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Citation for published version:

Digital Object Identifier (DOI):
10.1103/PhysRevB.87.104106

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Physical Review B: Condensed Matter and Materials Physics

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Molecular dynamics and object kinetic Monte Carlo study of radiation-induced motion of voids and He bubbles in bcc iron

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(Received 10 December 2012; published 12 March 2013)

We show that voids adjacent to radiation damage cascades can be moved in their entirety by several lattice spacings. This is done using molecular dynamics cascade simulations in iron at energies of 1–5 keV. An equation describing this process is added to an object kinetic Monte Carlo (OKMC) code to allow study of the mechanism at longer time scales. The mechanism produces an enhancement of void diffusion by two orders of magnitude from $1 \times 10^{-22} \text{ cm}^2/\text{s}$ to $3 \times 10^{-20} \text{ cm}^2/\text{s}$. Repeating the study on He bubbles shows that the movement is damped by the presence of helium in the void.

DOI: 10.1103/PhysRevB.87.104106 PACS number(s): 61.82.Bg, 61.72.Qq, 61.80.Az

I. INTRODUCTION

Steels under high-energy neutron bombardment such as experienced in fission and fusion reactors are known to develop voids and bubbles over time. These have seriously deleterious effects, and a clear understanding of how these defects affect the properties of a material is crucial to designing better steels. Molecular dynamics can be used to gain insight into the atomistic level interactions between radiation cascades and defects such as voids and bubbles. The long-term effects of the atomic mechanisms can be observed using object kinetic Monte Carlo (OKMC) and such a multiscale approach is becoming increasingly popular.

A study of very low-energy cascades of just a few eV impacting on voids has been performed by Dubinko and Lazarev.\textsuperscript{1} It showed that very low doses can lead to the dissolution of voids by focusons striking them. This is consistent with experimental observation\textsuperscript{2} that the voids can be reduced in size by irradiation under favorable conditions.

At higher energy, Pu et al.\textsuperscript{3} studied stability of helium bubbles struck by radiation cascades. They find that the stability of the bubble depends on its He/vac ratio. Clusters with significantly more He than vacancies tend to absorb vacancies from the cascade. Conversely, clusters with a low density of He tend to lose vacancies during the cascade, approaching the stable ratio of near 1:1.

Simulations by Parfitt and Grimes\textsuperscript{4} study the behavior of helium bubbles in uranium dioxide under radiation-induced cascades. An atom close to the bubble is given 10 keV of energy to simulate a radiation event, and the resulting cascade observed. They find that where the hot ‘melt’ region of the cascade overlaps the bubble, there is an increased chance of helium gas being emitted into the region. As the melt region cools, the helium is trapped in the lattice. Their study concludes that this emission is mostly not ballistic, but rather is due to the disorder of the bubble walls allowing low-energy diffusion paths into the lattice. Some ballistic emission of gas is also observed at shorter time scales. Most of the gas remains confined to the bubble. They expect the mechanism of increased surface diffusion to be dominant for larger bubbles.

Further properties of He in iron during cascades (but not in the presence of voids or bubbles) have been obtained by Lucas and Schäublin.\textsuperscript{5} They find that the presence of substitutional He tends to reduce the number of Frenkel pairs produced. Interstitial He increases lattice stress and favors the formation of self-interstitial atoms (SIA) as well as stabilizing SIA clusters. They observe increased diffusion of He at temperature, which explains how bubbles can form rapidly.

An interesting mechanism for interstitial cluster formation during cascades in pure α-iron has been observed by Calder et al.\textsuperscript{6} They find that some cascades emit particles moving faster than the cascade wave front, which causes secondary cascades ahead of the main one. Collision of the high-density primary wave with the core of the secondary cascades can lead to nucleation sites for interstitial clusters. This is due to atoms being forcefully pushed into the secondary cascade’s low-density core region.

Under certain irradiation conditions, voids can form lattices.\textsuperscript{7} This effect has been reproduced by Heinisch and Singh\textsuperscript{8} in kinetic Monte Carlo (KMC) simulations. Their study uses clusters of crowdions formed by irradiation and moving mostly one dimensionally along close-packed planes to explain the alignment. Voids off the lattice are exposed to a greater flux of crowdions and so tend to shrink, whereas voids on lattice are shielding each other along the close-packed planes. The simulations do produce a void lattice successfully although the lattice does not show the same refinement as experimentally observed.

