Atomistic Simulations of the Defect Chemistry and Self-Diffusion of Li-ion in LiAlO$_2$

N. Kuganathan $^{1,2,*}$, J. Dark $^2$, E.N. Sgourou $^{3,4}$, Y. Panayiotatos $^4$ and A. Chroneos $^{1,2}$

$^1$ Department of Materials, Imperial College London, London W7 2AZ, UK  
$^2$ Faculty of Engineering, Environment and Computing, Coventry University, Priory Street, Coventry CV1 5FB, UK  
$^3$ Solid State Physics Section, University of Athens, Panepistimiopolis Zografos, 157 84 Athens, Greece  
$^4$ Department of Mechanical Engineering, University of West Attica, 12210 Athens, Greece  
* Correspondence: n.kuganathan@imperial.ac.uk or ad0636@coventry.ac.uk

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Abstract: Lithium aluminate, LiAlO$_2$, is a material that is presently being considered as a tritium breeder material in fusion reactors and coating material in Li-conducting electrodes. Here, we employ atomistic simulation techniques to show that the lowest energy intrinsic defect process is the cation anti-site defect (1.10 eV per defect). This was followed closely by the lithium Frenkel defect (1.44 eV per defect), which ensures a high lithium content in the material and inclination for lithium diffusion from formation of vacancies. Li self-diffusion is three dimensional and exhibits a curved pathway with a migration barrier of 0.53 eV. We considered a variety of dopants with charges $^+$1 (Na, K and Rb), $^{+2}$ (Mg, Ca, Sr and Ba), $^{+3}$ (Ga, Fe, Co, Ni, Mn, Sc, Y and La) and $^{+4}$ (Si, Ge, Ti, Zr and Ce) on the Al site. Dopants Mg$^{2+}$ and Ge$^{4+}$ can facilitate the formation of Li interstitials and Li vacancies, respectively. Trivalent dopants Fe$^{3+}$, Ni$^{3+}$ and Mn$^{3+}$ prefer to occupy the Al site with exoergic solution energies meaning that they are candidate dopants for the synthesis of Li (Al, M) O$_2$ (M = Fe, Ni and Mn) compounds.

Keywords: LiAlO$_2$; defects; Li diffusion; dopants; atomistic simulation

1. Introduction

The ever-increasing demand for energy generated, initiated further research into alternate methods of renewable energy production and storage methodologies for the power production. In that respect, Li-based ceramics are technologically important and considered by the community for applications ranging from nuclear fusion to batteries [1–8]. LiAlO$_2$, a lithium based ceramic material, has been considered with the prospect of being used as a tritium breeder material in magnetic confined nuclear fusion reactors after promising initial research regarding the physical and thermodynamic properties of the material as well as the potential tritium breeding ratio (TBR) [8,9]. In addition, this material has great potential as a coating material in Li-conducting electrodes [10].

Previous studies have highlighted the importance of atomic scale calculation to gain insights on the defect processes (intrinsic defect processes, doping and diffusion) of oxides for energy applications [11–15]. Previous experimental and theoretical studies determined a wide range of activation energies for Li self-diffusion in $\gamma$-LiAlO$_2$ (0.50–1.26 eV) [16–22]. Additionally, there are no systematic studies of the intrinsic defect processes and doping in this material. In the present study, we have employed atomistic simulations to investigate the structure, intrinsic defect processes, Li self-diffusion and the introduction of dopants.
2. Computational Methods

The present computational study was based on classical pair-wise potential calculations to describe LiAlO₂ via the General Utility Lattice Program (GULP) code [23,24]. In the present approach, the total energy (lattice energy) is determined by long range (i.e., Coulombic) and short range [i.e., electron-electron repulsive and attractive intermolecular forces (van der Waals forces)]. The latter were modelled using Buckingham potentials (refer to the supplementary information). The van der Waals forces arising from the spontaneous formation of instantaneous dipoles are very important as the formation energy results are sensitive to those forces. The present modelling approach takes into the account of Van der Waals forces as a function of the interatomic distance (r) [23,24]. Two-body Buckingham potential mentioned in the supplementary information consists of two parts. The first part of the equation represents the Pauli repulsion (electron-electron) and the second part denotes the van der Waals interaction. Ionic positions and lattice parameters were relaxed using the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm [24]. Convergence criteria dictated that in relaxed configurations, forces on each atom was <0.001 eV/Å. To introduce point defects in the lattice we used the Mott-Littleton methodology [25] similarly to recent work [26–28]. It is established that although the present methodology may overestimate the defect formation enthalpies at the dilute limit, the trends will be unchanged [29]. In a thermodynamic perspective defect parameters can be described by comparing the real (i.e., defective) crystal to an isobaric or isochoric ideal (i.e., non-defective) crystal. Defect formation parameters can be interconnected through thermodynamic relations [30,31], with the present atomistic simulations corresponding to the isobaric parameters for the migration and formation processes [32–34].

