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Phys. Rev. Materials **6**, 083602 — Published 15 August 2022

DOI: [10.1103/PhysRevMaterials.6.083602](https://doi.org/10.1103/PhysRevMaterials.6.083602)

A Statistical Perspective on Embrittling Potency for Intergranular Fracture

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Abstract

Embrittling potency is a thermodynamic metric that assesses the influence of solute segregation to a grain boundary on intergranular fracture. Historically, studies have reported embrittling potency as a single, scalar value, assuming a single segregation site of importance at a grain boundary and a particular cleavage plane. However, the topography of intergranular fracture surfaces is not generally known *a priori*. Accordingly, this study presents a statistical ensemble approach to compute embrittling potency, where many free surface permutations are systematically considered to model fracture of a grain boundary. The result is a statistical description of the thermodynamics of grain boundary embrittlement. As a specific example, embrittling potency distributions are presented for Cr segregation to sites at two Ni $\langle 111 \rangle$ symmetric tilt grain boundaries using atomistic simulations. We show that the average embrittling potency for a particular grain boundary site, considering an ensemble of free surface permutations, is not equal to the embrittling potency computed using the lowest energy pair of free surfaces. A mean grain boundary embrittlement is proposed, considering both the likelihood of formation of a particular free

surface and the probability of solute occupancy at each grain boundary site, to compare the relative embrittling behavior of two distinct grain boundaries.

Keywords

Grain Boundary Segregation, Embrittlement, Atomistic Simulation, Probability Distribution

1. Introduction

Atomic solutes in metals are prone to segregation where, depending on energetic favorability, they may migrate toward grain boundaries (GBs) [1] or other regions of free volume [2,3]. Solute segregation can be intended and beneficial, such as in the case of grain size stabilization in nanocrystalline metals [4–6], or it can have deleterious effects on material properties, such as in the case of hydrogen embrittlement [7,8]. A variety of material property changes result from segregation [2,9,10], but we focus specifically on GB embrittlement in this work, where solute segregation changes the work of GB separation. GBs are known to be energetically favorable pathways for fracture [11,12]; thus, embrittlement caused by solute segregation can have a profound impact on the strength and ductility of an alloy [13].

McLean provided a thermodynamic description of GB solute segregation in binary systems by defining an isotherm that relates the concentration of solutes segregated to a GB to bulk concentration and a solute segregation energy [14]. The McLean isotherm assumed that segregation energies to each GB site were equal. However, different atomic sites at a GB have different geometric and chemical configurations. White, Stein, and Coghlan [15,16] showed evidence that solutes do not segregate to all GB sites equally and defined a different isotherm to account for site-specific segregation energetics. Building upon GB segregation energy, the thermodynamic concept of embrittling potency was proposed by Rice and Wang [13]. Embrittling potency for a GB was defined as the difference in energy between solute segregation to the GB and solute segregation to a corresponding free surface. Furthermore, Rice and Wang argued that embrittlement was related to the reduction of the work of separation of a GB due to the presence of the segregant. The amount by which the

ideal work of separation was changed by solute segregation could be written as $\Delta E_b \Gamma$, where ΔE_b is the embrittling potency, and Γ is the number of segregants per unit area on the interface (coverage). Hence, embrittling potency served as a scalar measure of the impact of a solute on GB fracture. Several authors have since studied site-specific embrittling potency for various metals [7,17–22]. For example, Razumovskiy et al. [20,22] studied embrittlement in a variety of Ni-based alloys using first-principles methods. These authors used their findings on GB embrittling potency to tailor compositions for Ni-based superalloys with higher mechanical strength. Recently, Aksoy et al. [21] introduced a method for comparing embrittling potencies between different symmetric tilt GBs (STGBs) embedded within a theoretical polycrystal. By defining a statistical method of analysis which calculates GB site occupancy within a polycrystalline environment and performing atomistic simulations of solute segregation to STGBs and their corresponding free surfaces (FS), Aksoy et al. identified critical grain boundaries with outstanding embrittling behavior in the case of S segregation in a Ni polycrystal.

In previous works that focused on embrittling potencies, a prevalent underlying assumption was made that GB fracture occurred on a predetermined, flat cleavage plane [7,13,17–21,23]. The consequence of this assumption is that a single value of the embrittling potency was computed for each atomic site. However, GB fracture can be tortuous, producing atomically rough surfaces depending on the boundary and loading conditions [18,23–26]. For example, tensile tests of H-embrittled Inconel from Seita et al. showed tortuous intergranular fracture surfaces resulting from a mixture of brittle and ductile fracture [25]. Atomistic simulations from Yamakov et al. [26] showed that unstable crack propagation in intergranular fracture can result in micro-void coalescence producing

a variety of non-minimum energy surface topographies. Molecular dynamics simulations from the work of Spearot et al. [24] and Bitzek et al. [27] clearly demonstrated jagged cleavage planes with surface roughness between 3-6 Å normal to the GB. Such observations imply that a free surface pair can form that does not correspond to the static minimum energy free surfaces for that lattice orientation. Unfortunately, the influence of intergranular fracture surface topography on embrittling potency has not been discussed or evaluated at length; although, one recent study of H embrittlement used a small set of atomically flat free surfaces [28].

In this work, we examine embrittling potency of solutes at GBs from a thermodynamic and statistical perspective. Our approach considers an ensemble of possible free surface topographies resulting from an implied tortuous intergranular crack propagation. We illustrate this approach via calculations of Cr segregation to individual GB sites in Ni. Ni-Cr is chosen as model alloy system since the atomic mismatch between Ni and Cr is negligible, limiting the effects of compositional expansion and the associated elastic release of stresses around the segregating solute [29]. Additionally, the origin of segregation in this alloy system is well documented in previous studies [20,22,29,30], making it an ideal candidate for this study. Atomistic calculations of segregation across the ensemble of free surfaces produce a distribution of embrittling potencies for each GB site, showing that some GB sites exhibit strengthening or embrittling behavior depending on the free surface topography. Analyzing the new ensemble data requires a different set of tools; thus, we propose an ensemble average of the embrittling potencies that accounts for the probability density associated with free surface formation, based on calculations of equilibrated free surface energies. Additionally, we propose a mean GB embrittlement which considers the ensemble

average embrittling potencies, the probability of site occupancy by a solute, and the area of the GB unit cell. The mean GB embrittlement condenses the embrittling potency distributions into a single, scalar value representative of the full ensemble that can be used to compare two different GBs when the embrittling potency distributions are themselves unique.

We note that this work does not consider the dynamics of crack propagation along a GB. Rather, we present a thermodynamic and statistical perspective on the calculation of embrittling potency. We assume fracture conditions where solutes do not move from their site position between GB and FS models, although local relaxations are allowed via energy minimization within the framework of molecular statics calculations. Thus, the results described below are best applied to cases of crack initiation.

