PCCP



PAPER View Article Online



Cite this: Phys. Chem. Chem. Phys., 2016, 18, 13974

Accepted 11th April 2016
DOI: 10.1039/c6cp01525e

Received 5th March 2016,

www.rsc.org/pccp

Understanding the anomalous behavior of Vegard's law in $Ce_{1-x}M_xO_2$ (M = Sn and Ti; $0 < x \le 0.5$) solid solutions†

Tinku Baidya,‡*^{ab} Parthasarathi Bera,*^c Oliver Kröcher,^{bd} Olga Safonova,^b Paula M. Abdala,^e Birgit Gerke,^f Rainer Pöttgen,^f Kaustubh R. Priolkar^g and Tapas Kumar Mandal^h

The dependence of the lattice parameter on dopant concentration in $Ce_{1-x}M_xO_2$ (M = Sn and Ti) solid solutions is not linear. A change towards a steeper slope is observed around $x \sim 0.35$, though the fluorite structure (space group Fm3m) is preserved up to x = 0.5. This phenomenon has not been observed for $Ce_{1-x}Zr_xO_2$ solid solutions showing a perfectly linear decrease of the lattice parameter up to x = 0.5. In order to understand this behavior, the oxidation state of the metal ions, the disorder in the oxygen substructure and the nature of metal-oxygen bonds have been analyzed by XPS, 119 Sn Mössbauer spectroscopy and X-ray absorption spectroscopy. It is observed that the first Sn-O coordination shell in $Ce_{1-x}Sn_xO_2$ is more compact and less flexible than that of Ce-O. The Sn coordination remains symmetric with eight equivalent, shorter Sn-O bonds, while Ce-O coordination gradually splits into a range of eight non-equivalent bonds compensating for the difference in the ionic radii of Ce^{4+} and Sn^{4+} . Thus, a long-range effect of Sn doping is hardly extended throughout the lattice in $Ce_{1-x}Sn_xO_2$. In contrast, for $Ce_{1-x}Zr_xO_2$ solid solutions, both Ce and Zr have similar local coordination creating similar rearrangement of the oxygen substructure and showing a linear lattice parameter decrease up to 50% Zr substitution. We suggest that the localized effect of Sn substitution due to its higher electronegativity may be responsible for the deviation from Vegard's law in $Ce_{1-x}Sn_xO_2$ solid solutions.

Introduction

CeO₂ has been extensively studied in the last few years with respect to its manifold functionalities. It is an attractive material for various catalytic, electronic, optical, electrical, electrochemical, gas sensing, and corrosion resistant applications. ^{1–14} CeO₂ based materials have been useful in auto-exhaust catalysis, hydrogen

production, H₂-O₂ recombination, or electrodes in fuel cells. 1-7,15-19 In auto-exhaust catalysis, CeO2 has been recognized as an important material because of its unique redox properties. Due to their excellent oxygen storage/release properties, CeO₂ based oxides have been used as excellent supports for noble as well as transition metals promoting various catalytic reactions. The oxygen storage capacity (OSC) of CeO2 plays a unique role in mediating crucial oxygen exchange processes during catalytic reactions in auto-exhaust catalysis. 1,20 In the efforts to enhance the OSC of CeO₂ by various cation substitution reactions, the tetravalent cations such as Ti4+, Sn4+, and Zr4+ have been found to be effective in creating more labile oxygen through structural changes in the Ce_{1-x}M_xO₂ solid solutions. 4,21,22 Such substitutions of isovalent as well as aliovalent transition metals into the CeO₂ matrix enhance the reducibility, thermal stability, oxygen storage capacity and catalytic activities of CeO2.4 The abovementioned solid solutions have been used as supports for noble metals like Pt, Pd, Rh, and Ru leading to a better catalytic activity for a variety of reactions at comparatively low temperatures.4

Hegde and coworkers have intensively investigated the structural and redox properties of $Ce_{1-x}M_xO_2$ (M = Sn, Ti and Zr) solid solutions both experimentally and theoretically.^{23–27} It has been

^a Solid State and Structural Chemistry Unit, Indian Institute of Science, Bangalore 560012. India

^b Paul Scherrer Institut, 5253 Villigen, Switzerland

^c Surface Engineering Division, CSIR — National Aerospace Laboratories, Bangalore 560017, India. E-mail: partho@nal.res.in

^d Ecole Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland

^e SNBL, European Synchrotron Radiation Facility, 38043 Grenoble, France

f Institut für Anorganische und Analytische Chemie, Universität Münster, Corrensstrasse 30, 48149 Münster, Germany

g Department of Physics, Goa University, Taleigao Plateau, Goa 403206, India

h Department of Chemistry and Centre of Nanotechnology, Indian Institute of Technology Roorkee, Roorkee 247667, India

 $[\]dagger$ Electronic supplementary information (ESI) available. See DOI: 10.1039/c6cp01525e

[‡] Present address: Department of Mechanical and Aerospace Engineering, University of California, 9500 Gilman Drive, San Diego, CA 92093, USA. Email: tinku26@gmail.com

Paper

demonstrated that the substitution of isovalent cations in CeO_2 creates significant distortions in the anion substructure, splitting eight equidistant oxygen neighbors in fluorite type CeO_2 into 4+4 neighbors with different distances. The extent of dissolution of foreign cations forming solid solutions is confirmed by a linear change of the lattice parameter of the parent oxide as postulated by Vegard's law. However, careful examination of the lattice parameters at different concentrations of Ti, Zr and Sn have demonstrated unusual behavior. Although all these substituents are tetravalent, the change in the lattice parameter with the increase of Ti or Sn concentration deviates from linearity, while for Zr it is perfectly linear up to x=0.5. This observation certainly questions the universal validity of Vegard's law in the present solid solutions and has motivated us to investigate this phenomenon in more detail.