3. Results

3.1. Crystal Structure, Intrinsic Defect Processes and Li Diffusion

The LiAlO₂ crystal structure considered in the present study is the tetragonal γ-phase (Figure 1) with lattice parameters a = b = 5.1687 Å, c = 6.2679 Å, and α = β = γ = 90° [35]. This crystal structure consists of LiO₄ and AlO₄ tetrahedra. They are inter-connected via edge and corner sharing to form three dimensional channels. The calculated structural parameters are given in Table 1 and it can be seen that there is excellent agreement with experiment [35] validating the potentials (refer to Table S1 in the Supplementary Materials) used in this study.

![Figure 1. Crystal structure of LiAlO₂ (space group P4₁2₁2).](image)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Exp</th>
<th>Calc</th>
<th>Δ(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a = b (Å)</td>
<td>5.1687</td>
<td>5.1403</td>
<td>0.55</td>
</tr>
<tr>
<td>c (Å)</td>
<td>6.2679</td>
<td>6.3860</td>
<td>1.88</td>
</tr>
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<td>α = β = γ (°)</td>
<td>90.00</td>
<td>90.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Volume [V = a × b × c (Å³)]</td>
<td>167.4498</td>
<td>168.7370</td>
<td>0.77</td>
</tr>
</tbody>
</table>

Table 1. Calculated structural parameters for LiAlO₂ (space group P4₁2₁2) as compared to the experimental determinations [35].
Next, we considered the process of calculating the defect energy required for the key intrinsic defect processes such as Frenkel, Schottky and anti-site in LiAlO$_2$. Frenkel and Schottky defect formation energies were calculated by combining individual point defects. Frenkel energy is the sum of vacancy and interstitial energies. Schottky energy was calculated by adding appropriate vacancy energies and subtracting cohesive energy. Relative energetics of these defect processes are useful in predicting the electrochemical behaviour of LiAlO$_2$. The reaction energies for the intrinsic defect processes can be represented by using the established Kröger-Vink notation [36].

$$\text{Li Frenkel} : \text{Li}^X_{\text{Li}} \rightarrow V'_{\text{Li}} + \text{Li}^*$$  \hspace{1cm} (1)

$$\text{Al Frenkel} : \text{Al}^X_{\text{Al}} \rightarrow V''_{\text{Al}} + \text{Al}^{**}$$  \hspace{1cm} (2)

$$\text{O Frenkel} : \text{O}^X_{\text{O}} \rightarrow V''_{\text{O}} + \text{O}^*$$  \hspace{1cm} (3)

$$\text{Schottky} : \text{Li}^X_{\text{Li}} + \text{Al}^X_{\text{Al}} + 2\text{O}^X_{\text{O}} \rightarrow V'_{\text{Li}} + V''_{\text{Al}} + 2V''_{\text{O}} + \text{LiAlO}_2$$  \hspace{1cm} (4)

$$\text{Li}_2\text{O Schottky} : 2\text{Li}^X_{\text{Li}} + \text{O}^X_{\text{O}} \rightarrow 2V'_{\text{Li}} + V''_{\text{O}} + \text{Li}_2\text{O}$$  \hspace{1cm} (5)

$$\text{Li}/\text{Al antisite (isolated)} : \text{Li}^X_{\text{Li}} + \text{Al}^X_{\text{Al}} \rightarrow \text{Li}''_{\text{Li}} + \text{Al}^{**}_{\text{Al}}$$  \hspace{1cm} (6)

$$\text{Li}/\text{Al antisite (cluster)} : \text{Li}^X_{\text{Li}} + \text{Al}^X_{\text{Al}} \rightarrow \{\text{Li}''_{\text{Li}} : \text{Al}^{**}_{\text{Al}}\}^X$$  \hspace{1cm} (7)

The reaction energies are reported in Figure 2. The lowest intrinsic defect energy process was calculated to be the cation mixing (anti-site) in which Li and Al exchange their atomic positions. This defect was noted in various oxide materials experimentally and theoretically [37–47]. The primary reasons for this defect include experimental conditions for the preparation of as-prepared compounds and cycling of as-prepared materials particularly in battery applications. Using the Pechini sol–gel process, Dominko et al. [48] synthesised a pure orthorhombic phase of Li$_2$MnSiO$_4$. Later, Politaev et al. [39] observed a small amount of cation exchange (Li-Mn) defect in their experimental preparation (solid-state reactions in hydrogen at 950–1150 $^\circ$C) of monoclinic phase of Li$_2$MnSiO$_4$. Nyten et al. [38] observed a shift in the potential plateau between first and second cycles in Li$_2$FeSiO$_4$ and suggested that this is due to the structural rearrangement associated with the interchange of some of the Li and Fe ions. The second lowest reaction energy is calculated for the Frenkel defect and this process is higher only by 0.34 eV than the anti-site defect. This would signify the concentration of lithium vacancies and interstitials, inferring this material is suitable for use as a tritium breeder. Other defects exhibit high reaction energies meaning that they are not significant in this material.

