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Chemical Activity of Oxygen Vacancies on Ceria: A combined Experimental and Theoretical Study on CeO2 (111)

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ABSTRACT: The chemical activity of oxygen vacancies on well-defined, single-crystal CeO2(111)-surfaces is investigated using CO as a probe molecule. Since no previous measurements are available, the assignment of the CO stretch frequency as determined by IR-spectroscopy for the stoichiometric and defective surfaces are aided by ab-initio electronic structure calculations using density functional theory (DFT).

Ceria, one of the most easily reducible materials, provides the basis for numerous applications ranging from heterogeneous catalysis (vehicle emission control, ethanol steam reforming, water gas shift reactions) over solar thermal hydrogen production from water and to the generation of CO from CO2. In all cases, the oxidation reduction cycle (Ce4+/Ce3+ cations – and the associated oxygen vacancies) is the most determining factor. The importance of this material has triggered numerous experimental and theoretical studies, in particular aiming at elucidating the properties of oxygen vacancies. Earlier pictures proposing the two electrons left behind by removing an oxygen atom from the surface to be localized at the surrounding cations (i.e., presence of two Ce3+ ions adjacent to the defect, see the Ce cations labeled 1 in Fig. 1) had to be revised later and recent theoretical work has revealed that the remaining electrons are actually localized at some distance away from the vacancy. Though it is intuitive to believe that the density of oxygen vacancies may affect the chemical activity of ceria, direct experimental evidence is lacking and even the determination of O-vacancy densities represents a major experimental challenge. The most frequently used approach to determine the concentration of this active site crucial for all redox reactions occurring on this surface is to use carbon monoxide (CO) as a probe molecule and to deduce the nature of the cations it binds to from the frequency of the CO stretch vibration. Typically, the CO stretch frequency shows substantial variations upon changes in the charge state of the metal ion the CO is bound to. In the case of Cu ions, for example, the difference in stretch frequency between CO bound to Cu2+ and Cu+ amounts to about 70 cm⁻¹. Although a large set of data exists for CO adsorbed on ceria powders, the extent of this shift for Ce-cations is still being discussed quite controversially. For a reliable identification of the shifts urgently reference data for ceria single crystal surfaces are needed.

EXPERIMENTAL

The IRRAS measurements were conducted in an UHV apparatus combining a state-of-the-art FTIR spectrometer (Bruker Vertex 80v) with a multi-chamber UHV system (Prevac). The CeO2(111) single crystal (SurfaceNet) surface was prepared by repeated cycles of sputtering with 1 keV Ar⁺ and annealing at 800 K for 15 min in an O2 atmosphere of 1×10⁻⁵ mbar for forming a stoichiometric surface, or alternatively without O2 to create a reduced one. Exposure to CO at sample temperatures typically below 75 K was achieved by backfilling the IR chamber up to 10⁻¹ mbar. The base pressure during acquisition of IR spectra was ~8×10⁻¹¹ mbar.

Fig. 1. Ball-and-stick model of the CeO2(111) surface (top view) with (a) top- or (b) subsurface oxygen vacancy (V_o). The position of the vacancy is indicated by the dashed circle. Color: red – top surface oxygen, light red – subsurface oxygen, white – Ce4+, blue – Ce3+. The IR-MOPC measurements were conducted in an UHV apparatus combining a state-of-the-art FTIR spectrometer (Bruker Vertex 80v) with a multi-chamber UHV system (Prevac). The CeO2(111) single crystal (SurfaceNet) surface was prepared by repeated cycles of sputtering with 1 keV Ar⁺ and annealing at 800 K for 15 min in an O2 atmosphere of 1×10⁻⁵ mbar for forming a stoichiometric surface, or alternatively without O2 to create a reduced one. Exposure to CO at sample temperatures typically below 75 K was achieved by backfilling the IR chamber up to 10⁻¹ mbar. The base pressure during acquisition of IR spectra was ~8×10⁻¹¹ mbar.
Results

