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Molecular Dynamics Modelling of Radiation Damage in Normal, Partly Inverse and Inverse Spinels

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Abstract

The radiation response of perfect crystals of MgAl$_2$O$_4$, partially inverted MgGa$_2$O$_4$ and fully inverse MgIn$_2$O$_4$ were investigated using molecular dynamics. Dynamical cascades were initiated in these spinels over a range of trajectories with energies of 400 eV and 2 keV for the primary knock-on event. Collision cascades were set up on each of the cation and anion sublattices and were monitored up to 10 ps. Simulations in the normal MgAl$_2$O$_4$ spinel for the 2 keV energy regime resulted in similar defect structures as obtained at the post-threshold 400 eV energies, with little clustering occurring. The predominant defect configurations were split interstitials and cation antisites. For the inverse spinels, a much wider variety of lattice imperfections was observed. More defects were also produced due to the formation of interstitial-vacancy cation chains and oxygen crowdions.

Key words: collisions cascades, spinel, defect formation

PACS: 61.72.Ji, 61.80.-x, 61.82.Ms

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Magnesium aluminate MgAl$_2$O$_4$ spinel is a ceramic oxide that has demonstrated strong radiation resistance to amorphisation and to the formation of defect clusters and dislocation loops [1] [2]. Recent irradiation experiments conducted by Matsumura et al. revealed disorder occurring over the cation sublattices, but the crystal structure remained stable in the damaged area [3] after 1 MeV Ne$^+$ irradiation to a dose of 4.5x10$^{20}$ ions/m$^2$.

The term spinel generally refers to the class of compounds with the molecular formula AB$_2$O$_4$, where A and B are generally either divalent and trivalent respectively or tetravalent and divalent [4]. Within the cubic closed-packed array of anions, the structure holds two types of interstices: octahedral and tetrahedral. Eight formula units make up a unit cell of spinel, containing a total of 64 tetrahedral interstices and 32 octahedral interstices between the anions [5]. In the perfect normal spinel, MgAl$_2$O$_4$, all Mg$^{2+}$ ions occupy one eighth of the tetrahedral positions, and the Al$^{3+}$ ions occupy half of octahedral interstices with six oxygen nearest neighbours, as shown in Fig.1. The remaining interstices are able to accommodate defect species.

Cationic interchange is one of the factors that explains the radiation tolerance of spinels. This occurs when one divalent cation (Mg$^{2+}$) initially at a tetrahedral site replaces a trivalent cation (Al$^{3+}$) at an octahedral site and vice-versa, resulting in the formation of a pair of antisite defects. Cation antisites are known to have the lowest formation energy as compared to other types of defects in spinels because the degree of local distortion of the crystalline structure resulting from their formation is minimal [6] [7]. The extent of cation disordering is quantified by the inversion parameter $i$, which is the
fraction of tetrahedral sites occupied by trivalent cations; \( i \) ranges from 0 for the normal spinel to 1 for a fully inverse spinel, and relates to the formula \([\text{Mg}_{1-i}\text{Al}_i](\text{Mg}_{2-i}\text{Al}_i\text{O}_4)\), where square brackets indicate the tetrahedral cation sites and curved brackets the octahedral sites.

In this work, we use molecular dynamics (MD) to simulate the damage response of three spinel compounds, namely normal magnesium aluminate, \( \text{MgAl}_2\text{O}_4 \), half inverse magnesium gallate \( \text{MgGa}_2\text{O}_4 \) \((i = 0.5)\) and the fully inverse magnesium indate \( \text{MgIn}_2\text{O}_4 \) \((i = 1)\). In reality the inversion parameter of \( \text{MgAl}_2\text{O}_4 \) is only close to zero at low temperatures and for natural spinel samples that have been able to equilibrate over geological time scales [8] [9]. Synthetic materials exhibit a broad range of inversion parameters from \( \approx 0.2 - 0.4 \) [10].

We compare the structural changes in these three compounds and investigate the most common types of defects for energies of 400 eV and 2 keV. The simulation is set up by imparting a certain amount of kinetic energy to an atom or ion at the centre of the simulation cell. This atom, known as the primary knock-on atom (PKA), displaces other target atoms along its trajectory and creates a disturbed region consisting of point defects. Details such as the number and distribution of the residual defects together with the mechanisms for defect recombination or aggregation can be extracted from the simulations.

