Vacancy clustering and prismatic dislocation loop formation in aluminum

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The formation of prismatic dislocation loops is an important factor leading to radiation damage of metals. However, the formation mechanism and the size of the smallest stable loop has remained unclear. In this work, we use electronic structure calculations with millions of atoms to address this problem in aluminum. Our results show that there is a cascade of larger and larger vacancy clusters with smaller and smaller energy. Further, the results show that a seven vacancy cluster on the (111) plane can collapse into a stable prismatic loop. This supports the long-standing hypothesis that vacancy clustering leads to prismatic loops, and that these loops can be stable at extremely small sizes. Finally our results show that it is important to conduct calculations using realistic concentrations (computational cell size) to obtain physically meaningful results.

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The embrittlement of metals subjected to radiation is a long-standing problem in various applications including nuclear reactors. As the irradiation dose increases above a certain threshold, a significant population of prismatic dislocation loops (dislocation loops whose Burgers vector has a component normal to their plane) has been experimentally observed to arise in metals [1–5]. It is widely believed these prismatic loops form through the clustering of vacancies that are generated randomly by irradiation [6]. Specifically, the vacancies diffuse and eventually cluster on specific planes. Once there is a large enough planar cluster, the atoms on the two faces collapse on to each other leaving behind a prismatic dislocation loop.

However, the formation mechanism and the size of the smallest stable loop remain unclear: there is no direct experimental observation of the process, and the theoretical investigations are inconclusive. Recent molecular dynamics simulations [7] support the hypothesized mechanism for iron, but these calculations used the Finnis-Sinclair empirical atomistic potentials whose validity is uncertain in situations involving changing atomic bonds [8]. In contrast, calculations for aluminum using quantum mechanical density-functional theory [9, 10] show that di-vacancies – a complex of two vacancies – are either energetically unfavorable if they are aligned along the ⟨110⟩ direction or barely favorable with negligible binding energy if aligned along ⟨100⟩. If two vacancies can barely bind, it seems doubtful that they can be stable and grow to form clusters that can turn into prismatic loops. While these DFT methods are far more accurate, the computational effort is extremely large and consequently these calculations were limited to less than a hundred atoms. This corresponds to an unphysically high concentration of vacancies. Furthermore, the results are in variance with experiments [11, 12] that are indicative of a high binding energy of di-vacancies and a significant concentration of di-vacancies, especially at elevated temperatures.

We study vacancy clustering and prismatic loops by performing electronic structure calculations using orbital-free density-functional theory (OFDFT) [13]. Specifically, the kinetic energy functional is modelled using the Thomas-Fermi-Weizsacker family of functionals with λ = 1/6. We use the modified form of Heine-Abarenkov pseudopotential [14] for aluminum to model the external field created by the nuclei and core electrons. The exchange-correlation effects are treated using a local density approximation [15, 16]. These functionals have been shown to correctly predict the bulk and vacancy properties of aluminum [17, 18].

A challenge in studying defects in solids, and especially vacancies, is their extremely small concentrations which is typically a few parts per million in aluminum [19]. Therefore any realistic calculation of vacancies and their interaction has to involve millions of atoms. Unfortunately, performing electronic structure calculations with such numbers of atoms remained beyond reach till the recent development of the quasi-continuum orbital-free density-functional method (QC-OFDFT) [18]. This method has enabled for the first time the calculation of the electronic structure using OFDFT of samples with millions of atoms subjected to arbitrary boundary conditions. Importantly, the method is completely seamless, does not require any ad hoc assumptions, uses OFDFT as its only input and enables convergence studies of its accuracy.

The independent unknowns of the QC-OFDFT analysis are the nuclear positions, and the electronic fields comprising of electron-density and electrostatic potential. As in the conventional QC approach, the nuclei positions are interpolated from the positions of representative nuclei, Figure 1(a). Near the defect core, all nu-
Finally, the Cauchy-Born predictor is sampled on an auxiliary mesh as the coarse mesh. The electronic fields exhibit subatomic oscillations and require a fine mesh to accurately capture these oscillations. In regions away from the defect cores these oscillations are locally periodic on the length scale of the lattice. Hence, a first guess to the electronic fields is computed from a periodic unit cell calculation using the underlying Cauchy-Born lattice deformation. We refer to this guess as the Cauchy-Born predictor. Though the Cauchy-Born predictor is an accurate representation of the electronic fields in regions far away from the defect, they greatly differ close to the defect cores. Thus, the electronic fields are represented as a sum of the Cauchy-Born predictor and corrector fields. The corrector fields are computed on a mesh that exhibits subatomic resolution near the defect core, where the electronic fields differ greatly from the Cauchy-Born predictor, and coarsens away from the defect core, where the corrector fields exhibit slow variation on the scale of the lattice, Figure 1(b). We refer to this computational mesh as the intermediate mesh. Finally, the Cauchy-Born predictor is sampled on an auxiliary mesh that resolves an integration lattice unit cell in each finite element with exceedingly fine resolution, Figure 1(c). This auxiliary mesh is introduced for purposes of representing the Cauchy-Born predictor and does not introduce any degrees of freedom into the calculation. The degrees of freedom of the QC-OFDFT analysis are the corrector fields on the intermediate mesh and the positions of representative atoms on the coarse mesh which are computed from a variational principle. These fields completely describe the electronic structure and the relaxed atomic structure of the material system.

