Combined effect of Mg and vacancy on the generalized planar fault energy of Al

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By first-principles calculations, a systematic study has been undertaken to investigate the combined effect of vacancies and Mg impurities on the generalized planar fault energy (GPFE) of pure Al. It was predicted that introduction of a single vacancy at the stacking fault plane can decrease the GPFE and enhance twinning propensity of Al. Furthermore, vacancies were shown to exhibit a natural/activated Suzuki segregation feature towards the intrinsic/extrinsic stacking fault of Al, respectively. Along with an increasing vacancy content localized near the deformation plane a noticeable decrease in the intrinsic stacking fault energy $\gamma_{\text{ISFE}}$ was obtained, which potentially may induce more split-up of dislocations into partials. However, a further increase of twinnability (based on the predicted GPFE curves) with increasing vacancy concentration was not found. More interestingly, we discovered a coupled Suzuki segregation behavior of Mg solutes and vacancies towards the intrinsic stacking fault of Al. A systematic analysis of the twinnability parameter $\tau_a$ on basis of the modified GPFE curves under the influence of Mg solutes and vacancies clearly demonstrates the general difficulty of deformation twinning of Al-Mg alloys. Moreover, a decreased $\gamma_{\text{ISFE}}$ with Mg alloying as well as with the presence of vacancies may serve, at least partially, to explain the high work-hardening rate and the formation of band structures in Al-Mg alloys processed by severe plastic deformation (SPD).

1. Introduction

The generalized planar fault energy (GPFE), representing the energy penalty of rigid crystal displacements within the fcc (111) planes along the $<11\bar{2}>$ direction, has in recent years been widely recognized as an important descriptor for the deformation mechanisms of both coarse and nano grained fcc metals [1–6]. The GPFE (or $\gamma$ surface) has four extrema: the unstable stacking fault energy $\gamma_{\text{USFE}}$, the intrinsic stacking fault energy $\gamma_{\text{ISFE}}$, the unstable twinning fault energy $\gamma_{\text{UTFE}}$ and the twinning fault energy (two-layer micro twin) energy $\gamma_{\text{TTEF}}$, where the latter is also referred to as the extrinsic stacking fault energy $\gamma_{\text{ESFE}}$ [1,7]. Due to the difficulties in accurate experimental measurement of these extrema [8], atomistic simulations have been widely employed to determine the GPFE, aiming to explore the atomistic mechanisms to properly analyze and interpret deformation behaviors in metals and alloys. A combination of GPFE and associated models can be used to predict and interpret many dislocation-related phenomena, e.g. emission of partial dislocations [19], critical twinning stress [3], Peierls stress [10], plastic deformation regimes [16], etc. For instance, following earlier works [2,11,12], a twinnability analysis based on the GPFE is possible, using the extensively used criterion $\tau_a$ which is defined as a function of the number and strength of active twinning systems in terms of $\gamma_{\text{USFE}}$, $\gamma_{\text{ISFE}}$ and $\gamma_{\text{TTEF}}$:

$$\tau_a = \left[1.136 - 0.151 \frac{\gamma_{\text{ISFE}}}{\gamma_{\text{TTEF}}} \frac{\gamma_{\text{USFE}}}{\gamma_{\text{TTEF}}} \right]$$

(1)

Herein, a larger $\tau_a$ means stronger twinning tendency. By calculating $\tau_a$ via GPFE, an increasing propensity of Cu–Al to deform preferentially by twinning with increasing Al content has been predicted [2]. Specifically, based on relevant criteria with respect to, e.g. $\gamma_{\text{USFE}}/\gamma_{\text{TTEF}}$ and $\gamma_{\text{ISFE}}/\gamma_{\text{TTEF}}$ postulated by Rice [13], Tadmor and Hai [14], the GPFE curve can be utilized to evaluate the competition between different deformation regimes like full slip and twinning. In addition, intrinsic energy barriers characterized by $\gamma_{\text{USFE}}$, $\gamma_{\text{ESFE}}/\gamma_{\text{TTEF}}$ and $\gamma_{\text{ISFE}}/\gamma_{\text{TTEF}}$ are widely utilized for analyzing the activation of stacking fault, twinning, and full slip.
Recently, Jo et al. [6] revisited the plastic deformation mechanisms of crystalline materials by using the effective energy barriers (EEBs) on basis of GPFE, which take into account the effect of shear directionality. Apart from this, a hierarchical, multi-scale theory in conjunction with GPFE was put forward by Kibey et al. [3] to predict the critical twinning stress in Al. Furthermore, as demonstrated in Refs. [10,15], a coupling of GPFE and generalized Peierls-Nabarro model is able to predict the Peierls stress, which characterizes the mobility of dislocations. All these important works have proven and established the fundamental importance of GPFE in plastic deformation of crystalline materials.

Generally, most previous works have put a focus on predicting GPFE of perfect pure metals, e.g. Al, Cu, Ni [1,5,16], or the effect of solute alloying on GPFE [4,17–19]. Nevertheless, rare attention has been expended in the influence of defects, especially vacancies, upon GPFE. Vacancies, which to some extent are always present, are crucial in many material processes and reactions at the atomistic level, i.e. diffusion [20], precipitation [21,22], segregation [23], etc. and thereby have long been known to strongly influence the properties of materials. For instance, a significant high vacancy concentration generated via severe plastic deformation (SPD) can promote grain boundary segregation of Mg solutes, thus suppressing grain growth and maintaining an ultra-fine grained (UPG) state of materials [23]. Hence, a fine control of the roles that vacancies play in these fundamental material processes can assist the engineering of special materials with desired properties.

