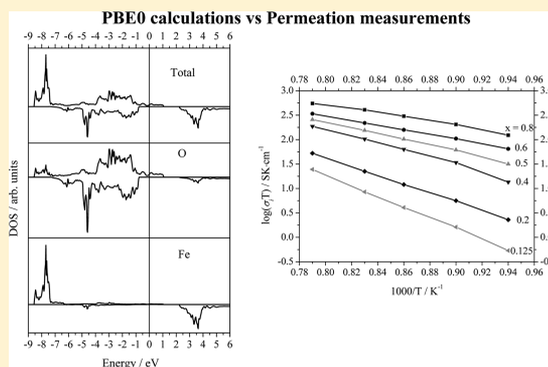


Comparison of Permeation Measurements and Hybrid Density-Functional Calculations on Oxygen Vacancy Transport in Complex Perovskite Oxides

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Supporting Information

ABSTRACT: The oxygen vacancy ($V_{\text{O}}^{\bullet\bullet}$) formation enthalpy (ΔH_{F}) was determined from the effective activation energies obtained in the permeation measurements on a series of solid solutions $\text{La}_{0.98-x}\text{Sr}_x\text{Co}_{0.2}\text{Fe}_{0.8}\text{O}_{3-\delta}$ ($x = 0.125-0.8$) and compared with the results of *ab initio* calculations at finite temperatures based on the hybrid (PBE0) and GGA+U calculations, as well as previous thermogravimetric experiments. The Mulliken atomic charges, magnetic moments, and the Fe- $V_{\text{O}}^{\bullet\bullet}$ and Sr- $V_{\text{O}}^{\bullet\bullet}$ distances are analyzed. The strong dependence of formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ on the Fe oxidation state due to variation in Sr and $V_{\text{O}}^{\bullet\bullet}$ concentrations is discussed.



1. INTRODUCTION

(La,Sr)(Co,Fe) $\text{O}_{3-\delta}$ (LSCF) is known as one of the best materials for oxygen permeation membranes^{1,2} and cathodes in solid oxide fuel cells.³ This is due to the fact that such complex perovskite oxides show high mixed ionic and electronic conductivities (MIEC) and, hence, high oxygen permeability. The highest oxygen flux is generally observed for dense ceramic membranes having a perovskite structure ABO_3 at temperatures above 800 °C (ref 4). This relies on a high concentration of mobile oxygen vacancies as well as a high electronic conductivity. Since the electronic conductivity in perovskites is usually orders of magnitude larger compared to the ionic conductivity, the latter is the limiting transport mechanism, so that permeability (assuming fast surface reaction) can be expressed as

$$j_{\text{O}_2} = \frac{RT}{16F^2L} \sigma_i \ln \frac{p'_{\text{O}_2}}{p''_{\text{O}_2}} \quad (1)$$

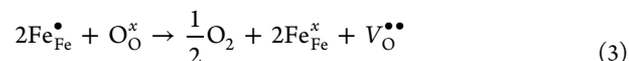
where R is the gas constant, T temperature, F Faraday constant, L membrane thickness, σ_i ionic conductivity, and p'_{O_2} and p''_{O_2} the oxygen partial pressure on both sides of the membrane.

High ionic conductivity can be achieved by maximizing diffusivity and particularly concentration of oxygen vacancies. It is well-known that Sr^{2+} ion substituting for La^{3+} is compensated by electronic defects (holes) or oxygen vacancies. The oxygen vacancies prevail over electronic defects at high temperatures.⁵

Such vacancy formation, e.g., in LaFeO_3 (LF), can be described (in Kröger-Vink notations⁶) as



With increasing temperature and/or decreasing oxygen partial pressure, additional vacancies are formed due to partial reduction of the B-site cation, that is, Fe in a solid solution $\text{La}_{1-x}\text{Sr}_x\text{FeO}_{3-\delta}$ (LSF) decreases its oxidation state from partially 4+ ($\text{Fe}_{\text{Fe}}^{\bullet}$) closer to 3+ (Fe_{Fe}^x)



Since Co shows higher reducibility compared to Fe, maximum oxygen fluxes in LSCF can be found for Sr- and Co-rich compounds.¹ On the other hand, excessively high concentration of oxygen vacancies inherently reduces the stability of the crystal lattice, so that pure strontium cobaltite does not have the desired perovskite structure and, hence, permeability vanishes. It turned out that 20 mol % of Fe is sufficient to stabilize the perovskite structure of $\text{SrCo}_{0.8}\text{Fe}_{0.2}\text{O}_{3-\delta}$ in air. However, in nitrogen atmosphere, i.e., decreased oxygen partial pressure, the perovskite transforms into a brownmillerite phase,⁷ which has an ordered arrangement of oxygen vacancies,

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resulting again in drastically decreased vacancy mobility. Partial substitution of Sr with Ba helps to avoid the brownmillerite formation in $\text{Ba}_{0.5}\text{Sr}_{0.5}\text{Co}_{0.8}\text{Fe}_{0.2}\text{O}_{3-\delta}$,⁷ which shows the highest oxygen fluxes observed so far.^{8,9} But still, this compound is close to the stability limits, with regard to crystal structure¹⁰ and reactivity with acid gases. At temperatures below 840 °C an additional hexagonal phase forms reducing oxygen fluxes.^{11–13} Moreover, carbonates are easily formed in the presence of CO_2 , again drastically reducing the permeability.¹⁴ Therefore, LSCF with limited Sr- and Co-content, although already discovered 30 years ago, has again attracted increasing interest as oxygen transport membrane material; significant oxygen fluxes could be achieved using thin supported membrane layers in $\text{La}_{0.6}\text{Sr}_{0.4}\text{Co}_{0.2}\text{Fe}_{0.8}\text{O}_{3-\delta}$.

The two parent compounds LF and SrFeO_3 (SF) are characterized by the existence of Fe ion in the two different oxidation states, Fe^{3+} ($3d^5$) and Fe^{4+} ($3d^4$), respectively. Moreover, LF shows the electronic conductivity increasing with the temperature whereas SF is a metallic compound (at least for small δ). The lattice structure changes from orthorhombic for LF to cubic for SF at room temperature.¹⁵ It would then not be surprising to observe complex magnetic,¹⁶ electrical,¹⁷ and phonon properties¹⁸ for the solid solutions of LF and SF. It is already known from the literature¹⁹ that the $V_{\text{O}}^{\bullet\bullet}$ formation enthalpy for the solid solution LSF changes in a step-wise manner at $\delta = x/2$, where the jump between the oxidation states of Fe ($2+/3+$ and $3+/4+$) occurs.