Deviations from Vegard's law in solid solutions of ionic salts and ionic oxides have previously been discussed in the literature. 29,30 The main explanations are related to limited solubility and microscopic cluster formation. Similar deviations from Vegard's type behavior have been observed for thin films of various solids. 31,32 The present study is focused on the understanding of the anomalous behavior of the lattice parameter with the dopant concentration in $Ce_{1-x}Sn_xO_2$ solid solutions. The detailed characterization of these solutions has been done by X-ray powder diffraction (XRD), UV-Vis spectroscopy, X-ray photoelectron spectroscopy (XPS), ^{119}Sn Mössbauer spectroscopy and X-ray absorption spectroscopy (XAS).

Experimental details

 $Ce_{1-x}Sn_xO_2$ (x=0.1-0.5) oxides were prepared by the solution combustion method using ceric ammonium nitrate ((NH₄)₂-Ce(NO₃)₆·6H₂O), tin oxalate (SnC₂O₄), and glycine (C₂H₅NO₂) as precursors. The detailed method of preparation has been reported earlier.²³ Similarly, $Ce_{1-x}Zr_xO_2$ oxides have been prepared by using stoichiometric amounts of ceric ammonium nitrate, zirconium nitrate (Zr(NO₃)₄·xH₂O) and glycine.²⁷ Using a similar method, $Ce_{1-x}Ti_xO_2$ (x=0.1-0.4) oxides were prepared by taking stoichiometric amounts of ceric ammonium nitrate, titanyl nitrate (TiO(NO₃)₂), and glycine.²⁵

Transmission electron microscopy (TEM) images were collected using a Hitachi HD-2700CS electron microscope with a cold field emission gun operated at 200 kV, a CEOS corrector system and an EDXS detector. For Ce and Sn elemental analysis, respective CeL α and SnL α emission lines were considered.

All bulk samples of the $Ce_{1-x}M_xO_2$ (M = Sn, Ti and Zr) solid solutions were characterized by powder XRD employing a PANalytical X'Pert PRO X-Ray diffractometer operated under CuK α radiation of 1.5418 Å wavelength at 40 kV and 30 mA in the 2θ range $20-80^\circ$.

UV-Vis diffuse reflectance spectra (DRS) of $Ce_{1-x}M_xO_2$ (M = Sn) solid solutions were recorded using a Shimadzu UV-2450 UV-Vis spectrophotometer in the wavelength range of 200–800 nm. For the UV-Vis DRS measurements, samples were mixed with BaSO₄ in the sample to the BaSO₄ ratio of 1:100.

The recorded reflectance data were converted to absorbance using the Kubelka–Munk (K–M)³³ function. For the estimation of band gaps, a direct transition was considered in a Tauc plot³⁴ with $(\alpha h \nu)^2$ vs. $h\nu$.

The XPS of $Ce_{1-x}M_xO_2$ (M = Sn and Ti) solid solutions were recorded using a SPECS spectrometer non-monochromatic AlK α radiation (1486.6 eV) as an X-ray source operated at 150 W (12.5 kV and 12 mA). All the spectra were obtained with a pass energy of 25 eV and a step increase of 0.05 eV. The CasaXPS program was employed for curve-fitting of the Ce3d core level spectra into several components with Gaussian–Lorentzian peaks after Shirley background subtraction. Peak positions, spin–orbit splitting, doublet intensity ratios, and full width at half maximum (FWHM) were fixed as given in the literature. 35

A Ca^{119m}SnO₃ source was used for the Mössbauer spectroscopic experiments. The measurements were carried out at ambient temperature. A palladium foil of 0.05 mm thickness was used to reduce the Sn K X-rays concurrently emitted by this source. The samples were enclosed in small PMMA containers at a thickness corresponding to about 10 mg Sn per cm². Fitting of the data was done by using the Normos-90 program package.³⁶

X-ray absorption spectra at Sn and Ce K-edges were measured at SNBL (BM01B, European Synchrotron Radiation Facility, Grenoble, France) using the Si(111) double crystal monochromator. The sizes of the X-ray beam in the horizontal and vertical directions were 8 mm and 0.6 mm, respectively, giving the total beam intensity on the sample of about 10¹⁰ ph per s. Three ionization chambers were used and they were filled with Ar and Kr to produce the absorption of 20% for I_0 , 50% for I_t and I_{ref} , where I_0 , I_t and I_{ref} are intensities of incident, transmitted and reference beams, respectively. An optimal amount of sample powders was mixed with cellulose and pressed into pellets of 1.3 cm in diameter to measure the spectra in the transmission mode. The XAS were acquired in the continuous scanning mode. The as-recorded Ce and Sn K edge XAS data were reduced following the standard procedures in the Demeter program package.37

Ce $\rm L_3$ -edge X-ray absorption near edge spectroscopy (XANES) measurements were performed at SuperXAS beamline of the Swiss Light Source (PSI, Villigen, Switzerland). The incident photon beam was selected using a Si(111) double crystal monochromator and the rejection of higher harmonics and focusing were achieved using a Si-coated collimating mirror at 2.8 mrad and a Rh-coated toroidal mirror at 2.8 mrad. The beamline was calibrated using a V foil (V K-edge at 5465 eV). The resulting spectra were reduced following standard procedures in the Demeter program package. 37

Results

TEM-EDXS studies

All $Ce_{1-x}M_xO_2$ (M = Sn, Ti, Zr) solid solutions crystallize in a single-phase with the fluorite structure. All of them are exceptionally homogeneous down to the nanometer scale, most probably at the atomic level. A typical example of the elemental maps

Ce_1 Sn 1 Sn 2 Sn 2

Fig. 1 Elemental maps of Ce and Sn obtained by TEM analysis in different regions (900k magnification; image size -0.0001×0.0001 mm).

of Ce and Sn in $Ce_{0.5}Sn_{0.5}O_2$ obtained by TEM-EDXS is shown in Fig. 1.