Ion diffusion is a key issue in many energy related materials, including cathode materials for batteries. The current simulation approach has the ability to calculate Li-ion diffusion pathways together with activation energies. As experimental determination of Li-ion diffusion is quite challenging, simulation results would be useful to assist in the investigation of experimental data. The results reveal that there is a single Li local hop with the jump distance of 3.09 Å. The activation energy for this hop is calculated to be 0.53 eV. Long range diffusion pathway was constructed by connecting local Li hops. Figure 3a shows the three dimensional long range diffusion pathway with a non-linear pattern. This shows that Li-ion conduction in this material is high. In a previous theoretical study based on the molecular dynamics by Jacobs et al. [18], it is shown that Li-ion diffusion is three dimensional though the pathway is not clear enough from their calculated snapshots derived at different temperatures. Density functional theory simulations were also employed to calculate the Li-ion diffusivity in γ-LiAlO$_2$ [21,22]. In those calculations, Li hopping mechanisms and activation energies for different Li-Li separations are reported. However, the information regarding the Li migration pathway (linear or non-linear) is unclear. Our simulation clearly presents the diffusion pathway that can be directly compared in experiments.
The activation energy calculated here (0.53 eV) is within the range of previous theoretical studies (0.5–0.63 eV) [18,21,22] and lower than previous experimental (NMR spectroscopy and conductivity measurements) work by Idris et al. (0.70 eV) [17]. More direct experimental work (e.g., secondary ion mass spectrometry, SIMS) would be required to experimentally pinpoint the activation energy of Li self-diffusion. As was previously demonstrated, simple atomistic models can yield reliable activation energies of diffusion as compared to experiment with mechanisms compatible with DFT [49].

The calculated activation energy exceeds 0.5 eV and it is therefore anticipated that LiAlO$_2$ will only be a mediocre ion conductor. In Table 2 we compare the activation energies of migration of recently investigated Li, Na, Mg ion conducting oxides considered for battery applications. There is a good agreement between our calculations and experiments for some oxide materials. As mentioned earlier, experimental investigation of activation energy is often difficult. Current simulations, in addition to giving activation energy information, have the added bonus of elucidating diffusion pathways that can be useful in the interpretation of experimental data.
This section details the processes of monovalent, divalent, trivalent and tetravalent doping into the lithium aluminate lattice. Monovalent dopants (R = Na, K and Rb) and divalent dopants (M = Mg, Ca, Sr and Ba) were considered on the Al site in order to increase the Li content in the form of Li interstitials in LiAlO$_2$ according to the following reaction equations:

$$2Li_2O + R_2O + 2Al^{1+}_{Al} \rightarrow 2R^{2+}_{Al} + 4Li^{+} + Al_2O_3 (8)$$

$$Li_2O + 2MO + 2Al^{1+}_{Al} \rightarrow 2M^{2+}_{Al} + 2Li^{+} + Al_2O_3 (9)$$

This efficient engineering strategy can increase the lithium density in lithium-based ceramics and tune mechanical, electrical and optical properties of materials such as lithium aluminate to further increase the applicability for use as a tritium breeding material. Solution enthalpies calculated for monovalent dopants are highly endothermic (>4.5 eV) suggesting that the doping process is quite difficult (refer to Figure 4a). Solution enthalpy gradually increases with ionic radius. High solution enthalpies are due to the larger radii of cations than that of Al$^{3+}$ (0.53 Å). In the case of divalent cations, Mg$^{2+}$ is promising as the solution enthalpy of MgO is −5.66 eV (refer to Figure 4b). The highly exothermic solution enthalpy is due to the ionic radius of Mg$^{2+}$ (0.57 Å) which is closer to the ionic radius of Al$^{3+}$. The other three dopants exhibit endoergic solution enthalpies, as their ionic radii significantly deviate from the ionic radius of Al$^{3+}$.

![Figure 4](image)

**Figure 4.** Solution enthalpies of (a) R$_2$O (R = Na, K and Rb) with respect to the R$^+$ ionic radius and (b) MO (M = Mg, Ca, Sr and Ba) with respect to the M$^{2+}$ ionic radius in LiAlO$_2$.