Recently, we have considered the oxygen vacancy properties in $\text{La}_{1-x}\text{Sr}_x\text{Co}_{0.25}\text{Fe}_{0.75}\text{O}_{3-\delta}$ using the density functional theory (DFT) for calculating defect formation enthalpy as a function of nonstoichiometry and temperature.²⁰ The calculated defect formation enthalpy is very close to that from a similar DFT study on LSF due to low content of Co.²¹ However, a comparison of the results of these calculations and the thermogravimetric measurements revealed that the calculated formation enthalpy may significantly differ from the measured one at low temperature and high concentrations of Sr although the oxidation entropies suggested no such significant differences. From the experimental point of view, the formation enthalpy may be affected by the oxygen partial pressure and, thus, the Fe oxidation state at fixed Sr content.¹⁹ From the calculation point of view, the magnetic moment on Fe is sensitive to the exchange-correlation functional, affecting the formation enthalpy. It has been shown indeed within the DFT+U calculations and generalized gradient approximation (GGA) by Ritzmann et al.²² that the inclusion of the strong correlation effects significantly affects the calculated formation energy of $V_{\text{O}}^{\bullet\bullet}$. However, one should keep in mind that the results of such GGA+U calculations depend very much on the choice of Hubbard-U parameter.

The $V_{\text{O}}^{\bullet\bullet}$ migration energy was calculated in LF using the GGA+U approach,^{22,23} and in LF, LSF within the standard GGA approach,²⁴ both studies applied the so-called NEB method as implemented in the VASP code.²⁵ It was concluded²⁴ that the defect migration energy is not essentially influenced by the average Sr concentration while the *local* configuration of Sr and/or La in the critical triangle in the saddle point of migration affects the barrier height. Thus, the formation energy of $V_{\text{O}}^{\bullet\bullet}$ and its strong dependence on Sr doping is at the focus of the comparison with the experimental data and represents a key milestone for the DFT calculations. In addition, *hybrid* DFT calculations were shown to be very effective in treatment of the electronic structure of correlated

materials and, in particular, perovskites,^{26–29} as well as in calculations of $V_{\text{O}}^{\bullet\bullet}$ formation energy therein.³⁰ We, therefore, utilize in the present study not only the GGA+U calculations, but also such the hybrid DFT calculations; in order to deeper analyze the defect formation energies in LSF with different Sr concentrations.

Our calculations were motivated by the permeation experiments on the role of Sr doping in oxygen transport. In Section 2 the experimental technique, sample preparation are described. Section 3 is devoted to the computation methods including the computational parameters and calculations of the $V_{\text{O}}^{\bullet\bullet}$ formation energies. Results and Discussion are given in Section 4, subdivided into three subsections: oxygen permeation measurements, the calculated bulk properties and a comparison between the calculated $V_{\text{O}}^{\bullet\bullet}$ properties and experimental results. In Section 5 we present the main conclusions.

2. EXPERIMENTAL SECTION

$\text{La}_{0.98-x}\text{Sr}_x\text{Co}_{0.2}\text{Fe}_{0.8}\text{O}_{3-\delta}$ ($x = 0.2, 0.4, 0.6, 0.8$) perovskite powders were prepared by spray pyrolysis of the aqueous nitrate solutions containing the corresponding metal nitrates. After spray-drying the powders were calcined at 1200 °C for 5 h and examined by X-ray diffraction (XRD) using a Bruker, D4 Endeavor to ensure phase purity and by inductive coupled plasma–optical emission spectroscopy (ICP-OES) to verify the desired stoichiometry. Lanthanum under-stoichiometry was chosen in order to prevent possible La_2O_3 phases, which lead to sample failure due to high hygroscopy. In case of $x = 0.4$ another powder with stoichiometric La content was prepared, processed, and tested identically. There was no influence of the small La under-stoichiometry on oxygen permeation and, hence, ionic conductivity.

All powders were treated by ball milling in ethanol, using 3 mm ZrO_2 balls, to achieve an average particle size (d_{50}) of 2–4 μm . Samples for oxygen permeation measurements were uniaxially dry pressed and sintered at 1200 °C for 5 h achieving 90% of theoretical density and subsequently ground to an outer diameter of 14.7 mm.

Oxygen permeation measurements of the membranes were conducted in air/Ar gradients with a constant flow rate of 250 mL/min of air as feed gas and 50 mL/min of Ar as sweep gas at varying temperatures. The disc-shaped samples with thickness of 1 mm were sealed with gold rings to the gasket of the quartz glass reactor at 1000 °C. Inner diameter of gold seals is 13 mm leading to 1.33 cm^2 membrane area. The permeated gas was analyzed by a mass spectrometer (Omnistar, Pfeiffer Vacuum).

3. COMPUTATIONAL SECTION

The DFT calculations in the present study are performed using both the plane wave (PW) basis set as implemented in VASP5.2 code^{25,31} and the linear combination of atomic orbitals (LCAO) as implemented in CRYSTAL09 code.³² We will compare the results of DFT+U method³³ and hybrid PBE0³⁴ exchange-correlation functional (hereafter, functional) important for the proper treatment of the electronic structure of strongly correlated systems such as LSF.^{22,35} We give emphasis to the analysis of the electronic density of states (DOS), atomic charges, the *z*-projection of the magnetic moments (hereafter magnetic moments) on Fe, and the Fe–O bond lengths in perfect as well as defective LSF in the calculations, using both the DFT+U and hybrid PBE0 functionals.

We use, hereafter, the following abbreviations for the two solid solution compositions used in the present study for the analysis of main results in perfect and defective systems: LSF12.5 (12.5% of Sr) for $\text{La}_{0.875}\text{Sr}_{0.125}\text{FeO}_{3-\delta}$ and LSF50 (50% of Sr) for $\text{La}_{0.5}\text{Sr}_{0.5}\text{FeO}_{3-\delta}$.

It is known from XRD measurements^{15,16,36} of the crystalline structure of LSF12.5 and LSF50 that the former is orthorhombic at room temperature and rhombohedral at higher temperatures whereas the latter is rhombohedral at low and room temperature but becomes cubic at high temperatures. Thus, for a better comparison of these two compounds we used the rhombohedral symmetry for both LSF12.5 and LSF50.

3.1. Parameters for Density Functional Calculations.

In the LCAO calculations, the Gaussian basis set and relativistic pseudopotentials (PPs) of a free La atom with 46 electrons was taken from the PPs library of the Stuttgart/Cologne group.³⁷ The small-core relativistic effective core pseudopotential replacing 28 electrons and the Gaussian basis set for Sr were taken from ref 38. Notice that the basis set for Sr in our study included the 3d-electron exponent, since in the previous study on SrTiO_3 ,³⁹ this basis set reproduced the phonon properties better than without this exponent. The all-electron basis set 8–411G⁴⁰ was used for O atom whereas the all-electron basis set for Fe was taken from ref 41. In order to avoid spurious interactions between the diffuse functions and the core functions of neighboring atoms, the basis set diffuse exponents smaller than 0.15 bohr⁻² were removed. To optimize the basis set, we used Powell's conjugate-directions minimization method⁴² without calculations on the total-energy derivatives as implemented in the basis set optimization code OPTBAS.⁴³ The bound-constrained optimization was performed for the exponents with 0.1 bohr⁻² lower bound. The exponents of Gaussian basis functions in noncontracted basis functions were optimized in the calculations with PBE0 functional for rhombohedral LSF50. Notice that this basis set optimization method was already successfully used for LaCoO_3 ³⁵ and SrTiO_3 .³⁹ In Supporting Information (Tables S1–S4) we give the basis set for LSF50 used in the present study.