XRD studies

PCCP

The detailed structural characterizations and Rietveld refinements are presented in earlier reports. 23,25,27 The dependence of the lattice parameter ν s. the Sn concentration in the $\mathrm{Ce}_{1-x}\mathrm{Sn}_x\mathrm{O}_2$ solid solution is shown in Fig. 2a. The lattice parameter decreases monotonically until the Sn concentration reaches 35 at% followed by a linear decrease but with a steeper slope. Similar behavior occurs for the Ti substitution with a change of the slope at \sim 30 at%. In contrast, in $\mathrm{Ce}_{1-x}\mathrm{Zr}_x\mathrm{O}_2$ solid solution, the lattice parameter decreases with a constant linear slope up to at least 50 at% of Zr (Fig. 2c), clearly indicating that Sn^{4+} and Ti^{4+} are responsible for the observed anomalies. In the following discussion, we will focus only on a comparison of Sn and Zr substitutions.

UV-Vis DRS studies

The changes in the electronic structure of the $Ce_{1-x}Sn_xO_2$ oxides determined by UV-Vis spectra are presented in Fig. 3. It demonstrates that absorption edge is not significantly shifted, although, a major change is observed at \sim 300 nm. The new electronic transition peak fills up the valley with increasing Sn concentration, which is a clear indication of electronic transition due to charge transfer from O2p to Sn5s. Thus, there is a change in the metal–oxygen bonding nature gaining more covalency with increasing Sn concentration. However, the band gaps (2.81–2.92 eV) in these materials are slightly lower than pure CeO_2 and largely remain unaltered with progressive Sn-substitution (see Fig. S1, ESI†). A small change in the band gaps of the solid solutions can be originated from the changes in the nature of metal–oxygen bonding.

XPS studies

According to XRD, the fluorite-type structure remains preserved for both $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ solid solutions though they have different behaviors towards lattice parameter νs . Sn/Zr concentration. Consequently, the important question arises about

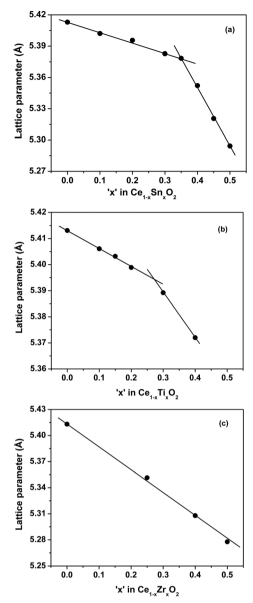


Fig. 2 Lattice parameter vs. substituent concentration in (a) $Ce_{1-x}Sn_xO_2$, (b) $Ce_{1-x}Ti_xO_2$ and (c) $Ce_{1-x}Zr_xO_2$ solid solutions.

the oxidation states of the constituent elements, especially for Ce. For this purpose, detailed XPS studies have been carried out.

The Ce3d core level spectra of $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ oxides are presented in Fig. 4(a and b). The spectral envelops indicate that the oxides contain both Ce^{4+} and Ce^{3+} species and they can be curve-fitted into several component peaks related to Ce^{4+} and Ce^{3+} , along with the corresponding satellites. Fig. 5 represents a typical curve-fitted Ce3d core level spectrum of $Ce_{0.65}Sn_{0.35}O_2$. In the figure, peaks labeled as v correspond to $3d_{5/2}$ photoemissions, whereas u peaks stand for related $3d_{3/2}$ photoemissions. The $u^{\prime\prime\prime}$ peak is relatively well separated from the rest of the spectrum and is characteristic of the presence of tetravalent cerium. In the case of $Ce_{0.65}Sn_{0.35}O_2$, $v^{\prime\prime\prime}$ and $u^{\prime\prime\prime}$ spin–orbit peaks at 898.0 and 916.5 eV, respectively, with 18.5 eV separation are assigned for the primary photoionization from

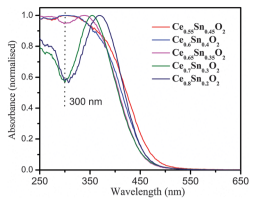
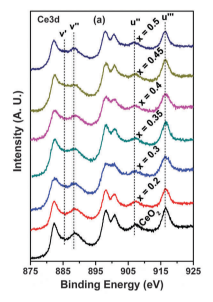


Fig. 3 UV-Vis spectra of Ce_{1-x}Sn_xO₂ oxides with different Sn concentrations.



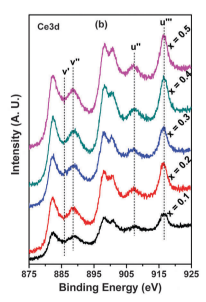
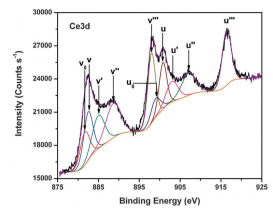


Fig. 4 XPS of Ce3d core level spectra in the solid solutions: (a) $Ce_{1-x}Sn_xO_2$ and (b) $Ce_{1-x}Zr_xO_2$.



Curve-fitted Ce3d core level spectrum of Ce_{0.65}Sn_{0.35}O_{2-δ}

Ce⁴⁺ with the Ce3d⁹4f⁰O2p⁶ final state. Lower binding energy states of v"-u" (888.6 and 907.1 eV) and v-u (882.5 and 900.8 eV) correspond to the shake-down satellite features of Ce3d⁹4f¹O2p⁵ and Ce3d⁹4f²O2p⁴ final states, respectively. ³⁸⁻⁴³ Peaks labeled as v_0 , v' and u_0 , u' at 881.6, 885.1 and 899.1, 903.2 eV, respectively, are related to the Ce^{3+} final states. It is to be noted that v'-u' spin orbit doublet peaks correspond to main photoionization from the Ce3d⁹4f¹O2p⁶ final state, whereas lower binding energy v₀-u₀ peaks are assigned to characteristic shake-down satellites of the Ce3d⁹4f²O2p⁵ final state. Relative surface concentrations of Ce⁴⁺ and Ce^{3+} species present in the $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ solid solutions have been calculated from curve-fitted spectra that are summarized in Table 1. Peak areas (A) of Ce4+ and Ce3+ components are commonly used to estimate their relative concentrations (C) in the oxides using the following equations:³⁸

$$A_{Ce^{4+}} = A_{v} + A_{u} + A_{v''} + A_{u''} + A_{v'''} + A_{u'''} + A_{u'''}$$
 (1)

$$A_{\text{Ce}^{3+}} = A_{v_0} + A_{u_0} + A_{v'} + A_{u'} \tag{2}$$

$$C_{\text{Ce}^{3+}} = \frac{A_{\text{Ce}^{3+}}}{A_{\text{Ce}^{4+}} + A_{\text{Ce}^{3+}}} \tag{3}$$