2 Methodology

2.1 Potential

The ionic spinel crystal was modelled as an ordered array of point masses with formal charges. Electronic contributions were not considered explicitly. The interactions were calculated by a pairwise additive potential energy function comprising of the standard Buckingham potential [11] for short range interactions and the usual Coulomb potential for the long range electrostatic field. In these dynamical cascades, especially in the high energy regimes, the repulsive behaviour of colliding ionic cores at very small separation is taken into account by joining the potential profile to the Ziegler-Biersack-Littmark (ZBL) potential [12]. This is achieved by using a splining function \( g(r_{ij}) \), where \( r_{ij} \) denotes the pair separation between ions \( i \) and \( j \), to ensure continuity of the first and second derivatives for these functions. We have also implemented a smooth cutoff of the Buckingham potential at an interatomic distance greater than \( r_2 \). The potential energy function, \( \phi(r_{ij}) \) and the splining function are given by Eq. (1) and Eq. (2) respectively, where \( f_i \) are fitted constants.
Pair interactions (i and j)

<table>
<thead>
<tr>
<th></th>
<th>$A_{ij}$ (eV)</th>
<th>$\rho_{ij}$ (Å)</th>
<th>$C_{ij}$ (Å$^6$eV)</th>
<th>$r_0$ (Å)</th>
<th>$r_1$ (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg–O</td>
<td>1279.69</td>
<td>0.29969</td>
<td>–</td>
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</tr>
<tr>
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<td>0.21916</td>
<td>32.0</td>
<td>0.2</td>
<td>1.05</td>
</tr>
<tr>
<td>O–Al</td>
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<td>0.3013</td>
<td>–</td>
<td>0.15</td>
<td>0.65</td>
</tr>
<tr>
<td>O–Ga</td>
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<td>0.3019</td>
<td>–</td>
<td>0.15</td>
<td>0.65</td>
</tr>
<tr>
<td>O–In</td>
<td>1595.65</td>
<td>0.3296</td>
<td>7.402</td>
<td>0.45</td>
<td>1.10</td>
</tr>
</tbody>
</table>

Table 1
Potential parameters for magnesium aluminate, semi inverse magnesium gallate, and inverse magnesium indate spinels.

\[
\phi(r_{ij}) = \begin{cases} 
\text{ZBL} & r_{ij} < r_0; \\
\frac{g(r_{ij})}{r_0} & r_0 \leq r_{ij} < r_1; \\
A_{ij}\exp\left(-\frac{r_{ij}}{\rho_{ij}}\right) - \frac{C_{ij}}{r_{ij}^6} + V(r_{ij}) & r_1 \leq r_{ij} < r_2; \\
V(r_{ij}) & r_{ij} \geq r_2; 
\end{cases} 
\]

(1)

\[g(r_{ij}) = e^{(f_1 + f_2 r_{ij} + f_3 r_{ij}^2 + f_4 r_{ij}^3 + f_5 r_{ij}^4 + f_6 r_{ij}^5)}.\]

(2)

In Eq. (1), $V(r_{ij})$ denotes the Coulomb potential, and potential parameters $A_{ij}$, $\rho_{ij}$ and $C_{ij}$ reported in Table 1, were determined by a multistructure-fitting scheme to describe correctly the inversion in these spinels [13] [14]. The cation-cation interactions are purely Coulombic, such that the $A_{ij}$ and $\rho_{ij}$ parameters are zero. The potential gives a lattice parameter of 8.12 Å for MgAl$_2$O$_4$, 8.31 Å for the half-inverse MgGa$_2$O$_4$ and 8.85 Å for the inverse MgIn$_2$O$_4$. These values are slightly higher than those reported experimentally in [15].

2.2 Simulation setup

The simulations were performed in crystalline structures that reflect the degree of disorder that can be observed experimentally for each material: $i = 0$ for MgAl$_2$O$_4$, $i = 0.5$ for MgGa$_2$O$_4$ and $i = 1$ for MgIn$_2$O$_4$. For MgGa$_3$O$_4$ and MgIn$_2$O$_4$, this was achieved by using a Monte-Carlo energy minimisation technique [16] to generate a 2x1x1 supercell from which the whole lattice was generated.