In the PW calculations we used the projector augmented wave (PAW) method⁴⁴ and scalar-relativistic pseudopotentials for 46 electrons on La, 12 electrons on Fe, 28 electrons on Sr, and 2 electrons on O atoms. The exchange-correlation functional was due to Perdew, Burke, and Ernzerhof (PBE).⁴⁵ The cutoff energy was fixed at 520 eV throughout all the calculations. The electron occupancies were determined with the Gaussian method using a smearing parameter 0.1 eV. Correspondingly, the rotationally invariant Dudarev's form⁴⁶ for the Hubbard correction was used in the PBE+U calculations. However, the advantage of this form is due to the difference $U_{\text{eff}} = U - J$, of the Hubbard parameter U and the exchange parameter J . We use, hereafter, the abbreviation PBE+ U_{eff} in discussion of the results obtained. We used the two values of U_{eff} 4.5 and 6.5 eV, to compare not only the bulk properties but also the $V_{\text{O}}^{\bullet\bullet}$ formation enthalpy. A ferromagnetic structure was assumed throughout all the calculations in VASP and CRYSTAL.

In the perovskite-type rhombohedral structure (space group (SG) 167) the primitive unit cell contains 10 atoms, thus, twice the number of atoms in the perovskite-type cubic structure (5 atoms, SG 221).³⁵ In the present study, we substituted one La atom for one Sr atom in the unit cell containing 10 atoms in order to consider the bulk properties for the content of Sr 50% and to guarantee the highest possible symmetry (SG 155) in

this case. In contrast, the content of Sr 12.5% needs an increase of the unit cell up to 40 atoms, at least, and substituting one La atom for one Sr atom therein. We used the supercell approach⁴⁷ for the calculations of LSF12.5 and a single $V_{\text{O}}^{\bullet\bullet}$ in both LSF12.5 and LSF50. The 40 atoms supercell was

generated with the help of the transformation matrix $\begin{pmatrix} 211 \\ 121 \\ 112 \end{pmatrix}$ for

the translational vectors. A single $V_{\text{O}}^{\bullet\bullet}$ corresponds in such a supercell to the nonstoichiometry $\delta = 0.125$. In addition, the 80 atom supercell (the $2 \times 2 \times 2$ extension of the rhombohedral primitive unit cell) with two Sr ions was used to simulate the case of lower nonstoichiometry, $\delta = 0.0625$ in LSF12.5. In the present study, we used the FINDSYM program of Hatch and Stokes⁴⁸ in order to identify the corresponding SG and Wyckoff positions of all atoms in the supercells. Notice that the number of symmetry operations is reduced in CRYSTAL calculations when making the supercells for the non-symorphic SGs. If the La atom substitutes for the Sr atom in the supercells for LSF12.5, the SG 146 is obtained in the CRYSTAL calculations.

The Monkhorst–Pack scheme⁴⁹ for $8 \times 8 \times 8$ k -point mesh in the Brillouin zone was applied to the 10 atoms unit cell, $3 \times 3 \times 3$ to extended 40 atoms and $4 \times 4 \times 4$ to extended 80 atoms supercells. The tolerance factors of 7, 7, 7, and 14 for the Coulomb and exchange integrals calculations were used in the CRYSTAL calculations. Furthermore, the forces for the self-consistent cycles were optimized until the energy difference reached 10^{-7} eV and were smaller than 0.01 eV/Å for the lattice structure optimization (in both perfect and defective crystals) in the CRYSTAL and VASP calculations, respectively.

3.2. Defect Formation Enthalpy. Calculating the enthalpy of $V_{\text{O}}^{\bullet\bullet}$ formation is relevant for a comparison with the results of permeation measurements. The defect formation enthalpy is given by

$$\Delta H_{\text{F}}(T) = E_{\text{tot}}^{V_{\text{O}}^{\bullet\bullet}} - E_{\text{tot}}^{\text{P}} + \Delta H_{\text{vib}}(T) + \frac{1}{2}\mu_{\text{O}_2}^0(T) \quad (4)$$

where superscripts p and $V_{\text{O}}^{\bullet\bullet}$ indicate perfect and defective crystals, E_{tot} total electron energies, and T the temperature. All the energies are given per supercell. The standard chemical potential of oxygen $\mu_{\text{O}_2}^0$ is given by half the total energy of oxygen molecule plus the vibrational, rotational, and translational contributions, to include the temperature effect. In our previous study⁵⁰ we have demonstrated that the calculations of the properties of O_2 with the hybrid PBE0 functional in the LCAO method lead to the very accurate results for the binding energy (5.30 eV), equilibrium distance (1.20 Å), as well as rotational (2.11 K) and vibrational (2478.60 K) temperatures. Therefore, the chemical potential of O_2 in the LCAO calculations reads

$$\mu_{\text{O}_2}^0(T) = E_{\text{tot}}^{\text{O}_2} + E^{\text{O}_2} + kT \quad (5)$$

where $E_{\text{tot}}^{\text{O}_2}$ the total electronic energy of O_2 , E^{O_2} the thermal energy which includes the translational, rotational, and vibrational contributions to the chemical potential of O_2 as calculated within an ideal gas model in the CRYSTAL code, and k Boltzmann's constant. It means that the calculations of ΔH_{F} with the PBE0 functional in the LCAO method do not need use of any experimental data, in strong contrast to the PW calculations with standard DFT and/or PBE+ U_{eff} functionals. We included the vibrational contribution in the solid phase $\Delta H_{\text{vib}}(T)$ in the present study by taking data from our previous study on $(\text{La,Sr})\text{Co}_{0.25}\text{Fe}_{0.75}\text{O}_3$,²⁰ as calculated within the

harmonic approximation and the direct method⁵¹ for calculating phonon frequencies.

As is known, the standard DFT functional in the PW approach leads to errors for the binding energy and/or equilibrium distance of O₂.⁵² There are several suggestions in the literature regarding how to correct this effect.^{53,54} We rely on the approach suggested by Finnis et al.⁵⁵ which is based on the experimental data on oxide heats of formation, thereby avoiding a calculation of the total energy of an oxygen molecule. Thus, the oxygen chemical potential in our PBE+U_{eff} calculations of ΔH_F reads

$$\mu_{\text{O}}^0(T) = E_{\text{tot}}^{\text{AO}} + \frac{1}{m}E_{\text{vib}}^{\text{AO}}(T^0) - E_{\text{tot}}^{\text{A}} - \frac{1}{l}E_{\text{vib}}^{\text{A}} - \Delta G^{\text{AO}}(T^0) + \Delta H_{\text{O}}(T) \quad (6)$$

where superscripts AO and A stand for the reference binary oxide, and its metal, respectively; in this we have used MgO, T⁰ is equal in our case to standard state temperature 298.15 K, ΔG^{AO} is the standard Gibbs free energy of formation, taken from a thermodynamic database,⁵⁶ i.e., ΔG^{MgO} = 601.6 kJ/mol. ΔH_O(T) is the difference in enthalpies at the temperature of interest and standard state, which is also taken from the thermodynamic database.⁵⁶ E_{vib} is the vibrational contribution in addition to the internal energy. The prefactors 1/l and 1/m represent the ratios of the number of atoms in the primitive unit cell to that in the supercell for binary oxide and metal, respectively, which are both 1/27 for our case.