In the course of SPD at ambient temperature, vacancies are massively created due to intensive interactions of dislocations under tremendous plastic strain [23,24]. However, to the best of the authors’ knowledge, no complete and thorough theoretical work has so far been reported to address the effect of vacancy on the GPFE of Al and its alloys except for Lu and Kaxiras [25], Asadi et al. [26], both of which have confirmed theoretically that vacancies have a decreasing effect on the ISFE in Al. Using a semi-discrete variational Peierls-Nabarro model in conjunction with GPFE determined by first-principles calculations, Lu and Kaxiras [25] predicted a lubricating effect ascribed to the introduction of vacancies on the dislocation motion of Al. These interactions are considered to have strong effects upon the properties of materials [25]. It is thereby timely as well as scientifically interesting to quantify the effect of vacancy upon the GPFE of Al and its alloys. Al-Mg alloys possessing a good compromise of properties (e.g. light weight, mechanical strength, corrosion resistance etc. [23,27]) are taken as the model system in the present work. Experimentally, Mg solutes and vacancies are reported to be strongly coupled in an Al matrix [24], thereby the combined effect of vacancies and Mg solutes upon GPFE of Al needs to be considered.

In the first part of this work, we will study the influence of a single vacancy on the GPFE of Al. Next, the focus will be put on the effects of vacancies, in terms of local concentration and local arrangement near the deformation plane, upon the GPFE curve of Al. Hereafter, the effect of Mg-vacancy pairs and complexes upon GPFE of Al is quantified. In the last part, the special deformation behaviors of Al-Mg alloys subjected to various SPD techniques, e.g. twinningality, increased work hardening rate, planar slip of dislocations, etc. are discussed in view of the calculated effects of Mg solutes, vacancies or their combination on the GPFE of Al.

2. Computational methodology

Density functional theory with plane wave basis functions as implemented in the highly-efficient Vienna ab initio simulation package (VASP) [28,29] was utilized for all the calculations in the present work. The core region were treated with the frozen-core projected augmented wave (PAW) method [30,31], and the exchange–correlation functions were described with the Generalized Gradient Approximation (GGA) of Perdew–Burke–Ernzerhof (PBE) [32]. A cutoff of 350 eV employed in the present work can yield sufficiently converged results. k-point sampling of $13 \times 7 \times 1$ based on the Monkhorst-Pack scheme [33] together with the linear tetrahedron method including Bloch corrections [34] was adopted for the integration of reciprocal-space energy in the Brillouin zone (BZ). The convergence criterion of $10^{-6}$ eV and $10^{-4}$ eV were employed for the termination of electronic self-consistency and ionic loop, respectively, in the relaxation process.

A 72-atom supercell as illustrated in Fig. 1(a) was constructed in order to find the energetically favorable distributions of Mg solutes and vacancies in the Al matrix. By putting Mg solutes and vacancies at different positions (see Fig. 1(a) and Table 1), the binding energies $E_b$ of the vacancy-vacancy and Mg-vacancy pairs at different neighboring state in the Al matrix can be evaluated in terms of the following equations,

$$E_b(X - X) = -(E(Al_{170}X_2) + E(Al_{172}) - 2E(Al_{171}X))$$  \hspace{0.5cm} (2)

$$E_b(X - Y) = -(E(Al_{170}XY) + E(Al_{172}) - E(Al_{171}X) - E(Al_{171}Y))$$  \hspace{0.5cm} (3)

where e.g. $E(Al_{172})$ is the total electronic energy of $Al_{172}$ as calculated by VASP, $X$ is a vacancy, and $Y$ is a Mg solute atom. Thus, Eq. (2) gives $E_b$ for vacancy-vacancy pairs, while Eq. (3) for Mg-vacancy pairs. A positive $E_b$ value corresponds to a favorable binding of the pairs.

Table 1 lists the predicted $E_b$ of the various pairs at different neighboring distances in the Al matrix. It is found that vacancies always like to stay as 2nd nearest neighbor (2nn) in the Al matrix. The repulsion between 1st nearest neighbor vacancies (1nn) and attractions as 2nn predicted by the present work are in good agreement with the calculations by Karling and Wahnström [35]. As for Mg-vacancy pairs, 4th nearest neighbors (4nn) are energetically more preferable. It is worth noting that 1nn Mg-vacancy pairs has an $E_b$ of $-0.0148$ eV, which is in good agreement with the experimental value of $-0.01 \pm 0.04$ eV [20] and close to the LDA-produced value of $-0.02$ eV by Wolverton [20].

In order to calculate GPFE curves, a slab model containing a perfect stacking sequence of 12 (111) planes (ABCABCABCABC) with total 96 atoms as well as a vacuum spacing of 15 Å along the (111) direction was constructed, as shown in Fig. 1(b). Through a rigid displacement of half the crystal with respect to the other along the <112> direction within (111) planes (see Fig. 1(b)), we can obtain the whole GPFE curve by calculating the energies for different configurations along the displacement path. To investigate the entire GPFE curve, two separate shear deformations of the slab model along the <112> direction have to be enforced. Note that in each operation, the final displacement distance was $a_0/\sqrt{6}$, being the Burgers vector of a partial dislocation, with $a_0$ as the lattice constant of Al. As a start, a stacking fault was created through displacing the atomic layers numbered with 1–6 in one-half of the crystal (cf. Fig. 1(b)). In a second move, based on the stacking fault configuration, layers numbered from 1 to 5 in the lower part of the crystal were shifted in the opposite direction. To attain the equilibrium state of different configurations, the atomic positions in the slab model were solely allowed to relax in the direction perpendicular to the stacking fault plane, using a conjugate-gradient relaxation algorithm. The fault energies of the different configurations along the displacement path were calculated using Eq. (4) [36,37]:

$$E(\overline{f}) = \frac{E_{\text{faulted}}(\overline{f}) - E_{\text{perfect}}}{A}$$  \hspace{0.5cm} (4)

In which, $E_{\text{faulted}}(\overline{f})$ is the total energy of the supercell deformed...
by a fault vector $\mathbf{f}$. $E_{\text{perfect}}$ corresponds to the total energy of the perfect stacking slab, and $A$ is the cross-section area of slab.

