_	0
1Ce _{Sn} 2V _O	2.095	+3	_	1
2Ce _{sn} 1V _O	1.268	+4, +4	_	0
2Ce _{Sn} 2V _O	4.510	+3, +3	1.67	2
$3Ce_{Sn}2V_{O}$	1.918	+3, +4, +3	—	2

calculations, we adopted a rutile-like SnO_2 structure with a 3 × 3 × 3 supercell,⁴⁴ containing 54 and 108 atoms of Sn and O, respectively, in its pristine form. Thus, we created a variety of point defects inside it. Our experiments indicated substitutional doping of Ce atoms at the Sn sites and the presence of oxygen vacancies. The investigated point defects, shown in Fig. 9, are taken into account from these experimental insights.

Panel (a) shows a single Ce atom doped into the SnO₂ at the Sn sites, namely $1Ce_{Sn}$. This simulation reveals the absence of a magnetic moment, indicating that all Ce valence electrons are shared in covalent bonds, characterizing the +4 oxidation state. Panel (b) depicts a point defect with a single oxygen vacancy surrounding a Ce-ion substituting a Sn site, namely, $1Ce_{Sn}1V_O$. Again, the simulation results in the oxidation state +4 for the Ce ion. Thus, a single-vacancy neighboring the Ce-ion does not suffice to modify its oxidation state. However, after adding two neighboring oxygen vacancies to a Ce impurity ($1Ce2V_O$), as depicted in Fig. 9(c), we found an unpaired localized valence electron in the Ce-ion, resulting in a magnetic moment characteristic of Ce³⁺. Thus, the addition of oxygen vacancies to Ce-

doped SnO_2 was shown to be a determining factor to tune the oxidation state of Ce-ions.

In our investigation, we also consider complexes comprised of two and three Ce substitutional impurities surrounded by one and two oxygen vacancies, as depicted in Fig. 9(d)–(f). The $2Ce_{sn}1V_O$ complex (panel d) results in the oxidation state +4 for all ions. However, the $2Ce_{sn}2V_O$ complex has all Ce ions in the oxidation state +3. Thus, for the complexes, our simulations also indicate the importance of oxygen vacancies for determining the Ce oxidation state. However, different to single Ce impurities, one oxygen vacancy per cerium atom results in Ce³⁺ ions. Finally, we show in panel (f) a complex with three Ce substitutional impurities and two vacancies. Here, the central Ce atoms present a +4 oxidation state, whereas the other ones are in the +3 oxidation state. This simulation indicates the possibility of Ce³⁺ and Ce⁴⁺ co-existing in the same complex. Table 3 presents these results more concisely.

Thus, our *ab initio* simulations demonstrate that the presence of oxygen vacancies is a determining factor for the oxidation state of the doped Ce impurity. In this sense, according to our calculations, in the absence (presence) of the oxygen vacancies the preferential oxidation state for cerium atoms is +4 (+3). Furthermore, it is worth noting that the possibility of Ce³⁺ and Ce⁴⁺ co-existing in the sample depends on the Ce/V_o ratio, whereas an increased amount of oxygen vacancies favors the oxidation state Ce³⁺. Regarding the experimental result, when the Ce doping content is increased, the concentration of Ce³⁺ ions decreases. This drives the decrease of oxygen vacancies, reducing the presence of the neighboring oxygen vacancies around the cerium ions, in agreement with previous work, where they have shown that this occurs mainly on the particle shell.^{31,37}



Fig. 10 The local density of states (LDOS) for Sn (1), O (2) and Ce (3) for the respective structure displayed in each column: (a) $1Ce_{Sn}$, (b) $1Ce_{Sn}1V_{O}$ and (c) $1Ce_{Sn}2V_{O}$.



Fig. 11 The local density of states (LDOS) for Sn (1), O (2) and Ce (3) for the respective structure displayed in each column: (a) $2Ce_{sn}1V_O$, (b) $2Ce_{Sn}2V_O$ and (c) $3Ce_{Sn}2V_O$. In the last row, we display the LDOS separately by highlighting each Ce-ion.



