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**Molecular dynamics simulations of oxygen diffusion in GdBaCo$_2$O$_{5.5}$**

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The mechanisms of oxygen diffusion in GdBaCo$_2$O$_{5.5}$ compound are investigated by molecular dynamics simulations. The results confirm that diffusion is mainly bidimensional with oxygen moving in the (a,b) plane while diffusion along the c axis is much more difficult. Between 1000 and 1600 K, the activation energy for diffusion is about 0.6 eV, close to experimental values. Going deeper inside the oxygen diffusion mechanism, we see that this diffusion occurs mainly in the cobalt planes while most of the oxygen vacancies are kept in the Gd planes. Analysis of oxygen motions show that Gd planes can be seen as source-sink for the oxygen vacancies rather than as fast pathways. © 2010 American Institute of Physics, doi:10.1063/1.3504250

The main issue in solid oxide fuel cells (SOFC) technology is to lower the operating temperature$^1$ in the range 500–750 °C. Due its high activation energy process, the cathode becomes a major source of losses at low temperature.$^2$ Recent experiments show the suitability of cathode SOFCs.$^3, 4$ To understand the particularities of this promising material, we study here the oxygen diffusion in GdBaCo$_2$O$_{5.5}$ using molecular dynamics (MD) simulations. GdBaCo$_2$O$_{5.5}$+x is a double perovskite compound with Gd and Ba cations alternating on the A site along the c axis, leading to a tetragonal symmetry at high temperature. It generally contains oxygen vacancies and mixed valency on cobalt site. The diffusion of oxygen species is far from being understood in these compounds even if the presence of oxygen vacancies even if the presence of oxygen vacancies in the Gd plane, as found from neutron diffraction,$^5$ suggested preferential diffusion in this atomic plane. The system was first equilibrated at 1000, 1200, 1400, and 1600 K and at zero pressure for 10 000 time steps (with a time step of 2 fs) in the isothermal-isobaric (NPT) ensemble with Nosé–Hoover thermostat. Then 100 000 time steps allowed to find the equilibrium lattice parameters. For the set of potentials no. 1, the tetragonal high temperature structure was maintained with cell parameters close to experimental values, for the set no. 2, a dramatic increase in the c cell parameter was observed leading to a collapse of the structure. In this case, we thus used a

$$\varphi_{ij}(r) = A_{ij} \exp\left(-\frac{r}{\rho_{ij}}\right) - \frac{C_{ij}}{r^6},$$  

where $r$ is the distance between the atoms $i$ and $j$, and $A_{ij}$, $\rho_{ij}$, and $C_{ij}$ are potential parameters specific to each ion pair whose values are given in Table I. Two sets of potentials were used which were previously validated in MD simulations of cobaltite compounds. For the set no. 1, we used the potentials from Fisher et al.$^9$ for Ba–O, Co–O, O–O interactions, and we determined the parameters for gadolinium-oxygen pair by fitting structural data of Gd$_2$O$_3$. For the second set, we directly used the interatomic potentials from Rupasov et al.$^{10}$ Busker et al.$^{11}$ and Minervini et al.$^{12}$ The potentials were tested by performing energy minimizations with the GULP software$^{13}$ at a temperature of 0 K on a 4 $\times$ 4 $\times$ 6 supercell with vacancies located in the Gd plane. As results, the calculated cell $a=b$ and $c$ parameters expressed for a pseudotetragonal double perovskite cell, were 3.76(1) Å and 7.58(1) Å for the potentials set no. 1, and 3.87(3) Å and 7.71(6) Å for potentials set no. 2. These values are close to those extrapolated to 0 K from high temperature measurements (a=b=3.85337 Å and c=7.5318 Å).

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<table>
<thead>
<tr>
<th>Interaction</th>
<th>$A$ (eV)</th>
<th>$\rho$ (Å)</th>
<th>$C$ (eV Å$^{-6}$)</th>
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</thead>
<tbody>
<tr>
<td>Ba$^{2+}$...O$^{2-}$</td>
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<td></td>
<td></td>
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<tr>
<td>Set 1$^a$</td>
<td>1214.4</td>
<td>0.3537</td>
<td>0.0</td>
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<tr>
<td>Set 2$^b$</td>
<td>905.70</td>
<td>0.3976</td>
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<td>Gd$^{3+}$...O$^{2-}$</td>
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<tr>
<td>Set 1$^c$</td>
<td>1458.38</td>
<td>0.3522</td>
<td>0.0</td>
</tr>
<tr>
<td>Set 2$^d$</td>
<td>1885.75</td>
<td>0.3399</td>
<td>20.34</td>
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<tr>
<td>Co$^{3+}$...O$^{2-}$</td>
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<tr>
<td>Set 1$^e$</td>
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<tr>
<td>Set 2$^f$</td>
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<tr>
<td>O$^{2-}$...O$^{2-}$</td>
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<tr>
<td>Set 1$^g$</td>
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<td>0.149</td>
<td>43.0</td>
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<tr>
<td>Set 2$^h$</td>
<td>9547.96</td>
<td>0.2192</td>
<td>32.0</td>
</tr>
</tbody>
</table>

$^a$Fischer et al.$^8$

$^b$Busker et al.$^{10}$

$^c$This study.