About 12% Ce exists in +3 oxidation state on the surface of pure CeO₂ in line with the literature reports. 44-46 With increasing

Table 1 Concentration of Ce^{3+} ion in $Ce_{1-x}M_xO_2$ (M = Sn and Zr) as calculated by curve-fitting of Ce3d core level spectra

$Ce_{1-x}Sn_xO_2$	$Ce^{3+}/(Ce^{3+} + Ce^{4+})$ (%)	$Ce^{3+}/(Ce + Sn)$ (%)			
CeO ₂	12	_			
$Ce_{0.8}Sn_{0.2}O_2$	22	17.6			
$Ce_{0.75}Sn_{0.25}O_2$	24	18.0			
$Ce_{0.7}Sn_{0.3}O_2$	26	18.2			
$Ce_{0.65}Sn_{0.35}O_2$	30	19.5			
$Ce_{0.6}Sn_{0.4}O_2$	27	16.2			
$Ce_{0.55}Sn_{0.45}O_2$	30	16.5			
$\mathrm{Ce_{0.5}Sn_{0.5}O_{2}}$	29	14.5			
$Ce_{1-x}Zr_xO_2$	$Ce^{3+}/(Ce^{3+} + Ce^{4+})$ (%)	$Ce^{3+}/(Ce + Zr)$ (%)			
$Ce_{0.9}Zr_{0.1}O_2$	24	21.6			
$Ce_{0.8}Zr_{0.2}O_2$	24	19.2			
$Ce_{0.7}Zr_{0.3}O_2$	18	12.6			
$Ce_{0.6}Zr_{0.4}O_2$	17	10.2			
$Ce_0 {}_5Zr_0 {}_5O_2$	14	5.0			

PCCP

Sn concentrations, the concentration of Ce^{3+} (Ce^{3+} /(Ce^{4+} + Ce^{3+})) gradually increases to 30% in $Ce_{1-x}Sn_xO_2$. Although, an apparent increase in the relative concentration of Ce^{3+} occurs, the relative concentration of Ce^{3+} with respect to the total number of atoms (Ce + Sn) have been found at an almost constant value of $\sim 18\%$. However, the relative surface concentration of Ce^{3+} with respect to the total one (Ce + Zr) in $Ce_{1-x}Zr_xO_2$ increases initially at 10% Ce^{3+} occurs and then gradually decreases with increasing Ce^{3+} concentration.

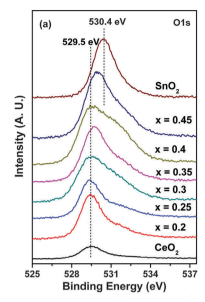
Sn3d and Zr3d core level spectra of samples from the $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ solid solutions are presented in Fig. S2 (ESI†). In Fig. S2 (ESI†), $Sn3d_{5/2,3/2}$ core level peaks at 486.1 and 494.4 eV observed in $Ce_{1-x}Sn_xO_2$ correspond to Sn^{4+} . However, if Sn is present with mixed oxidation states (Sn^{4+} and Sn^{2+}) in these oxides XPS cannot distinguish between these two oxidation states because of the very small core level binding energy difference. This issue is addressed in more detail by ¹¹⁹Sn Mössbauer spectroscopy (*vide infra*). $Zr3d_{5/2,3/2}$ core level peaks in the $Ce_{1-x}Zr_xO_2$ solid solutions as presented in Fig. S2 (ESI†) remain constant at 182.3 and 184.6 eV, indicating the presence of only the +4 oxidation state of Zr in these oxides. This is expected because Zr^{4+} is normally irreducible in such oxidic materials.

O1s core level spectra have also been analyzed carefully to check for a change of the nature of the oxygen species in the solid solutions due to the difference in electronegativity of Sn and Zr as substituents. Fig. 6 shows the O1s core level spectra of $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ samples. Interestingly, the peak positions of the O1s core levels in $Ce_{1-x}Sn_xO_2$ gradually shift towards higher binding energy. For example, the O1s core level binding energies in pure CeO₂ and SnO₂ are 529.5 and 530.4 eV, respectively (Fig. 6a). Therefore, while they form solid solutions, the O1s binding energy increases gradually with increasing Sn substitution. The change is indeed significant and detectable. Thus, the binding energy is shifted from 529.5 eV in CeO₂ to 530.0 eV in Ce_{0.55}Sn_{0.45}O₂. On the other hand, no significant shift in the O1s binding energy position in the $Ce_{1-x}Zr_xO_2$ samples is observed, which remains almost constant at 529.6 eV (Fig. 6b). This shows that the presence of Sn in CeO2 makes Ce/Sn-O bonding more covalent as compared to the Zr substitution.

Mössbauer spectroscopic studies

Since XPS is unable to explicitly distinguish the $\rm Sn^{4+}$ and $\rm Sn^{2+}$ states in the solid solution of $\rm Ce_{1-x}Sn_xO_2$, $^{119}\rm Sn$ Mössbauer spectroscopy is very helpful to separate them. Fig. 7 shows the experimental spectra along with transmission integral fits. The corresponding fitting parameters are listed in Table 2. For comparison, the corresponding samples of $\rm SnO_2$ and $\rm SnO$ (with a contamination of $\rm SnO_2$) have also been measured. Both $\rm Sn$ species can be clearly distinguished by their isomer shifts in $\rm ^{119}\rm Sn$ Mössbauer spectroscopy. The lone-pair character of divalent $\rm Sn$ is additionally expressed in a strong quadrupole splitting parameter.