We used also the so-called *ghost* basis set as implemented in the CRYSTAL code.³² It assumes that the basis set of a missing O atom remains in the vacancy with a zero core charge. This means that the electron density of two electrons stemming from a missing oxygen atom is divided between the oxygen site and neighboring cations. Thus, use of a ghost basis set provides a reasonable initial guess for the electronic density distribution and in addition allows us to determine the effective electronic charge in the vacancy position.

4. RESULTS AND DISCUSSION

4.1. Oxygen Permeation Measurements. In order to properly compare the results of permeation measurements and present hybrid PBE0 and PBE+U_{eff} calculations, we discuss here the procedure used to deduce the V_O^{••} formation and migration energy from the experimental profiles. We suppose that the high oxygen permeation in perovskites is due to the conduction mechanism of oxygen ions diffusing through oxygen vacancies in the crystal lattice. According to eq 1, the oxygen permeation rate is proportional to the ionic conductivity at constant thickness, temperature, and oxygen partial pressure gradient.

Normalization of the permeation rate by the geometrical (thickness) and experimental (ln(p_{O₂'₂)/p_{O₂'₁)) terms results in the *specific permeability*, first introduced by Kharton⁵⁷ (assuming fast surface reaction)}}

$$J_{\text{O}_2} = \frac{j_{\text{O}_2} \cdot L}{\ln \frac{p_{\text{O}_2}'^2}{p_{\text{O}_2}''^2}} = \frac{R}{16F^2} \sigma_i T \quad (7)$$

According to the Nernst–Einstein-relation, the product of ionic conductivity and temperature is

$$\sigma_i T = \frac{4F^2 [V_{\text{O}}^{\bullet\bullet}] D_V}{RV_m} \quad (8)$$

where V_m is the molar volume of the compound, [V_O^{••}] the oxygen vacancy concentration, and D_V the vacancy diffusion coefficient.

Combining eqs 7 and 8 results in

$$J_{\text{O}_2} = \frac{1}{4V_m} [V_{\text{O}}^{\bullet\bullet}] D_V \quad (9)$$

Since the temperature dependence of the molar volume of LSCF is negligible compared to that of [V_O^{••}] and D_V, the effective activation energy found from the slope of Arrhenius plots of the specific permeability equals a *sum* of the oxygen vacancy formation energy and migration energy (for low [V_O^{••}], see discussion below).

Figure 1 shows experimental results for a series of solid solutions La_{0.98-x}Sr_xCo_{0.2}Fe_{0.8}O_{3-δ} with increasing Sr content, x

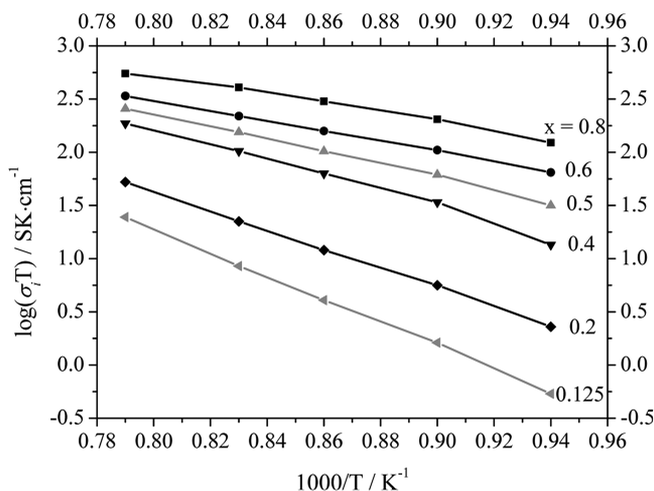


Figure 1. Arrhenius plot of ionic conductivity times temperature of La_{0.98-x}Sr_xCo_{0.2}Fe_{0.8}O_{3-δ}. Black lines represent experimental data for x = 0.2, 0.4, 0.6, and 0.8. Gray lines are interpolated for x = 0.125 and 0.5. Typical error bar is 5%.

= 0.2, 0.4, 0.6, and 0.8. The curves shown are calculated σ_iT from oxygen permeation measurements in the temperature range of 1073–1273 K. For x = 0 the permeation rate is assumed to be zero due to negligible oxygen vacancy concentration. Using this experimental data set, it is possible to interpolate the specific permeability for each temperature. The resulting lines for x = 0.125 and 0.5 are also plotted in Figure 1. Table 1 shows the respective effective activation

Table 1. σ_iT@1200 K (in SK/cm) for La_{0.98-x}Sr_xCo_{0.2}Fe_{0.8}O_{3-δ} and Corresponding Effective Activation Energies E_A (in eV) in the Temperature Range 1073–1273 K According to Figure 1

x	σ _i T	E _A
0.125	0.89	2.15
0.2	1.35	1.74
0.4	1.93	1.41
0.5	2.15	1.17
0.6	2.29	0.95
0.8	2.60	0.88

energies. We compare the formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ found from measured effective activation energies by subtracting the corresponding $V_{\text{O}}^{\bullet\bullet}$ migration energies from the DFT calculations:⁵⁸ 0.75 and 0.84 eV for LSF12.5 and LSF50, respectively. This leads to a sharp decrease of the predicted $V_{\text{O}}^{\bullet\bullet}$ formation energies as Sr concentration grows: from 1.40 eV down to 0.33 eV.

The equilibrium constant for reaction given by eq 3 depends on temperature via the standard reaction (vacancy formation) enthalpy

$$\ln K = \frac{-\Delta H^0}{RT} + \frac{-\Delta S^0}{R}$$

and is related to the defect concentrations by

$$K = \frac{\sqrt{p_{\text{O}_2}} [V_{\text{O}}^{\bullet\bullet}] [\text{Fe}_{\text{Fe}}^x]^2}{[O_{\text{O}}^x] [\text{Fe}_{\text{Fe}}^x]^2} = \frac{\sqrt{p_{\text{O}_2}} [V_{\text{O}}^{\bullet\bullet}] (1 - [\text{Sr}'_{\text{La}}] + 2 \cdot [V_{\text{O}}^{\bullet\bullet}])^2}{(3 - [V_{\text{O}}^{\bullet\bullet}]) ([\text{Sr}'_{\text{La}}] - 2 \cdot [V_{\text{O}}^{\bullet\bullet}])^2} \quad (10)$$

However, the temperature dependence of K does not always directly correspond to the temperature dependence of the vacancy concentration (as extracted from E_a of σ_{ion} in Table 1 and the vacancy migration barrier). This simple relation holds only for $[V_{\text{O}}^{\bullet\bullet}] \ll [x/2]$ when

$$K \approx \frac{\sqrt{p_{\text{O}_2}} [V_{\text{O}}^{\bullet\bullet}] (1 - [\text{Sr}'_{\text{La}}])^2}{3 \cdot [\text{Sr}'_{\text{La}}]^2} \propto [V_{\text{O}}^{\bullet\bullet}] \quad (11)$$