2. Methods

2.1. Material Selection, EAM Potential, and Software

The selected material system consists of bulk Ni as the solvent with Cr solutes. We choose this system since there is less than a 5% difference between the atomic radii of Ni and Cr. Thus, embrittlement can be attributed to chemistry only since Cr substitution will not create significant mechanical strain in the lattice for the solvent/solute pair. Additionally, the segregation behavior of Cr in Ni has been well-documented by Startt et al. [29,30] using *ab initio* calculations, providing a point of validation for the results of our calculations. We use the LAMMPS molecular dynamics software [31,32] with an embedded-atom method (EAM) potential designed by Zhou et al. [33]. While originally designed as an Fe-Ni-Cr potential, this potential was fitted to accurately simulate the formation of different material

phases and provide stacking-fault energies which aligned with experimental results. Stacking-fault energies and material phase emulation are essential in the modeling of grain boundaries and segregation. While other potentials were also considered for study [34,35], the Zhou et al. potential was chosen as the best option. The mean and breadth of the distributions that are presented in Section 3 are likely to be sensitive to the interatomic potential; however, the fundamental concepts of the existence of an embrittling potency distribution for each GB site and the proposed ensemble average are valid regardless of the choice of interatomic potential.

We use Ovito [36] to visualize the resulting structures and identify specific atoms at and surrounding the GBs for segregation and embrittlement analysis. Within a unit cell area on the GB plane, we identify and number unique atom sites based on centrosymmetry [37].

2.2. Grain Boundary Models

We select a pair of complementary symmetric tilt grain boundaries (STGBs) with $[111]$ misorientation axis for analysis: the $\Sigma 7 \{213\}$ and $\Sigma 21 \{415\}$ STGBs. These GBs are chosen because the adjoining lattice regions exhibit similar characteristics [38]. To construct the STGBs, we create two regions, “top” and “bottom”, in a simulation cell with periodic boundary conditions in all three cartesian directions as shown in Fig 1a. The misorientation axis $M = [111]$ is defined along the x-axis, parallel to the GB plane, while the normal to the GB plane N in the top lattice region is defined as $[\overline{4}15]$ for the $\Sigma 21$ STGB and $[\overline{2}13]$ for the $\Sigma 7$ STGB. We populate both regions with Ni atoms in a face-centered cubic (FCC) crystal structure oriented according to those crystallographic directions.

The dimensions of the top and bottom regions must be integer multiples of the repeating distance of the lattice determined from the crystallographic orientation to satisfy periodic boundary conditions. In this study, we choose repeating distances to be $8 \times 7 \times 6$ units in the x, y, and z directions, respectively. Hence, the $\Sigma 21 \{415\}$ model has the dimensions $48.7 \times 66.2 \times 91.8 \text{ \AA}$, and the $\Sigma 7 \{213\}$ model has the dimensions $48.7 \times 114.7 \times 53.0 \text{ \AA}$. These dimensions are significantly larger than the cutoff distance for the EAM potential used in this study; thus, they are sufficient for convergence of atomistic calculations of segregation energies.

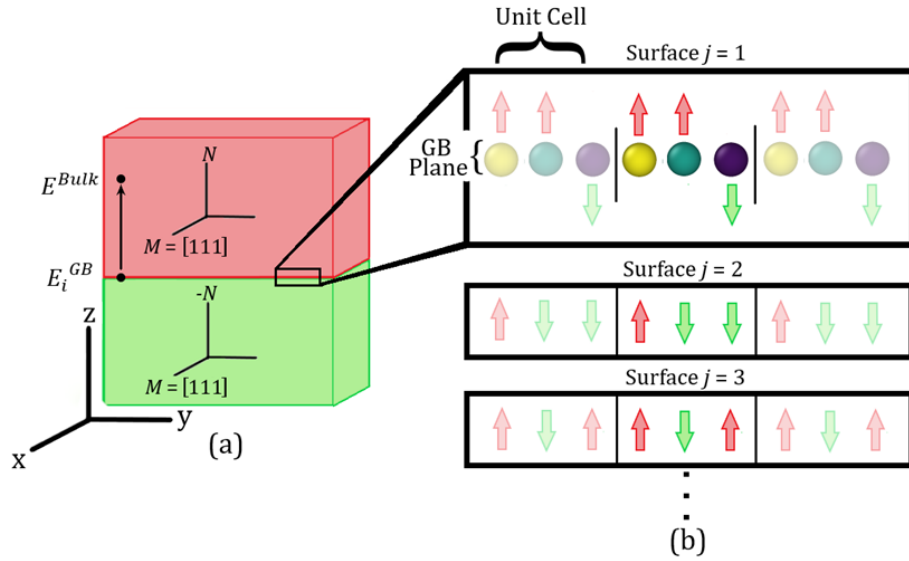


Figure 1: (a) Lattice orientations for both top (red) and bottom (green) regions illustrating the GB construction. Cr is substituted into a GB site and energy minimization is performed to obtain E_i^{GB} . Then, Cr is swapped from the GB into the bulk and energy minimization is performed to obtain E^{Bulk} . (b) Example of the free surface topography permutation process with a fictitious GB with 3 atoms at the GB plane in one repeating unit cell. Decisions for an atom to move up or down during fracture are made within a single GB unit cell, and then those decisions are repeated in all other repeating units.

To find the minimum energy configuration of each GB, we perform a gamma-surface construction. Within this construct, we translate the top lattice region in the x, y, and z directions relative to the bottom lattice region, with a conjugate gradient molecular statics (energy minimization) calculation performed after each translation. The total translation distance in the x and y directions is equal to the repeating distance, while the translation distance in the z direction is $2 \text{ \AA}$, roughly the width of a layer of atoms in the z direction. Since the z translation may cause atoms to overlap each other, atoms from the bottom region are deleted if they are within $1 \text{ \AA}$ of atoms in the top region. 20 increments in the x and y direction and 10 increments in the z direction provide 4000 initial configurations for determination of the lowest energy GB structure.

2.3. Free Surface Models

We initially assume that fracture occurs along the GB plane as illustrated in Fig. 1b. Atoms along the GB plane in a single unit cell are partitioned into either the top or bottom region, and the partitions are repeated periodically for each unit cell on the GB plane. For example, referring to the fictitious GB in Fig. 1b, the ensemble contains a permutation ($j = 2$) where a pair of unrelaxed free surfaces (FS) is formed by moving atoms within a unit cell in an up-down-down pattern. The top and bottom regions are then separated by $20 \text{ \AA}$ and conjugate gradient molecular statics is used to determine the relaxed free surface topography. Energy minimization is performed with no external pressure. Generating all permutations with this method results in $32 (2^5)$ free surface permutations for the $\{415\}$ GB, and $512 (2^9)$ for the $\{213\}$ GB. We call this set of fracture permutations the “periodic GB ensemble”. Note, this procedure implicitly assumes that solute concentration does not

change at the GB during fracture, and that the solute does not move significantly from its site location, although it may relax locally during energy minimization.