All samples of the $Ce_{1-x}Sn_xO_2$ solid solutions exclusively show Sn(iv) resonances. The isomer shift shows a small but continuous decrease from 0.13(1) mm s⁻¹ in $Ce_{0.9}Sn_{0.1}O_2$ to



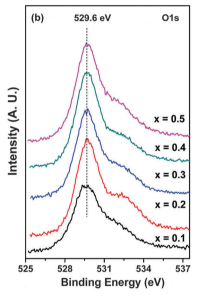
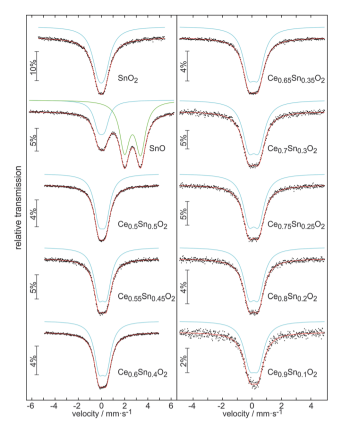


Fig. 6 $\,$ XPS of O1s core levels in the solid solutions: (a) Ce $_{1-x} \rm Sn}_x \rm O_2$ and (b) Ce $_{1-x} \rm Zr}_x \rm O_2$.

 $0.06(1)~{\rm mm~s^{-1}}$ in ${\rm Ce_{0.5}Sn_{0.5}O_2}$, indicating a weak decrease of the electron density at the Sn nuclei. This is in close agreement with the end member of the solid solution, rutile type ${\rm SnO_2}$, which shows an isomer shift of $-0.02~{\rm mm~s^{-1}}$. All spectra show a slightly increased line width parameter, a consequence of the statistical distribution of the Sn atoms. The local non-cubic site symmetry of the Sn atoms (a consequence of the Sn–O substructure distortions discussed below) is expressed in a weak quadrupole splitting of $\sim 0.65~{\rm mm~s^{-1}}$.

XANES and EXAFS studies

In order to explore the possible presence of Ce^{3+} in the bulk of Sn substituted CeO_2 , Ce L_3 -edge XANES spectra were recorded in all the samples along with CeO_2 and $Ce(NO_3)_3$ as standards of Ce^{4+} and Ce^{3+} , respectively. Fig. 8 shows CeL_3 -edge XANES in



 119 Sn Mössbauer spectra of various samples from the $Ce_{1-x}Sn_xO_2$ solid solutions along with spectra of SnO₂ and SnO samples.

Table 2 Fitting results from ¹¹⁹Sn Mössbauer spectra of various samples from the solid solution Ce_{1-x}Sn_xO₂ along with spectra for SnO₂ and SnO samples: isomer shift δ ; quadrupole splitting ΔE_{Ω} ; experimental line width Γ

Sample	$\delta \ ({\rm mm \ s^{-1}})$	$\Delta E_{\rm Q} \ ({\rm mm\ s^{-1}})$	$\Gamma \text{ (mm s}^{-1}\text{)}$	ratio (%)
SnO ₂	-0.02(1)	0.54(1)	1.26(1)	100
SnO	2.68(1)	1.38(1)	1.02(1)	70(1)
	0.00(1)	0.53(1)	0.96(1)	30(1)
$Ce_{0.5}Sn_{0.5}O_2$	0.06(1)	0.59(1)	0.97(1)	100
$Ce_{0.55}Sn_{0.45}O_2$	0.09(1)	0.68(1)	1.00(1)	100
$Ce_{0.6}Sn_{0.4}O_2$	0.10(1)	0.65(1)	0.97(1)	100
$Ce_{0.65}Sn_{0.35}O_2$	0.10(1)	0.64(1)	0.95(1)	100
$Ce_{0.7}Sn_{0.3}O_2$	0.12(1)	0.69(1)	0.99(1)	100
$Ce_{0.75}Sn_{0.25}O_2$	0.12(1)	0.68(1)	0.95(1)	100
$Ce_{0.8}Sn_{0.2}O_2$	0.12(1)	0.64(1)	0.92(1)	100
$Ce_{0.9}Sn_{0.1}O_2$	0.13(1)	0.60(1)	0.94(1)	100

the two end members, Ce_{0.8}Sn_{0.2}O₂ and Ce_{0.5}Sn_{0.5}O₂, as typical examples. Both compounds exhibit similar features as compared to the CeO2 reference and no shift in the edge position is observed, while the Ce³⁺ standard shows a massive edge shift towards lower energies (\sim 2 eV). The results do not exclude the presence of small amounts (below 5%) of Ce3+ in the bulk of CeO2 48-51 and also in the Ce1-xSnxO2 solid solutions. However, they clearly indicate that Ce³⁺ concentration in the bulk of solids is lower than that on the surface determined by XPS under vacuum.

EXAFS analysis at the Sn K- and Ce K-edges was performed to study the changes in the local structure around these ions

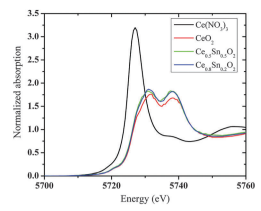


Fig. 8 XANES spectra at the Ce L₃-edge in Ce_{1-x}Sn_xO₂ solid solutions together with the reference samples.

with increasing Sn concentration. The magnitude of Fourier transform (FT) of the EXAFS data along with fitted curves is presented in Fig. 9 and 10, respectively. The coordination number, bond distances and mean square disorder in bond distances are listed in Table S1 (ESI†).

As can be seen from Fig. 9, there is hardly any change in either the position or the height of the first peak in magnitude of the FT of Sn K-edge EXAFS. The first peak corresponds to the Sn-O coordination shell and its unchanged behavior essentially indicates that the local structure around the Sn atoms remains quite stable with increasing tin content. A second peak at about 3.5 Å appears in the FT spectra with increasing Sn concentration. In Ce_{0.8}Sn_{0.2}O₂ the second shell peak is hardly visible but a clear peak can be seen in Ce_{0.5}Sn_{0.5}O₂. The absence of the peak in FT magnitude is generally ascribed to the presence of local disorder around the absorbing atom. This disorder may result in a destructive interference of EXAFS signals originating from the scattering of two different types of neighbors (Sn and Ce) placed at slightly different distances from the absorbing ion.

