Oxygen Nonstoichiometry in (Ca$_2$CoO$_3$)$_{0.62}$(CoO$_2$): A Combined Experimental and Computational Study

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ABSTRACT: The oxygen nonstoichiometry in the misfit calcium cobaltite (Ca$_2$CoO$_3$)$_{0.62}$(CoO$_2$) has been studied experimentally and by density functional theory (DFT) calculations. The standard oxidation enthalpy $\Delta H^0_{\text{ox}}$ of oxygen deficient (Ca$_3$CoO$_5$)$_{0.62}$(CoO$_2$) was measured directly using simultaneous thermogravimetry and differential scanning calorimetry. $\Delta H^0_{\text{ox}}$ was found to be in agreement with the prediction from a previously published defect chemical model based on purely thermogravimetical analysis. A series of samples with different oxygen vacancy concentration was prepared by annealing in air, followed by rapid quenching. Room-temperature Raman spectroscopy showed a sharp mode at 700 cm$^{-1}$ decreasing in intensity with increasing vacancy concentration. We discuss this observation as evidence for oxygen vacancies being preferably formed within the central layer of the Ca$_2$CoO$_3$ subsystem. DFT calculations demonstrated that the calculated electronic structure is sensitive to the chosen model of the crystal structure. Still, for all investigated models, the standard formation enthalpy of oxygen vacancies within the Ca$_2$CoO$_3$ moiety was much lower than that for a site within the CoO$_2$ layer, in agreement with the presented experimental data.

1. INTRODUCTION

The family of misfit cobaltites has attracted significant scientific attention in recent years because of their interesting electronic properties, the intriguing crystal structure, and the interplay between the two.† The common building block of misfit cobaltite structures is a layer of edge-sharing CoO$_6$ octahedra (hereafter denoted as the cobalt oxide layer, COL), which is alternately stacked along the c axis with layers of rock salt symmetry (rock salt layer, RSL), thereby forming a natural superlattice along the c axis.‡ Interesting electronic properties of these materials result from charge transfer between the two subsystems, leading to a mixed valent cobalt state within the COL.§

In this paper, we study the misfit calcium cobalt oxide (CCO), where the COL is separated by Ca$_2$CoO$_3$ as the rock salt subsystem. While the a- and c-axes of both the COL and RSL subsystem are identical, the b-axes are not, and the ratio of $b_{\text{COL}}$ and $b_{\text{RSL}}$ leads to a structural modulation along b. The most precise notation for CCO is thus (Ca$_2$CoO$_3$)$_q$(CoO$_2$) with $q = b_{\text{COL}}/b_{\text{RSL}} = 0.62 \pm 0.03$,§,‡— but the alternative notation “Ca$_3$Co$_4$O$_{13}$” is frequently found in the literature as well. CCO doped with various elements has been intensively studied as a thermoelectric material suitable for high-temperature energy harvesting and exhibits one of the highest figures of merit among all oxide materials (e.g., see ref 7). At high temperatures, the electronic properties of many oxide materials are influenced by reduction: thermal creation of effectively positively charged oxygen vacancies and the accompanying change in the concentration of electronic charge carriers. Using the Kröger–Vink notation,§ the formation of an oxygen vacancy, charge compensated by annihilation of two hole type carriers, can be written as

$$O_2^\text{-} + 2e^\text{-} + 2h^\text{+} = v^\text{++} + \frac{1}{2}O_2$$

(1)

While there has been some discussion about the presence and degree of oxygen nonstoichiometry in the structurally related compound Na$_3$CoO$_2$,9—12 it is generally agreed that
CCO can exhibit a significant amount of oxygen vacancies. Previous studies have shown the reversible creation of oxygen vacancies in CCO at high temperatures and atmospheres with a low oxygen partial pressure $pO_2$ (e.g., see refs 4, 13, and 14). Writing the chemical formula as $Ca_3Co_{3.92}O_{9.34}$, Shimoyama et al. showed that CCO can accommodate oxygen vacancies up to $\delta = 0.20$ without decomposition. On the basis of an X-ray diffraction refinement study, Ling et al. concluded that oxygen vacancies are exclusively formed in the central Co–O layer of the RSL subsystem. Oxygen vacancies inside the RSL may adjust the charge carrier concentration without disturbing the conduction path of carriers inside the COL by acting as scattering centers. The presence of oxygen vacancies could therefore be a possible reason for reported discrepancies in the electronic properties of nominally identical samples (e.g., see ref 7). Furthermore, oxygen vacancies may also affect the internal stress between the two subsystems, thereby causing the different structural modulations observed experimentally.