Next, we considered trivalent dopants (X = Ga, Fe, Co, Ni, Mn, Sc, Y and La) on the Al site according to the following reaction equation:

$$X_2O_3 + 2Al^{1+}_{Al} \rightarrow 2X^{3+}_{Al} + Al_2O_3 (10)$$
Figure 5 shows the solution enthalpies of $X_2O_3$. The results present nickel as the favourable isovalent dopant on the Al site, having the lowest solution enthalpy of $-2.03 \text{ eV}$ to integrate into the lattice. Solution enthalpies for both Fe$^{3+}$ and Mn$^{3+}$ are also negative implying that these two dopants are also worth examining experimentally. The favourable nature of potential candidate dopants (Fe$^{3+}$, Ni$^{3+}$ and Mn$^{3+}$) can be due to their ionic radii which are closer to the ionic radius of Al$^{3+}$ (0.53 Å). However, the dopant Co$^{3+}$ exhibits a positive solution enthalpy though its ion radius is closer to that of Al$^{3+}$ indicating other factors should be responsible. The dopants Ga$^{3+}$ and Sc$^{3+}$ exhibit almost zero solution enthalpies. Other dopants are not favourable as their solution enthalpies are endoergic. From Mn$^{3+}$ to La$^{3+}$ solution enthalpy gradually increases with ionic radius.

Finally, tetravalent dopants (Y = Si, Ge, Ti, Zr and Ce) were considered on the Al site. This doping process will introduce additional Li vacancies in LiAlO$_2$ assisting vacancy mediated self-diffusion according to the following equation:

$$2YO_2 + 2Al_{\text{Al}}^X + 2Li_{\text{Li}}^X \rightarrow 2Y_{\text{Al}}^{X^*} + 2V_{\text{Li}}^{\prime} + Al_2O_3 + Li_2O \quad (11)$$

We report the solution energies in Figure 6. The results present germanium as the favourable tetravalent dopant on the aluminium site, having the lowest solution enthalpy of 1.08 eV to integrate into the lattice. The second most stable dopant is Zr with the solution enthalpy of 1.54 eV. The ionic radii of these two dopants are closer to the ionic radius of Al$^{3+}$. However, the high positive solution enthalpy of Ti ($\sim 14 \text{ eV}$) is unclear though its ionic radius is somewhat closer to that of Al$^{3+}$. Furthermore, this dopant cannot be doped under any condition. Both Si and Ce exhibit high solution enthalpies ($\sim 4 \text{ eV}$) due to the mismatching of their ionic radii with that of Al$^{3+}$.
Finally, tetravalent dopants (Y = Si, Ge, Ti, Zr and Ce) were considered on the Al site. This doping process will introduce additional Li vacancies in LiAlO$_2$ assisting vacancy mediated self-diffusion according to the following equation:

\[
2 \text{YO}^{2+} + 2 \text{Al}^{4+} + 2 \text{Li}^{1+} \rightarrow 2 \text{Y}^{4+} + 2 \text{Li}^{1+} + \text{Al}^{3+} + \text{Li}^{1+}
\]  

We report the solution energies in Figure 6. The results present germanium as the favourable tetravalent dopant on the aluminium site, having the lowest solution enthalpy of 1.08 eV to integrate into the lattice. The second most stable dopant is Zr with the solution enthalpy of 1.54 eV. The ionic radii of these two dopants are closer to the ionic radius of Al$^{3+}$. However, the high positive solution enthalpy of Ti (~14 eV) is unclear though its ionic radius is somewhat closer to that of Al$^{3+}$. Furthermore, this dopant cannot be doped under any condition. Both Si and Ce exhibit high solution enthalpies (~4 eV) due to the mismatching of their ionic radii with that of Al$^{3+}$.

**Figure 6.** Solution enthalpies of YO$_2$ (Y = Si, Ti, Ge, Zr and Ce) with respect to the Y$^{4+}$ ionic radius in LiAlO$_2$.

### 4. Conclusions

In this computational modelling investigation, we have employed static atomistic simulation techniques to study the defect processes (intrinsic disorder mechanisms and doping) and Li self-diffusion in LiAlO$_2$. The lowest energy intrinsic defect process is the Li-Al anti-site implying that a small concentration of Li on the Al site and Al on the Li site will be present. The long range Li-ion diffusion is three-dimensional with an activation energy of migration of 0.53 eV. Numerous dopants were considered and it is calculated that the solution of GeO$_2$ will increase the concentration of the Li vacancies that are required to act as vehicles of diffusion. Furthermore, Mg$^{2+}$ on the Al site would increase the concentration of Li interstitials that are necessary for the enhancement in the capacity of NaAlO$_2$. It is anticipated that the present atomistic simulation work will encourage the further experimental investigation of LiAlO$_2$.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/1996-1073/12/15/2895/s1, Table S1: Interatomic potential parameters used in the atomistic simulations of LiAlO$_2$.

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