The Coulomb sums were calculated using the Distributed Parallel Multipole Tree Algorithm (DPMTA) which is a fast multipole method developed by
Rankin [17]. In the implementation of the algorithm, we choose not to impose periodic boundary conditions but instead to use a charge-neutral cube and fix the three outer atomic layers. We ensured that the cascades did not interact significantly with the cell boundaries by using sufficiently large cells. Consequently, the 400 eV simulations were carried out using a cubic cell of size $\sim 50 \AA$ containing 12331 ions, whereas the 2 keV simulations were performed in a system of size $\sim 100 \AA$ containing 97,669 ions. Before the dynamical cascades were initiated, the system was relaxed for up to 10 ps. The cascades were carried out at an initial temperature of 0 K. The knock on event was set up by giving extra kinetic energy to the PKA along a specific crystallographic direction and the simulation was monitored up to 10 ps, the time by which most defects had relaxed.

3 400 eV cascades

We simulated 400 eV irradiation events since simulations showed that the higher energy cascades produce sub-cascades with similar defect structures. Because of differences in the atomic masses and displacement energies of the separate sublattices, 9 simulations were performed for each of the O$^{2-}$, Mg$^{2+}$ and B$^{3+}$ PKAs where B = Al, Ga or In. The residual defects for each PKA reported in Table 2 are averaged and are representative of the damage per simulation. Interstitials, vacancies and antisite defects are denoted using the simplified Kröger-Vink notation [18]. In the table a split interstitial is counted as two interstitials and a vacancy, not a single defect.

3.1 MgAl$_2$O$_4$

Some results concerning cascades in normal spinel have been previously reported in [7]. The maximum number of displaced ions typically occurred around 100–150 fs after the PKA was set in motion. Most displaced ions recombined to lattice sites within the first picosecond leaving only a few surviving defects. For the Mg PKA, the predominant defects occurring were the Mg split interstitial Mg$^{\ddag}_i$–V$^{\ddag}_Mg$–Mg$^{\ddag}_i$, and the isolated Mg vacancy V$^{\ddag}_Mg$. In normal spinel, cation split interstitials centred about a tetrahedral Mg vacancy were common. This configuration is usually aligned along $<110>$ so that the two interstitials could fill the space of the neighbouring unoccupied octahedral interstices. Such a situation is illustrated in Fig 2(a) with a split Mg interstitial separated from an isolated vacancy with one Al and one Mg antisites. Table 2 shows that the Mg PKA created a total of 1 cation antisite per simulation for this energy. Out of the 9 trajectories, 3 ended up without any damage to the crystal when O was the primary knock-on atom. The remaining cascades
Table 2
The residual defects obtained at 400 eV for the three spinels. The defects tabulated are antisites, interstitials and vacancies respectively, and the bold characters refer to the predominant defect species of each type.

<table>
<thead>
<tr>
<th>Compound</th>
<th>PKA</th>
<th>MgB</th>
<th>B_{Mg}</th>
<th>M_{B}</th>
<th>O_{B}^i</th>
<th>V_{Mg}''</th>
<th>V_{O}^-</th>
<th>V_{B}''</th>
</tr>
</thead>
<tbody>
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<td>Mg^{2+}</td>
<td>0.67</td>
<td>0.44</td>
<td>2.0</td>
<td>0.22</td>
<td>0.22</td>
<td>2.22</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>O^{2-}</td>
<td>0.11</td>
<td>0.11</td>
<td>0.00</td>
<td>1.11</td>
<td>0.00</td>
<td>0.00</td>
<td>1.11</td>
</tr>
<tr>
<td></td>
<td>Al^{3+}</td>
<td>0.56</td>
<td>1.11</td>
<td>1.22</td>
<td>0.89</td>
<td>0.33</td>
<td>0.67</td>
<td>0.89</td>
</tr>
<tr>
<td>MgGa_2O_4</td>
<td>Mg^{2+}</td>
<td>1.33</td>
<td>0.67</td>
<td>1.56</td>
<td>1.44</td>
<td>2.00</td>
<td>2.22</td>
<td>1.44</td>
</tr>
<tr>
<td></td>
<td>O^{2-}</td>
<td>0.22</td>
<td>0.33</td>
<td>1.00</td>
<td>3.11</td>
<td>0.78</td>
<td>1.11</td>
<td>3.11</td>
</tr>
<tr>
<td></td>
<td>Ga^{3+}</td>
<td>0.44</td>
<td>1.00</td>
<td>3.33</td>
<td>1.89</td>
<td>2.56</td>
<td>2.78</td>
<td>1.89</td>
</tr>
<tr>
<td>MgIn_2O_4</td>
<td>Mg^{2+}</td>
<td>0.33</td>
<td>0.22</td>
<td>1.33</td>
<td>1.78</td>
<td>1.56</td>
<td>1.11</td>
<td>1.78</td>
</tr>
<tr>
<td></td>
<td>O^{2-}</td>
<td>0.00</td>
<td>0.11</td>
<td>0.33</td>
<td>3.22</td>
<td>1.11</td>
<td>0.33</td>
<td>3.22</td>
</tr>
<tr>
<td></td>
<td>In^{3+}</td>
<td>0.33</td>
<td>0.89</td>
<td>0.89</td>
<td>3.11</td>
<td>2.11</td>
<td>0.33</td>
<td>3.11</td>
</tr>
</tbody>
</table>