The oxidation–reduction thermodynamic behavior of a material is usually studied by means of thermogravimetry or coulometric titration (e.g., see refs 17 and 18), and the extraction of equilibrium coefficients and standard entropies and enthalpies relies on the validity of several assumptions. These assumptions include the negligible temperature dependence of the thermodynamic parameters and not least the applicability of the considered defect chemical model. It is therefore desired to employ a technique allowing the direct determination of parameters for the thermodynamic reduction (or oxidation) processes. This is done here by an instrument combining thermogravimetry and differential scanning calorimetry (TG-DSC), in which the enthalpy change is measured directly and simultaneously with the respective mass change upon sample reduction or oxidation. When the chemical reaction involved is identified, the enthalpy change of the reaction is readily obtained from the TG-DSC analysis.

The incommensurate structure of misfit compounds makes it difficult to describe them computationally, as relatively big unit cell approximations and high computational power are required. Previous attempts to model the electronic structure of CCO have led to results contradicting experimental evidence or have been shown to be sensitive toward the chosen unit cell.

Recently, we have presented a defect chemical model to describe the oxygen nonstoichiometry in CCO, which is based on the assumption that oxygen vacancies are formed only next to Co sites within the central Co–O layer in the rock salt subsystem. In this paper, we report our results of a combined experimental and computational study of oxygen nonstoichiometry in CCO. Density functional theory (DFT) calculations and Raman spectroscopy give further evidence that oxygen vacancies are mainly formed in the central Co–O layer of the rock salt subsystem. The oxidation thermodynamics in CCO are studied directly by TG-DSC, and the results are compared to values obtained from DFT and the defect chemical modeling.

2. EXPERIMENTAL METHODS

Powder and pellet samples of CCO with a $Ca/Co$ ratio of 3/3.95 were fabricated as reported elsewhere. The change in oxygen content of a pellet was determined by means of thermogravimetry (CI Electronics MK2 microbalance). The sample and the counterweight were attached to the arms of the balance using platinum wires. The sample was suspended in an outer alumina tube in a vertical tubular furnace and the counterweight was hung in a glass tube kept at room temperature. The apparatus was connected to an in-house built gas mixer of a type as described elsewhere. The weight change was monitored when varying the oxygen partial pressure ($pO_2$) from 1 to $10^{-4}$ atm by diluting $O_2$ with He until equilibrium was obtained. All measurements were performed for temperatures and $pO_2$ where CCO is stable. The observed weight change can thus be solely assigned to a variation of the oxygen stoichiometry. In practice, a resolution of $\sim0.01$ mg was obtained, which translates to a resolution in the oxygen nonstoichiometry $\delta$ in $Ca_3Co_{3.95}O_{9.34}$ of $\sim0.0003$ for our sample masses. All weights were corrected for estimated buoyancy arising from the asymmetrical temperature and volume of the sample and counterweight.

To study the oxidation thermodynamics of oxygen deficient CCO, we use an instrument with simultaneous thermogravimetry and differential scanning calorimetry (Netzsch STA 449 C Jupiter) connected to a gas mixer similar to that used for the pure thermogravimetric measurements. The DSC signal is a measure of the heat flow from the sample in comparison to an empty reference crucible. When the sample undergoes a reaction associated with an enthalpy change (e.g., a phase transition or chemical decomposition), the DSC curve shows a peak, where the peak area corresponds to the respective change in enthalpy. Because CCO is stable at the temperatures and $pO_2$ used in the TG-DSC experiments, the observed signals can be associated with the formation and oxidation of oxygen vacancies. The DSC peak area, divided by the corresponding mass change, can thus be identified with the oxidation enthalpy $\Delta H_{\text{ox}}$ of oxygen deficient CCO.

Instrument and crucible calibration was done right before the measurement of the sample by using reference samples with structural phase transitions in the investigated temperature range. As for the TG experiment, the $pO_2$ was varied by diluting $O_2$ in He. At each temperature, three oxidation and reduction cycles were recorded to increase the statistical significance, and only the exothermic oxidation signal was used for peak integration.

Ceramic samples of CCO were prepared with different oxygen nonstoichiometries for Raman spectroscopy measurements. These samples were obtained by annealing in air at different temperatures well below the decomposition temperature of CCO until compositional equilibrium was reached and then rapidly ($\approx15$ s) cooled to room temperature. The surface was polished after quenching.