Fig. 12 The total spin density polarization (SDP) for the main type of defect 2Ce_{sn2}V_O with FM (a), and AFM (b) configurations. We depicted the spin-resolved charge density differences (blue – spin-up, and green – spin-down) and isosurface value 0.005 eV Å⁻¹. The total density of states (DOS) for the main type of defects $2 Ce_{\text{Sn2}}V_{\text{O}}$ with FM (c), and AFM (d), configurations.

Here, we calculated the formation energy $E_{\rm f}$ with the following expression:⁴⁵

$$E_{\rm f} = E_{\rm D} - E_{\rm P} + n_{\rm Sn}\mu_{\rm Sn} + n_{\rm O}\mu_{\rm O} - n_{\rm Ce}\mu_{\rm Ce}$$
(5)

where $E_{\rm D}$ and $E_{\rm P}$ are the total energies of the supercells with and without defects, respectively; $\mu_{\rm Sn}$, $\mu_{\rm O}$ and $\mu_{\rm Ce}$ are the elemental chemical potentials; $n_{\rm Sn}$, $n_{\rm O}$ and $n_{\rm Ce}$ are the number of tin atoms replaced by cerium atoms, the number of oxygen vacancies, and the number of cerium atoms in the supercell, respectively.

According to our results, the formation energies are in the following order: $1Ce_{sn}1V_{O}$ ($E_{f} = 1.65 \text{ eV}$) < $3Ce_{sn}2V_{O}$ ($E_{f} = 1.93$ eV) < 1 $Ce_{Sn}2V_O$ ($E_f = 2.09 eV$) < 2 $Ce_{Sn}2V_O$ ($E_f = 4.51 eV$) < 1 Ce_{Sn} $(E_{\rm f} = 7.23 \text{ eV})$. The 1Ce_{sn}1V_O is the lowest $E_{\rm f}$, suggesting the most probable defect to occur in our experimental measurements. Also, Fig. 10 depicts the local density of states (LDOS) for 1Ce, 1Ce1Vo, and 1Ce2Vo, where we note that the effect of Ce impurities and surrounding oxygen vacancies, originates from the impurity electronic states between the conduction and the valence bands. In addition, the d-Sn has a stronger contribution at the valence band, and also, p-O and f-Ce. Another consequence to introduce oxygen vacancies in the system is the emergence of the s-Sn impurity states in O and Sn LDOS, which increases with the introduction of more O-vacancies, see the top and center panels of Fig. 10. As long as we are concerned with the Ce oxidation state, it is clear from panels (a-3), (b-3) and (c-3)that we need two oxygen vacancies to reduce the Ce oxidation from +4 to +3, in agreement with the equation:

$$\operatorname{Sn_2}^{4+}\operatorname{Ce_2}^{4+}\operatorname{O_8}^{2-} + 2e^- \to \operatorname{Sn_2}^{4+}\operatorname{Ce_2}^{3+}\operatorname{O_7}^{2-} \Box_{V_0} + \operatorname{O}^{2-}.$$
 (6)

Eqn (6) suggests that the formation of the unstable Ce^{3+} requires two electrons for the reaction. This can be confirmed by the only fully occupied majority spin-up states without its minority counterpart Fig. 10(c-3), which is also accompanied by one peak of the corresponding p-O orbital just below of the Fermi energy Fig. 10(c-2). On the other configurations (Fig. 10(a and b)) Ce ions prefer to stay at valence +4 and the no polarized states signature is also exhibited by the corresponding p-O orbitals (*cf.* Fig. 10(a-2) and (b-2) with (a-3) and (b-3)).

Also, the increase of the cerium content was analyzed $(2Ce_{sn}2V_O \text{ and } 3Ce_{sn}2V_O)$. As displayed in Fig. 11, in the case of two Ce ions, both display the same oxidation state. Therefore, it shows that each Ce ion needs to be correlated, at least, with two oxygen vacancies to exhibit +3 valence, as a preferential state. Therefore, only Fig. 11(b-3) and (c-3), show two atoms with Ce-f orbitals full spin-up polarized. Furthermore, this last effect is accompanied by the p-O orbitals, as can be seen by *cf.* Fig. 11(b-2) and (c-3).