Fig. 2. Typical defects observed at 400 eV in MgAl_2O_4 for a Mg^{2+} PKA initiated along ⟨135⟩ (a), an O^{2-} PKA along ⟨173⟩ (b), an Al^{3+} PKA along ⟨175⟩ (c), an Al^{3+} PKA directed along ⟨123⟩ (d) an Al^{3+} PKA along ⟨173⟩ (e). Interstitials are denoted by spheres and vacancies are shown as cubes. The green colour refers to the ternary cation B (here B = Al); Mg defects are coloured blue and O defects red. Cones represent antisites (purple - M_{GB}; grey - B_{Mg}). The same colour scheme applies for all the three spinels.

Fig. 2. Typical defects observed at 400 eV in MgAl_2O_4 for a Mg^{2+} PKA initiated along ⟨135⟩ (a), an O^{2-} PKA along ⟨173⟩ (b), an Al^{3+} PKA along ⟨175⟩ (c), an Al^{3+} PKA directed along ⟨123⟩ (d) an Al^{3+} PKA along ⟨173⟩ (e). Interstitials are denoted by spheres and vacancies are shown as cubes. The green colour refers to the ternary cation B (here B = Al); Mg defects are coloured blue and O defects red. Cones represent antisites (purple - M_{GB}; grey - B_{Mg}). The same colour scheme applies for all the three spinels.

Typically resulted in 1 anion split interstitial O_{B}'' - V_{O}^- - O_{B}'' with an isolated V_{O}^- (see Fig. 2(b)). The Al PKA produced more Al antisites as compared to Mg antisites. The other defect structures obtained were cation antisites, Mg and O split interstitials as shown in Fig. 2(c), and a mixed Mg-Al split interstitial centred about a Mg vacancy (Fig. 2(d)). Two simulations resulted in a defect configuration including an Al interstitial trapped between 2 adjacent Al vacancies. Such a situation is illustrated in Fig. 2(e). Cation split inter-
stitials centred about an Al vacancy did not occur. Oxygen split interstitials could reform with neighbouring vacancies if the axis of the split interstitial was in the direction of the O vacancies, leading to a one-dimensional diffusion mechanism. The migration energy for this transition to occur was found to be 0.29 eV. Conversely if the O split interstitial had its axis perpendicular to that joining its central vacancy to the neighbouring O vacancy as shown in Fig. 2(b), then an extra 0.63/0.67 eV was required for the configuration to align along the favourable direction before migrating (these energies refer to at least two mechanisms for the split interstitial to rotate). Al antisites could easily pin the O split interstitial to prevent the latter from migrating (see Fig. 2(c)). The two defects have a positive binding energy of 1.65 eV.

3.2 Half inverse MgGa₂O₄

Figure 3 shows the damage in the half inverse \((i = 0.5)\) magnesium gallate when choosing a Mg\(^{2+}\), O\(^{2−}\) and Ga\(^{3+}\) PKA. In this crystal, half of the occupied tetrahedral interstices were taken up by the Ga\(^{3+}\) ions. In the normal spinel, this site was occupied solely by the Mg ion and was the only favourable site where cation split interstitials could form. Cation split interstitials centred about a tetrahedral Ga vacancy were observed in the magnesium gallate spinel (see pure Ga split interstitial in Fig. 3(a)).