$^d$Minervini, Zacate, and Grimes.$^{11}$

$^e$Rupasov et al.$^9$

$^f$Author to whom correspondence should be addressed. Electronic mail: guilhem.dezanneau@ecp.fr.
DL_POLY isotropic option during NPT simulations for which, the $a=b$ and $c$ parameters are allowed to vary in a non-independent way. Then, 100 000 time steps were performed in the canonical ensemble (NVT) with Nosé–Hoover thermostat, keeping the lattice constants at their previously determined equilibrated values, and finally the diffusion properties were studied in the microcanonical ensemble (NVE), for $2 \times 10^6$ time steps (up to $5 \times 10^6$ time steps). The calculated thermal expansion was 15.2 $\times 10^{-6}$ K$^{-1}$ for set no. 1, while it was only $8.8 \times 10^{-6}$ K$^{-1}$ for set no. 2. The first value is very close to the one experimentally found by Tarancon et al. of 16.4 $\times 10^{-6}$ K$^{-1}$ for GdBaCo$_2$O$_{5+x}$. This result would indicate that set no. 2 is probably not fully adapted for such simulations.

The oxygen diffusivity was determined from the evolution of the mean square displacement as a function of time at different temperatures using the Einstein’s law. The values of $D$ are represented in the Arrhenius plot of Fig. 1. We also plot the oxygen tracer diffusion coefficient as determined from Tarancon et al.$^{12}$ These two coefficients could be directly compared, neglecting the evolution of oxygen composition with temperature. We also present in Fig. 1 the chemical diffusion coefficients as determined from relaxation measurements to which was applied a correction. Indeed, the chemical diffusion coefficient was divided by the thermodynamic factor set to 75, corresponding to the value obtained from the oxygen content in GBCO in the range 950–1050 °C.$^{15}$ From our simulations, the oxygen transport can be described by an Arrhenius law with an activation energy of 0.67 eV and 0.56 eV, respectively, for sets 1 and 2. These values reasonably compare with the value of 0.60 eV found from isotope exchange.$^{14}$ They are also close to the values of oxygen chemical diffusion activation energies of 0.66 eV and 0.77 eV found, respectively, by Taskin et al.$^{16}$ and Choi et al.$^{17}$ from mass and conductivity relaxation measurements. Nevertheless, all the diffusion coefficients presented in Fig. 1 spread over several orders of magnitude for a same temperature, underlining the difficulty to determine experimentally the oxygen diffusion coefficient.

MD simulations also allow a deeper understanding of diffusion mechanism. Figure 2 compares the mean square displacement of oxygen along the $c$-axis and in the $a$-$b$ plane at 1400 K. Oxygen atoms mostly diffuse in the $a$-$b$ plane with a diffusivity of $1.1 \times 10^{-7}$ cm$^2$ s$^{-1}$. This diffusivity is almost ten times higher than the one found along the $c$-axis of $1.3 \times 10^{-8}$ cm$^2$ s$^{-1}$. This anisotropic diffusion can be explained by the absence of oxygen vacancies in the barium planes, making the efficient diffusion of oxygen vacancies across Ba planes difficult.

Barium planes logically do not contain oxygen vacancies. As foreseen from previous neutron diffraction studies, vacancies are mainly located in the gadolinium planes ($\sim$95%), but significant amount of them is also in the cobalt planes ($\sim$5%). When studying the individual displacements of oxygen atoms, we observe that they spend most of their time in the cobalt planes and occasionally hop into gadolinium planes (where they stay only for a few picoseconds while they can remain up to some nanoseconds in cobalt planes). Figure 3 presents the oxygen density (a) in (001) cobalt planes and (b) in (100) cobalt planes. Figure 3(b) confirms that oxygen atoms in barium planes stay at the same position and do not diffuse in the neighboring cobalt planes. Besides, a count of the oxygen hops shows that oxygen atoms make roughly three times more jumps in the cobalt planes than going back and forth or cross a gadolinium plane. This would mean that diffusion mainly occurs in the Co plane and that vacancies in the Gd planes are mainly trapped. Their presence is nevertheless necessary to make one oxygen escaping from time to time from the Co plane.
leaving a highly diffusing oxygen vacancy in the Co plane. The first intuition that vacancies in Gd planes lead to an enhanced diffusion is then probably not fully correct. Actually, the diffusion in perovskite compounds involves nearest-neighbor jumps between oxygen positions and can thus not be explained by the only consideration of vacancies in Gd planes. From these simulations, we indeed show that Gd planes serve as source-sink for oxygen vacancy rather than as real pathways.

In conclusion, MD simulations predict that oxygen transport in GdBaCo$_2$O$_{5.5}$ is strongly anisotropic with an activation energy of around 0.6 eV in the temperature range of 1000–1600 K. Oxygen atoms mainly diffuse along the cobalt planes with short hops into gadolinium planes and very seldom cross a barium plane. Meanwhile, most of oxygen vacancies are kept into gadolinium planes while some of them are moving along the cobalt planes to allow oxygen diffusion.

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8PL POLY code: www.ccp5.ac.uk/DL POLY/.