This study looks at the interaction of voids and bubbles with cascades and how this affects their center of mass. Simulations show a mechanism of atomic injection into the void by the density wave front of the cascade. This leads to a new void being formed at the cascade core because the atoms absorbed by the original void are not available for recombination. The mechanism is explored atomistically in molecular dynamics. A simple equation is fitted to model the motion observed in the molecular dynamics simulations. This equation is then used as an event in the OKMC simulations, along with other thermal processes, to allow the long-term evolution of the system to be explored.

II. METHOD

The molecular dynamics study used a version of MOLDY\textsuperscript{9} modified for variable time-step cascade simulations as detailed previously.\textsuperscript{10} The Ackland et al.\textsuperscript{11} embedded atom potential was used for Fe-Fe interactions. The Juslin and Nordlund pair
potential\textsuperscript{12} was used for Fe-He and the Beck\textsuperscript{13} pair potential for He-He. The Beck-He-He potential was splined to the Ziegler-Biersack-Littmark (ZBL)\textsuperscript{14} universal potential at short range. Helium is a closed-shell atom, and we do not expect any chemical bonding, so the pair potential formalism is sensible.

The study consists of triggering radiation cascades in \(\alpha\)-iron near to a void or bubble. To create stable voids or bubbles in a bcc iron lattice, a single atom was removed near the center of the system. The system was then relaxed and the atom with the highest potential energy removed. Iterating this process produces a stable void with faceted sides aligned to stable lattice directions. Stability was tested by running the sample at 900 K for 20 ps and observing no changes.

Bubbles were produced by placing helium atoms onto the vacant sites in the void. The system was then quenched using low-temperature molecular dynamics (MD) with a high thermostat value to limit the velocity of the helium atoms. Once the helium has reached a stable configuration, the system may be thermally equilibrated with standard thermostating.

Cascades were initiated by giving a substantial kinetic energy to one iron atom near to the defect cluster; this atom is called the primary knock-on atom (PKA). Cascade energies of 1, 2.5, and 5 keV were considered. For 1- and 2.5-keV cascades, a box dimension of 40\(a_0\) (128 000 atoms) was used, with 65\(a_0\) (549 250 atoms) for 5 keV. We use the \(NVT\) canonical ensemble (constant number, volume, cell shape, temperature, and zero total momentum) with periodic boundaries.

The final atomic configurations were analyzed by comparison to a perfect lattice aligned to the defective lattice.\textsuperscript{10} The Wigner-Seitz cell around each perfect lattice point is checked for atoms, allowing easy identification of interstitials, vacancies, and substitutional defects. Defects were considered to be clustered if they were within second-nearest-neighbor distance of each other.

The data were further processed by comparing the nearest neighbors of every atom before and after the collision, marking atoms with at least two new neighbors. This identifies the area that has been affected by the cascade, as well as being sensitive to correlated motion of atoms. Additionally, it provides an effective method for visualizing the cascade by joining a line from the initial to final location of each atom (cf. Calder et al.).\textsuperscript{6} The region covered by these lines is the part of the simulation “melted” by the cascade and the lines tend to form connected paths due to replacement-collision events. By taking the list of atoms that have changed neighbor, and filtering it so that only members that have moved at least one lattice site remain, an estimate of cascade volume is obtained. The cascade radius is calculated from this volume.

The point around which the cascade is centered is identified by the center of mass of those listed atoms with at least four neighbors also in the list. This eliminates small subclusters and replacement-collision chains that would otherwise skew the estimate.

\section*{III. Results}
Simulations of a cascade interacting with a void or bubble in a periodic cubic supercell of \(\alpha\)-Fe were performed under various conditions. Most PKAs were given initial velocities close to (111) direction towards the void, with the slight deviation to avoid perfect replacement-collision chains. This high-symmetry direction was chosen in order to localize the cascade\textsuperscript{15} and give easily comparable data across different simulations.

To explore if the effects observed are dependent on the crystal alignment, some cascades were also performed in the \(\langle 111 \rangle\) direction and in random directions. Simulations were performed at 1, 2.5, and 5 keV on voids/bubbles of sizes 30 and 100, with the bubbles at ratio 1:1 for vacancy:helium. One set of data is also performed at 300 K to explore temperature effects (the various simulations along with their damage yields are summarized in the Supplemental Material\textsuperscript{16}).