On the other hand, for conditions with large $[V_{\text{O}}^{\bullet\bullet}]$ (i.e., high T and/or less positive ΔH^0), the increase of K with increasing temperature (decreasing T^{-1}) mainly corresponds to the term $([\text{Sr}'] - 2[V_{\text{O}}^{\bullet\bullet}])$ in the denominator becoming small, while $[V_{\text{O}}^{\bullet\bullet}]$ in the numerator approaches $[\text{Sr}']/2$ and is almost constant. Thus, while K increases with decreasing $1/T$, $[V_{\text{O}}^{\bullet\bullet}]$ levels off. In between these two regimes, the slope of σ_{ion} versus $1/T$ gradually decreases. Indeed the curves in Figure 1 indicate some bending, in particular, for materials with high Sr content. Unfortunately, the temperature range covered in the measurement is not large enough to exploit this nonlinearity for fitting the complete defect model (and thus ΔH^0) as defined by eq 11. Nevertheless, one has to keep in mind that ΔH^0 values estimated from Figure 1, assuming validity of the approximate eq 11 might underestimate the real ΔH^0 values.

4.2. Bulk Properties Obtained in Density Functional Calculations. The results of PBE0 and PBE+ U_{eff} calculations of bulk properties for LSF50 and LSF12.5 are compared with the experimental data in Tables 2–3. Obviously, the experimental lattice parameters in the rhombohedral LSF50 are better reproduced with the hybrid PBE0 functional than with the PBE+ U_{eff} one (Table 2). The latter approach typically overestimates the lattice parameters (if the U_{eff} parameter is not directly fitted). The calculated lattice parameters for LSF12.5 with PBE0 are close to experiments for the rhombohedral phase (Table 3). Since the substitution of La for Sr in the hypothetical rhombohedral LF with the SG 167 reduces the symmetry of the LSF50 crystal to the SG 155, the oxygen atoms are split and occupy two Wyckoff positions in its 10 atoms unit cell, namely, d for O1 and e for O2. In the 40 atom unit cell of LSF12.5 (SG146) there are 8 oxygen atoms occupying the Wyckoff position b with different coordinates.

Table 2. Calculated Lattice Parameters ($a = b, c$, in Å) Using the Hexagonal Setting, Atomic Charge (q , in e), Magnetic Moment (μ_{Fe} in μ_{B}) of Fe and the Fe–O Distance in Å in the 10 Atom (Stoichiometric in Oxygen) Unit Cell of LSF50 (SG 155) for PBE0 in the LCAO Calculations and PBE+ U_{eff} Functional in the PW Calculations^a

property	LCAO		PW, PBE+ U_{eff}		experiment
	PBE0		$U_{\text{eff}} = 4.5$ eV	$U_{\text{eff}} = 6.5$ eV	
a,b	5.54		5.58	5.59	5.51 ^b
c	13.42		13.48	13.49	13.42 ^b
q					
Sr	1.92		1.58	1.58	
La	2.65		2.11	2.12	
Fe	2.24		1.85	1.90	
O1	−1.50		−1.23	−1.26	
O2	−1.51		−1.24	−1.27	
μ_{Fe}	4.10		4.06	4.22	3.8 ^c
$d_{\text{Fe-O}}$	1.96 [1.97]		1.98 [1.99]	1.98 [2.00]	1.96 ^c

^aSee the text for details. Notice that the atomic charge values for the PBE+ U_{eff} correspond to the Bader topological analysis^{60,61} whereas those for PBE0 are due to the Mulliken population analysis.⁶² The values in square brackets correspond to the Fe–O2 distance in LSF50. ^bRef 59. ^cNeutron diffraction measurements at room temperature for CO/CO₂ quenched samples with high deviation from oxygen stoichiometry.⁶³

Table 3. Calculated Lattice Parameters ($a = b, c$, in Å) Using the Hexagonal Setting, Atomic Charge (q , in e), Magnetic Moment (μ_{Fe} in μ_{B}) of Fe and the Fe–O Distance in Å in the 40 Atom (Stoichiometric in Oxygen) Unit Cell of LSF12.5 (SG 146) for PBE0 in the LCAO Calculations^a

property	LCAO		experiment
	PBE0		
a,b	5.61		5.61 ^b
c	13.46		13.63 ^b
q			
Sr	1.92		
La	2.65		
Fe	2.26		
O	−1.60		
μ_{Fe}	4.29		
$d_{\text{Fe-O}}$	2.00		1.96–2.00 ^c

^aDue to large number of atoms in the supercell and low symmetry in the CRYSTAL calculations, only the average values for q and μ_{Fe} are given. Notice that the atomic charge values are due to the Mulliken population analysis.⁶² ^bTaken from ref 15 at 1023 K. ^cTaken from ref 16 for orthorhombic phase.

Therefore, only the average values for q and μ_{Fe} are given in Table 3. The two split oxygen atoms in LSF50 differ by the distance to Sr and Fe atoms. So, the oxygen atom O1 [O2] has a distance from the Sr atom of 2.55 [2.46] Å in the PBE+ U_{eff} calculations, irrespective of the choice of U_{eff} (6.5 or 4.5 eV). The values for the distance between O1[O2] and Sr atoms are slightly larger in the PBE0 calculations, being 2.57 [2.76] Å. It is also seen from the analysis of atomic charges for LSF50 (Tables 2 and 3) that the LCAO calculations with the PBE0 functional suggest a somewhat more ionic picture for LSF50 than the PW calculations with the PBE+ U_{eff} functional.

Also, the magnetic moment of Fe (μ_{Fe}) from the PBE0 calculations is slightly larger than that from the PBE+ U_{eff} calculations, at least for LSF50 and $U_{\text{eff}} = 4.5$ eV. The $d_{\text{Fe-O}}$ values from PBE+ U_{eff} calculations are overestimated, whereas