Note, this method produces free surfaces in pairs for symmetric tilt GBs. For each free surface where an atom site is partitioned into the top group, there exists a complementary free surface where the opposite is true. Each free surface should have the same free energy and segregation energy as its mirror duplicate; thus, we keep data resulting from these duplicate surfaces to verify the consistency of the simulation results.

Of course, each GB could have more possible free surface permutations than those in the periodic GB fracture ensemble. To limit the total number of molecular statics calculations that are necessary for every site in the GB, we initially assume that fracture occurs along the GB plane and that decisions are periodic for all GB unit cells. We hypothesize that the ensemble resulting from these decisions is representative of the ensemble of all possible free surfaces. To test this hypothesis, in Section 3.2., we include some atoms from one atomic layer above the GB plane in the free surface permutation process to understand how this expanded set changes the distribution of FS energy. We call this set of permutations the “periodic expanded ensemble”.

In addition, to verify how free surface topography neighboring the analyzed GB unit cell can affect segregation and embrittlement, we create a third set of fracture permutations. In this new set, a single GB unit cell is permuted in the same way as the periodic GB ensemble; however, periodicity is not enforced across the entire GB plane. Instead, the remainder of the GB adopts an atomically flat fracture topography. Hence, we call this ensemble the “flat neighboring ensemble”. For the $\Sigma 7$ {213} STGB, since the GB cell is composed of three

identical units, we include translations of the flat fracture region in the y direction, shifting the starting location of the permuted region. Ultimately, for both STGBs, the average embrittling potency is compared against the periodic GB ensemble.

2.4. Segregation Energy and Embrittling Potency

The segregation energy of a solute to a GB or FS, $\Delta E_{seg,i}^{GB/FS}$, is defined as the difference between the energy of a system with a solute atom at a substitutional site at the GB or FS, $E_i^{GB/FS}$, and the energy of a system with a solute atom at a substitutional site within the bulk, E^{Bulk} [16],

$$\Delta E_{seg,i}^{GB/FS} = E_i^{GB/FS} - E^{Bulk} . \quad (1)$$

Here, i denotes the site type. Negative segregation energy implies that solute segregation to the GB or FS is energetically preferred.

Figure 1 illustrates how these values are acquired. We initially substitute a Cr atom into the desired Ni GB site location. Following an energy minimization, we compute the potential energy of the system as E_i^{GB} . Then, we swap the GB site Cr with a Ni atom from within the bulk and perform another energy minimization to calculate E^{Bulk} .

For calculation of $\Delta E_{seg,i}^{FS}$, we must include an additional step. As before, we substitute a Cr atom into the desired surface site and perform an energy minimization to calculate E_i^{FS} . Then, we swap the surface Cr with a Ni atom within the bulk. Unlike before, we restrict the relaxation of all atoms except for those within a sphere around the bulk Cr of radius equal to the interatomic cut-off distance of 5.7 Å. Then, we calculate the energy again as E^{Bulk} . The purpose of ordering the substitutions and locally minimizing the bulk Cr is to limit model-

scale reconstruction of the free surface during energy minimization. If the free surface reconstructs, then the free surface permutation of interest would not be the same as the free surface that results from the energy minimization. This would lead to inconsistencies in the calculation of the embrittling potency. Ultimately, the surfaces for which we prevent reconstruction have much higher FS energies than the minimum energy free surfaces (difference greater than $200 \text{ mJ}/\text{m}^2$), and thus will end up being weighted lower when averages are performed in Section 4, so local minimization should not notably impact the results of the study.

Once we have segregation energies for the GB and each FS, embrittling potency can be calculated. The embrittling potency of a particular GB site $\Delta E_{b,i}$ is defined using the Rice-Wang formulation [13],

$$\Delta E_{b,i} = \Delta E_{seg,i}^{GB} - \Delta E_{seg,i}^{FS}. \quad (2)$$

A positive $\Delta E_{b,i}$ denotes embrittling, whereas a negative $\Delta E_{b,i}$ denotes strengthening. Note that Eq. (1) computes a single value of $\Delta E_{seg,i}^{GB}$ for each GB site, but each GB site exists on many possible sampled free surfaces (Fig. 1), providing a distribution of values for $\Delta E_{seg,i}^{FS}$. Hence, each GB site will have a distribution of embrittling potencies $\Delta E_{b,i}$ computed via Eq. (2).

3. Results

The objective of this work is to show how distributions of embrittling potencies computed using different free surface topographies can contribute to an ensemble average, and hence a more complete understanding of solute embrittlement. To do so, we must gather the following information: solute segregation energies to the GBs (Section 3.1), embrittling

potencies for each free surface permutation for the STGBs (Section 3.2), and relaxed FS energies, which will define weighting in the ensemble average (Section 3.3.). Note, for completeness, we present embrittling potency distributions for all Ni GB sites, even if they are anti-segregation sites that would never be occupied by the Cr solute. In the ensemble average (Section 4), we employ the McLean isotherm [14–16] which effectively addresses the coupling between GB site occupancy and embrittlement.

3.1. Grain Boundary Segregation Energies

Figure 2a shows the minimum energy $\Sigma 7 \{213\}$ and $\Sigma 21 \{415\}$ STGB structures. GB structural units are labeled according to the nomenclature presented by Wang et al. [39]. Figure 2b shows maps of segregation energies for a Cr solute to sites within these STGBs. The most energetically favorable segregation sites at the GB plane are sites 2, 5, and 8 for the $\Sigma 7 \{213\}$ boundary, and site 2 for the $\Sigma 21 \{415\}$. Interestingly, these sites are associated with the F structural units in the case of the $\Sigma 7 \{213\}$ STGB while they are associated with the E structural unit in the $\Sigma 21 \{415\}$ STGB. On the other hand, the largest anti-segregation sites are sites 1, 4, and 7 for the $\Sigma 7 \{213\}$ STGB, and site 4 for the $\Sigma 21 \{415\}$ STGB. In both GBs, these anti-segregation sites are associated with the F structural unit and are positioned near pockets of free volume within the GB structure.

The solute segregation energy map for the $\Sigma 7 \{213\}$ STGB illustrates a pattern of energies that repeats in unison with the structural unit description, which is expected. The regular geometric pattern ensures that each FE structural unit block is identical to its neighbor, resulting in the triplet data shown in Fig. 2b for the $\Sigma 7 \{213\}$ STGB.