Room-temperature Raman measurements were performed with a Dilor XY triple grating Raman spectrometer in backscattering configuration using the unpolarized 514.5 nm laser line of a Coherent Innova 90 Ar$^+$/Kr$^+$ mixed-gas laser. The samples measured were placed with the compacting direction (mainly the c axis) parallel to the incident laser beam. To avoid heating of the sample, the power of the laser was kept below 20 mW, and the beam was focused onto the surface of the samples with a spot size of $\approx100$ nm in diameter.

3. THEORETICAL AND COMPUTATIONAL METHODS

We adopted a computational methodology similar to that successfully used before by Rébola et al. Spin-polarized periodic DFT calculations were performed using the VASP.
The plane wave basis set was limited to a 600 eV energy cutoff for valence electrons while the atomic cores were included via the projector augmented wave (PAW) method. The Brillouin zone was sampled by a Γ-centered grid of 3×2×3 k points (placed less than 0.035 Å/Å from each other in reciprocal lattice direction). The exchange and correlation effects were embodied by the Perdew–Burke–Ernzerhof (PBE) generalized gradient functional. Simultaneous relaxation of ionic positions and lattice parameters was performed for the defect-free monoclinic structures starting from the experimental values and without assuming a certain oxygen position for the vacancy in the defective (V_{def}) supercell.26,27 We computed the standard formation enthalpy \( \Delta H_{\text{form}}(p_0,T = 0) \) for oxygen vacancies from this by
\[
\Delta H_{\text{form}}(p_0,0) = H_{\text{def}}(p_0,0) - H_{\text{perf}}(p_0,0) + \frac{1}{2}H_{O_2}(p_0,0) + 2\mu_e
\]
where \( H_{\text{perf}}(p_0,0) \) and \( H_{\text{def}}(p_0,0) \) are the enthalpies for the pristine and defective structures, respectively, and \( H_{O_2}(p_0,0) \) is that of oxygen gas. All enthalpies are taken at standard pressure \( p_0 = 1 \) bar and temperature \( T = 0 \) K and assumed to correspond to the DFT total energies. The system charge \( n \) (here \( n = 2 \) for an oxygen vacancy with a double positive effective charge) was simulated by adjusting the total number of electrons in the cell and at the same time adding a compensating jellium background to avoid diverging Coulomb contributions. The electron potential \( \mu_e \) was defined as
\[
\mu_e = E_F + \gamma = E_F + (V_{\text{def}} - V_{\text{perf}})
\]
where \( E_F \) is the Fermi level in the pristine material. \( \gamma \) is a correction term which accounts for the energy of the jellium-like potential, here considered as the shift in the average electrostatic potentials at a bulklike lattice site far from the vacancy in the defective (\( V_{\text{def}} \)) and pristine (\( V_{\text{perf}} \)) supercell.26,27

Equation 3 could be extended to obtain \( \Delta G_{\text{form}} \) at finite temperatures and oxygen partial pressures. However, as this is usually based on several assumptions (e.g., that thermal contributions of the solid are negligible compared to that of the gas; \( G(p,T) = G(p_0,0) \)), the association between experiment and calculation will in this work be done at 0 K by comparing the computational formation enthalpy of an oxygen vacancy to the experimental results.

4. RESULTS AND DISCUSSION
4.1. Thermogravimetry and Defect Chemical Model. The measured oxygen nonstoichiometry \( \delta \) in CaxCo3yO9.34−\( \delta \) as a function of \( pO_2 \) and temperature \( T \) is shown in Figure 1. The data is identical to those published in ref 21 but plotted as CaxCo3yO9.34−\( \delta \) instead of (CaxCo3−\( \delta \)).24 (CoO2) (with \( \Delta = 0.62\times\delta \)) to not presuppose a certain site for oxygen vacancy...
The formation. We use the mass at 350 °C and $p_{O_2} = 1$ atm as a reference for $δ = 0$.

The defect chemical model to describe oxygen non-stoichiometry and other relevant electronic defects in CCO is explained in detail elsewhere, but we present the main features here.

Because of the incommensurate stacking of the $\text{Ca}_2\text{CoO}_3$ and the $\text{CoO}_2$ layers, a mixed-valent state for the Co ions within the COL is inherently present. As a perfect reference state, we define that all Co atoms in the rock salt layer are in a $3^+$ state, which results in a mixed valent state with an average valency of $4 - q = 3.38$ in the CoO₂ layer, in order to satisfy overall charge neutrality.

When choosing a localized representation, an electron charge carrier can be identified with a Co$^{3+}$ ion, and thus carries an effective charge of $-(1 - q)$, while a hole carrier (or Co$^{4+}$ ion) has an effective charge of $+q$. In the following, we use a modified Kröger–Vink notation$^{29}$ to describe defects with fractional effective charges and subscripts RSL or COL to denote the specific location of each defect.