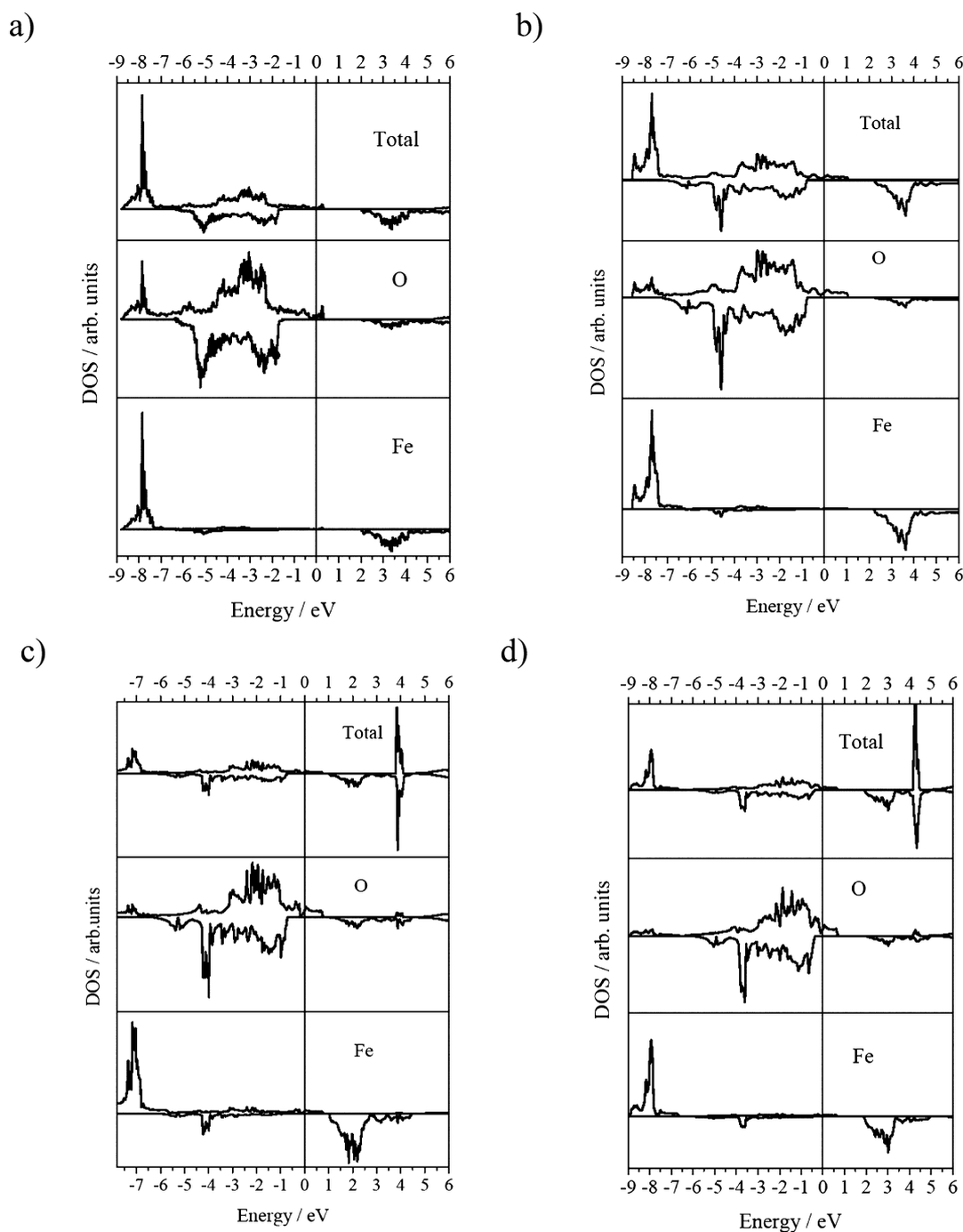


Figure 2. Total and projected density of states (DOS) for stoichiometric in oxygen ($\delta = 0$) LSF12.5 (a) and LSF50 (b), both for the hybrid PBE0 functional, PBE+ U_{eff} functional with $U_{\text{eff}} = 4.5$ eV (c) or $U_{\text{eff}} = 6.5$ eV (d), all in the FM state. The Fermi energy is taken as zero and indicated by the vertical solid line. The DOS was recalculated with the tetrahedron method in the PBE+ U_{eff} calculations. The DOS for spin-down electrons is represented with negative values.

the PBE0 values are more consistent with the experiments. (A closer comparison of our results and those reported by Ritzmann et al.²² using the PBE+ U_{eff} approach for LSF50 suggested very similar μ_{Fe} but larger atomic charges for Fe in our case (1.85 e vs 1.70 e) even though very similar values for U_{eff} (4.5 eV vs 4.3 eV) were applied in both studies. The difference could be attributed to the symmetry of LSF50: rhombohedral in our calculations and cubic in ref 22. As expected, $d_{\text{Fe-O}}$ is larger in LSF12.5 than in LSF50 which is also confirmed in our PBE0 calculations and previous PBE+ U_{eff} calculations.²²

We compare below the bulk properties calculated here for LSF with those for LF and use it as a reference material for the analysis of changes in oxidation state of Fe in defective and perfect LSF. The stoichiometric LF was calculated in the LCAO method with the PBE0 functional only. The antiferromagnetic structure and orthorhombic symmetry (SG 62) were assumed. The lattice constants for LF ($a = 5.58$, $b = 5.59$, $c = 7.90$ Å) are in a very good agreement with the experimental values.^{17,59} The magnetic moment μ_{Fe} is high ($4.28 \mu_{\text{B}}$) in this compound in agreement with the Fe d^5 high spin configuration and the oxidation state of Fe 3+. The moment reduction from

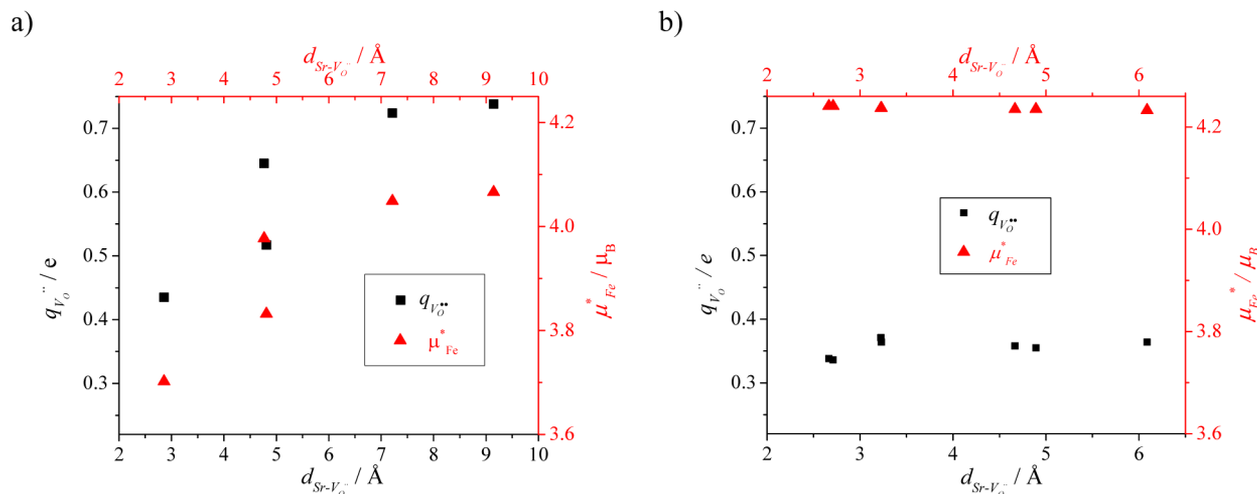


Figure 4. Mulliken charge of V_{O}^{2+} and magnetic moment of Fe as functions of $d_{\text{Sr}-V_{\text{O}}^{2+}}$ for LSF12.5: (a) $\delta = 0.125$ and (b) $\delta = 0.0625$.

supercells. However, these supercells contain different numbers of nonequivalent oxygen ions occupying different Wyckoff positions. Respectively, as the first step, we identified the Wyckoff positions for O ions in the supercells using the computer code FINDSYM.⁴⁸ Second, we calculated the formation enthalpies of all nonequivalent oxygen ions, in order to find those having smallest and largest formation enthalpies at finite temperature. Notice again that the LCAO calculations with the PBE0 functional have an advantage of using the *ghost* basis set in the vacancy position and allow one to estimate the Mulliken charge of V_{O}^{2+} ($q_{V_{\text{O}}^{2+}}$), the distance of Fe closest to V_{O}^{2+} ($d_{\text{Fe}-V_{\text{O}}^{2+}}$) and its magnetic moment μ_{Fe}^* .