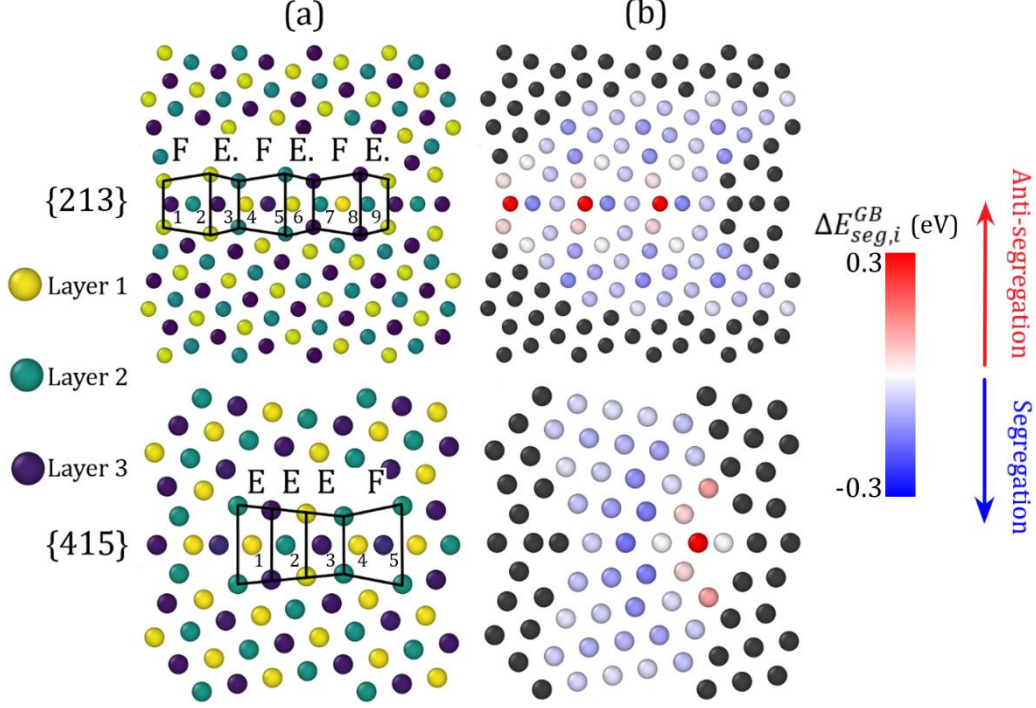


Figure 2: (a) Minimum energy structure of the $\Sigma 7$ {213} and $\Sigma 21$ {415} STGBs in Ni. Grain boundary sites are numbered for reference and colored by {111} atomic layer to identify structural units. The structural unit nomenclature was developed by Wang et al. [39]. (b) Map of Cr solute segregation energies for $\Sigma 7$ {213} and $\Sigma 21$ {415} STGBs. Units are in eV. Negative values indicate favorable solute segregation while positive values indicate anti-segregation. Grey atom sites are not evaluated.

Segregation energies provide sufficient information to determine equilibrium concentrations at the GB. According to the site-specific interpretation of the McLean isotherm [16], the atomic percent of sites i occupied by solutes, X_i^{GB} , can be determined via,

$$X_i^{GB} = \frac{X^{Bulk} \exp\left(-\frac{\Delta E_{seg,i}^{GB}}{k_b T}\right)}{1 - X^{Bulk} + X^{Bulk} \exp\left(-\frac{\Delta E_{seg,i}^{GB}}{k_b T}\right)} \quad (3)$$

Where X^{Bulk} is the atomic percent of solutes in the bulk lattice, k_b is the Boltzmann constant, and T is the absolute temperature, taken to be $T = 300 \text{ K}$ in this work to emulate room temperature conditions. We broadly use the term “site occupancy” when describing

solute occupation of sites at the grain boundary. Site occupancy is often taken to mean either “the probability that a site i is occupied by a solute” or “the atomic percent of sites i occupied by a solute”. We use these contexts interchangeably throughout this work.

The choice of X^{Bulk} should be small, as the atomistic simulations in this work are in a dilute limit where solute-solute interactions are not considered. Wagih and Schuh reported that the limit for dilute studies lies at $X^{Bulk} \cong 0.02$ or 2% [40] for Mg segregation in Al. Beyond this limit, real dilute segregation behaviors diverge from the non-dilute approximations due to solute-solute interactions. Al-Mg is not identical to Ni-Cr; however, the interatomic potential for Al-Mg [41] used by Wagih and Schuh has a similar cutoff distance to the Ni-Cr potential used in this study, and they both show energetic preference for solvent-solvent interactions over solute-solute interactions. Hence, it is reasonable to assume that the underlying cause of non-dilute behavior, solute-solute interaction, is similar for both cases and we take $X^{Bulk} = 2\%$ for our calculations. Note, there is nothing that precludes the extension of our method to the non-dilute case, but that is beyond the scope of the present study.

Frequently, studies which discuss GB width look toward geometric indicators such as centrosymmetry or CNA to determine which sites are part of the GB and which sites belong in the bulk [42,43]. However, our method is not concerned with structural unit geometry, but instead with the thermodynamic impact of a segregant on GB fracture. Hence, we define GB width through a segregation energy cutoff for the purpose of studying embrittlement. Atoms along $\langle 110 \rangle$ crystallographic directions outward from the GB plane are included in the analysis until the segregation energy diminishes to a value between -0.025 and 0.0 eV, since

sites with segregation energy within this range will have $X_i^{GB} < 5$ at.% (computed using Eq. 3), which we consider to be negligible. Figure 3 demonstrates for the three unique $\Sigma 7$ {213} STGB sites how we implement this criterion along $\langle 110 \rangle$ crystallographic paths. Based on the data, this criterion provides a GB width of approximately 6 Å on either side of the GB plane. An identical criterion is appropriate for determining GB sites for the $\Sigma 21$ {415} STGB.

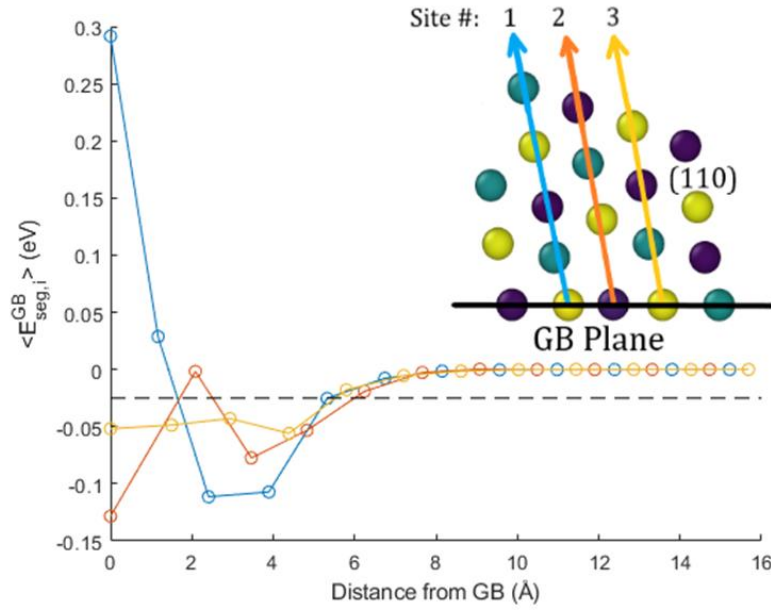


Figure 3: Segregation energy versus distance from the GB plane along $\langle 110 \rangle$ directions for the $\Sigma 7$ {213} STGB. The cutoff criterion is indicated above the dashed black line. Atoms in the figure inlay are colored by different {111} planes, as shown in Fig. 2a.