However, in the case of Ce K-edge EXAFS (Fig. 10) the position, the height as well as the shape of the Ce-O correlation peak (first peak) changes with increasing Sn concentration. These changes indicate that the oxygen coordination around Ce gets adjusted to the increasing number of tin neighbors in the second coordination shell. In pure CeO₂, every cerium ion is surrounded symmetrically by 8 oxygen atoms. With the increase of Sn concentration, these 8 oxygen neighbors around Ce become non-equivalent in a way that some Ce-O bonds become longer and the others shorter. The changes in oxygen coordination are plotted in Fig. 11 (also see Table S1, ESI†). Interestingly, the bond distribution changes gradually from 6 shorter +2 longer Ce-O distances to 4 longer +4 shorter Ce-O distances when the Sn concentration increases above 40%. There is also a small change in the shorter Ce-O bond length. With increasing tin concentration, the shorter Ce-O bond decreases from 2.3 to 2.25 Å while the longer one decreases from 2.56 Å to around 2.5 Å. This behavior can be explained by the difference in the ionic radii of Ce4+ and Sn4+ ions in 8 fold coordination. Ce4+ has an ionic radius of 0.97 Å while that of Sn4+ is about 0.81 Å (Shannon radii⁵²). The difference in ionic radii creates a strong

PCCP

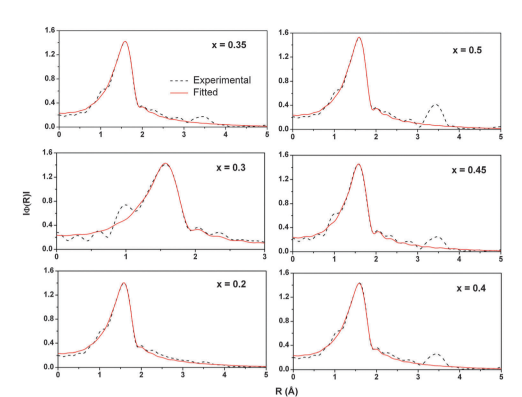


Fig. 9 Magnitude of Fourier transform and its fitting for the Sn K-edge EXAFS in $Ce_{1-x}Sn_xO_2$ solid solutions.

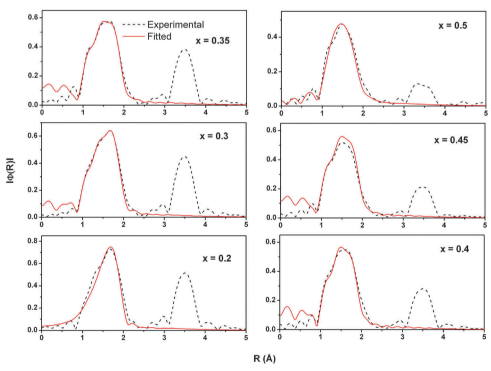


Fig. 10 Magnitude of Fourier transform and its fitting for the Ce K-edge EXAFS in $Ce_{1-x}Sn_xO_2$ solid solutions.

disorder in the local structure around the cerium ions leading to a large distribution of Ce–O bond distances. In contrast, the oxygen coordination around Sn does not split into short and long bonds. All 8 Sn–O bond lengths remain close to 2.051 Å

and only show a small isotropic expansion with the increase of Sn content.

The disorder around the Ce ions is also visible from the behavior of the peak corresponding to the second coordination

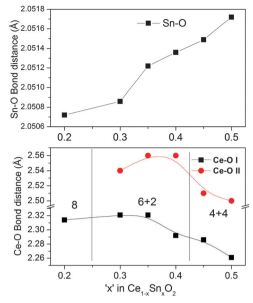


Fig. 11 Different Ce-O and Sn-O bond distances in the Ce_{1-x}Sn_xO₂ solid solutions

shell in the FT of Ce K-edge EXAFS. With increasing Sn concentration, it gradually becomes less intensive, gets larger and changes from a single peak to a doublet. In Ce_{0.8}Sn_{0.2}O₂, most of the second shell neighbors of Ce are Ce atoms, thus giving rise to a single peak in the FT spectra of Ce-edge EXAFS. With increasing Sn concentration, the number of Sn neighbors grows, Ce-O bonds become non-equivalent resulting in the large variety Ce-Sn and Ce-Ce bond distances, resulting in a doublet peak structure.

In contrast, according to the literature, 27 EXAFS analysis shows that the oxygen coordination shell in the Ce_{1-x}Zr_xO₂ solid solutions is divided into 4 + 4 neighbors in the entire substitution range up to x = 0.5.

Discussion

This study compares the structure of $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ solid solutions. Despite similar ionic radii, oxidation state, and coordination number 8 (0.81 Å for Sn⁴⁺ and 0.84 Å for Zr⁴⁺), and a fluorite-type structure, substitution of tin and zirconium shows a distinctly different behavior of the lattice parameter as a function of x. There is a pronounced deviation from Vegard's law in the case of $Ce_{1-x}Sn_xO_2$.

In order to understand this phenomenon, the oxidation states of the metals have been analyzed by XPS, 119Sn Mössbauer spectroscopy and Ce L3-edge XANES spectra. These techniques have confirmed that both Sn and Zr exist in the +4 oxidation state. XPS studies have indicated the presence of a significant amount (12%) of Ce³⁺ on the surface of pure ceria, which increases up to 20–30% for $Ce_{1-x}Sn_xO_2$ and $Ce_{1-x}Zr_xO_2$ solid solutions. This is suspected to be an overestimation of surface Ce³⁺ concentration and does not reflect the bulk concentration. Since XPS measurements have been carried out under high vacuum, which promotes the formation of Ce3+ due to low oxygen partial pressure, the

concentration of Ce³⁺ in the bulk structure can be much smaller. Indeed, Ce L₃- edge XANES measurements carried out in air demonstrate that bulk Ce3+ concentration in all solids is smaller and rather independent of the concentration of the Sn substituent. For this reason, the role of oxygen vacancies in a sudden change of the lattice parameter at x = 0.35 for $Ce_{1-x}Sn_xO_2$ solid solutions can be ruled out.