3. Results

3.1. Mono-vacancy effect on GPFE and Suzuki segregation of vacancies in Al

The effect of a vacancy upon the GPFE curve of Al was investigated through introducing a single vacancy ($\text{Al}_{95}\text{Va}_1$) in the stacking fault plane (layer 0 in Fig. 1(b)) residing in site 1 (see Fig. 1(e)). In this situation, an overall concentration of 1.08 at.% and a layer concentration of 12.5 at.% of vacancies is produced. The mono-vacancy affected $g_{\text{USFE}}$, $g_{\text{ISFE}}$, $g_{\text{UTFE}}$, and $g_{\text{TFE}}$ of Al are summarized in Table 2, and the corresponding mono-vacancy induced GPFE curve of Al is displayed in Fig. 2, in comparison with the GPFE curves of Al$_{95}$Mg$_1$ as well as pure Al [38]. One can see from Fig. 2 that mono-vacancy has a more prominent decreasing effect on $g_{\text{ISFE}}$ of Al than that of mono-Mg solute [38], i.e. from 142.4 mJ/m$^2$ by a fault vector $\mathbf{f}$. $E_{\text{perfect}}$ corresponds to the total energy of the perfect stacking slab, and $A$ is the cross-section area of slab.

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![Fig. 1](image1.png)

Fig. 1. (a) Indication of the positions for substitution of Mg and vacancy as different neighbors in Al matrix. Typical atomic configurations adopted in the present calculations including the perfect stacking (b), intrinsic stacking fault (c), and twinning fault configurations (d), a 15 Å thickness of vacuum was implemented in all the slab models. The displacement path of the GPFE along the $<11\overline{2}>$ direction is also illustrated. (e) Atomic positions (indicated as 1−12) for possible Mg or vacancy substitutions to evaluate its effect on the GPFE of Al.

![Fig. 2](image2.png)

Fig. 2. Mono-vacancy effect on the GPFE curves of Al, in comparison with the GPFE curves of Al$_{95}$Mg$_1$ as well as pure Al [38]. One can see from Fig. 2 that mono-vacancy has a more prominent decreasing effect on $\gamma_{\text{ISFE}}$ of Al than that of mono-Mg solute [38], i.e. from 142.4 mJ/m$^2$

<table>
<thead>
<tr>
<th>Occupations</th>
<th>Neighbors</th>
<th>Va-Va</th>
<th>Mg-Va</th>
</tr>
</thead>
<tbody>
<tr>
<td>1−2</td>
<td>1nn</td>
<td>−0.0642</td>
<td>−0.0148</td>
</tr>
<tr>
<td>1−3</td>
<td>2nn</td>
<td>0.0173</td>
<td>0.0031</td>
</tr>
<tr>
<td>1−4</td>
<td>3nn</td>
<td>0.0049</td>
<td>−0.0040</td>
</tr>
<tr>
<td>1−5</td>
<td>4nn</td>
<td>0.0159</td>
<td>0.0187</td>
</tr>
<tr>
<td>1−6</td>
<td>5nn</td>
<td>0.0038</td>
<td>−0.0103</td>
</tr>
</tbody>
</table>

Table 1 Binding energy $E_b$ (eV) for the Mg-Mg, vacancy-vacancy, and Mg-vacancy pairs as function of their separation distance in the Al matrix. A positive value indicates a favorable binding state. The occupations of Mg or vacancy in the Al matrix as different neighbors are as indicated in Fig. 1(a).

Table 2 Mono-Mg and mono-vacancy effect on unstable stacking fault energy $\gamma_{\text{USFE}}$, stacking fault energy $\gamma_{\text{ISFE}}$, unstable twinning fault energy $\gamma_{\text{UTFE}}$ and twinning fault energy $\gamma_{\text{TFE}}$ of Al. The energies are in mJ/m$^2$.

<table>
<thead>
<tr>
<th>System</th>
<th>$\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{ISFE}}$</th>
<th>$\gamma_{\text{UTFE}}$</th>
<th>$\gamma_{\text{TFE}}$</th>
<th>$\gamma_{\text{ISFE}}/\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{UTFE}}/\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{UTFE}}-\gamma_{\text{USFE}}$</th>
<th>$S_t$ (twinning)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Al}_{96}$</td>
<td>177.4</td>
<td>142.4</td>
<td>226.5</td>
<td>135.4</td>
<td>0.803</td>
<td>1.277</td>
<td>49.1</td>
<td>0.898</td>
</tr>
<tr>
<td>$\text{Al}_{95}\text{Mg}_1$</td>
<td>168.0</td>
<td>135.0</td>
<td>211.6</td>
<td>119.1</td>
<td>0.804</td>
<td>1.260</td>
<td>43.6</td>
<td>0.904</td>
</tr>
<tr>
<td>$\text{Al}_{95}\text{Va}_1$</td>
<td>156.2</td>
<td>121.3</td>
<td>196.0</td>
<td>102.5</td>
<td>0.777</td>
<td>1.255</td>
<td>39.8</td>
<td>0.909</td>
</tr>
</tbody>
</table>

* Data from Ref. [38].
to 121.3 mJ/m$^2$, as will do for the other three extremum energy points. The vacancy mediated decrease of $\gamma_{SSF}$ in Al was also demonstrated in the theoretical works by Lu and Kaxiras [25] and Asadi et al. [26], but neither of them investigated an entire vacancy-influenced GPFE curve. Based on the analysis of GPFE curves, $\mathrm{Al}_{59}\mathrm{Va}_1$ was calculated to exhibit a lowered $\gamma_{SSF}/\gamma_{USFE}$, i.e. 0.777 (see Table 2), relative to pure Al, indicating that vacancies may make the emission of partial dislocations easier. Mg ($\mathrm{Al}_{59}\mathrm{Mg}_1$) has previously been predicted to facilitate twinning due to an increased $\tau_a$ and decreased $\gamma_{SSF}/\gamma_{USFE}$, compared with pure Al [38]. In this work, we find that the introduction of vacancies can also increase the twinnability $\tau_a$ of Al ($\tau_a = 0.909$), and this promoting role is even higher than that of Mg solutes ($\mathrm{Al}_{59}\mathrm{Mg}_1$, $\tau_a = 0.904$).