For 300-K simulations, little change was seen in the results apart from a slight reduction of Frenkel pairs, and so further temperature studies were not pursued, although higher temperatures may have more effect. Results along the \(\langle 111 \rangle\) direction were found to be consistent with similar studies of the \(\langle 111 \rangle\) direction. The radii and damage of the cascades were also consistent across randomly launched trajectories.

Normally, voids move by capture and emission of point defect, but interestingly some of our simulations show that the void is moved by several angstroms, far more than absorption of the defects created by the cascade would allow. The process appears to have an upper and lower interaction radius, close to the size of the cascade, as shown in Figs. 1 and 2.

A typical cascade displaces atoms around itself creating a density “wave” moving spherically outwards, with a depleted region at its core. Normally, in a perfect lattice this wave will collapse back as the energy dissipates, causing recombination of vacancies and interstitials at the core (Fig. 3). The defect yield (dpa) from a cascade normally considers only point defects surviving this recombination.

However, when a preexisting void is adjacent to the cascade, a more complex mechanism may take place. The density wave injects atoms into the preexisting void, where they find sites to occupy and so do not return as the wave collapses back. This results in the original void being filled, and a new void of equal size appearing at the cascade core. The outcome of this process is equivalent to the movement of the entire void by several lattice spacings. This behavior was reproducible for

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure1.png}
\caption{Graph showing movement of void center of mass from starting position \(\Delta d\) cluster (filled squares) and bubble (empty triangles) vs distance to PKA (\(d\)) for a 30 defect cluster hit at 1 keV. It can be seen that the void can move significantly towards the PKA site for certain cascade distances. The presence of helium in the void greatly damps this effect.}
\end{figure}
voids under all conditions considered, providing the cascade overlaps the void.

Figures 1 and 2 show how the distance moved by the center of the void depends on the separation between void and cascade. The cascade will typically begin a few lattice spacings from the starting position of the PKA. At very short distances, the PKA can pass directly through the void, which is why there is a minimum interaction threshold for PKA distance.

Data suggest interaction thresholds of 12 and 25 Å with peak interactions around 18 Å for 1 keV (Fig. 1). For 2.5 keV, the interaction starts at 18 Å, lasting until 40 Å and has peak at 31 Å. For 5 keV, the interaction starts around 12 Å and ends at 38 Å and had peak interaction at 31 Å (Fig. 2).

For 1-keV cascades on bubbles of size 30, almost no motion of the cluster is observed even for high cascade overlap. The slight movement seen (Fig. 1) is due to the side of the bubble nearest the cascade absorbing a few vacancies, shifting the center of the cluster. On average, the bubble grew by one vacancy and almost all the helium remained clustered in it.

For a void of size 100 impacted at 5 keV, significant movement of the void towards the cascade was observed (Fig. 2). Void size tended to decrease slightly on average to 84 vacancies. This change is largely due to the void being split, as shown in Fig. 4(a), which occurs in three simulations. Ignoring the split samples void size tends to be around 88 after cascade interaction.

Similar results to the 1-keV bubble case were observed for a 5-keV cascade impacting a helium bubble of size 100. Figure 4(b) shows a typical bubble after impact. The helium reduces the mobility of the cluster (Fig. 2) and the helium remains clustered in the bubble. Cascade radius is unaffected, although the number of Frenkel pairs formed seems to increase slightly (Table I). Occasional shrinkage by up to 5 vacancies is observed at longer distance due to interstitial absorption from the cascade periphery.

The standard deviation of the vacancies in the voids and bubbles was analyzed to explore the compactness of the cluster. For each void, the standard deviation of the vacancies in the void is calculated, and then this number is averaged over different voids. The standard deviation for clusters that move more than 1 Å during the cascade is compared to the value for a perfect cluster (see Table II).
Results shows that the bubbles remain compacted, while the voids become more diffuse. Since all the defects are within second-nearest-neighbor distance of each other in a cluster, it is expected that they will collapse to a faceted void due to vacancy attraction. In kinetic Monte Carlo models, it is standard to assume that there is a capture radius of at least this distance\(^{17-20}\) between vacancies, leading to collapse to a compact void.