In this notation, the oxidation of an oxygen vacancy (eq 1) can be written as

$$v^\bullet_0 + \frac{1}{2}O_2(g) + 2e^{(1-q)}/ = O^\circ_0 + 2h^\bullet_{\text{COL}}$$

with the equilibrium coefficient

$$K_{Ox} = \frac{[O^\circ_0][h^\bullet_{\text{COL}}]^2}{[v^\bullet_0][e^{(1-q)}]^2} = \exp\left(-\frac{\Delta G_{Ox}}{k_BT}\right)$$

$$= \exp\left(-\frac{\Delta H_{Ox}}{k_BT}\right) \exp\left(\frac{\Delta S_{Ox}}{k_B}\right)$$

$$\Delta G_{Ox}, \Delta H_{Ox}, \text{and } \Delta S_{Ox} \text{ are the Gibbs energy, enthalpy, and entropy of the oxidation reaction, respectively.}$$

Equivalently, one can consider eq 5 in the reverse direction. The reaction then describes the formation of an oxygen vacancy, in analogy to the process studied by DFT (eq 2). The corresponding thermodynamic parameters for oxygen vacancy formation are then the negative of the respective parameters of the oxidation reaction $\Delta G_{\text{form}} - \Delta G_{Ox}, \Delta H_{\text{form}} = -\Delta H_{Ox}$ and $\Delta S_{\text{form}} = -\Delta S_{Ox}$.

The experimental $δ$ shows signs of saturation at low $p_{O_2}$, where the saturation level increases with temperature. We therefore suggested a thermal excitation of hole type charge carriers within the COL and a compensating reduction of Co$^{3+}$ to Co$^{2+}$ in the rock salt layer$^{21}$

$$\text{Co}^{x}_{\text{Co},RSL} + e^{(1-q)}/ = \text{Co}^{x}_{\text{Co},COL} + h^\bullet_{\text{COL}}$$

(7)

$$K_D = \frac{[\text{Co}^{x}_{\text{Co},RSL}][h^\bullet_{\text{COL}}]}{[\text{Co}^{x}_{\text{Co},COL}][e^{(1-q)}/]}$$

(8)

Furthermore, the overall electroneutrality requires

$$q(\text{Co}^{x}_{\text{RSL}}) + (1 - q)(e^{(1-q)}/) = q[h^\bullet_{\text{COL}}] + 2q(v^\bullet_0)$$

(9)

A numerical fit of the thermogravimetric data to these equations describes the experimental results only qualitatively (e.g., $R^2 = 0.84$ at 675 °C) (Figure 1a). We therefore included an additional, plausible restriction in ref 21. We postulated that oxygen vacancies can be formed only on oxygen sites next to Co$^{3+}$ ions in the rock salt layer, rationalized by a weaker chemical bond between those two species as compared to a Co$^{3+}$ site. The resulting model allows fitting the thermogravimetric data to the equilibrium constants of the two equations and describes the data very well ($R^2 = 0.997$ at 675 °C) (Figure 1b). The obtained thermodynamic parameters for charge disproportionation (eq 8) and oxidation (eq 6) follow an Arrhenius-type behavior with the oxidation enthalpy $\Delta H_{Ox}$ determined to be $-53 \pm 2 \text{ kJ/mol}$ (equivalent to a formation enthalpy of $\Delta H_{\text{form}} = 0.57 \pm 0.02 \text{ eV per oxygen vacancy}$).

4.2. Raman Spectroscopy. Room-temperature Raman spectra of ceramic CCO samples with different oxygen content are shown in Figure 2. The phonon modes found in the investigated spectral range from 250–800 cm$^{-1}$ can be ascribed to vibrational motion of oxygen, rather than the heavier Ca and Co atoms. The $δ$ values for the respective samples were estimated from Figure 1b. The spectra were normalized to the intensity of the mode at 630 cm$^{-1}$. We note, however, that a normalization over the whole spectral range yields qualitatively
similar results. Each spectrum is dominated by three modes at 300, 540, and 630 cm\(^{-1}\). In addition, the spectrum is built up of a range of broad, overlapping modes, which are in general agreement with a Raman study of CCO single crystals.\(^{30}\) The assignment of these modes is challenging because of the complicated structure of CCO and beyond the scope of this paper. By comparing our results to the reported Raman spectrum of Na\(_2\)CoO\(_\text{2}\)\(_\text{−}\)\(_1\)\(_\text{−}\)\(_1\), which has a similar layer of edge-sharing CoO\(_6\) octahedra, we note that the latter is dominated by modes at 475 and 570 cm\(^{-1}\) at room temperature, which have been assigned to the in-plane E\(_1\)g and out-of-plane A\(_1\)g modes, respectively.\(^{31,32}\) In analogy, we therefore assign the dominant modes found in CCO at 540 and 630 cm\(^{-1}\) to the E\(_1\)g and A\(_1\)g modes, respectively. The energy shift as compared to NCO is caused by slightly different lattice parameters and different interlayer strain between the two subsystems.