The results of PBE0 and PBE+ U_{eff} calculations for V_{O}^{2+} in LSF12.5 and LSF50 (using both 40 atoms and 80 atom supercells) are presented in Table 4 for the two oxygen vacancies having smallest and largest formation energies, whereas Figure 3 shows the dependence of $q_{V_{\text{O}}^{2+}}$ and μ_{Fe}^* on $d_{\text{Fe}-V_{\text{O}}^{2+}}$ for all calculated vacancies in the 40 atoms supercells. Also, Figure 4 shows the dependence of $q_{V_{\text{O}}^{2+}}$ and μ_{Fe}^* on $d_{\text{Sr}-V_{\text{O}}^{2+}}$ for LSF12.5 with $\delta = 0.125$ and $\delta = 0.0625$.

We have observed that the distances $d_{\text{Fe}-V_{\text{O}}^{2+}}$ and $d_{\text{Sr}-V_{\text{O}}^{2+}}$ are correlated with $q_{V_{\text{O}}^{2+}}$ and μ_{Fe}^* in LSF12.5 with $\delta = 0.125$: both $q_{V_{\text{O}}^{2+}}$ and μ_{Fe}^* increase considerably with $d_{\text{Fe}-V_{\text{O}}^{2+}}$ (Figure 3a) and $d_{\text{Sr}-V_{\text{O}}^{2+}}$ (Figure 4a). Thus, the electron population (charge inside vacancy) $q_{V_{\text{O}}^{2+}}$ lies in the region between 0.43 and 0.74 e, as the $d_{\text{Fe}-V_{\text{O}}^{2+}}$ distance increases from 1.77 to 1.91 Å and $d_{\text{Sr}-V_{\text{O}}^{2+}}$ distance increases from 2.86 to 9.15 Å, respectively (Figures 3a and 4a). The value of μ_{Fe}^* also shows the correlation as discussed: it is smallest (3.70 μ_{B}) for the shortest $d_{\text{Fe}-V_{\text{O}}^{2+}}$ and largest (4.24 μ_{B}) for the longest $d_{\text{Fe}-V_{\text{O}}^{2+}}$ in LSF12.5. That is, there is a considerable dispersion (0.5 μ_{B}) of magnetic moments in different Fe locations and surroundings in this case of $\delta > x/2$, i.e., involving Fe^{2+} ions.

On the other hand, in the defective supercells of LSF50 with $\delta = 0.125$ and LSF12.5 with $\delta = 0.0625$ the values of parameters $q_{V_{\text{O}}^{2+}}$ and μ_{Fe}^* are less sensitive to the local position of V_{O}^{2+} within the supercell than in the above-discussed LSF12.5 with $\delta = 0.125$. It is reflected in the small changes of the properties in LSF50 with $\delta = 0.125$ shown in Figure 3b as functions of

$d_{\text{Fe}-V_{\text{O}}^{2+}}$. In both the hybrid PBE0 and PBE+ U_{eff} calculations for LSF50 with $\delta = 0.125$ and LSF12.5 $\delta = 0.0625$, the total magnetic moment of the supercells increases in the presence of V_{O}^{2+} . Also, its value is independent of the functional. The maximum/minimum values of $q_{V_{\text{O}}^{2+}}$ in LSF50 $\delta = 0.125$ and LSF12.5 with $\delta = 0.0625$ are 0.26/0.24 e (Figure 3b) and 0.34/0.37 e (Figure 4b), respectively, demonstrating more delocalized behavior of electrons left behind by a missing oxygen atom, in comparison with LSF12.5 with $\delta = 0.125$. Interestingly, both LSF50 with $\delta = 0.125$ and LSF12.5 with $\delta = 0.0625$ are characterized by the condition $\delta \leq x/2$.

The distance $d_{\text{Fe}-\text{O}}$ in defective LSF50 changes in comparison with stoichiometric material, i.e., in such supercells it is not only the Fe ions closest to V_{O}^{2+} that are affected by the presence of V_{O}^{2+} . For example, the shortest/longest distance $d_{\text{Fe}-\text{O}}$ in the case of V_{O}^{2+} having largest formation energy in LSF50 is 1.85/2.00 Å and the Fe magnetic moment is 3.82/4.35 μ_{B} . Here also the Sr ions play an important role. The closer the Sr ion the longer the distance with Fe, i.e., if $d_{\text{Fe}-\text{O}} = 1.85/2.00$ Å, then the Sr ion has the distance of 2.68/2.51 Å, respectively.

We present in Table 4 the V_{O}^{2+} formation enthalpy ΔH_{F} as given by eq 4 calculated at 1200 K (which is also the temperature closest to the operation temperature of the cathode of the solid oxide fuel cells) for the two functionals used. The calculated values of ΔH_{F} are compared with those obtained using the results of permeation measurements of the present study and thermogravimetric measurements from the literature. In order to make the comparison at realistic temperatures, the inclusion of vibrational contribution to ΔH_{F} is necessary. The vibrational contribution in the solid phase ΔH_{vib} was taken from our previous study on (La,Sr)- $\text{Co}_{0.25}\text{Fe}_{0.75}\text{O}_{3-\delta}$.²⁰ ΔH_{vib} at $T = 1200$ K found with the standard PBE functional in ref 20 for the content of Sr 50% is -0.31 eV for the oxygen nonstoichiometry $\delta = 0.125$. Interestingly, the same contribution for another perovskite (BaZrO_3) was obtained using the CRYSTAL code with the PBE0 functional. Notice that the calculated oxidation entropy in ref 20 agrees very well with the measured one.⁶⁵ Such calculations of phonon contributions require ground state structure; otherwise, the imaginary frequencies are present and the calculation of free energies is not possible. Because of very time-consuming calculations of phonon vibrations in large supercells, the

estimated value of -0.31 eV was also used for LSF12.5 with $\delta = 0.0625$. The overall contribution of the temperature effects, consisting of the vibrational contribution and kT -term in eqs 4 and 5, is approximately -0.06 eV at 1200 K which is slightly larger than suggested in ref 22 (-0.02 eV at 973.15 K).