In a recent work, we used QC-OFDFT to study di-vacancies in aluminum [18], where we found a very strong cell-size (concentration) effect. Specifically, we found that (110) di-vacancies were repulsive for small cell-sizes, in agreement with previous calculations [9, 10], and the same di-vacancies were attractive for larger cell-sizes corresponding to realistic concentrations, with binding energies of 0.19 eV in agreement with experimental measurements [11, 12]. This work, showed that electronic structure calculations do not rule out vacancy clustering in aluminum. Therefore, we examine this mechanism further in the current work.

We begin by examining the binding energies of various quad-vacancies formed from a pair of di-vacancies. The number of possible quad-vacancies that may be formed from a pair of di-vacancies is very large. Thus we restrict our analysis to configurations such that each vacancy has at least two other vacancies as nearest or second nearest neighbors. We shall justify this choice subsequently. This criterion results in 9 distinct configurations (up to symmetry), 6 of which are planar vacancy clusters and 3 of which are non-planar. These configurations are listed in Table I.

The vacancy cluster binding energy of a \( n \)-vacancy cluster is defined as
\[
E^{\text{bind}}_{nv} = nE_v^f - E_{nv}^f,
\]
where \( E_v^f \) denotes the formation energy of a single vacancy and \( E_{nv}^f \) the formation energy of the \( n \)-vacancy cluster. The vacancy cluster binding energies for the 9 configurations of quad-vacancies are tabulated in Table I. Figure 2(a) shows the contours of electron-density for the quad-vacancy cluster with the highest binding energy. This corresponds to configuration No. 5 in Table I, which denotes a planar quad-vacancy on (110) plane. Binding energies of each of these vacancy clusters listed in Table I are computed using a computational cell consisting of a million atoms. This corresponds to realistic vacancy concentrations of a few parts per million [19]. The boundary conditions for all simulations are chosen such that the electronic fields decay to bulk values on the boundaries of the sample. Numerical parameters were chosen to keep the error in the formation energy due to discretization and coarse-graining to be less than 0.01 eV.

It is interesting to observe that all the quad-vacancies considered have positive binding energies. Further, they also have binding energies larger than twice the computed di-vacancy binding energy of 0.19 and 0.23 eV for (110) and (100) di-vacancies respectively. This indicates that pairs of di-vacancies are attractive in all cases. These results suggest that quad-vacancy formation is an energetically feasible process and that vacancies prefer to condense rather than split into mono or di-vacancies. This observation also justifies our restriction to 9 quad-vacancy configurations.

The cell-size used to simulate defects effectively sets the concentration of the defects. To understand the effect of vacancy concentration on the feasibility of vacancy clustering, we study the cell-size effect on quad-vacancy binding energy for the first configuration in Table I. This configuration represents a square shaped quad-vacancy on (100) plane, whose electronic structure is shown in Figure 2(b). Figure 3 shows a strong dependence of the vacancy binding energy on the cell-size. The quad-vacancy which is energetically favorable for large cell-sizes becomes unstable for small cell-sizes. This a result of the long-ranged elastic and electrostatic effects in the presence of these defects. This cell-size dependence shows that vacancy clustering which is feasible at low and realistic vacancy concentrations becomes unfavorable at high concentrations.

The results in Table I also show that the configurations with the highest binding energy (No. 5 and 6) are planar quad-vacancy clusters on {110} and {111} planes. Therefore, we performed simulations on larger vacancy clusters on {110} and {111} planes, again using cell-sizes with a million atoms. On the {111} plane, we studied an
FIG. 1: (a) Coarse computational mesh used to interpolate nuclei positions away from the fully-resolved defect core. (b) Intermediate computational mesh used to represent the corrector fields. It has subatomic resolution in the defect core, and coarsens away from the defect core. (c) Fine auxiliary mesh used to sample the Cauchy-Born predictor fields within an integration lattice unit cell, $D_e$, in each element $e$.