Suzuki segregation is an interesting phenomenon describing that impurities or vacancies are prone to segregate to stacking faults. Most previous work has focused on the Suzuki interaction between impurities and stacking fault [39,40]. However, the Suzuki segregation of vacancies have been scarcely investigated apart from Ref. [41], in which a positive binding energy between vacancies and fault plane and has a spatial distribution in the vicinity of the fault plane.

Spatially distributed vacancies to the stacking fault; a negative value denotes the tendency for vacancies to segregate toward the stacking fault. Calculations reveal that $E_{\text{int}}$ is not restricted to the stacking fault plane and has a spatial distribution in the vicinity of the fault plane. Spatially distributed $E_{\text{int}}$ can produce a spatial concentration distribution of vacancies near the stacking fault. This layer-by-layer concentration profile of vacancies at finite temperature $T$ can be calculated by employing the following equation [42]:

$$c(n) = \frac{1}{1 + \frac{1}{c_0} \exp \left( \frac{E_{\text{int}}}{k_B T} \right)}$$  \hspace{1cm} (6)

where $c_0$ is the nominal overall concentration of vacancies, $T$ is the temperature, and $k_B$ is the Boltzmann constant.

A quantified segregation of vacancies towards intrinsic and extrinsic stacking faults in Al based on Eq. (5) is presented in Fig. 3(a). The intrinsic stacking fault planes correspond to the layers labelled “0” and “1” in Fig. 1(c). It is observed in Fig. 3(a) that, similar to the Suzuki segregation of Mg in Al [38], $E_{\text{int}}$ is noticeable only at a few atomic layers in the vicinity of the stacking fault plane and then diminishes at larger distances away from the fault plane. As indicated in Fig. 3(a), the effect of an intrinsic (extrinsic) stacking fault on a vacancy only extends two (three) layers. The negative $E_{\text{int}}$ near the stacking fault suggests the Suzuki segregation of vacancies either intrinsic or extrinsic. One can further find that the interaction between a vacancy and the extrinsic stacking fault is stronger than that with the intrinsic counterpart, which indicates that the vacancy is more effective in modifying the energy of extrinsic stacking faults. In spite of Suzuki segregation in Al, vacancies would behave differently in the process of approaching the fault region of intrinsic and extrinsic stacking faults. As shown in Fig. 3(a), for a vacancy getting near to an intrinsic stacking fault, energy barrier barely exists. This is consistent with the predictions by Ref. [41], which is unlike the slightly activated Suzuki segregation of Mg in Al predicted in a previous work [38]. However, in the case of approaching the extrinsic stacking fault region, vacancies would exhibit activated Suzuki segregation behavior due to the two large energy barriers on both sides of the stacking fault (layers −1 and 1 in Fig. 3(a)). We thereby conclude that the kinetic barrier is higher for vacancies to diffuse toward extrinsic stacking faults despite their higher thermodynamic stability.

The corresponding spatial concentration profile of vacancies in the vicinity of a stacking faults which was evaluated using Eq. (6) at room temperature $T = 300$ K is displayed in Fig. 3(b). A local enrichment of vacancies can be observed both for intrinsic and extrinsic stacking faults. Due to a stronger interaction energy $E_{\text{int}}^i$, the highest vacancy concentration at the extrinsic stacking fault plane is much higher than that of the intrinsic counterpart. Interestingly, it is found that the spatial vacancy concentration near the intrinsic stacking fault exhibits a Gaussian-like distribution (cf. Fig. 3(b)), while the distribution is oscillatory around the extrinsic stacking fault. Furthermore, we should keep in mind that Eq. (6) is only valid at low $c_0$, since interactions between vacancies would become important and shall take a decisive role in controlling the local distributions near the stacking fault defect.

3.2. Vacancy concentration effect on GPFE curve of Al

As noted earlier, it is energetically favorable for vacancies to stay as 2nn in Al. Therefore, to study the effect of varying vacancy concentration on GPFE in Al, we have created models with vacancies symmetrically distributed as 2nn in the vicinity of the stacking fault plane. A variety of models with 3($\mathrm{Al}_{39}\mathrm{Va}_3-1$, $\mathrm{Al}_{39}\mathrm{Va}_3-2$) and 5($\mathrm{Al}_{39}\mathrm{Va}_5-1$, $\mathrm{Al}_{39}\mathrm{Va}_5-2$) vacancies were considered, corresponding to an overall concentration of 3.1 at.% and 5.2 at.%. This is much higher than the typical equilibrium vacancy concentration in bulk materials, but can be used to evaluate the effect of different vacancy concentrations on the GPFE of Al. The detailed distribution of vacancies for these models can be found in Table 3 (Column of “Occupation”) and Fig. 1(e).

Fig. 4 shows the calculated GPFE curves of these models, and the corresponding GPFE data are collected in Table 3. We first note that the vacancy concentration effect on GPFE is strongly dependent on the local configurations and concentrations of vacancies in the stacking fault region. As discussed above, a single vacancy residing in the stacking fault plane can enhance twinning propensity of Al. However, no increased twinnability can be observed for the models with 3 or 5 vacancies, due to the reduced $\tau_a$ compared to that of Al (cf. Table 3). $\mathrm{Al}_{39}\mathrm{Va}_3-1$ and $\mathrm{Al}_{39}\mathrm{Va}_5-1$ have relatively lower GPFE than $\mathrm{Al}_{39}\mathrm{Va}_3-2$ and $\mathrm{Al}_{39}\mathrm{Va}_5-2$, which can be ascribed to the co-plane distribution of vacancies near the deformation plane in $\mathrm{Al}_{39}\mathrm{Va}_3-1$ and $\mathrm{Al}_{39}\mathrm{Va}_5-1$. The evolution of GPFE versus vacancy concentration are displayed in Fig. 5(a) and (b). As can be seen, the increase of vacancy concentration leads to a sustained decrease of $\gamma_{SSF}$, with accordingly $\mathrm{Al}_{39}\mathrm{Va}_5-1$ has the lowest $\gamma_{SSF}$, i.e. 125.8 mJ/m$^2$, which may facilitate the nucleation of leading partial dislocations and thus enhance plasticity. A maximum decrease of $\gamma_{SSF}$ down to 105.6 mJ/m$^2$ was obtained for $\mathrm{Al}_{39}\mathrm{Va}_3-1$. No further decrease of $\gamma_{SSF}$ was observed with an increasing vacancy concentration from 3.1 at.% to 5.2 at.% (as shown in Fig. 3(b)). A lowered $\gamma_{SSF}$ would increase the width of the stacking fault ribbon...
connecting the dissociated partial dislocations, leading to the more splitting-up of full dislocations into partials. The modified separation distance between the partials induced by vacancies calls for some caution when using transmission electron microscopy data to predict $\gamma_{\text{USFE}}$, since the accidental introduction of vacancies due to Suzuki segregation could result in inaccurate results.