### IV. OKMC RATES

In this work, we wrote an object kinetic Monte Carlo (OKMC) code to study how voids repeatedly exposed to radiation cascades move, on a time scale far longer than could be achieved in molecular dynamics. In addition to thermally induced vacancy and interstitial cluster motion, we include defect creation from cascade debris, and allow vacancy clusters to jump to cascade sites. These latter two rates are dose dependent, and their effect is modeled using data gained from the molecular dynamics simulations.

Object kinetic Monte Carlo is similar to normal kinetic Monte Carlo of defects, but allows clusters of defects to be treated as objects. Clusters can have their own individual diffusion rates, dissociation rates, and interaction events that affect the cluster as a whole, instead of dealing with the constituent point defects. This method is more efficient than KMC as the number of bodies in the simulation is greatly reduced. Other methods such as accelerated molecular dynamics\(^{21}\) also allow the time scale to be increased beyond molecular dynamics time scales, and have the advantage of not requiring predefined rate tables. However, given the time scales that we wished to explore and the availability of data for rates, including input gained from the molecular dynamics simulations, OKMC was chosen.

<table>
<thead>
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<th>Type</th>
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<th>(E_d) (eV)</th>
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</tr>
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<td>(V_2)</td>
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<tr>
<td>(I_4)</td>
<td>0.43</td>
<td>1.26</td>
</tr>
<tr>
<td>(L_{m,n}) Immobile</td>
<td>(E_m^{n+1} + E_f^I + (E_b^I - E_f^I)g(n))</td>
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</tr>
</tbody>
</table>

The OKMC code was written and used to explore the significance of long-term evolution of the void motion. The short-time-scale jump events identified in the molecular dynamics are added as an equation in the OKMC and are not directly atomistically simulated. Jump events are simulated by moving the void to the cascade core if it is within the interaction radius of the cascade. These jumps can then be simulated alongside other mechanisms such as radiation-induced Frenkel pairs and thermal dissociation. Locally, the motion induced by the cascade is ballistic in nature, as the void is essentially moved by the expanding wave of interstitials. However, when considered over longer time scales, the motion is diffusive in that the void makes jump at a given rate, undergoing a random walk. The diffusion rate for such a process can be calculated as \(D = \frac{x^2}{6t}\), where \(x\) is the average jump distance and \(t\) is the average time between jumps. This is not thermally activated diffusion, rather it is activated by proximity to a cascade.

The code implements the standard resident-time algorithm, also called the kinetic Monte Carlo or Gillespie algorithm.\(^{22}\) The established mechanism for void diffusion is by point defect absorption and emission events (later referred to as standard diffusion) and so our new mechanism’s diffusion is compared to this process. The OKMC model includes vacancy clusters, interstitial clusters, defect creation from cascade debris, and direct cascade-void interactions. It allows diffusion of small clusters and thermal dissociation of point defects from clusters following the energy barriers given by the Fu et al.\(^{23}\) study as used in Björkas’s et al.\(^{24}\) model (see Table III). The capture radii of clusters follows the model of Ortiz and Caturia\(^{17}\):

\[
r_{L,V} = Z_I V \left[ \left( \frac{3n\Omega}{4\pi} \right)^{1/3} + r_0 \right],
\]

where \(r_{L,V}\) are the capture radius for interstitial and vacancy clusters, respectively. \(Z_I = 1.15\) is the bias factor accounting for the increased strain field observed for interstitials. \(\Omega\) is the atomic volume and \(r_0 = 3.3\) Å.

Open absorbing boundaries were used for this study, simulating a single void at the center of a single grain. Radiation cascades were introduced into the sample with the maximum interaction radius taken from the data sets.
in the current study (Figs. 1 and 2). The cascade debris is calculated by the Bacon et al.\cite{Beck95} empirical formula for iron: \(N_f = 5.67E_{0.779}\), where \(N_f\) is the number of Frenkel pairs and \(E\) is the damage energy in keV. Cascades were implemented in a shell-like manner, with vacancies randomly placed in a smaller radius, and surrounded by a shell of interstitials. Cluster interactions were carefully performed to take into account the correct change in cluster position for absorption and emission events, as this will affect the diffusion.

If the cascade is sufficiently close to the void, the void is shifted to the center of the cascade region as seen in the MD simulations. By checking the interaction volume of the cascade for the largest void within distance and updating according to

\[
\text{if } r_{\text{cascade}} + r_{\text{void}} > d_{\text{cascade-void}} \text{ then } \bar{r}_{\text{void}} = \bar{r}_{\text{cascade}}.
\]

\(r_{\text{cascade}}\) is the cascade radius, \(r_{\text{void}}\) the void radius, \(d_{\text{cascade-void}}\) is the distance from the cascade center to the void center, and \(\bar{r}_{\text{void}}/\bar{r}_{\text{cascade}}\) is the position vector for the void/cascade. This mechanism was disabled to simulate the standard diffusion.