Fig. 3. The pictures show the damage after 5 ps for a PKA energy of 400 eV in the gallate spinel. (a) Mg\(^{2+}\) PKA along \((123)\) resulting in the formation of pure O, Ga, and Mg split interstitials, one Ga–O vacancy pair and cation antisites; (b) O\(^{2−}\) PKA along \((175)\); (c) Ga\(^{3+}\) PKA along \((210)\) forming a distinct ‘ring’ structure; (d) Mg\(^{2+}\) PKA along \((751)\); (e) Ga\(^{3+}\) PKA along \((173)\) creating disorder on the cation sublattice.

For the Mg\(^{2+}\) PKA, the common defects consisted of O, Mg, Ga and mixed Mg–Ga split interstitials centred about either a Mg vacancy or a Ga vacancy. Fig. 3(a) exemplifies the final damage obtained with a Mg PKA. The simu-
lations initiated by O$^{2−}$ PKAs produced more damage on the O sublattice, the defects being mainly O split interstitials or O crowdions oscillating one-dimensionally in the $<110>$ directions. Fig. 3(b), gives an example where the O interstitial settled at a tetrahedral interstice between 3 O vacancies and 1 Ga vacancy. For the Ga$^{3+}$ PKA more cation defects were obtained. Typical defects were interstitial–vacancy ‘crowdion’ chains, with the interstitials trapped at unoccupied interstices between the vacancies. Fig. 3(c) illustrates a ‘ring’ structure made up of 3 Mg$^{i}$, 2 V$^{m}$ Ga and 1 V$^{m}$ Ga. Closer inspection revealed that the 3 Mg interstitials settled at tetrahedral interstices between the octahedral vacancies. This defect formed around the 1 ps time mark following the cascade initiation. Over the whole set, more defects were seen in this compound as compared to the normal spinel. This difference occurs because of extended split interstitials on the cation sublattice, anion crowdions, and also the tendency of an isolated O vacancy to push the nearest neighbour tetrahedral cation slightly off its site resulting in an extra vacancy and interstitial.

3.3 Inverse MgIn$_2$O$_4$

Table 2 shows a smaller number of antisite defects produced in the inverse indate spinel as compared to the normal and half-inverse spinel. In this case, the irradiated crystals contained mainly anion and cation split interstitials and crowdions as in the half inverse spinel. The cation split interstitials were centred about an In vacancy, the latter being an initially occupied tetrahedral site. Figure 4 illustrates some of the final structures for 3 simulations.

The average number of interstitials created by a Mg$^{2+}$ PKA for a 400 eV cascade in magnesium indate is 1.33 for Mg, 1.78 for O and 1.56 for In. The interstitials formed Mg$^{i}$–V$^{m}$ In–In$^{i}$, O$^{i}$–V$^{m}$ O–O$^{i}$ and In$^{i}$–V$^{m}$ In–In$^{i}$ structures and O crowdions. No split interstitials around an Mg octahedral vacancy were found just as no split interstitials were observed around an octahedral Al site in normal MgAl$_2$O$_4$. The O$^{2−}$ PKAs produced damage mainly on the oxygen sublattice. An average of 3.2 O interstitials were created per trajectory as compared to 0.3 and 1.1 for Mg and In interstitials respectively. For the 9 trajectories considered only 1 In antisite was observed. For an O PKA initiated along (123) the irradiated lattice ended as shown in Fig. 4(b), with the formation of 1 O divacancy defect, a cation chain, an isolated Mg vacancy and an O split interstitial. The cation chain was created around 1.3 ps with the interstitials occupying octahedral interstices. Near the chain an O interstitial is trapped at a tetrahedral interstice between two O vacancies. Similar defects were obtained when the PKA was In$^{3+}$. In this case, the ballistic phase involved more displaced ions. For a PKA orientation of (731), the final damage is presented in Fig 4(c). In this case the O crowdion structure hops around during the period of the simulation and does not settle to a stable configu-
ration. Over the whole set, O defects were predominant as reported in Table 2.

Fig. 4. Some defect structures in magnesium indate for 400 eV using (a) Mg$^{2+}$ PKA along ⟨210⟩, (b) O$^{2−}$ directed along ⟨123⟩ and (c) In$^{3+}$ PKA along ⟨731⟩.