To elucidate the magnetic properties of the Ce-doped SnO_2 systems, we study the local magnetic centers and spin-resolved charge densities for the lowest energy defect configuration to obtain the spin density polarization (SDP). In these calculations, we fix the initial magnetic moments for the ferromagnetic (FM) or antiferromagnetic (AFM) alignments. Fig. 12(a) and (b) show the SDP of the $2\text{Ce}_{\text{Sn}}2\text{V}_{\text{O}}$ defect. For the FM case, although

the SDP nearby Ce1 becomes mainly negative (green-colored region) and the SDP of Ce2 is mainly positive (blue colored region). The negative SDP around the first vacancy ($1Ce_{sn}1V_O$) overcomes the opposite positive SDP around the second vacancy ($2Ce_{sn}2V_O$), favoring a net resultant polarization. Meanwhile, for the AFM configuration, the SDP nearby each cerium atom, and the oxygen atoms nearby each Ce impurity, display the same opposite spin polarization.

Moreover, the spin-resolved total density of states was plotted for both configurations (see Fig. 12(c) and (d)). A total energy difference between both magnetic configurations $\Delta E_{\text{mag}} = E_{\text{AFM}} - E_{\text{FM}} = 1$ meV. The small positive value suggests that FM is more favorable than the AFM state for this type of defect, as observed in the experimental part for Ce³⁺-ions, where the paramagnetic response is dominant.

IV. Conclusions

Undoped and cerium-doped (0 to 30 mol%) SnO2 NPs with high crystallinity have been successfully synthesized by a polymer precursor method. A mean crystallite size of 6 nm has been determined for the undoped SnO₂ NPs, meanwhile, a small size of 2 nm was determined for the 30 mol% Ce-doped. Samples showing the formation of a single phase rutile-type structure have been obtained for doping amounts from 0 to 10 mol%, and the formation of a secondary CeO_2 phase was determined for samples with cerium concentration above 10 mol%. Roomtemperature ferromagnetism associated with oxygen vacancies in the single SnO₂ sample was found. Meanwhile, for the cerium doped impurity, the FM is increased for the 1 mol% Ce doped sample, which was assigned to the long-range interaction of cerium ions with nonzero magnetic moment (4f¹) mediated by the F-centers. On the other hand, we found a high $M_{\rm S}$ of 0.27 \times 10^{-3} emu g⁻¹ in the 10 mol% of SnO₂ NPs, this fact can be associated to ceria NPs. The coexistence of Ce³⁺ and Ce⁴⁺ has been corroborated from the Ce L3-edge XANES data analysis, with a decreasing tendency of the Ce^{3+}/Ce^{4+} ratio as the Ce content increases. This fact is in good agreement with the magnetic measurement. The XANES was theoretically simulated via the FEFF code, our results show that the oxygen vacancies surrounding the cerium ions are the key to obtaining the +3 state in our samples as determined from the experimental part and corroborated by ab initio calculations. It was confirmed theoretically that for low concentrations of doping, the Ce atoms need to be correlated at least with two oxygen vacancies to exhibit the +3 state, as a preferential oxidation state, and after the Ce content in the host sample increases, the Ce⁴⁺ ions take over the Ce³⁺ ones, since after a critical doping level, Ce atoms can no longer coordinate with more oxygen vacancies. The presented results pave the way for tuning the magnetic properties of oxides through the control of the oxidation state of impurities.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by CONCYTEC – FONDE-CYT within the framework of the call E038-01 [contract No. 07-2019-FONDECYT-BM-INC. INV]. LVL thanks PNPD/CAPES for grant number 88887.319028/2019-00. L. Cabral thanks FAPESP (Fundação de Amparo à Pesquisa do Estado de São Paulo) for grant number 2018/20729-9. M. P. Lima thanks FAPESP (grant number 17/02317-2). AM thanks the FAPESP (grant number 2013/12993-4). JAHC want to thank the Brazilian agencies CNPq [grants number 301455/2017-1, 443652/2018-0] and FAPDF [grant number 00193.0000151/2019-20] for the financial support. The authors thank to the Brazilian Synchrotron Light Laboratory (LNLS) for the beamtime granted (proposal 20160072) to carry out the X-ray absorption measurements.

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