### 3.3. The effect of combining Mg solutes and vacancies on the GPFE curves of Al

As discussed in the Introduction, during the course of severe plastic deformation, massive defects, especially vacancies, would be produced, which are generally coexisting with Mg solutes in Al-Mg alloys. The interaction between Mg solutes and vacancies might affect the GPFE curves of Al, motivating the present investigation of the combined effect of Mg solutes and vacancies on the GPFE of Al. To address this problem more systematically, we distinguish between the effect of Mg-vacancy pairs and Mg-vacancy complexes.

#### 3.3.1. Mg-vacancy pairs

Mg-vacancy pairs were first introduced in the stacking fault plane with different neighboring distance between Mg and vacancy: 1nn (Al$_{94}$Mg$_1$Va$_1$-1nn), 3nn (Al$_{94}$Mg$_1$Va$_1$-3nn), and 4nn (Al$_{94}$Mg$_1$Va$_1$-4nn). The resulting GPFEs are listed in Table 4 and the corresponding GPFE curves are presented in Fig. 6.

#### Table 3

<table>
<thead>
<tr>
<th>System</th>
<th>Occupation</th>
<th>$\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{ISFE}}$</th>
<th>$\gamma_{\text{UTFE}}$</th>
<th>$\gamma_{\text{TFE}}$</th>
<th>$\gamma_{\text{ISFE}}/\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{UTFE}}/\gamma_{\text{USFE}}$</th>
<th>$\gamma_{\text{UTFE}}-\gamma_{\text{USFE}}$</th>
<th>$t_a$ (twinnability)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$_{96}$</td>
<td></td>
<td>177.4</td>
<td>142.4</td>
<td>226.5</td>
<td>135.4</td>
<td>0.803</td>
<td>1.277</td>
<td>49.1</td>
<td>0.898</td>
</tr>
<tr>
<td>Al$_{93}$Va$_3$-1</td>
<td>Va$_1$(1,5,7)</td>
<td>131.0</td>
<td>105.6</td>
<td>171.3</td>
<td>106.3</td>
<td>0.806</td>
<td>1.307</td>
<td>40.3</td>
<td>0.887</td>
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<td>Al$_{93}$Va$_3$-2</td>
<td>Va$_1$(1,8,6)</td>
<td>140.1</td>
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<td>208.2</td>
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<td>Al$_{91}$Va$_5$-1</td>
<td>Va$_1$(1,5,7,9,11)</td>
<td>125.8</td>
<td>109.8</td>
<td>172.0</td>
<td>107.4</td>
<td>0.821</td>
<td>1.367</td>
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</tr>
<tr>
<td>Al$_{91}$Va$_5$-2</td>
<td>Va$_1$(1,5,7,10,12)</td>
<td>152.9</td>
<td>125.5</td>
<td>196.4</td>
<td>117.1</td>
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</table>

Fig. 3. (a) The interaction energy $E_{\text{int}}^n$ (Eq. (5)) between single vacancy and intrinsic (dotted circles) or extrinsic (open circles) stacking faults in Al, as a function of distance to the stacking fault, in number of atomic layers $n$ (Fig. 1(c) and (d)). (b) The corresponding concentration profile of vacancies $c_{Va}$ evaluated from Eq. (6) at temperature $T = 300$ K.

Fig. 4. Vacancy concentration effect on the GPFE curves of Al. For the detailed vacancy distributions in the vicinity of the fault plane, please refer to Table 3.

As an example, the reduction in $\gamma_{\text{USFE}}$ from pairs is more significant than that of separate Mg solutes and vacancies. Besides, a correlation between the interaction energy of Mg-vacancy pairs and their effect on the GPFE of Al can be derived by comparing Tables 4 and 1; the shorter distance between Mg and vacancy, the larger reduction of the GPFE of Al. The 4nn Mg-vacancy pair with largest positive binding energy displays the smallest reduction in GPFE of Al. The correlation between the interaction energy of Mg-vacancy pairs...
and $\tau_a$ suggests that the more positive the binding energy in Mg-vacancy pairs, the higher propensity for twinning.