According to the defect chemistry model,^{19,66} there are two different regimes which should be considered with respect to nonstoichiometry δ and doping x : $\delta \leq x/2$ and $\delta > x/2$. These regimes control the Fe oxidation state, the $V_{\text{O}}^{\bullet\bullet}$ formation energies, and, thus, the oxygen transport in LSF under different conditions. Thus, the formation enthalpy ΔH_{F} obtained in the thermogravimetric measurements corresponds to different oxidation states: Fe 2+/3+ (large formation enthalpy) and 3+/4+ (low formation enthalpy)^{19,66} as were obtained under different experimental conditions controlled by the oxygen partial pressure. The magnitude of ΔH_{F} corresponding to given oxidation state is nearly a constant, only weakly dependent on dopant concentration x , provided $\delta \leq x/2$. We present in Table 4 the experimental values of ΔH_{F} extrapolated from the permeation data discussed in Section 4.1 and for $\text{La}_{0.9}\text{Sr}_{0.1}\text{FeO}_{3-\delta}$ and $\text{La}_{0.4}\text{Sr}_{0.6}\text{FeO}_{3-\delta}$ compositions (close to our calculated compositions) taken from the from the thermogravimetry study^{19,66} for the above-mentioned two regimes. As mentioned above, the results of permeation measurements demonstrate strong decrease of ΔH_{F} with Sr doping (1.40 vs 0.33 eV) due to changes of the Fe oxidation state. Our calculations with $\delta = 0.125$ for LSF50 satisfy the criterion $\delta \leq x/2$ whereas $\delta = 0.0625$ for LSF12.5 lies at the border between the two Fe oxidation states (2+/3+ and 3+/4+, i.e., $\delta = x/2$).

The calculated values of ΔH_{F} given in Table 4 for two extreme vacancy configurations for LSF50 with $\delta = 0.125$ (the average oxidation state 3.25) as obtained in the PBE0 calculations lie in the range 0.04–0.27 eV (the average value 0.16 eV), in reasonable agreement with the permeation measurements ($\Delta H_{\text{F}} = 0.33$ eV) but smaller than in the thermogravimetry experiments (0.99 eV). On the other hand, the defect formation enthalpy $\Delta H_{\text{F}} = 0.54$ –1.09 eV (the average value 0.83 eV) for LSF12.5 with $\delta = 0.0625$ (average oxidation state is 3.0) in the PBE0 calculations reasonably agrees not only with the value of 1.40 eV from the permeation measurements, but also with 1.20 eV from the thermogravimetry measurements. Thus, our calculations reproduce the general trend of a moderate decrease of the $V_{\text{O}}^{\bullet\bullet}$ formation energy with the Sr concentration increase (and vice versa). In contrast, the opposite regime $\delta > x/2$ for LSF12.5 with $\delta = 0.125$ (average oxidation state is 2.875) is characterized by the average calculated value of $\Delta H_{\text{F}} = 2.07$ eV, which is more than by 1 eV larger in a comparison with the above-discussed $\delta = x/2$. This results from the presence of 12.5% of Fe^{2+} in this compound and high energy price for their formation from Fe^{3+} . Note that the calculated average value of 2.07 eV should be compared with the average of experimental energies of 1.2 and 5.0 eV (Table 4), i.e., 3.1 eV since a sequential release of the two electrons from the oxygen vacancy occurs in the two stages, with $0 \leq \delta \leq x/2$ and $x/2 < \delta$.

In Figure 5 we also present the variation in ΔH_{F} as a function of $d_{\text{Sr}-V_{\text{O}}^{\bullet\bullet}}$ for LSF12.5 with $\delta = 0.0625$. It is worth mentioning that the variation in ΔH_{F} is larger for LSF12.5 with $\delta = 0.0625$ (where $V_{\text{O}}^{\bullet\bullet}$ can be very close to or far from Sr) than LSF50 with $\delta = 0.125$ (where $V_{\text{O}}^{\bullet\bullet}$ will always be close to Sr), cf., the low variations in the formation energy of $V_{\text{O}}^{\bullet\bullet}$ in ref 21.

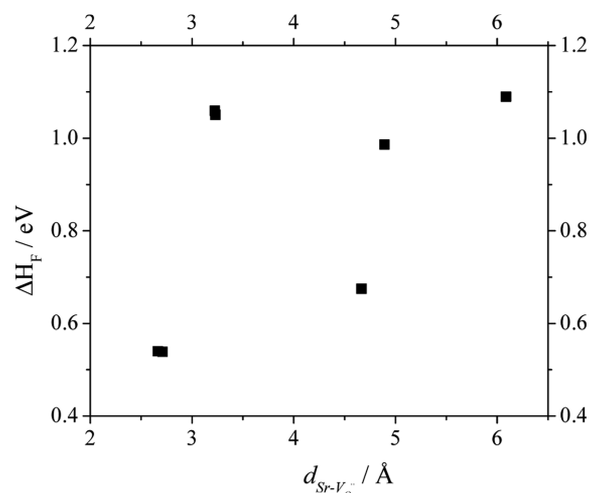


Figure 5. Formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ (ΔH_{F} at 1200 K) as a function of $d_{\text{Sr}-V_{\text{O}}^{\bullet\bullet}}$ for LSF12.5 with $\delta = 0.0625$. The chosen values of ΔH_{F} correspond to $q_{V_{\text{O}}^{\bullet\bullet}}$ —and μ_{Fe}^* -values in Figure 4b.

The analysis of Fe magnetic moments in Table 4 shows that in the oxidation state Fe 3+ (LSF12.5, $\delta = 0.0625$) they are closest to those in pure LF (4.28 μ_{B}), with the same oxidation state, whereas for other compounds the magnetic moments are reduced, due to increased fraction of Fe^{4+} or Fe^{2+} (both having 4 unpaired spins but different configurations: d^4 and d^6 , respectively).

Note that the absolute values of ΔH_{F} in the PBE+ U_{eff} calculations (Table 4) differ considerably from the results of permeation measurements, and the hybrid PBE0 calculations. This brings us the conclusion that the hybrid PBE0 functional provides us with the most accurate description of bulk LSF and $V_{\text{O}}^{\bullet\bullet}$ behavior therein.⁶⁷

5. CONCLUSIONS

The DFT calculations based on the hybrid PBE0 functional (with the standard exact exchange contribution of 25%) and the LCAO basis set were combined with the permeation measurements to analyze the formation enthalpy of oxygen vacancies in complex perovskites like $(\text{La,Sr})\text{FeO}_{3-\delta}$ with different Sr doping. The use of hybrid functionals allows us to reproduce the bulk properties of these solid solutions and their behavior as semiconductors. The finite band gap is important to calculate properly the formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ in LSF which is in agreement with previous experimental studies on the formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ in LSF. Moreover, the LCAO basis set allowed us to determine the electronic densities within the vacancy (due to additional so-called ghost basis set at the vacancy site) and find the correlation between the charge of $V_{\text{O}}^{\bullet\bullet}$, the $\text{Fe}-V_{\text{O}}^{\bullet\bullet}$ distance, and the magnetic moment on Fe. It is demonstrated that a careful comparison of the calculation results with the experiments requires the knowledge of the oxidation state of Fe under experimental conditions and thus the relation between nonstoichiometry δ and doping x . We have confirmed that the formation enthalpy of $V_{\text{O}}^{\bullet\bullet}$ significantly decreases with the Sr doping, which greatly accelerates oxygen transport through permeation membranes and SOFC cathodes.

■ ASSOCIATED CONTENT

● Supporting Information

Exponents (bohr⁻²) and contraction coefficients. 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 interest.

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standard DFT calculation. ΔH_{F} in this case is reduced significantly (in comparison with data in Table 4), varying in the range from 0.38 to 1.07 eV at $U_{\text{eff}} = 4.5$ eV.