As Sn⁴⁺ is more electronegative than Zr⁴⁺, the Sn-O bond becomes more covalent in nature than the Zr-O bond (Pauling electronegativities are: 1.12 for Ce, 1.96 for Sn, and 1.33 for Zr). The higher covalency of the -Sn⁴⁺-O²⁻- bond is also evident from the increase of O1s core level binding energy in the Ce_{1-x}Sn_xO₂ solid solutions. A gradual color change from pale white of CeO2 to intense yellow color in solid solutions with increasing Sn content (Fig. S3, ESI†) gives an indication of higher covalency, probably due to charge transfer from O2p to Sn5s states. The absorbance data are consistent with the above fact probably due to increasing charge transfer (see Fig. 3) as indicated by the filling of valley with increasing Sn concentration. Therefore, all 8 oxygen atoms are drawn closer to the Sn atom, which is indicated by equal and shorter Sn-O bonds. As the oxygen shell around the Sn atom is shrinked and becomes more localized, it may not be able to create sufficient distortion on the Ce atom coordination shell until the Sn concentration reaches a critical value, x = 0.35. On the other hand, the coordination number of Ce changes from 8 (up to x = 0.2) to 6 + 2 (up to x = 0.35) to 4 + 4 (above x = 0.35) with increasing Sn concentration. The interesting feature is that the Ce-O bond length remains almost unchanged up to x = 0.2 and then slightly changed up to $x \le 0.35$ (see Fig. 11a). The reason can be that contraction of the SnO₈ shell may facilitate the nearby CeO₈ shells keeping the Ce-O bond length/shell volume almost unchanged in the cell with a reduced lattice parameter. Thus, the lattice parameter of $Ce_{1-x}Sn_xO_2$ decreases at a shallow slope up to 35 at% Sn substitution and above this concentration, the Sn-O bond becomes predominant throughout the lattice, forcing the local structure around cerium to change from 6 + 2 to 4 + 4, which allows the lattice parameter to decrease steeply up to 50 at% Sn substitution. In comparison, the Zr-O bond is more ionic in nature. The 4 + 4 coordination of the Ce-O shell leads to isotropic shrinking on the lattice parameter in the whole range of Ce_{1-x}Zr_xO₂ solid solution up to x = 0.5. Thus, Sn-O local coordination, as a consequence of higher electronegativity of Sn, leads to the deviation from Vegard's law for $Ce_{1-x}Sn_xO_2$ solid solutions.

Conclusions

The substitution of Ce⁴⁺ by the tetravalent cations such as Ti⁴⁺ and Sn^{4+} in $Ce_{1-x}M_xO_2$ solid solutions does not result in a linear change of the lattice parameter up to x = 0.5. In both cases the lattice parameter decreases linearly with a shallow slope up to $\sim 35\%$ substitution, followed by a steeper slope for higher concentrations. In contrast, Zr substitution results in a perfectly linear change in the lattice parameter up to 50%.

The role of oxygen vacancies for this difference can be ruled out, because Zr and Sn remain in +4 oxidation state in all solids and the bulk concentration of Ce3+ is small and rather independent of the dopant content. On the other hand, we observe significant differences in the oxygen coordination shell around the metal as a function of dopant concentration. In $Ce_{1-r}Sn_rO_2$, the oxygen coordination shell around the Ce atoms splits into two sub-shells with shorter and longer bonds, while Sn-O coordination remains symmetric. In contrast, for Ce_{1-x}Zr_xO₂, both Ce–O and Zr–O shells split into different sub-shells and oxygen atoms are cooperatively adjusted to compensate for the size mismatch between Ce and Zr. The different lattice parameter vs. concentration behavior between Sn and Zr substituted CeO₂ can be explained by higher electronegativity of Sn as compared to Zr, supported by XPS of the O1s core level resulting in the deviation from Vegard's law for $Ce_{1-x}Sn_xO_2$ solid solutions.

Acknowledgements

We sincerely thank Professor M. S. Hegde, Solid State and Structural Chemistry Unit, Indian Institute of Science, Bangalore, India for access to the experimental facilities for this work. We thank Dr Frank Krumeich from ETH Zurich for TEM-EDXS measurements.

References

- 1 A. Trovarelli, Catalysis by Ceria and Related Materials, Imperial College Press, London, 2002.
- 2 A. Trovarelli and P. Fornasiero, Catalysis by Ceria and Related Materials, Imperial College Press, London, 2nd edn, 2013.
- 3 J. Kašpar, P. Fornasiero and M. Graziani, Catal. Today, 1999, 50, 285.
- 4 P. Bera and M. S. Hegde, RSC Adv., 2015, 5, 94949.
- 5 M. S. Hegde and P. Bera, Catal. Today, 2015, 253, 40.
- 6 P. Bera and M. S. Hegde, Catal. Surv. Asia, 2011, 15, 181.
- 7 P. Bera and M. S. Hegde, J. Indian Inst. Sci., 2010, 90, 299.
- 8 C.-S. Oh, C.-I. Kim and K.-H. Kwon, J. Vac. Sci. Technol., A, 2001, 19, 1068.
- 9 F.-C. Chiu and C.-M. Lai, J. Phys. D: Appl. Phys., 2010, 43, 075104.
- 10 C. O. Avellaneda, M. A. C. Berton and L. O. S. Bulhões, Sol. Energy Mater. Sol. Cells, 2008, 92, 240.
- 11 M. Mogensen, N. M. Sammes and G. A. Tompsett, Solid State Ionics, 2000, 129, 63.
- 12 H. Inaba and H. Tagawa, Solid State Ionics, 1996, 83, 1.
- 13 M. R. Mohammadi and D. J. Fray, Sens. Actuators, B, 2010,
- 14 X. Zhong, Q. Li, J. Hu and Y. Lu, Corros. Sci., 2008, 50, 2304.
- 15 P. Y. Sheng, W. W. Chiu, A. Yee, S. J. Morrison and H. Idriss, Catal. Today, 2007, 129, 313.
- 16 B. Hariprakash, P. Bera, S. K. Martha, S. A. Gafoor, M. S. Hedge and A. K. Shukla, Electrochem. Solid-State Lett., 2001, 4, A23.
- 17 S. Y. Yao, W. Q. Xu, A. C. Johnston-Peck, F. Z. Zhao, Z. Y. Liu, S. Luo, S. D. Senanayake, A. Martínez-Arias, W. J. Liu and J. A. Rodriguez, Phys. Chem. Chem. Phys., 2014, 16, 17183.