### 3.3.2. Mg-vacancy complexes

The Mg-vacancy complex effect on the GPFE of Al was investigated with multi Mg and multi vacancies as ordered in the vicinity of stacking fault plane. The construction of energetically favorable Mg-vacancy complexes needs simultaneous consideration of vacancy-vacancy, Mg-vacancy interactions. Coplanar vacancy-Mg-vacancy and Mg-vacancy-Mg triplets not being in the fault plane are one kind of Mg-vacancy complex that affects the GPFE of Al. Four models with notation S1, S2, S3, and S4 were constructed (see the detailed Mg and vacancy site occupations in Table 5) to quantify the triplet effects. Fig. 7(a) shows the GPFE curves under the effect of vacancy-Mg-vacancy and Mg-vacancy-Mg triplets, and the corresponding GPFE values are collected in Table 5. Clearly, both vacancy-Mg-vacancy (S1, S2) and Mg-vacancy-Mg (S3, S4) triplets lower the GPFE of Al (cf. Fig. 7(a)). In comparison with Al$_{95}$Mg$_1$ [38], S1 and S2 have further reduced GPFE, showing that the coupled segregation effect of Mg atoms and vacancies to the fault plane is not surprising since both Mg and vacancies exhibit a Suzuki segregation feature as predicted previously. In addition, Mg atoms have a larger atomic radius than Al. Mg residing in the stacking fault plane would create lattice distortions which would drive vacancies to segregate to the stacking fault plane so as to lower the elastic strain energy. On the other hand, lattice contraction created by vacancy segregation at the stacking fault would attract the Mg atoms to diffuse to the stacking fault. The further reduction of $\gamma_{ISFE}$ induced by Mg-vacancy pairs indicates that a coupled segregation of Mg and vacancy to stacking fault is more energetically favorable than having only Mg or vacancy segregation along the stacking fault.

### Table 4

The effect of Mg and vacancy pairs on unstable stacking fault energy $\gamma_{USFE}$, stacking fault energy $\gamma_{ISFE}$, unstable twinning fault energy $\gamma_{UTFE}$ and twinning fault energy $\gamma_{TFE}$ of Al. Both Mg and vacancy are placed in the stacking fault plane. The subscript number in the "Occupation" column corresponds to the atomic positions as indicated in Fig. 1(e) for possible Mg or vacancy substitutions. The energies are in mJ/m$^2$.

<table>
<thead>
<tr>
<th>System</th>
<th>Occupation</th>
<th>$\gamma_{USFE}$</th>
<th>$\gamma_{ISFE}$</th>
<th>$\gamma_{UTFE}$</th>
<th>$\gamma_{ISFE}/\gamma_{USFE}$</th>
<th>$\gamma_{UTFE}/\gamma_{USFE}$</th>
<th>$\gamma_{TFE}/\gamma_{USFE}$</th>
<th>$\tau_a$ (twinnability)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$_{96}$</td>
<td>–</td>
<td>177.4</td>
<td>142.4</td>
<td>226.5</td>
<td>135.4</td>
<td>0.803</td>
<td>1.277</td>
<td>49.1</td>
</tr>
<tr>
<td>Al$_{94}$Mg$_1$Va$_1$-1nn</td>
<td>Mg$_1$Va$_2$</td>
<td>135.0</td>
<td>107.6</td>
<td>174.4</td>
<td>91.9</td>
<td>0.797</td>
<td>1.292</td>
<td>39.5</td>
</tr>
<tr>
<td>Al$_{94}$Mg$_1$Va$_1$-3nn</td>
<td>Mg$_1$Va$_3$</td>
<td>148.7</td>
<td>116.2</td>
<td>184.8</td>
<td>92.1</td>
<td>0.782</td>
<td>1.243</td>
<td>36.1</td>
</tr>
<tr>
<td>Al$_{94}$Mg$_1$Va$_1$-4nn</td>
<td>Mg$_1$Va$_4$</td>
<td>153.8</td>
<td>119.5</td>
<td>189.2</td>
<td>92.3</td>
<td>0.777</td>
<td>1.230</td>
<td>35.4</td>
</tr>
</tbody>
</table>
Table 5
Mg and vacancy complex effect on unstable stacking fault energy $\gamma_{USFE}$, stacking fault energy $\gamma_{ISFE}$, unstable twinning fault energy $\gamma_{UTFE}$ and twinning fault energy $\gamma_{TFE}$ of Al. The energies are in mJ/m$^2$. The subscript number in the “Occupation” column corresponds to the atomic positions as indicated in Fig. 1(e) for possible Mg or vacancy substitutions.

