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Pair Distribution Function Analysis of Structural Disorder by Nb$^{5+}$ Inclusion in Ceria: Evidence for Enhanced Oxygen Storage Capacity from Under-Coordinated Oxide

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Supporting Information Placeholder

ABSTRACT: Partial substitution of Ce$^{4+}$ by Nb$^{5+}$ is possible in CeO$_2$ by co-inclusion of Na$^+$ to balance the charge, via hydrothermal synthesis in sodium hydroxide solution. Pair distribution function analysis using reverse Monte Carlo refinement reveals that the small pentavalent substituent resides in irregular coordination positions in an average fluorite lattice, displaced away from the ideal cubic coordination towards four oxygens. This results in under-coordinated oxygen, which explains significantly enhanced oxygen storage capacity of the materials of relevance to redox catalysis used in energy and environmental applications.

Cerium dioxide, or ceria, has numerous applications in heterogeneous catalysis. The reversible reducibility of Ce$^{4+}$ ions and high mobility of oxide ions in the solid-state give rise to significant oxygen storage properties that make the material an ideal redox-active support for a wide range of catalytic processes such as in automotive emission control, low temperature water-gas shift catalysis for hydrogen purification and in selective oxidation catalysis for conversion of organics, as well as an oxide-conducting solid-electrolyte. The current focus on ceria-based materials is aimed at further improvement of oxygen storage capacity for new and emerging energy and environmental applications and there is a continued need for new synthetic routes to ceria materials with favorable oxygen storage properties. Inclusion of substituent metals can produce drastic improvements in the redox properties of ceria by introducing strain, increasing the oxide vacancy concentration and/or improving high temperature stability. Many transition-metal cations have been reported as substituents into ceria and often these are considerably smaller than Ce$^{4+}$ and have a strong preference for octahedral geometry.

In this communication we consider the preparation of substituted ceria by a low-temperature hydrothermal route where the inclusion of the pentavalent niobium leads to thermally robust materials with oxygen storage capacities much greater than pure ceria. The choice of a pentavalent substituent ion is counterintuitive, since its presence alone would imply the need for oxygen excess for charge balance, but we show how Na$^+$ as co-substituent provides a mechanism for charge balance. Previous work on niobium-containing ceria is limited, compared to the extensive studies made of other substituent cations, but includes materials for catalysis, with rather small amounts of Nb present (typically 5 atom %) or in fact as phase separated oxides, but no structural study has been performed. We use reverse Monte Carlo (RMC) analysis of neutron total scattering data to obtain atomistic structural models for the materials and compare them with unsubstituted ceria: this reveals new information about the extent of oxide-ion disorder in this important class of materials that shows how local structure may be dramatically disrupted by inclusion of substituent cations, despite retaining an average cubic fluorite structure.

Materials were prepared by the hydrothermal reaction of (1-x)CeCl$_2$·7H$_2$O, xNbCl$_5$ and H$_2$O$_2$ in NaOH solution at 240 °C (see Supporting Information for full details). For samples with x ≤ 0.30 all powder XRD peaks from the materials can be indexed to a face-centered fluorite unit cell (Figure S1). When x = 0 the refined cell parameter a is 5.415(1) Å, close to the NIST-determined value for highly crystalline CeO$_2$(5.411651(1) Å). As x increases, a linear decrease in a is observed to 5.3699(6) Å for x = 0.30 (Figure 1a), explained by the inclusion of the smaller Nb into the fluorite lattice. Further to this, the F$_{2g}$ phonon mode in the Raman spectra, associated with eight-coordinate M-O bond vibrations, displays a slight shift to lower wavenumber with increasing x, from 464 cm$^{-1}$ to 459 cm$^{-1}$, consistent with shorter M-O bonds (Figure 1b); although the effect of decreasing crystal domain size, as observed by Scherrer analysis of powder XRD patterns (Figure 1a) and by HREM images (Figure 1d-f), may also contribute. The intensity of Raman peaks at 540 cm$^{-1}$ and 600 cm$^{-1}$ (labelled α and β, respectively), attributed to second-order phonon modes caused by oxide vacancies in Ce$^{4+}$/Ce$^{3+}$ and Nb$^{5+}$, increases as a function of x. Ce L$_{3}$- and Nb K-edge X-ray absorption near-edge structure (XANES) spectra (Figures S3 and S4) give no indication for reduction of the metals from their highest oxidation-states (+4 and +5 for Ce and Nb, respectively). Instead, charge balance in these materials was found to be achieved by adventitious co-substitution of Na from the hydrothermal reaction mixture, to give general formula (Ce$_{1-x}$Nb$_x$)$_3$Na$_y$O$_{2.6}$ (where x ≤ 0.30 and y ≥ ~0.30). Elemental analysis by ICP-OES shows that the amounts of Na and Nb increase such that the proportion of Na, y, is always at least a third of the Nb content (~x) (Figure 1c), the...
minimum level of Na substitution required to achieve charge balance. The excess Na substituted into the structure allows oxide vacancies, and Na incorporation into complex cerium oxides has been previously shown to generate oxide-deficient fluorite type materials with notably high low-temperature reducibility.\textsuperscript{15}

Figure 1. (a) Refined lattice parameter (black) and Scherrer crystallite size (red) from powder XRD of (Ce\textsubscript{1-x}Nb\textsubscript{x})\textsubscript{1+y}Na\textsubscript{2-y}O\textsubscript{2-δ} as a function of Nb content, x, with lines of best fit. Error bars smaller than data points. (b) Raman spectra with a 514 nm excitation laser. (c) ICP-OES measured Nb and Na content as a percentage of total metal content (remainder Ce). (d-f) HR-TEM micrographs with nominal values of x of 0, 0.10 and 0.20, respectively. All micrographs are presented at the same scale for comparison.