Note, this GB width criterion still includes some sites with positive segregation energy within the GB, which would have $X_i^{GB} \cong 0$. This includes sites 1, 4, and 7 in the $\Sigma 7$ {213} STGB and site 4 in the $\Sigma 21$ {415} STGB. For completeness, we include these sites in the embrittling potency calculations; however, they will not impact the GB work of separation since they will never be occupied by solute. In Section 4, we address this by weighting GB site embrittling potencies by GB site occupancy.

3.2. Embrittling Potency Distributions

Segregation energy and embrittling potency calculations are performed for all GB sites identified in Fig. 2b; however, in this section, we limit our demonstration of embrittling potency distributions to the numbered atoms at the GB plane because these atoms contain the extrema of GB segregation energies. Figs. 4a and 4b show embrittling potency distributions for GB sites at the $\Sigma 7$ {213} and $\Sigma 21$ {415} STGBs, respectively. The box-whisker plots show condensed distributions of embrittling potencies for each site type. The red line denotes the mean, the box denotes the interquartile range, and the whiskers denote the data beyond the third quartile and before the first quartile. Red crosses denote statistical outliers.

Each box-and-whisker corresponds to a histogram which expands the distribution in detail. Examples of these histograms are shown in Fig. 4a for site 9 in the $\Sigma 7$ {213} STGB and in Fig. 4b for site 5 in the $\Sigma 21$ {415} STGB. Each histogram is fit with a skew-normal distribution,

$$F(\Delta E_{b,i}) = \frac{2}{\sigma} \frac{1}{\sqrt{2\pi}} \exp\left(-\frac{\left(\frac{\Delta E_{b,i} - \mu}{\sigma}\right)^2}{2}\right) \left[\frac{1}{2} \left(1 + \operatorname{erf}\left(\lambda \frac{\Delta E_{b,i} - \mu}{\sigma\sqrt{2}}\right) \right) \right], \quad (4)$$

where λ , μ , and σ are the skew, location, and shape of the distribution, respectively. The use of this distribution model to fit these results is inspired by the work of Wagih and Schuh [44], for which they show how closely a well-fit skew-normal distribution can approximate the density of states based on segregation energy. Skew-normal distributions are flexible distributions capable of capturing a variety of behaviors which have nearly normal behavior, so this function is equally applicable here.

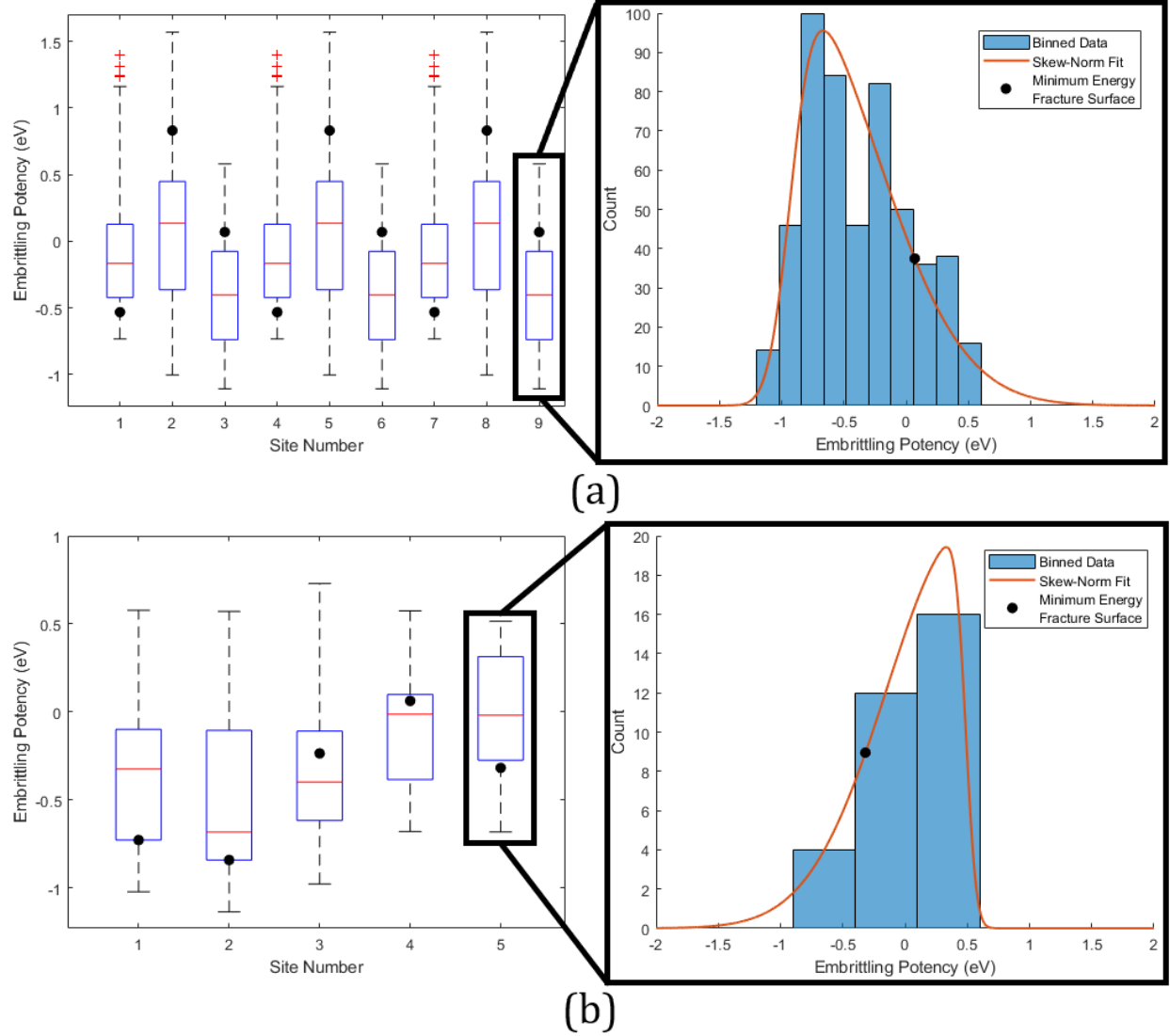


Figure 4: Distributions of embrittling potencies for (a) $\Sigma 7 \{213\}$ STGB sites with a detailed histogram for site 9 and (b) $\Sigma 21 \{415\}$ STGB sites with a detailed histogram for site 5. Distributions are represented compactly as box-and-whisker plots, with plus symbols denoting statistical outliers. The dots indicate embrittling potencies associated with the relaxed minimum energy free surfaces.

