Ce=O Terminated CeO₂

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Abstract: Multiply bonded lanthanide oxo groups are rare in coordination compounds and have not previously been reported for a surface termination of a lanthanide oxide. Here we report the observation of a Ce=O-terminated ceria surface in a CeO₂(111)-(√3×√3)R30° reconstruction of ≈3 nm thick ceria islands prepared on Pt(111). This is evidenced by scanning tunnelling microscopy (STM), low energy electron diffraction (LEED) and high-resolution electron energy loss spectroscopy (HREELS) measurements in conjunction with density functional theory (DFT) calculations. A Ce=O stretching frequency of 775 cm⁻¹ is observed in HREELS, compared with 766 cm⁻¹ calculated by DFT. The calculations also predict that the Ce=O bond is weak, with an oxygen vacancy formation energy of 0.85 eV. This could play an important role in the facile removal of lattice oxygen from CeO₂, accompanied by the reduction of CeIV to CeIII, which is a key attribute of ceria-based systems in connection with their unique catalytic properties.

Ceria is a common component of many heterogeneous catalysts employed for important industrial processes including CO oxidation, CO₂ hydrogenation, water–gas–shift, methane oxidation and methanol reforming.\(^1\)–\(^3\) Key to its catalytic function is the facile switching between CeIII and CeIV; the subsequent case of oxygen vacancy formation leading to excellent oxygen storage capacity as well as influencing the adsorption of active metal nanoparticles.\(^4\)–\(^5\) Moreover, oxygen adsorption and activation that results in the formation of peroxy (O₂⁻), superoxide (O₂⁻), and weakly bound oxygen (O²⁻) species has been linked to the dynamic exchange of lattice oxygen.\(^6\)–\(^8\) A recent study of a CeO₂-Rh inverse model catalyst concluded that oxygen spill-over from the metal also plays an important role in regeneration.\(^9\) Near-surface oxygen vacancies are clearly important in oxygen exchange, with calculations indicating that subsurface oxygen vacancies are stabilized at the CeO₂(111) surface.\(^10\)–\(^13\) Here we report the observation of a Ce=O-terminated (111) surface of ceria. Calculations suggest that this oxo species could play a crucial role in the mechanism of oxygen vacancy formation.

The lowest energy surface of CeO₂ is the stoichiometric O-terminated (111) surface with an ABC stacking of O-Ce-O trilayers.\(^14\) As with other reducible oxides, the surface reconstruction is dependent on the oxygen chemical potential\(^15\) (Supporting Information, Table S1). Here we focus on the CeO₂(111) (√3×√3)R30° reconstruction prepared as ultrathin islands on Pt(111). This reconstruction has been reported in an earlier study,\(^10\) although its precise origin and structure was not fully understood until now. The formation of this phase requires a slightly lower O₂ partial pressure for the oxidation stage compared with that of the (1×1) bulk terminated surface (1×10⁻⁷ mbar vs. 5×10⁻⁶ mbar). This results in changes to the morphology of ceria islands as well as the surface periodicity. Islands with the (√3×√3)R30° reconstructed surface, shown in Figure 1a, are imaged with atomic resolution (Figure 1b,c). The islands are significantly thicker than those observed for the (1×1) (Supporting information, Figure S1); 2.6–3 nm (9–10 O-Ce-O trilayers) as seen in Figure 1d, compared with 1-1.2 nm (3-4 O-Ce-O trilayers). Features in Figure 1c have a periodicity of 0.59±0.03 nm and are rotated 30° with respect to the (1×1) lattice, which corresponds to a (√3×√3)R30° termination.

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The \((\sqrt[3]{3} \times \sqrt[3]{3})R30^\circ\) reconstruction is also evidenced in the Low Energy Electron Diffraction (LEED) pattern (Supporting Information, Figure S1), is evidenced in the line profile shown in Figure 1e. This enhanced corrugation as well as the similarity to low temperature STM images of V\textsubscript{O} on V\textsubscript{2}O\textsubscript{5}(001)\textsuperscript{[17]} suggest that the features in Figure 1c correspond to Ce=O. Although multiply bonded lanthanide oxo species are rare, there are recent reports of Ce=O containing compounds\textsuperscript{[18,19]} One is Ce=O\textsubscript{(LOEt)}\textsubscript{2}-(H\textsubscript{2}O)]\textsubscript{2}MeC(O)NH\textsubscript{2}, where (LOEt)\textsubscript{2} is a Kl/C\textsubscript{228}ui oxygen tripodal ligand.\textsuperscript{[18]} The Ce=O bond length is 1.86 Å, with a stretching frequency of 684 cm\textsuperscript{-1}.\textsuperscript{[18]} There is a single report of a Ce=O oxo group that is not stabilized by H bonding or an alkali metal coordination, where the bond length is 1.84 Å.\textsuperscript{[19]} Further support for a Ce=O surface termination can be found in a comparison of filled and empty states STM images, shown in Figure 2. Filled (empty) state images sample valence (conduction) band states and hence are sensitive to O (Ce).\textsuperscript{[5,20]} Remarkably, the positions of the protrusions in the filled state images are a subset of those in the empty state images, indicating that O sits directly atop Ce (Figure 2c).