4 2 keV cascades

A total of 12 trajectories per compound was run for PKA energies of 2 keV. The simulations were equally divided among the PKA species resulting in 4 simulations for each sublattice. The average number of defects per simulation is reported in Table 3 and Fig. 5 shows the residual damage for the three spinels. Complex defect structures and clusters occurred in the inverse spinels for certain PKA trajectories, especially when the PKA was the ternary cation.

<table>
<thead>
<tr>
<th>Compound</th>
<th>PKA</th>
<th>MgB</th>
<th>BMg</th>
<th>Mg$^i$</th>
<th>O$'^i$</th>
<th>B$'''$</th>
<th>V$'^{''}$Mg</th>
<th>V$'^{''}$O</th>
<th>V$'^{''}$B</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgAl$_2$O$_4$</td>
<td>Mg$^{2+}$</td>
<td>2.25</td>
<td>1.75</td>
<td>7.75</td>
<td>6.75</td>
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<td>8.25</td>
<td>6.75</td>
<td>1.50</td>
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<td></td>
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<td>4.00</td>
<td>5.25</td>
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<tr>
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<td>15.75</td>
<td>4.00</td>
<td>15.75</td>
<td>17.75</td>
</tr>
</tbody>
</table>

Table 3
The table shows the average number of surviving defects for a 2 keV PKA. The numbers in each row are averages of 4 simulations for each PKA.
Fig. 5. Residual defects from a 2 keV event in the (a) normal spinel - cascade initiated by an Al PKA along $\langle 210 \rangle$, (b) gallate spinel resulting from a Mg PKA set up in the $\langle 751 \rangle$ direction, (c) inverse MgIn$_2$O$_4$ for a Mg PKA along $\langle 435 \rangle$. In general, the normal spinel ended with the least damage, while complex defect clusters were formed in the inverse spinels.

4.1 MgAl$_2$O$_4$

Damage resulting from 2 keV cascades in MgAl$_2$O$_4$ was characterised by an increased number, but similar defect structures, as the 400 eV cascades. For example, the O$^{2-}$ PKAs produced little damage on the cation sublattice with a small fraction of antisites defects as shown in Table 3. The cation PKAs created more disorder over their respective sublattices and also a considerable amount of anion defects. All the anion defects were of split interstitial form, while the cation defects could be observed to form split interstitials centred about an Mg vacancy and defect structures over the Al sublattice. In the latter case, defects with an Al ion trapped between two vacancies were observed, and a few ring structures were obtained consisting of 3 Al$^{\text{V}}$ and 3 V$^{\text{Al}}$ (see Fig. 5a). Detailed analysis of these ring defects using temperature accelerated dynamics [19] showed a formation energy of 10.24 eV and a barrier of 1.23 eV for the structure to annihilate. The ring defects alter only slightly the perfect lattice but absorb a considerable amount of energy in their creation. The existence of such defect clusters could contribute significantly to the radiation tolerance of spinel.

4.2 MgGa$_2$O$_4$

More antisites were generated in this compound as compared to the normal and fully inverse spinels. Substantial disorder on the cation sublattice formed, with interstitial-vacancy chains, where the interstitials were trapped at intrin-
sic vacancy interstices. Ring structures involving both cations were observed with many defect clusters. Fig. 5(b) results from a Mg PKA directed along ⟨751⟩. This cascade instantaneously divided into two distinct subcascades. The peak damage occurred around 200 fs with the formation of 237 interstitials, 237 vacancies and 9 antisites. Most of the point defects retracted within the next 1.5 ps leaving behind almost stable defect structures. One structure not observed for 400 eV PKAs is shown Fig. 5(b) with six cations sharing 4 octahedral vacancies. This structure preserves the coordination number of the neighbouring O ions, but the cations now occupy tetrahedral interstitial sites. Surviving defects were comparable for both Mg\(^{2+}\) and O\(^{2-}\) PKAs but the Ga\(^{3+}\) PKAs produced more damage.

4.3 MgIn\(_2\)O\(_4\)

Defects that were previously observed at 400 eV for the inverse spinel were produced by Mg\(^{2+}\) and O\(^{2-}\) PKAs, but the In PKAs created much disorder on the In sublattice with the formation of In interstitial–vacancy chains. An average of 15.8 In interstitials remained per trajectory for the In PKAs as compared to 8.5 and 9.0 for the Mg\(^{2+}\) and O\(^{2-}\) PKAs.