Each box-and-whisker in Fig. 4 contains a black dot indicating the embrittling potency for the indicated site associated with the minimum energy free surfaces. Regardless of the segregation site and GB considered, the value of embrittling potency associated with the minimum energy free surfaces does not lie at the mean of observations nor at the extrema,

but rather at an arbitrary value within the distribution. Hence, the use of the minimum energy free surface does not necessarily correlate to high, low, or even average embrittling potency when an ensemble of free surfaces is considered.

Figure 4 shows that GB sites can be embrittling or strengthening in nature depending on the intergranular fracture surface topography. Recall, the most energetically favorable sites for Cr segregations are 2, 5, and 8 for the $\Sigma 7$ {213} STGB, and site 2 for the $\Sigma 21$ {415} STGB. Each of these sites shows a spectrum of behavior depending on the topography of the free surface. For example, for the $\Sigma 7$ {213} STGB, sites 2, 5, and 8 show a mean positive embrittling potency (embrittling). However, many of the sampled free surfaces display a negative embrittling potency (strengthening) for these sites. Similarly, for the $\Sigma 21$ {415} STGB, site 2 shows a mean strengthening behavior; however, the box-whisker plot indicates that about a 25% of the sampled free surfaces show embrittling behavior. For all sites along the GB, the tails of the embrittling potency distributions show both embrittling and strengthening behavior. Studying a single free surface may capture either the embrittling or strengthening behavior but would ignore the spectrum of possible embrittling potencies resulting from the ensemble of possible free surfaces.

These distributions elucidate the embrittling nature of a particular GB site for a selection of intergranular crack surface topographies, but they are not useful in this form for comparing segregation-induced embrittlement between two GBs. Previously, Aksoy et al. [21] used a combination of site occupancy probability and embrittling potency to make meaningful comparisons between GBs. However, to meaningfully make conclusions about relative embrittlement of two GBs when considering an ensemble of free surfaces, we must

develop new tools which include the likelihood of free surface formation the probability of segregation to a particular GB site.

3.3. Free Surface Energies

Figures 5a and 5b show relaxed FS energies for all free surface permutations used to compute embrittling potency distributions for the $\Sigma 7$ {213} and $\Sigma 21$ {415} STGBs. These histograms demonstrate the wide variety of surface permutations tested in the generation of embrittling potency data. We uniquely determine the bin width of these histograms via the Freedman-Diaconis rule, which uses the interquartile range [45]. Since the pair of free surfaces generated are asymmetric with respect to the cleavage plane, the FS energies plotted are the average energies for both free surfaces.

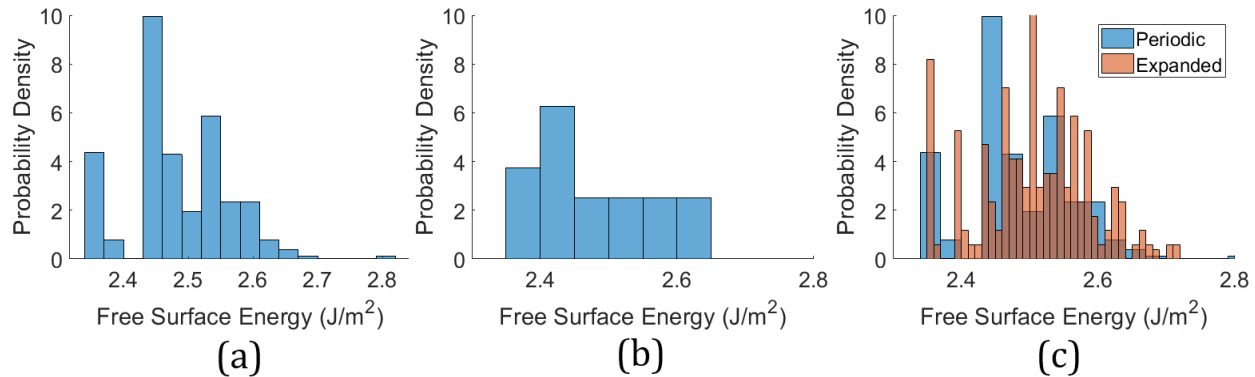


Figure 5: Histograms showing relaxed free surface energies for free surface topographies of the (a) $\Sigma 7$ {213} and (b) $\Sigma 21$ {415} STGBs. The distributions generally skew left towards lower FS energies. (c) Overlaid histograms of FS energies for fracture of the $\Sigma 7$ {213} STGB. The “Periodic” histogram includes the periodic GB ensemble shown in Fig 3a, while the “Expanded” histogram shows the periodic expanded ensemble that includes the GB plane atomic sites, as well as 3 atomic sites from the adjacent atomic plane above it. Note that the ranges of the FS energies are very similar.

For each GB, one pair of free surfaces stands out as the minimum energy surface topography. These minimum energy surfaces occur when all atoms at the GB plane are

assigned to either the top or bottom lattice during fracture. Hence, the minimum energy free surface does not split the GB plane as assumed in the works of Razumovskiy et al. [20] and Aksoy et al. [21], but rather delaminates the GB plane from the adjacent plane of atoms cleanly.

To show that our method of systematically generating free surfaces produces a representative ensemble of free surfaces, using the $\Sigma 7 \{213\}$ STGB as an example, we include FS energies from the periodic expanded ensemble. This increases the number of free surface permutations from 512 to 4096 (2^{12}), each of which would need to be evaluated to compute segregation for all GB sites. However, the addition of these new surfaces does not expand the range of relaxed FS energies, as shown in Fig. 5c; thus, free surfaces created using GB sites only are representative of the probable range of FS energies for these GBs.

4. Ensemble Averaging

To utilize the data computed from different free surface permutations, we present a method for condensing the tens of thousands of embrittling potencies computed in Section 3 into metrics that can be related back to fundamental concepts presented by Rice and Wang. A graphical overview of this approach is shown in Fig. 6. First, we average the embrittling potencies over the ensemble of free surfaces j for each GB site i , $\Delta E_{b,ij}$, to get ensemble averaged values for each site, $\langle \Delta E_{b,i} \rangle$. Then, an occupancy weighted average over the area of the GB unit cell is performed to obtain a mean embrittlement $\langle \Delta E_b^A \rangle$; this accounts for the coupling between occupancy and embrittlement and results in a mean contribution of solute segregation to embrittlement of the GB.