When creating oxygen vacancies in the structure, the most prominent change in the Raman spectra of Figure 2 is the suppression of the sharp mode at 700 cm\(^{-1}\) with increasing \(\delta\). Furthermore, the intensity of the mode at 290 cm\(^{-1}\) decreases slightly, while the spectral weights of the other modes remain roughly unchanged (see Figure 2c). Also, the positions of the peaks shift slightly to higher wavenumbers with increasing \(\delta\), thereby indicating a variation of the internal stress between the two layers. For example, the peak of the most intense mode shifts gradually from 625 cm\(^{-1}\) for \(\delta = 0\) to 633 cm\(^{-1}\) for \(\delta = 0.15\) (Figure 2b).

If we assume a significant formation of oxygen vacancies within the CoO\(_2\) layer, several modes connected to the vibration and rotation of the CoO\(_6\) octahedra would be affected by the removal of oxide ions, both by changing the respective energy as well as the intensity. In particular, such changes should be visible for the E\(_1\)g and A\(_1\)g modes, likely at 540 and 630 cm\(^{-1}\). Because only the modes at 290 and 700 cm\(^{-1}\) show a significant change upon the formation of oxygen vacancies, we conclude that no significant oxygen nonstoichiometry is present within the CoO\(_2\) layer.

We therefore assign both modes at 290 and 700 cm\(^{-1}\) to vibrations within the rock salt layer. A mode similar to the one at 700 cm\(^{-1}\) with the same annealing behavior was observed in Co-doped ZnO (at 690 cm\(^{-1}\)) with different concentrations of oxygen vacancies and identified as a local disordered vibrational mode of a Zn–O–Co cluster.\(^{33}\) In that material, oxygen vacancies are preferentially formed close to Co dopant sites in Zn–O, so that the mode at 690 cm\(^{-1}\) loses its spectral weight when the sample is annealed in an oxygen poor atmosphere and recovers upon annealing in air. In analogy, we therefore assign the mode observed in CCO at 700 cm\(^{-1}\) with a local vibration of a Co–O–Co... cluster within the rock salt subsystem. When oxygen vacancies are created on those oxygen sites, the spectral weight of the corresponding mode is expected to decrease, as was observed experimentally.

For \(\delta = 0.15\), the relative spectral weight at 700 cm\(^{-1}\) is already reduced to \(\leq 25\%\) of its value for \(\delta = 0\) (Figure 2c), indicating that the associated vibrating Co–O–Co... cluster extends over several Co–O units. The formation of oxygen vacancies could induce a structural clustering of the RSL, as discussed in detail in Computational Results, which impedes the vibration of Co–O–Co... clusters associated with the Raman-active mode at 700 cm\(^{-1}\). We emphasize that the mode at 700 cm\(^{-1}\) is not a defect mode but a Raman active mode present in the perfect structure and suppressed upon defect formation. In summary, the results obtained from Raman spectroscopy suggest that oxygen vacancies are mainly formed on sites within the central Co–O layer inside the rock salt subsystem.

### 4.3. Direct Investigation of Oxidation Thermodynamics

The defect chemical model presented in Thermogravimetry and Defect Chemical Model contains a set of assumptions and simplifications. It is thus desired to measure the oxidation enthalpy of CCO in a more direct way, allowing for a critical assessment of the predictions made from this model. Simultaneous thermogravimetry and differential scanning calorimetry allows an unambiguous measurement of the enthalpy change connected to the exchange with oxygen.

As an example, the raw data of a TG-DSC oxidation experiment of CCO at 1050 K is shown in Figure 3. After 10 min, when both the TG and DSC signals were stable, the oxygen partial pressure \(p\text{O}_2\) within the measurement chamber was changed from \(\approx 0.1\) to 1 atm. The sample weight increased and the DSC signal showed an exothermic peak. Both observations can be assigned to the oxidation of oxygen vacancies in CCO. We note that the exact knowledge of the \(p\text{O}_2\) and thus the oxygen vacancy concentration in both states is not required to extract the molar oxidation enthalpy as only the differential \(\Delta H/\Delta m\) is needed for the calculation. When both signals were stable again, the atmosphere was changed back to a \(p\text{O}_2\) of 0.1 atm. The measurement was repeated twice to increase the statistical significance of the obtained result.