TABLE I: Vacancy cluster binding energies for quad-vacancies formed from a pair of di-vacancies. All possible quad-vacancies such that each vacancy has two other vacancies as nearest or second nearest neighbors are considered.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Positions of vacancies</th>
<th>Vacancy binding energy (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>planar {100} {(0,0,0), (a/2,a/2,0), (a,0,0), (a/2,-a/2,0)}</td>
<td>0.52</td>
</tr>
<tr>
<td>2</td>
<td>planar {100} {(0,0,0), (a/2,a/2,0), (a,0,0), (3a/2,a/2,0)}</td>
<td>0.50</td>
</tr>
<tr>
<td>3</td>
<td>planar {100} {(0,0,0), (a/2,a/2,0), (a,0,0), (a,a,0)}</td>
<td>0.48</td>
</tr>
<tr>
<td>4</td>
<td>planar {100} {(0,0,0), (a,0,0), (0,a,0), (a,a,0)}</td>
<td>0.48</td>
</tr>
<tr>
<td>5</td>
<td>planar {110} {(0,0,0), (0,a/2,a/2), (a,0,0), (a,a/2,0)}</td>
<td>0.56</td>
</tr>
<tr>
<td>6</td>
<td>planar {111} {(0,0,0), (0,a/2,a/2), (a/2,a/2,0), (a/2,a,a/2)}</td>
<td>0.55</td>
</tr>
<tr>
<td>7</td>
<td>non-planar {(0,0,0), (0,a/2,a/2), (a/2,0,a/2), (a/2,a/2,0)}</td>
<td>0.53</td>
</tr>
<tr>
<td>8</td>
<td>non-planar {(0,0,0), (a,0,0), (a/2,a/2,0), (a/2,0,a/2)}</td>
<td>0.51</td>
</tr>
<tr>
<td>9</td>
<td>non-planar {(0,0,0), (a,0,0), (a/2,a/2,0), (0,a/2,a/2)}</td>
<td>0.50</td>
</tr>
</tbody>
</table>

FIG. 2: (a) Contours of electron-density around a planar quad-vacancy (Configuration No. 5 in Table 1) on (110) plane in a million atom sample. This planar quad-vacancy has the highest binding energy among the various quad-vacancies considered. (b) Contours of electron-density around a planar quad-vacancy (Configuration No. 1 in Table 1) on (100) plane in a million atom sample.

hexagonal cluster with 7 vacancies, and found two stable configurations. One of the stable configuration is a non-collapsed state with a vacancy cluster binding energy of 0.88 eV, with a maximum displacement of atoms of the order of 3.2% of the nearest neighbor distance. Note that this is larger than 7/2 times the di-vacancy binding energy (0.19 or 0.23 eV depending on orientation). This means that the hexagonal cluster is stable against dissociation into di-vacancies.

FIG. 3: Cell-size dependence of vacancy binding energy for the quad-vacancy given by the first configuration in Table I.
The second configuration is a prismatic loop where the atoms above and below the hexagonal vacancy disc collapse or move towards each other leaving a dislocation line at the boundary of the disc. Figure 4 shows the atomic positions and the contours of the electron-density on (001) and (111) planes of the collapsed prismatic loop. In particular, the dotted lines in Figure 4(a) depict the collapse of the planes resulting in the prismatic dislocation loop. The maximum displacement of atoms is around 44% of the nearest neighbor distance, the Burgers vector is $0.44[110]$ and the dislocation plane is (111). These results are consistent with experiments [20, 21]. Using transmission electron microscope (TEM), it was observed that prismatic loops form predominantly on a {111} plane with a 1/2{110} Burgers vector. Further, in these experiments, prismatic loops whose size is as small as 50 Å in diameter were observed. While these are larger than our hexagonal prismatic loop formed from 7 vacancies, it is impossible to detect a loop as it nucleates. Thus, the nucleation size of a prismatic loop was hitherto unknown. The computed vacancy cluster binding energy for the prismatic loop is 1.55 eV which means that not only is this structure stable against dissociation of di-vacancies, but that it is even more stable than the uncollapsed configuration.

On the (110) plane, we studied rectangular vacancy clusters with 6 and 9 vacancies. The computed binding energies for these vacancy clusters are 0.81 eV and 1.16 eV respectively. The maximum displacement of atoms in these vacancy clusters is around 4% of the nearest neighbor distance. These clusters did not display any bi-stability and collapse to prismatic loops.

These results point to four important facts. Firstly, the binding energy of vacancy clusters on {110} and {111} planes in aluminum increases with the size of the vacancy cluster. Also, these vacancy clusters are all stable, i.e., vacancies prefer to condense rather than split into mono or di-vacancies. To the best of our knowledge, this is the first numerical confirmation from an electronic structure perspective that vacancy clustering is energetically favorable. Secondly, we observe from direct numerical simulation that the hexagonal vacancy cluster on (111) plane collapses to form a prismatic loop. This establishes from electronic structure calculations that vacancy clustering and collapse of the planes surrounding the vacancy cluster is a possible mechanism for the nucleation of prismatic dislocation loops. Thirdly, our results point to the fact that vacancy clusters as small as 7 vacancies can collapse to form stable prismatic loops on {111} planes. Finally, our results show the importance of studying defects in solids at realistic concentrations.

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