We further test the validity of the Ce=O hypothesis through Density Functional Theory (DFT) calculations and High-Resolution Electron Energy Loss (HREELS) measurements. The Ce=O terminated model of the \((\sqrt[3]{3} \times \sqrt[3]{3})R30^\circ\) reconstruction, shown in Figure 3, was confirmed to be a local energy minimum by the absence of imaginary vibrational frequencies. A Ce-O-Ce bridge termination is also possible for the same Ce\textsubscript{2}O\textsubscript{3} stoichiometry of the uppermost trilayer (denoted TL1), which is more stable by \(\Delta E_{\text{Ce}-\text{O}-\text{bridge}} = -0.94\) eV (Supporting Information, Figure S3), although this structure has not been observed experimentally.

A likely explanation is that the Ce=O terminated phase is formed out of equilibrium, in other words it is a metastable structure formed from kinetically limited crystallization. Another example of this phenomenon is found in the case of TiO\textsubscript{2}(100)-(1 \times 3) reconstructions.\textsuperscript{[21]} In the simulated filled states STM image in Figure 2b, the threefold positions surrounded by Ce=O species appear grey where there is a Ce atom, and black at the location of the Ce defect of the Ce=O reconstruction.

In the simulated empty states image (Figure 2a), bright spots appear at the location of the Ce=O species, matching the experimental data well.
Calculated IR spectra for the Ce=O-terminated and unreconstructed CeO$_2$(111)-(1/√3) surfaces are shown in Figure 4. The most intense bands for the CeO$_2$(111)-(1/√3) surface appear at 538 and 526 cm$^{-1}$, assigned to F$_{1u}$-type vibrations where the near-surface O atoms move along the [111] direction (Fuchs–Kliewer modes). The band at 367 cm$^{-1}$ is assigned to the transversal stretch of the outermost oxygen atoms.$^{[22]}$ Fuchs–Kliewer modes are also observed for the Ce=O-terminated (√3×√3)R30° surface, at 516 and 529 cm$^{-1}$. The Ce=O surface also shows a distinctive band at 766 cm$^{-1}$, corresponding to the stretching vibration of the Ce=O double bond. For comparison, calculations for the O-bridge-(√3×√3)R30° reconstructed surface (Supporting Information, Figure S4) predict two similar F$_{1u}$-like modes, shifted to 537 and 527 cm$^{-1}$. The transversal vibration modes of the bridge O and the subsurface O located below it in TL1 (Supporting Information, Figure S3, A and B, respectively) appear at 502 and 586 cm$^{-1}$, respectively. The longitudinal stretching frequency of the bridging oxygen is 682 cm$^{-1}$ (not IR-active). Supporting Information, Table S2 contains a complete list of calculated frequencies.

HREEL spectra of the ceria islands are shown in Figure 5. The Ce=O stretch is expected at 680–780 cm$^{-1}$ from previous studies of cerium coordination compounds.$^{[18,23]}$ with H-bonding thought to give rise to the red shifted end of this spectrum. Our predicted frequency for surface Ce=O (766 cm$^{-1}$) is in line with these values. Measurements were recorded from a fresh region of the sample, corresponding to the CeO$_2$(111)-(√3×√3)R30° reconstruction (Figure 5a), and for comparison from an area exposed to the LEED beam where a (1×1) pattern was observed (Figure 5b).
HREEL spectrum of the (1 × 1) surface in Figure 5b shows significant loss features at 571, 1057 and 2107 cm⁻¹. The loss at 571 cm⁻¹ corresponds to the surface optical phonon mode of CeO₂(111)-(1 × 1).²⁴ The loss at 2107 cm⁻¹ is assigned to the CO-Pt(111) atom stretching frequency as seen in a control experiment for CO/Pt(111) (Supporting Information, Figure S5). The losses centered at 1057 cm⁻¹ are also seen in earlier data from CeO₂(111)-(1 × 1)/Pt(111),²⁴ being a combination of multiple loss peaks for the surface optical phonon and features associated with Pt₆Ce alloy on the surface. The component peaks are better resolved in the spectrum of the (3x√3)R30° reconstruction and are identified at 536 (ν₁), 775 (ν₂), 1014 (ν₃), 1144 (ν₄), 1247 (ν₅) and 2081 (ν₆) cm⁻¹. The only significant difference between the spectra in Figure 5a,b is the appearance of a loss at 775 cm⁻¹ for the reconstructed surface, which is very close to the calculated value for the Ce=O stretch frequency. As a further comparison, an HREEL spectrum from an un-reconstructed CeO₂(111)-(1 × 1) film (prepared at higher P₀₂) is displayed in Figure 5c; this matches qualitatively with that of Figure 5b and displays no sign of the Ce=O stretch. Small differences in the peak positions are likely due to variations in the film morphology and Pt support. Due to the higher oxygen pressure during preparation of the film in Figure 5c we would expect a smaller amount of residual PtCe surface alloy to be present, explaining the lower intensity of the peak at ≈1100 cm⁻¹.