The DSC signal in Figure 3 shows a very different equilibrium value for the two different atmospheric compositions (10% \(\text{O}_2\) diluted in He and pure \(\text{O}_2\)) because of variations in the heat capacity and heat conductivities of the different atmospheres, thereby changing the heat flow from the crucible in both conditions. To allow a meaningful integration of the DSC peak, we therefore recorded a DSC background of an empty crucible under similar experimental conditions. A typical background is shown in Figure 4a. Some experimental limitations have to be considered before background subtraction can be done: The switch in the atmospheric composition cannot be done instantaneously, but several flowmeters of the gas mixer have to be adjusted manually. The onset time of the DSC signal for both the sample and background shown in Figure 4a can therefore be determined...
with an accuracy of only approximately 15 s, as indicated by the two times \( t_i \) and \( t_{i+1} \). In practice, this inaccuracy corresponds to a horizontal shift of the background with respect to the DSC signal of the sample when subtracting the background. We subtracted the background for the two extreme conditions of an early and late onset. The two resulting peaks are shown in Figure 4b. Using a sigmoidal baseline, the area for the two peaks typically differs by 20%, indicating the statistical inaccuracy of the extracted data. The error in \( \Delta H/\Delta m \) due to different integration boundaries of both DSC and TG signals is in the order of 3% and therefore negligible as compared to that of background subtraction.

The resulting oxidation enthalpies in the investigated temperature range lie between \(-39 \) and \(-68 \) kJ/mol (corresponding to a formation enthalpy of \( \Delta H_{\text{form}} = 0.40-0.71 \) eV per oxygen vacancy for the reverse process) and do not show a clear temperature trend (Figure 5). The extracted values do not show a clear temperature dependency. The value predicted from the defect chemical model is indicated for comparison (dashed line).

We first studied the geometric structure of pristine CCO for different structural models. On the basis of these results, we calculated the formation energy of an oxygen vacancy placed in different crystallographic positions. Finally, we compared the obtained electronic properties for the different structures and computational methods.

As mentioned in Theoretical and Computational Methods, we restricted our calculations to the 3/5 (\( q = 0.60 \)) and 2/3 (\( q = 0.67 \)) unit cells to approximate the incommensurately modulated structure of CCO. The relaxed cells for the two structural models we used are shown in Figure 6.

4.4. Computational Results. In parallel to the experimental investigations, we addressed the oxygen nonstoichiometry in CCO computationally. In particular, we aimed to check the crystallographic restriction of oxygen vacancies within the RSL by calculating the formation enthalpies of oxygen vacancies placed at different oxygen sites.

The misfit between the two moieties (COL and RSL) can lead to local stress, which in turn can induce symmetry breaking along the \( b \) direction. The periodicity of the \( \text{Ca}_x\text{CoO}_3 \) layers may thus adapt to that of the COL, forming \( \text{Ca}_x\text{CoO}_3 \) clusters along the \( b \) axis rather than equidistant rock salt units. Some clustering was indeed found in experiments performed at low temperatures by Muguerra et al.\textsuperscript{15} and also in the first-
principles calculations of Rébola et al.\textsuperscript{19} although with different periodicities. Experiments have shown that at low temperatures the RSL forms disordered clusters of different sizes and shapes, while at $T = 400$ K, a disorder-to-order transition occurs, favoring a phase with a homogeneous RSL layer.\textsuperscript{34}

We found such clustering of the RSL along the $b$ axis for the 2/3 unit cell (Figure 6a,b) but a homogeneous (nonclustered) RSL in the case of the 3/5 unit cell (Figure 6c,d). However, in all our DFT optimizations we obtained a few local minima similar in total energy but with different symmetry of the RSL. For example, for the 3/5 model, the structure with a homogeneous RSL was found to be more stable than a clustered one by only 320 meV/cell. The determination of the precise form of the clustered phase goes beyond the scope of this study, as many quasiequivalent energy minima are expected. Therefore, we may in the following consider the 2/3 cell as a model for the clustered (low $T$) phase and the 3/5 cell as a model for the homogeneous RSL (high $T$).

In order to compute the preferred site of oxygen vacancies, we have calculated via eq 3 the formation enthalpy of a positively charged oxygen vacancy placed at different lattice sites in the structure. We considered three different positions for the oxygen vacancy: in the COL, within the outer Ca–O layer of the RSL, and in the inner Co–O layer of the RSL. For the considered 2/3 and 3/5 unit cells, the stoichiometry of CCO can be written as $\text{Ca}_3\text{Co}_3.75\text{O}_9$, $\text{Ca}_3\text{Co}_3\text{O}_{9.5}$, and $\text{Ca}_3\text{Co}_3\text{O}_9$ respectively, with $\delta = 0.25$ when one oxygen atom is removed.