The reducibility of the materials was initially assessed by H\textsubscript{2} temperature-programmed reduction (TPR, Figure 2a), showing that samples exhibit two H\textsubscript{2} uptakes – one at low temperature (<650 °C) and one at higher temperature, widely ascribed to surface and bulk reduction, respectively in ceria itself.\textsuperscript{16} The TPR profiles of the substituted materials are very different from the pure CeO\textsubscript{2} material, with a higher onset temperature of reduction, suggesting a different activation mechanism for the redox behavior. Although the bulk reduction remains approximately constant for all values of x, the degree of low temperature reduction increases dramatically as the level of substitution increases, Figure 2b, Table S2 so that ~60% of the Ce\textsuperscript{4+} in the x = 0.30 sample is reduced by 650 °C. \textit{In situ} powder XRD under simulated TPR conditions of (Ce\textsubscript{0.75}Nb\textsubscript{0.25})\textsubscript{1+y}Na\textsubscript{2-y}O\textsubscript{2-δ} reveals that the average fluorite structure is retained up to at least 800 °C, with a cell parameter increase that far exceeds expected thermal expansion\textsuperscript{17} (Figures 2c, S5), and an expanded lattice parameter of 5.4417(1) Å after cooling to room temperature, explained by the reduction of Ce\textsuperscript{4+} to Ce\textsuperscript{3+}, which has a 15-20% larger ionic radius. Subsequent heating in air recovers a fluorite which still has a slightly expanded lattice parameter, possibly due to incomplete re-oxidation, and a second heating cycle in 10% H\textsubscript{2}/N\textsubscript{2} again shows a drastic expansion as the Ce\textsuperscript{4+} is again reduced, demonstrating the cyclability of the reduction. Oxygen storage capacity (OSC) measurements were carried out to assess further the kinetically available oxygen in the materials, which is of direct applicability to their use as oxygen buffers in catalytic systems. Alternating pulses of CO and O\textsubscript{2} were passed over a sample (loaded with 0.5 wt% Rh, see Experimental Section) and the breakthrough time (the time taken for CO to be detected in the gas stream after passing over the sample), t, (Figures 2d, S6) of CO was used to calculate the OSC at a given temperature (Figure S7). These data, normalized to surface area to account for morphological effects, demonstrate that OSC per unit area is greatly improved by Nb incorporation (Figure 2e).

The coordination environment of Nb within the fluorite lattice so far remains unclear. Rietveld models in space group \textit{Fm}3\textit{m} with a mixed metal 4\textit{a} site fail to provide satisfactory agreement with neutron powder diffraction data collected from samples x = 0.10, 0.20, 0.30 (Figures S8-11, Table S3). Neutron total scattering data were therefore collected from samples with a nominal composition x = 0.20 (both as-made and fired at 500 °C for 5 hours) along with highly crystalline CeO\textsubscript{2} (NIST) and hydrothermally prepared Ce\textsubscript{1-x}Nb\textsubscript{x}O\textsubscript{2-δ}.
As a result of the displacement of Nb$^{5+}$ cations from the ideal cubic symmetry, a significant portion of the oxygen atoms become under-coordinated. This can be visualized in Figure 4, where the oxygen lattices from the refined RMC models are shown, colored by their coordination number. Any metal atoms found within 2.5 Å of an oxygen atom were considered to be coordinated to it. This value was chosen as it encompasses the expected M–O distances in CeO$_2$ and in NbO$_2$. In Figure 4 it can be seen that the vast majority of oxygen atoms in NIST ceria have the expected coordination number, whereas in the Nb-substituted samples a large proportion have a coordination number of 3 and 2. The percentages of under-coordinated oxygen are 17%, 23%, 46% and 45% in NIST CeO$_2$, hydrothermal Ce$_1$Na$_2$O$_{2.4}$, fired (Ce$_1$Na$_1$)$_1$-$\delta$Na$_{0.2}$ and as made

\[\delta = 0.20\] and \(\delta = 0.01\) (Figure 3a). Upon firing, the oxide atoms in this material were found close to their ideal sites, with the mean displacement of 0.13 Å. It is unclear if the perturbed oxide lattice in the as made material is a feature of the small particle size: Scherrer analysis of the XRD pattern of the \(x = 0.20\) as-made sample give a crystallite size of \(~10\) nm (Figure 1a), and a not insignificant quantity of the oxide ions will therefore presumably reside in a likely disordered environment on the particles’ surface. Analysis of bond distances resulting from the RMC analysis (Figure 3c) shows that the uniaxial displacement of the Nb atoms results in a 4 long (~2.7 Å) and 4 short (~2.0 Å) Nb–O distances in the most extreme cases (Figure S18), but there are also some undisplaced Nb atoms present in the refined configuration.

Figure 3. (a) RMC-fitted radial distribution functions obtained from total neutron scattering data. Data were collected on GEM or Polaris Instruments at ISIS Neutron Spallation Source. (b) 10x10x10 supercell from RMC fit to fired \(x = 0.20\) sample collapsed onto a single fluorite unit cell. (Ce,Na) atoms in blue, Nb atoms in red and O atoms in yellow. (c) and (d) Distributions of displacements of each atom type from their ‘ideal’ sites in each RMC model. (e) Resulting distribution of M–O bond distances in each RMC model.

ASSOCIATED CONTENT

Supporting Information
Detailed experimental procedures and additional characterization data. The Supporting Information is available free of charge on the ACS Publications website. The research data supporting this publication can be accessed at wrap.warwick.ac.uk

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REFERENCES


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$\text{(Ce}_{1-x}\text{Nb}_x)_{1-y}\text{Na}_y \text{O}_{2-\delta}$