- 18 P. Bera, A. Hornés, A. López Cámara and A. Martínez-Arias, Catal. Today, 2010, 155, 184.
- 19 M. Václavů, I. Matolínová, J. Mysliveček, R. Fiala and V. Matolín, J. Electrochem. Soc., 2009, 156, B938.
- 20 H. C. Yao and Y. F. Y. Yao, J. Catal., 1984, 86, 254.
- 21 A. Trovarelli, Catal. Rev.: Sci. Eng., 1996, 38, 439.
- 22 A. Trovarelli, Comments Inorg. Chem., 1999, 20, 263.
- 23 T. Baidya, A. Gupta, P. A. Deshpandey, G. Madras and M. S. Hegde, J. Phys. Chem. C, 2009, 113, 4059.
- 24 A. Gupta, M. S. Hegde, K. R. Priolkar, U. V. Waghmare, P. R. Sarode and S. Emura, Chem. Mater., 2009, 21, 5836.
- 25 T. Baidya, A. Gayen, M. S. Hegde, N. Ravishankar and L. Dupont, J. Phys. Chem. B, 2006, 110, 5262.
- 26 G. Dutta, U. V. Waghmare, T. Baidya, M. S. Hegde, K. R. Priolkar and P. R. Sarode, Chem. Mater., 2006, 18, 3249.
- 27 G. Dutta, U. V. Waghmare, T. Baidya, M. S. Hegde, K. R. Priolkar and P. R. Sarode, Catal. Lett., 2006, 108, 165.
- 28 L. Vegard, Z. Phys., 1921, 5, 17.
- 29 M. Castellanos and A. R. West, J. Chem. Soc., Faraday Trans. 1, 1980, 76, 2159.
- 30 P. Ganguly, N. Shah, M. Phadke, V. Ramaswamy and I. S. Mulla, Phys. Rev. B: Condens. Matter Mater. Phys., 1993, 47, 991.
- 31 F. Germini, C. Bocchi, C. Ferrari, S. Franchi, A. Baraldi, R. Magnanini, D. De Salvador, M. Berti and A. V. Drigo, J. Phys. D: Appl. Phys., 1999, 32, A12.
- 32 S. Seppänen, L. Hultman, J. Birch, M. Beckers and U. Kreissig, J. Appl. Phys., 2007, 101, 043519.
- 33 P. Kubelka and F. Munk, Z. Tech. Phys., 1931, 12, 593.
- 34 J. Tauc, R. Grigorovic and A. Vancu, Phys. Status Solidi, 1966, **15**, 627.
- 35 D. Briggs and M. P. Seah, Practical Surface Analysis by Auger and X-ray Photoelectron Spectroscopy, Wiley, New York, 1984.
- 36 R. A. Brand, Normos Mössbauer Fitting Program, Universität Duisburg, Duisburg, Germany, 2007.
- 37 B. Ravel and M. Newville, J. Synchrotron Radiat., 2005, **12**, 537.
- 38 C. Anandan and P. Bera, Appl. Surf. Sci., 2013, 283, 297.
- 39 A. Pfau and K. D. Schierbaum, Surf. Sci., 1994, 321, 71.
- 40 D. R. Mullins, S. H. Overbury and D. R. Huntley, Surf. Sci., 1998, 409, 307.
- 41 V. Chauvaut, V. Albin, H. Schneider, M. Cassir, H. Ardéléan and A. Galtayries, J. Appl. Electrochem., 2000, 30, 1405.
- 42 E. Beche, G. Peraudeau, V. Flaud and D. Perarnau, Surf. Interface Anal., 2012, 44, 1045.
- 43 E. J. Preisler, O. J. Marsh, R. A. Beach and T. C. McGill, J. Vac. Sci. Technol., B: Microelectron. Nanometer Struct.-Process., Meas., Phenom., 2001, 19, 1611.
- 44 M. Ricken, J. Nojlting and I. Riess, J. Solid State Chem., 1984, 54, 89.
- 45 R. J. Panlener, R. N. Blumenthal and J. E. Garnier, J. Phys. Chem. Solids, 1975, 36, 1213.
- 46 P. Dutta, S. Pal, M. S. Seehra, Y. Shi, E. M. Eyring and R. D. Ernst, Chem. Mater., 2006, 18, 5144.
- 47 P. E. Lippens, Phys. Rev. B: Condens. Matter Mater. Phys., 1999, 60, 4576.

Paper

- 48 P. Nachimuthu, W.-C. Shih, R.-S. Liu, L.-Y. Jang and J.-M. Chen, *J. Solid State Chem.*, 2000, **149**, 408.
- 49 A. M. Shahin, F. Grandjean, G. J. Long and T. P. Schuman, *Chem. Mater.*, 2005, **17**, 315.
- 50 C. Paun, O. V. Safonova, J. Szlachetko, P. M. Abdala, M. Nachtegaal, J. Sa, E. Kleymenov, A. Cervellino
- F. Krumeich and J. A. van Bokhoven, *J. Phys. Chem. C*, 2012, **116**, 7312.
- 51 C. Hennig, A. Ikeda-Ohno, W. Kraus, S. Weiss, P. Pattison, H. Emerich, P. M. Abdala and A. C. Scheinost, *Inorg. Chem.*, 2013, **52**, 11734.
- 52 R. D. Shannon, Acta Crystallogr., 1976, A32, 751.