<table>
<thead>
<tr>
<th>System</th>
<th>Occupation</th>
<th>$\gamma_{USFE}$</th>
<th>$\gamma_{USFE}$</th>
<th>$\gamma_{ISFE}$</th>
<th>$\gamma_{ISFE}$</th>
<th>$\gamma_{UTFE}$</th>
<th>$\gamma_{UTFE}$</th>
<th>$\gamma_{TFE}$</th>
<th>$\gamma_{TFE}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Al_{96}$</td>
<td>–</td>
<td>177.4</td>
<td>142.4</td>
<td>226.5</td>
<td>135.4</td>
<td>0.803</td>
<td>1.277</td>
<td>49.1</td>
<td>0.898</td>
</tr>
<tr>
<td>S1</td>
<td>Mg(1)Va(9,11)</td>
<td>160.1</td>
<td>126.7</td>
<td>203.6</td>
<td>108.1</td>
<td>0.791</td>
<td>1.272</td>
<td>43.5</td>
<td>0.901</td>
</tr>
<tr>
<td>S2</td>
<td>Mg(1)Va(10,12)</td>
<td>153.9</td>
<td>125.0</td>
<td>193.7</td>
<td>101.9</td>
<td>0.812</td>
<td>1.258</td>
<td>39.8</td>
<td>0.903</td>
</tr>
<tr>
<td>S3</td>
<td>Mg(9,11)Va(1)</td>
<td>156.3</td>
<td>124.9</td>
<td>202.1</td>
<td>106.1</td>
<td>0.799</td>
<td>1.293</td>
<td>45.8</td>
<td>0.893</td>
</tr>
<tr>
<td>S4</td>
<td>Mg(10,12)Va(1)</td>
<td>153.1</td>
<td>123.6</td>
<td>193.9</td>
<td>97.3</td>
<td>0.807</td>
<td>1.267</td>
<td>40.9</td>
<td>0.901</td>
</tr>
<tr>
<td>S5</td>
<td>Mg(9,10,11,12)Va(1)</td>
<td>155.7</td>
<td>128.9</td>
<td>206.7</td>
<td>108.3</td>
<td>0.828</td>
<td>1.328</td>
<td>51.0</td>
<td>0.877</td>
</tr>
<tr>
<td>S6</td>
<td>Mg(1)Va(9,11)</td>
<td>158.5</td>
<td>131.4</td>
<td>204.1</td>
<td>113.9</td>
<td>0.829</td>
<td>1.288</td>
<td>45.6</td>
<td>0.891</td>
</tr>
<tr>
<td>S7</td>
<td>Mg(1,4,10,12)Va(9,11)</td>
<td>166.3</td>
<td>140.8</td>
<td>221.0</td>
<td>120.9</td>
<td>0.846</td>
<td>1.329</td>
<td>54.7</td>
<td>0.875</td>
</tr>
<tr>
<td>S8</td>
<td>Mg(1,9,11)Va(4)</td>
<td>152.0</td>
<td>125.4</td>
<td>193.1</td>
<td>95.7</td>
<td>0.825</td>
<td>1.270</td>
<td>41.1</td>
<td>0.898</td>
</tr>
<tr>
<td>S9</td>
<td>Mg(1,10,12)Va(4)</td>
<td>157.8</td>
<td>125.0</td>
<td>201.3</td>
<td>102.4</td>
<td>0.792</td>
<td>1.276</td>
<td>43.6</td>
<td>0.900</td>
</tr>
<tr>
<td>S10</td>
<td>Mg(1,9,10,11,12)Va(4)</td>
<td>157.5</td>
<td>130.6</td>
<td>212.0</td>
<td>112.2</td>
<td>0.829</td>
<td>1.346</td>
<td>54.5</td>
<td>0.871</td>
</tr>
<tr>
<td>S11</td>
<td>Mg(1,9,11)Va(4,10,12)</td>
<td>138.0</td>
<td>113.9</td>
<td>195.2</td>
<td>105.3</td>
<td>0.825</td>
<td>1.415</td>
<td>57.2</td>
<td>0.850</td>
</tr>
<tr>
<td>S12</td>
<td>Mg(1,10,11)Va(4,10,12)</td>
<td>155.7</td>
<td>131.6</td>
<td>200.8</td>
<td>101.7</td>
<td>0.845</td>
<td>1.290</td>
<td>45.1</td>
<td>0.888</td>
</tr>
<tr>
<td>S13</td>
<td>Mg(1,10,12)Va(9,11)</td>
<td>165.1</td>
<td>132.0</td>
<td>212.7</td>
<td>107.1</td>
<td>0.800</td>
<td>1.288</td>
<td>47.5</td>
<td>0.895</td>
</tr>
<tr>
<td>S14</td>
<td>Mg(10,12)Va(1,9,11)</td>
<td>143.9</td>
<td>112.8</td>
<td>198.1</td>
<td>96.5</td>
<td>0.784</td>
<td>1.377</td>
<td>54.3</td>
<td>0.867</td>
</tr>
</tbody>
</table>

Fig. 7. Mg and vacancy complex effect on the GPFE curves of Al. The subscript numbers in each model are the atomic positions as indicated in Fig. 1(e) for possible Mg or vacancy substitutions.
introduction of the two extra vacancies is favorable for GPFE reduction. It is, however, a different case for S3 and S4. Compared with Al$_8$V$_4$, an introduction of two extra Mg atoms raises the GPFE of S3, while an increased $\gamma_{\text{USFE}}$ and reduced $\gamma_{\text{ISFE}}$, $\gamma_{\text{UTFE}}$, $\gamma_{\text{TFE}}$ were observed for S4. Besides, the calculated $\gamma_{\text{ISFE}}/\gamma_{\text{USFE}}$, i.e. $>0.790$ and $<0.905$ of S1 to S4, imply that these coplanar vacancy-Mg-vacancy and Mg-vacancy-Mg triplets do not produce high propensity for the formation of partial dislocations and twinning.

In the present work, ten more systems noted as S5-S14, listed in Table 5, were utilized to calculate the influence of other Mg-vacancy complexes on the GPFE of Al. Fig. 7(b-d) display the Mg-vacancy complex affected GPFE curves of Al with corresponding GPFE data summarized in Table 5. As can be seen, all these Mg-vacancy complexes have a tailoring effect on the GPFE of Al, including $\gamma_{\text{USFE}}$, $\gamma_{\text{ISFE}}$, $\gamma_{\text{UTFE}}$, and $\gamma_{\text{TFE}}$. However, no increased twinnability induced by these Mg-vacancy complexes was observed as compared to pure Al except for S9, which possesses a $\tau_a$ value of 0.900. A maximum decrease of $\gamma_{\text{USFE}}$ to 112.8 mJ/m$^2$ among these complexes was predicted for S14, which has one vacancy in the fault plane, and two 4nn Mg-vacancy pairs in layer 2 and -2. S11 with three 4nn Mg-vacancy pairs positioned in layer -2, 0 and 2 produced the lowest value of $\gamma_{\text{USFE}}$, i.e. 138.0 mJ/m$^2$ and a comparably low $\gamma_{\text{ISFE}}$, i.e. 113.9 mJ/m$^2$ in the complexes. This can be attributed to the fact that a 4nn Mg-vacancy pair is effective in altering the GPFE of Al, as discussed in Section 3.3.1. It is found that for complexes with vacancy residing in the stacking fault plane, a relatively lower $\gamma_{\text{USFE}}$ can be obtained, and a higher vacancy content can result in an even more reduction of GPFE (see S11, S14 in Table 5).

4. Discussion

To visualize and facilitate the discussion of the combined effect of Mg and vacancy upon the GPFE of Al, we propose an intrinsic stacking fault energy map (presented in Fig. 8) to show the variation of twinning parameter $\tau_a$ and $\gamma_{\text{USFE}}$ of different configurations. The results of all the investigated systems in this work, including Al, Al$_6$V$_3$, Al$_5$V$_4$, Al$_{13}$Mg$_7$V$_4$, Al$_8$Mg$_3$V$_4$, and Al$_{13}$Mg$_6$V$_4$, as well as Cu, Al$_6$Mg$_1$, Al$_3$Mg$_9$, from Ref. [38] are collected in this map. Additionally, two straight dot lines marking $\tau_a$ and $\gamma_{\text{USFE}}$ of pure Al have been added in the map, aiming to make direct comparison with Al.