In the present study, we first determined the stretch frequency of CO adsorbed on a fully-oxidized, single crystalline CeO$_2$(111) surface (“stoichiometric surface”) using infrared reflection absorption spectroscopy (IRRAS). The single sharp band seen in Fig. 2 indicates that ν$_1$ for CO adsorbed on the fully oxidized CeO$_2$(111) surface amounts to 2154 cm$^{-1}$, which differs by ~9 cm$^{-1}$ from the experimentally observed frequency of 2154 cm$^{-1}$. It can clearly be seen from Table 1 that the vibrational frequencies of CO adsorbed on the CeO$_2$(111) surface (“stoichiometric surface”) are unusually small, on the basis of previous assignments one would have expected shifts of up to 50 cm$^{-1}$. In addition, it is surprising to observe a blue-shift for CO bound to a defective site, as on other oxides like TiO$_2$ the increased binding energy at defect sites (which is also observe here) gives rise to a red-shift. In order to better understand the origins of this unexpected behavior and to assign the defect-induced ν$_1$, we have carried out first-principle density functional theory (DFT) calculations (see table 1, for details of the calculations see the SI). The results of the calculation allow to assign the defect-induced ν$_1$ to CO adsorbed at 6-fold coordinated Ce$^{4+}$ ions in the direct vicinity of the vacancy (site 1 in Fig. 1). An assignment to 7-fold coordinated Ce$^{3+}$ ions further away from the vacancy (site 2 in Fig. 1) can be ruled out.

The calculated vibrational frequency of CO bound to the 7-fold coordinated Ce$^{4+}$ on the stoichiometric surface amounts to 2145.2 cm$^{-1}$, which differs by ~9 cm$^{-1}$ from the experimentally observed frequency of 2154 cm$^{-1}$. It can clearly be seen from Table 1 that the vibrational frequencies of CO adsorbed on the CeO$_2$(111) surface (“stoichiometric surface”) are unusually small, on the basis of previous assignments one would have expected shifts of up to 50 cm$^{-1}$. In addition, it is surprising to observe a blue-shift for CO bound to a defective site, as on other oxides like TiO$_2$ the increased binding energy at defect sites (which is also observe here) gives rise to a red-shift. In order to better understand the origins of this unexpected behavior and to assign the defect-induced ν$_1$, we have carried out first-principle density functional theory (DFT) calculations (see table 1, for details of the calculations see the SI). The results of the calculation allow to assign the defect-induced ν$_1$ to CO adsorbed at 6-fold coordinated Ce$^{4+}$ ions in the direct vicinity of the vacancy (site 1 in Fig. 1). An assignment to 7-fold coordinated Ce$^{3+}$ ions further away from the vacancy (site 2 in Fig. 1) can be ruled out.

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Table 1. Calculated vibration frequencies V and adsorption energies E$_b$ of CO at various sites on the stoichiometric and reduced CeO$_2$(111) surface (one top-surface oxygen vacancy per supercell, experimental frequencies are listed for comparison).

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Adsorption Site</th>
<th>DFT v/cm$^{-1}$ (E$_b$/eV)</th>
<th>Exp. [cm$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free molecule</td>
<td></td>
<td>2122.5</td>
<td>2143.5</td>
</tr>
<tr>
<td>Ideal</td>
<td>Ce$^{4+}$</td>
<td>2145.2 (0.28)</td>
<td>2154</td>
</tr>
<tr>
<td>Reduced</td>
<td>Ce$^{4+}$ at OV (site 1)</td>
<td>2157.2 (0.40)</td>
<td>2169.6 (0.39)</td>
</tr>
<tr>
<td></td>
<td>Ce$^{4+}$ at site 2</td>
<td>2150.9 (0.27)</td>
<td>2143.1 (0.34)</td>
</tr>
<tr>
<td></td>
<td>Ce$^{4+}$ at site 3</td>
<td>2143.1 (0.34)</td>
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The calculated frequencies for adsorbed CO were corrected by multiplying the calculated values by a factor of $2143/2122.5$ or 1.0097, corresponding to the ratio of the experimental and calculated values for gas-phase CO.

defective surface sites shift to higher values, especially for CO adsorbed in a configuration allowing for direct interaction with the vacancy (site 1 in Fig. 1 or center of $V_O$).