The simulation initiated by an In PKA along ⟨752⟩ generated the most damage in this material. A total of 64 interstitials (56.2 % In, 17.2 % Mg, 26.6 % O), 64 vacancies and 15 antisites remained in the structure at the end of a 12 ps run. 33 out of the 36 In interstitials were trapped at unoccupied interstices: 28 were taken up at octahedral interstices and only 5 were seen at tetrahedral sites. In these complex environments, there is a strong tendency for the In interstitials to regain the coordination 6 with the anions, explaining the existence of these cation interstitial–vacancy chains.

Fig. 5(c) illustrates the resulting damage for a Mg\(^{2+}\) PKA directed along ⟨435⟩ in the inverse magnesium indate. Typical cation chains can be seen, while the O interstitials formed dancing crowdions, i.e. hop back and forth between the split interstitial and crowdion structures and do not attain a permanent stable configuration. However the cation crowdion structures that form remain stable over typical MD time scales.

4.4 Higher energy Cascades

A few calculations have also been carried out on larger systems but not in sufficient number to generate good statistics. For the normal aluminate spinel, calculations at a higher energy of 10 keV in a system of ~1 million ions resulted in similar defect structures as in the lower energy cascades. Cascade
branching occurred in the 10 keV simulations and spread over a larger volume of the computational cell. Figure 6 gives an example of residual damage from one such cascade triggered by a Mg$^{2+}$ PKA ending up with a total of 57 vacancies, 57 interstitials and 22 cation antisites. Cascade branching and the reannealing of defects is typical of all the spinel systems, in marked contrast to cascades reported in zircon [20] where large contiguous regions of the crystal structure are damaged and amorphised.

![Image](image_url)

**Fig. 6.** Damage resulting from a 10 keV cascade in magnesium aluminate spinel for a Mg$^{2+}$ PKA.

5 Discussions and Conclusions

The dynamical simulations resulted in primary defect structures for the three compounds consisting of cation split interstitials centred about a tetrahedral vacancy, oxygen split interstitials, cation antisites and vacancy pairs. The tabulated values show that more defects were created in the inverse spinels partly due to surviving crowdion chains of alternate interstitials and vacancies, for example the rightmost defect structure shown in Fig. 5c (this counts as 3 interstitials and 2 vacancies in our classification scheme). Although cascades initiated in the inverse and half-inverse spinels create more damage during the ballistic phase of the cascade, it is not clear how this damage would evolve in
the long term. For example, the defect structure shown magnified in figure 5b consists of a region of the crystal which has a local charge of +5 (an extra A and B ion). The long term survival of such a structure is questionable.

The results show clear differences between the different structures with a lower variety of defects created in normal spinel, especially at the lower energies. For example the low energy Mg PKA produced defects that were only a split Mg interstitial and an isolated vacancy while the O PKA produced only damage on the O sublattice for normal spinel. For the half inverse and inverse spinels, the Mg PKA produced more defects such as the Mg-B cation interstitial about a B ion vacancy and cation crowdion chains, while damage on the cation sublattice for the O PKA was larger than in normal spinel but still exceeded by that on the O sublattice. The B$^{3+}$ PKAs created about twice as many O defects in the half inverse spinel and about three times as many in the fully inverse spinel compared to normal. In the fully inverse spinel there was a tendency for the In interstitials to rearrange about octahedral sites and regain a co-ordination of 6 with the O ions. An O vacancy in the fully inverse case had also the effect of causing more local distortion, displacing slightly the neighbouring tetrahedrally-sited In ion.

The cascades in the spinels create defects which do not greatly change the overall crystal structure but which require a certain amount of energy to form. The cation anti-site defects are an example. These require only a small energy to form but are very numerous. On the other hand, the less frequently observed Al ‘ring’ defect in magnesium aluminate spinel requires an energy of 10.2 eV to form but involves only the symmetric switching of 3 Al cations to unoccupied tetrahedral sites. This ability to absorb energy, combined with sub-cascade branching which distributes the energy throughout the crystal, rather than a localised thermal spike creating a fully disordered region, means that the crystal structure is able to accommodate a cascade without undergoing radical distortion. Thus defects are more widely spread and not as clustered as in e.g. zircon [20].

References