4.1. Twinning propensity of Al-Mg alloys during plastic deformation

The systems exhibiting an increased twinnability compared to Al are marked out with a red rectangle in Fig. 8. Obviously, only a few configurations possess a higher twinning propensity than Al. It is demonstrated that the local concentrations and distributions of Mg and vacancy near the stacking fault plane play crucial roles in determining whether it can promote twinning in Al. As indicated in Fig. 8, although all these systems can result in an even more reduction of GPFE (see S11, S14 in Table 5).

Fig. 8. Two dimensional $\tau_a$-$\gamma_{\text{USFE}}$ map of all the investigated systems in the present work in comparison with Cu. The straight dot lines representing $\tau_a$ and $\gamma_{\text{USFE}}$ of Al are marked out. The increased twinnability as well as decreased $\gamma_{\text{USFE}}$ in comparison with Al are also as indicated.

4.2. Influence of reduced $\gamma_{\text{USFE}}$ in Al-Mg alloys on recrystallization

Al-Mg alloys subjected to SPD are reported to have a high dislocation density and high work-hardening rate [23,44,45], which have been attributed to the Mg alloying [23,44,45]. It is discussed that a suppressed dynamic recovery plays an important role in maintaining the high dislocation density and high work-hardening rate of Al-Mg alloys at larger strains [45]. As a matter of fact, the dynamic recovery suppression in SPD processed Al-Mg alloys can be partially attributed to the reduction of $\gamma_{\text{USFE}}$. It is widely known that to eliminate elastic energy, full dislocations in fcc metals and alloys usually split up into two Shockley partial dislocations through the dislocation reaction $\frac{1}{2}[110] \rightarrow \frac{1}{6}[121] + \frac{1}{6}[211]$ [51], with a stacking fault ribbon connecting the two partials. The width of the stacking fault ribbon depends on $\gamma_{\text{USFE}}$ and may significantly influence the dislocation mobility. Generally, the two partial dislocations have to be joined together before a dislocation can cross-slip to an adjacent plane [52]. Hence, in low $\gamma_{\text{USFE}}$ materials possessing wider stacking fault ribbon, cross-slip of full dislocations is relatively more difficult, leading to suppressed dynamic recovery and thus high work-hardening rate during plastic deformation [16]. As indicated in by the blue rectangle in Fig. 8, one can find that Mg solutes, vacancies, as well as Mg-vacancy pairs and/or complex can all reduce the value of $\gamma_{\text{USFE}}$ of Al, which is an indication that Al-Mg...
alloys in general would exhibit a lower YSFE than pure Al, being in good agreement with experimental results [53]. During SPD processing, more vacancies and Mg-vacancy pairs/complex can be introduced in Al-Mg alloys, which would further reduce YSFE, resulting in further suppressed dynamic recovery and thus enhanced work-hardening rate.

Al-Mg alloys with a high Mg content have been reported to show planar glide deformation characteristics [54,55]. As suggested by Hong et al. [56], YSFE imposes a crucial influence upon the planarity of dislocation movement. Usually in materials with high YSFE, for instance Al, dislocations have three-dimensional mobility and can easily cross slip [57], due to the small separation distances between the partial dislocations. Extensive cross slip of dislocations would make these materials exhibit a wavy glide behavior [56], which facilitates the formation of a heterogeneous distribution of dislocations, i.e., dislocation tangling, clustering or cell structures. A decrease in YSFE can help to bind the dislocations to their respective slip planes, thus promoting planar glide of dislocations. Besides, stacking faults connecting the partials will cover a geometrical region and have repulsions between each other. The width of the stacking fault region is inversely proportional to YSFE. With a smaller YSFE, a larger stacking fault region can be expected. The short range repulsions between these stacking fault regions would restrain the clustering of dislocations [56]. Consequently, a decrease of YSFE shall enhance the extent of planar slip of dislocations, which would facilitate the formation of band structures (slip band, deformation band) in the process of deformation. Thereby the case of formation of band structures in the SPD processed Al-Mg alloys can be partially explained by a decreased YSFE resulting from Mg alloying and vacancies. However, it has to be noted that the planar slip of dislocations in the SPD processed Al-Mg alloys is not only associated with the decreased YSFE, as the reduction of YSFE predicted in the current work is still limited. Since Al-Mg alloys possess a comparable shear modulus with Al, the promotion of planar slip behavior in the SPD processed Al-Mg alloys should also be attributed to the enhanced friction stress induced by the interactions between Mg solutes and dislocations [55,58–60], as well as atomic size mismatch, which need further investigations.

5. Conclusion

A systematic first-principles study has been conducted to investigate the combined effect of vacancies and Mg solutes on the GPFE of Al. It is found that a single vacancy residing in the stacking fault plane (plane concentration 12.5 at.%) can decrease the GPFE and increase twinnability of Al. As predicted, the natural Suzuki segregation of vacancy towards the intrinsic stacking fault gives rise to a Gaussian-like concentration distribution in the vicinity of intrinsic stacking fault in Al. It follows that an increasing vacancy concentration can produce a considerable decrease in YSFE of Al. However, no further increase of twinning propensity was observed when introducing more vacancies. One interesting phenomenon predicted in the present work is the coupled segregation of vacancies and Mg solutes towards intrinsic stacking fault in Al, i.e. the Suzuki segregation of Mg and vacancies to stacking fault are mutually stimulated. Finally, the twinnability analysis based on the modified GPFE of Al by introduction of vacancies and Mg atoms signifies the general difficulty of deformation twinning in Al-Mg alloys, consistent with the rarely reported deformation twin forming in coarse-grained Al-Mg alloys even subjected to various SPD strategies. Nevertheless, a reduction in YSFE attributed to Mg and vacancy introduction can, at least partially, explain the high work-hardening rate and the formation of band structures in SPD processed Al-Mg alloys.

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