The description of the localized Co($d$) states is often improved by the inclusion of a Hubbard-like correlation (GGA + $U$).\textsuperscript{35} However, our calculations with $U = 5$ eV, as used by Rébola et al. and similar to $U$ values used in DFT studies on CCO and the related compound $\text{Na}_3\text{CoO}_2$,\textsuperscript{19,36,37} did not give reasonable results and are therefore omitted from the present paper. The formation enthalpies as obtained by DFT are reported in Table 1. The obtained values for the same defect position differ significantly for the different models, indicating the sensitivity of the electronic structure to the chosen unit cell, as discussed below. Nevertheless, common for both models considered, we found the $\Delta H_{\text{f,form}}$ of oxygen vacancies formed in the RSL to be lower than that of vacancies in the COL.

The smallest formation enthalpy was calculated for the inner part of the RSL. This is hence the most stable site for $\text{v}_O^\text{**}$, being somewhat more favorable ($\geq 200$ meV) than a vacancy in the outer RSL. The defect in the COL is much less stable. We did not find any significant deviations in formation energy between vacancies placed at the different oxygen sites within the clustered rock salt layer (for the 2/3 cell).

The structural changes upon oxygen vacancy formation in the COL and RSL are shown in Figure 7 (for the 3/5 structure). For the $\text{v}_O^\text{RSL}$ defect, we found a strong rearrangement of the coplanar Co and O ions, which move away from the defect site, while both the Ca and the O ions lying in the outer part of the RSL undergo only negligible shifts (Figure 7b,d). The Co–Co distances change from 4.87 to 5.10 Å along the $a$-axis and from 4.68 to 5.49 Å along the $b$-axis. This distortion and the consequent structural stress on the RSL induce locally a clustering along the $b$ axis in all the models we have studied, resulting in a structure similar to that found by Rébola et al. for the corresponding pristine cell.

For the $\text{v}_O^\text{**}$ defect, the vacancy formation induces only minimal lattice rearrangement in the CoO$_2$ oxide layer, where the Co ions relax about 0.1 Å toward the vacancy site (Figure 7a,c). We therefore suggest that the larger relaxation around the vacancy may be one of the contributions that stabilize the vacancy defect in the RSL with respect to the COL. As these results were obtained with the 3/5 cell, we note that we qualitatively found the same results for the defects formed for the 2/3 unit cell. As the pristine structure in that case already showed clustering of the RSL, the lattice rearrangement upon formation of the $\text{v}_O^\text{**RSL}$ defect was less effective in stabilizing the defective structure, resulting in formation enthalpies for the 2/3 cell higher than those for the 3/5 cell.

As pointed out earlier, the 3/5 cell with a homogeneous RSL can be considered to model the high-temperature phase ($T > 400$ K) of CCO, thus mimicking the experimental conditions of the TG and TG-DSC measurements. Indeed, the oxygen vacancy formation enthalpy $\Delta H_{\text{f,form}}$ of 0.41 eV calculated by DFT for this cell compares favorably with the experimentally determined values (0.5–0.6 eV), suggesting that the defect formation process is appropriately described by our computational approach.

Identifying the energetically most favorable site of an oxygen vacancy to be inside the central Co–O layer of the RSL confirms the conclusion drawn from our Raman spectroscopy results and the structural study by Ling et al.\textsuperscript{8} Furthermore, our finding of high formation energies for an oxygen vacancy placed in the COL is in agreement with our previous calculations on $\text{Na}_3\text{CoO}_2$ and indicates that the layer of edge-sharing CoO$_6$ octahedra, the common structural element of the misfit cobaltite family, is generally stable toward oxygen loss.

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<tr>
<th>Defect Placed in Different Crystallographic Positions and for the Different Theoretical Models We Adopted</th>
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<tr>
<td>$\Delta H_{\text{f,form}}^{\text{v}_O^\text{**}}$ (eV)</td>
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<td>inner-RSL</td>
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<td>2/3 cell</td>
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<td>3/5 cell</td>
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Figure 7. Geometries as obtained by DFT of the (a) pristine COL, (b) pristine RSL, (c) COL-$\text{v}_O^\text{**}$, and (d) RSL-$\text{v}_O^\text{**}$ seen from the (001) direction. While no structural rearrangement is seen for the vacancy inside the COL (c), lattice relaxation occurs around the vacancy in the RSL (d), effectively stabilizing the defect.
As the defect formation enthalpies in Table 1 appear to be sensitive to the chosen model, we now inspect the electronic properties of the investigated models in more detail. The electronic structures for the pristine cells computed are shown in Figure 8. The obtained band structures are in general agreement with the calculations performed by Rébola et al. The states close to $E_F$ are mostly due to hybridized Co($d$)–O($p$) states. The COL states are split in the octahedral field to form t$_{2g}$–e$_g$ states. On the other hand, as expected for the rock salt coordination, the RSL Co($d$) states are not split into t$_{2g}$–e$_g$ states but appear spread over a broad energy range.