This spectroscopic evidence validates the conclusion from STM and DFT that there is a Ce=O termination of CeO₂(111). Ce=O bonds are expected to be weak based simply on their rarity in coordination compounds. This suggests the possibility that they could act as active sites for the formation of oxygen vacancies. To test this hypothesis, oxygen vacancy formation energies were calculated (Supporting Information, Table S3). For the Ce=O-terminated surface, we investigated the formation of oxygen vacancies in the upper three oxygen layers. We considered the five inequivalent types of oxygen atoms, labelled A–E in Figure 3. With (3x√3)R30° periodicity, a single oxygen vacancy corresponds to a vacancy concentration of Θ = 0.33. Θ is defined as the number of oxygen vacancies divided by the total number of atoms in a non-reduced and unreconstructed oxygen layer of the same cell (i.e. 3 atoms for (3x√3)R30° periodicity). We also investigated the same concentration of surface (s) and subsurface (ss) oxygen vacancies at the unreconstructed CeO₂(111) surface and a bridge-oxygen vacancy in the O-bridge-(3x√3)R30° reconstructed surface. At the unreconstructed CeO₂(111) surface with Θ = 0.33, the most stable vacancy is subsurface, with one Ce³⁺ ion in TL1 and one in TL2 (E_vac = 2.05 eV). This value is in good agreement with previous work as it lies in between the E_vac values calculated for subsurface vacancy concentrations of 0.25 and 0.50 (1.91 and 2.31 eV, respectively).²³ Removal of the terminating oxygen from the O-bridge-(3x√3)R30° reconstruction requires 1.81–2.02 eV, depending on the location of the excess charge. Vacancy formation on the Ce=O-terminated surface is heavily favored, with energies of ≈1 eV. The most stable vacancy is subsurface V⁴⁺ (cf. Figure 3, E_vac = 0.83 eV), but oxygen abstraction from the Ce=O species (V¹⁺) is also favorable, with virtually the same formation energy (E_vac = 0.85 eV). The filled states STM image in Figure 2b reveals several oxygen vacancies in the surface layer, imaged as dark holes. At the corresponding positions in the empty states image (highlighted with “X” in Figure 2c) there is an absence of Ce⁴⁺ features that are observed elsewhere on the surface. This can be understood on the basis of previous work on reduced CeO₂(111)²⁰ in which it was concluded that while Ce⁴⁺ gave rise to bright features, Ce³⁺ ions did not. The presence of Ce³⁺ below the O vacancies on the (3x√3)R30° reconstruction is consistent with the most stable excess charge distribution obtained upon oxygen abstraction from the Ce=O species (Supporting Information, Table S3, Figure S6). Moreover, neither Ce⁴⁺ below the vacancy in TL1 nor that in TL2 appear as a bright feature in the calculated empty states image. Furthermore, missing or attenuated features in the empty state image at Ce positions that are not located just below missing O sites, such as the large dark patches in Figure 2a and Figure S7, also correspond to Ce³⁺ ions in TL1 and are likely to result from deeper lying subsurface oxygen vacancies. The formation energy of subsurface oxygen vacancies, with one excess electron in TL1 and the other one in TL2, lie within a range of 0.85–1.08 eV (Table S3). The proposed structure for the Ce=O-terminated CeO₂(111)-(3x√3)R30° reconstruction, as well as the calculated low energy vacancy structures, allow us to satisfactorily explain the experimentally observed filled and empty state STM images.

In summary, a CeO₂(111)-(3x√3)R30° reconstruction of an ultrathin ceria film prepared on Pt(111) has been studied using surface imaging (STM), diffraction (LEED), and vibrational spectroscopy (HREELS). The results all point to the formation of a Ce=O termination, representing the first observation of a surface multiply bonded lanthanide oxo species. This interpretation is validated by DFT calculations of the filled and empty states STM images for the Ce=O model, which closely match the experimental images. Moreover, a Ce=O stretch frequency of 766 cm⁻¹ is predicted, which is in excellent agreement with the HREELS experimental value of 775 cm⁻¹. The calculations also predict that Ce=O species on CeO₂(111) could act as an active site for the formation of oxygen vacancies, which play a key role in ceria-based heterogeneous catalysis.

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Conflict of interest

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A combined experimental and computational study of the ($\sqrt{3} \times \sqrt{3}$)R30° reconstruction of the CeO$_2$(111) surface of an ultrathin ceria film is presented. The evidence collected indicates that the film is terminated by Ce=O groups.