These calculations beautifully corroborate our above assignment, namely that the blue-shift of 10 cm⁻¹ seen in the experiment when going from the stoichiometric to the reduced substrate has to be assigned to CO bound to Ce⁴⁺-sites in direct vicinity of the vacancy.

Calculations carried out for different CO-coverages (see SI), which in previous work has been found to affect both CO binding energies and stretch frequencies, revealed that the local CO concentrations have almost no effect on the CO binding energies. Since in previous assignments of IR-powder data a number of assignments has been reported for CO bound to a 6-fold coordinated Ce⁶⁺ (present only in a less stable defective surface configuration with the two excess electrons localized at nearest-neighbor Ce cations), we also considered this adsorption site. The adsorption energy was found to be 0.32 eV, i.e. lower compared to 6-fold coordinated Ce⁴⁺ (cf. Fig. 4a). However, this adsorption energy is slightly larger with respect to that on the 7-fold coordinated Ce⁷⁺ (cf. Fig. 4b).

**CONCLUSIONS**

Our experimental and theoretical observations provide a solid basis for the identification of stretch frequencies for CO bound to CeO₂(111) substrates. For CO bound to 7-fold coordinated (i.e., fully coordinated for a surface cation) Ce⁷⁺-cations (denoted Ce⁷⁺₉C where c stands for “coordinated”) within the stoichiometric surface, the stretch frequency, $ν_s$, amounts to 2154 cm⁻¹. The calculated adsorption energy of 0.28 eV for this species is in good agreement with the activation energy for desorption of 0.27 eV extracted from the temperature programmed desorption experiment.

Surprisingly, the CO species bound to defective parts of the surface exhibits a blue-shift (in contrast to other oxides). For Ce-ions in direct vicinity of the oxygen-vacancy, the corresponding blue-shift amounts to 10 cm⁻¹. We tentatively ascribe the increased adsorption energy and higher vibrational frequency observed for this species to electrostatic interactions resulting from a coupling of the CO dipole and the dipole of the top-surface-O-Ce double layer when the molecule moves inside the vacancy.

Based on the best agreement in the stretch frequency, we assign the band at 2163 cm⁻¹ to CO at Ce⁴⁺ ions at the Oₐ site. Note also that the number of available Ce⁶⁺₉C-sites is three times larger than that of the 3-fold hollow sites inside the vacancy. In fact, the spectra displayed in Fig. 3b would be consistent with the presence of a weak feature at higher frequencies, where theory would predict the position of the stretch frequency for CO adsorbed within the O-vacancies. As the calculated binding energy is very similar to the one calculated for adsorption on Ce⁷⁺ ions at the vacancy, one would expect that this site would be populated as well. Our results demonstrate that the coordination geometry can play a decisive role for weak chemical interactions on oxide surfaces. While on TiO₂ and many other oxides the frequency of adsorbed CO is mainly given by the charge state of the metal cation, the coordination geometry exhibits a more pronounced influence for CeO₂.

**ASSOCIATED CONTENT**

**Supporting Information**

Computational details, CO stretch frequency assignment for ceria powder, IR spectra of CO adsorbed on stoichiometric
and reduced CeO$_2$ with light incident along the [-211] direction, XPS data including analysis of the vacancy concentration on the stoichiometric and reduced surface, IRRAS desorption experiment on the stoichiometric and reduced surface (including determination of the activation energy of desorption). This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**
The authors declare no competing financial interests.

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**REFERENCES**

(1) Paier, J.; Penschke, C.; Sauer, J. Chemical reviews 2013, 113, 3949.