The trend of both diffusion mechanisms with temperature, dose rate, cascade energy, and system size was studied. The contribution from each process was calculated by separating out the jumps due to each mechanism. Studies were performed with default values: temperature of 500 K, dpa/s of 1, cubic cell of 100 Å and cascade energy of 10 keV on a void of 100 vacancies. Only one parameter was varied from default at a time, to gain information on system trends without performing an exhaustive search of the parameter space. Parameter ranges considered: temperature of 400–600 K, dpa/s of \(10^{-5}–10^{-2}\), cubic cell of 50–200 Å, and cascade energy of 1–20 keV. Under the default parameters, the standard diffusion was found to be \(1 \times 10^{-22} \text{ cm}^2/\text{s}\) while the diffusion due to the new mechanism is \(3 \times 10^{-20} \text{ cm}^2/\text{s}\), two orders of magnitude larger.

For fixed dpa rate, the standard diffusion rate is found to not be strongly dependent on cascade energy. The mechanism only depends on the number of free point defects and increasing cascade energy decreases the number of cascades to conserve dpa rate. However, the new mechanism is highly dependent on cascade energy because this sets the interaction radius.

System size does not strongly affect the new mechanism. It does affect the standard mechanism as the only initial sinks in the system are the void and cell walls. Increasing cell size gives defects more chance to be absorbed by the void instead of the walls and so effectively increases the void capture radius. This leads to a slight increase in diffusivity.

Rates for both mechanisms increase approximately linearly with dose rate so this parameter is not directly critical, although it is important in setting the relative time scale of the system as compared to dissociation events. Neither mechanism is strongly temperature dependent. The standard mechanism lacks temperature dependency because the diffusion of point defects is much faster than the time between cascades. Temperature will only become critical at lower \(T\) when diffusion time of point defects is comparable to the time between cascades.

The new mechanism is found to be dominant over the standard mechanism at all parameters considered apart from for energies of 2 keV or below, when the interaction radius becomes too small to have significant effect. Cascade energy is the most significant factor, with diffusion of the new mechanism rising to \(1 \times 10^{-19} \text{ cm}^2/\text{s}\) at 20 keV.

V. CONCLUSIONS

We have shown that the migration rates of voids in irradiated environments is much higher than expected from simple point defect migrations. Cascades occurring adjacent to voids may cause a jump of the entire void by several lattice spacings if the cascade is within appropriate interaction thresholds. This motion is due to the injection of atoms by the cascade into the original void, trapping them and leaving a new void at the cascade core. The effect becomes more significant as cascade energy increases (even at fixed dpa) because it depends on cascade radius. This jump mechanism has been shown in OKMC simulations to be a more significant mechanism than diffusion due to absorption and emission of point defects. The mechanism is suppressed when helium is present in the void.

The rate is equivalent to one 10.5-Å jump every 18.6 hours on average. This is still a much faster process than the standard diffusion mechanism. Although the overall void diffusion rate is low, the mechanism may still play a role in void mobility. Jumps displace the entire void by several lattice spacings. This may have impact on long-term void evolution processes, such as void lattice formation. Assuming the alignment method discussed in the Introduction, involving \(\langle 111 \rangle\) interstitial clusters,\cite{Juslin04} jumps may expose or shield voids from this flux of interstitials. These occasional jumps may help to fine tune the void lattice over long periods of time. Further study would be needed to verify this.

ACKNOWLEDGMENT

We thank D. J. Hepburn for useful discussions and the EPSRC Scottish CM-DTC for a studentship (G.J.G).


\[^{6}\text{A. F. Calder, D. J. Bacon, A. V. Barashev, and Y. N. Osetsky, Philos. Mag. 90, 863 (2010).}\]


\[^{12}\text{N. Juslin and K. Nordlund, J. Nucl. Mater. 382, 143 (2008).}\]

\[^{13}\text{D. E. Eckel, Mol. Phys. 14, 311 (1968).}\]
For some initial directions, the PKA can be channeled between atomic layers, causing the cascade to occur far from the void.