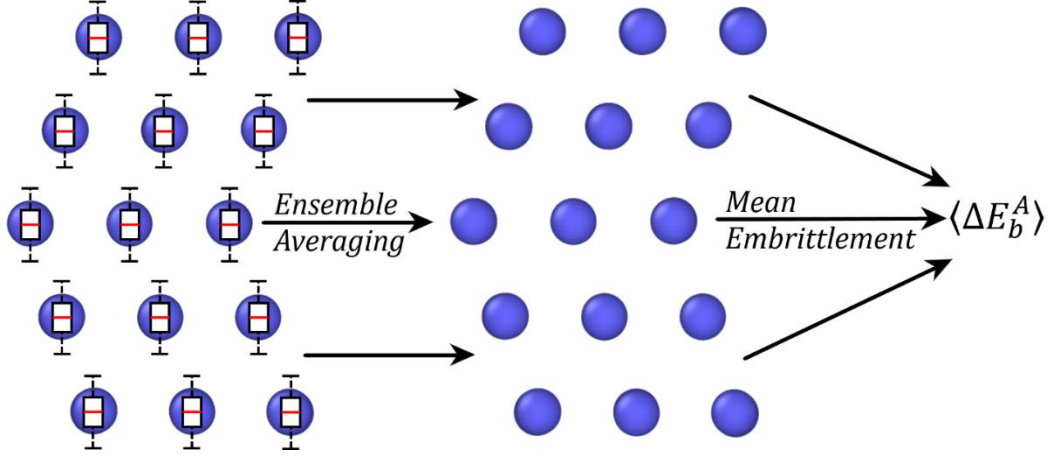


Figure 6: The process of creating the mean embrittlement. Each site at the GB begins with a distribution of embrittling potencies (left). Following an ensemble averaging procedure over many free surfaces, each site contains a single ensemble averaged embrittling potency (middle). Then, an average is performed over all sites considering solute occupancy to obtain a mean embrittlement.

4.1. Free Surface Weighting

To compute an average over the ensemble of the free surface permutations for a particular GB site, weights must be determined for the embrittling potency associated with each free surface. It is likely that free surface topographies with lower energy form more frequently during intergranular fracture. Thus, we assume that the weight for a particular embrittling potency depends on the relaxed energy of the corresponding free surface. We do not consider dissipative events that lead to small structural rearrangements, such as dislocation nucleation from a crack tip, nor do we consider the effects of lattice trapping.

To obtain weights for embrittling potencies associated with FS energies, we require a tunable, monotonically decreasing distribution function. We expect that free surfaces are less likely to form if they have higher energies [46]. Hence, a survival function derived from Weibull fracture statistics serves our needs. Weibull-style functions are frequently used to

describe distributions in studies of material strength and fracture [47–50], and its tunable size and shape parameters allow us to investigate the influence of including higher-energy free surfaces in the ensemble average for embrittlement.

We choose deliberately not to use the Boltzmann distribution, commonly used in statistical mechanics. While the NVT ensemble for which the Boltzmann distribution was derived has broad application, one of its core assumptions is that no work is or has been done on the system [51]. In our simulations, fracture is implied to occur between the GB and FS states, so work is considered to have been done on the system. Hence, the Boltzmann distribution is inappropriate for our approach.

Weibull fracture statistics, however, provides an appropriate lens through which we can examine our problem. In fracture statistics, elements of a material are envisioned as a series of nested chains and bundles, spanning in resolution from the continuum to the angstrom scale [52]. Failure of any one of these elements results in the failure of the entire structure, and failure is driven by an interplay between stress and the random distributions of kinetic energy among atoms, modelled by the Maxwell-Boltzmann distribution. One may consider the structures presented in this study as being composed of such a lattice of chain and bundle elements, and the “failure of the entire structure” is envisioned as crack initiation. From empirical observations, we expect the GB unit elements to be the dominant volumes of failure, so a model that accounts for the failure of all such elements in the structure could be statistically approximated by a model which only observes failure at the GB elements. Thus, we view the weakest-link model as an appropriate statistical mechanics method for modeling intergranular fracture on the nano scale.

From Bazant et al. [52], if we assume that the stress is uniform, we may represent the probability of failure as a cumulative distribution function (CDF) $P_f(\sigma; \lambda, k)$, where σ is the stress acting on the volume, and λ and k are fitting parameters. The survival function is simply $1 - P_f(\sigma; \lambda, k)$. While the survival function presented by Bazant et al. uses stress, thermodynamically, the fracture stress is proportional to the energy required to create two free surfaces from the GB model, i.e., $\sigma_f \propto 2E_j^{FS} - E^{GB}$. Recall from Fig. 5 that E_j^{FS} is the average free surface energy for both surfaces. Since the energy of the GB is a constant for each model, we can simplify the previous statement to $\sigma_f \propto E_j^{FS}$. Now, the survival function provides a monotonically decreasing probability of survival based on free surface energies and may be used as a weighting function for our study. Since a Weibull distribution is best suited for weakest-link models [52], we write the weighting function as,

$$w_j(E_j^{FS}; \lambda, k) = \exp \left[- \left(\frac{E_j^{FS}}{\lambda} \right)^k \right] \quad (5)$$

In this section, we choose a range of k values to compare, which physically represents different sets of free surface permutations, and for each k fit a corresponding λ such that $w_j = 0.99$ at the minimum energy free surface. This fitting is performed for each GB independently.

As an example, survival functions for the $\Sigma 7$ {213} STGB are shown in Fig. 7 overlaid on the histogram of the relaxed free surface energies. Lower k values represent ensembles that include higher-energy free surfaces.

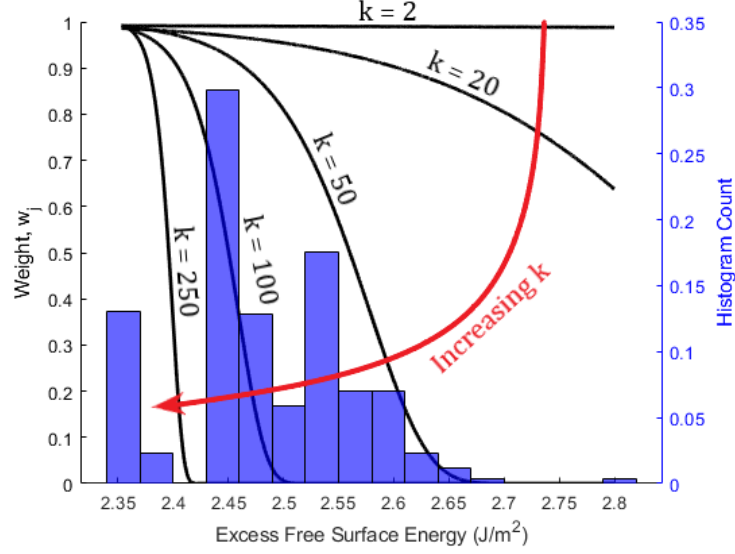


Figure 7: Examples of Weibull survival functions with different k values for the $\Sigma 7$ {213} STGB. The red arrow indicates increasing k . Higher k values result in lower weights for higher energy free surfaces.