The 2/3 cell (Figure 8a) is half-metallic, with a 0.75 eV gap in one of the spin channels, while the 3/5 cell (Figure 8b) is metallic, and the gap is shifted to higher energies, from about 0.25 eV above the Fermi energy. In both the 2/3 and 3/5 models, the t$_{2g}$ orbitals are fully occupied for one of the spin components and only partially occupied for the other. This implies a mixed valence scenario for the COL, in which most of the Co ions are in a +3 oxidation state with itinerant holes (representing Co$^{4+}$ states) contributing to the electrical conduction, as confirmed by Hall measurements.

On the other hand, the e$_g$ states are empty for both models. Furthermore, we find RSL states contribute to the total DOS at $E_F$ for both models, contradicting experimental results of Takeuchi et al., which find only COL states at the top of the valence band. We ascribe the differences in Figure 8 between the 2/3 and 3/5 cell to the different RSL symmetry in both models, in agreement with Rébola et al., suggesting that the density of states close to $E_F$ is strongly influenced by the formation of (Ca$_2$CoO$_3$)$_{62/3}$ clusters within the RSL.

Each of our models fits some of the experimental findings, but none is capable of reproducing the whole electronic structure of CCO. Therefore, we cannot conclude which of the studied models is the most realistic approximation to describe CCO. Extending the approximate unit cell along $b$ (5/8, 8/13 cells) may improve the agreement between computational and experimental results. However, such an increase in unit cell size would increase the computational effort significantly. This is in particular true for defect formation calculations, as multiples of the unit cell along the shortest lattice parameters ($a$ for CCO) need to be included to avoid interactions between periodic defect images. The accumulated magnitude of these and other sources of error is difficult to quantify without efforts beyond the scope of this work. Nevertheless, the good correspondence between the calculated and measured values of the formation energy indicates that the chosen atomistic models are indeed relevant for the physical system. Moreover and most importantly, we find the relative cost of forming oxygen vacancies at various sites to be robust with respect to all tested parameters, providing computational evidence that the main loss of oxygen takes place from the central Co–O layer of the RSL.

5. CONCLUSION

In summary, we have studied the position and thermodynamics of oxygen vacancies in (Ca$_2$CoO$_3$)$_{62/3}$(CoO$_2$) both theoretically and experimentally.

By simultaneous measurements of the weight and enthalpy change (TG-DSC), the molar oxidation enthalpy $\Delta H_{Ox}$ for temperatures between 950 and 1100 K was measured to be in the range of $-39$ to $-68$ kJ/mol (corresponding to a formation enthalpy $\Delta H_{form}$ of 0.40–0.71 eV per oxygen vacancy), confirming predictions made from a previously published defect chemical model ($\Delta H_{chem} = -53$ kJ/mol, corresponding to $\Delta H_{form} = 0.57$ eV).

Raman spectra of a series of CCO samples prepared with different oxygen vacancy concentrations $\delta$ were taken at room temperature. We interpreted the observed depletion of a sharp mode at 700 cm$^{-1}$ with increasing $\delta$ as an indication that oxygen vacancies are formed predominantly within the central Co–O layer of the Ca$_2$CoO$_3$ subsystem.

This was confirmed by DFT calculations on different structural models of CCO, which found the lowest oxygen vacancy formation enthalpy $\Delta H_{form}$ for the same site. Strong defect-induced lattice rearrangement of the Ca$_2$CoO$_3$ subsystem led to a computational $\Delta H_{form} = 0.41$ eV (corresponding to $\Delta H_{chem} = -40$ kJ/mol) in good agreement with the experimental values. The electronic properties calculated by DFT are sensitive to the employed structural model, thereby demonstrating the difficulties of a detailed computational analysis of misfit compounds. Nevertheless, this study showed how different techniques can be used to obtain fundamental and detailed understanding of defects and their effect on functional materials in general and the high-temperature properties of (Ca$_2$CoO$_3$)$_{62/3}$(CoO$_2$) in particular.

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