Utilizing different values of k allows us to answer the question: To what extent does inclusion of non-minimum energy free surfaces change the average embrittling behavior? As $k \rightarrow +\infty$, the survival function will approach the form of a step function, removing all free surface data except that of the free surface with the lowest formation energy. Likewise, as $k \rightarrow 1$, the survival function includes all free surface data from higher-energy free surfaces into the ensemble. In other words, lower values of k represent a larger ensemble, i.e., more possible free surface topographies, while larger values of k represent more limited ensembles, i.e., only lower-energy free surface topographies. While $k \rightarrow +\infty$ represents the idealized circumstance of equilibrium fracture, we expect from atomistic simulations [24,27] that higher-energy free surfaces may appear during intergranular crack propagation, requiring this data to be included in the analysis of solute embrittlement.

For the proceeding discussion, we assume $k = 250$. We explore how this might affect our results by considering the following question: what is the free surface energy range

included by a particular value of k ? For example, for $k = 250$, w_j is 0.05 at $E_j^{FS} = 2.41 \text{ J/m}^2$, so the included free surface energy range is only $[2.35, 2.41] \text{ J/m}^2$, or 0.06 J/m^2 . For the $\Sigma 7 \{213\}$ STGB, the area of a unit cell is about $69.7 \text{ \AA}^2$, so the energy range for different free surface topographies over the unit cell is $4.18 \times 10^{-20} \text{ J}$, or 0.26 eV . Lower k expand the free surface energy range, while higher k shrink this range. This brief example provides some insight into the energetics associated with different intergranular fracture surfaces that may form under thermal and mechanical loading.

4.2. Ensemble Averaged Embrittling Potency

With the FS weight w_j clearly defined, we now define an embrittling potency for each site $\langle \Delta E_{b,i} \rangle$ as a weighted average over the ensemble of many free surface permutations j . Initially, this is done regardless of the segregation or anti-segregation nature of the GB site. In this work, we refer to this term as the *Ensemble Average Embrittling Potency*,

$$\langle \Delta E_{b,i} \rangle = \frac{1}{\sum_j w_j} \sum_j \Delta E_{b,ij} w_j \quad (6)$$

Here, $\Delta E_{b,ij}$ is the embrittling potency of GB site i with free surface topography j . The weight w_j is the probability of formation for free surface topography j from Eq. (5). This provides each site with an embrittling potency that considers all sampled free surface topographies and their likelihoods of appearing during fracture. This approach allows for the identification of critical sites for embrittlement. Values of $\langle \Delta E_{b,i} \rangle$ are shown for the $\Sigma 7 \{213\}$ and the $\Sigma 21 \{415\}$ STGBs in Fig. 8a and 8b. Note that the ensemble average embrittling potency shows strengthening at the GB plane, indicating a reduced energetic preferences for

Cr to be at the free surface, which corroborates with first principles calculations by Startt et al. [29,30].

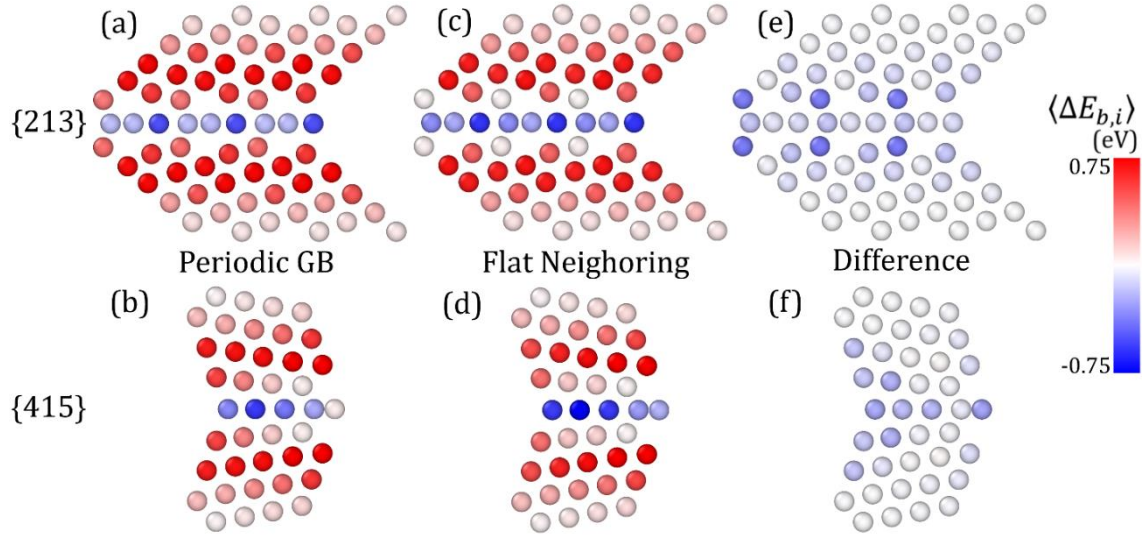


Figure 8: The ensemble average embrittling potency $\langle \Delta E_{b,i} \rangle$ in eV for each site i in the (a) $\Sigma 7 \{213\}$ STGB and the (b) $\Sigma 21 \{415\}$ STGB for the periodic GB ensemble. Sites at the GB plane have a negative $\langle \Delta E_{b,i} \rangle$ (strengthening) while all other sites surrounding the GB layer have a positive $\langle \Delta E_{b,i} \rangle$ (embrittling). In these examples, $k = 250$ for the purpose of weighting site embrittling potencies. (c) and (d) show the flat neighboring ensemble for $\Sigma 7 \{213\}$ and $\Sigma 21 \{415\}$ STGBs respectively. (e) and (f) show the difference between the flat neighboring and periodic GB ensembles for each site.

Recall, in the periodic GB ensemble, we repeat the free surface topography decisions for each unit cell in the model, assuming that the topography adjacent to the segregation region is unimportant. To evaluate the impact of that assumption, we construct the flat neighboring ensemble, where the tortuous segregation region lies between flat neighboring cleavage planes above (or below) the GB, as discussed in Section 2.3. Fig. 8c-8d show results for the flat neighboring ensemble. Lastly, Fig. 8e-8f show a comparative plot which takes the difference between the flat crack ensemble and original ensemble results. The greatest difference in magnitude is at the GB plane or the plane of atoms adjacent to the GB plane. Notably, the differences are mostly negative, suggesting that the flat neighboring ensemble

shows more strengthening behavior. Differences for some sites were as large as -0.2 eV. Ultimately, evaluation of this additional ensemble provides an error estimation for the embrittling potencies. This may warrant future studies devoted to understanding the impact of distant fracture surface topography on segregation energies and embrittling potencies.

It is important to understand the impact of different ensembles of free surface topographies (represented by different values of k) on the ensemble average embrittling potency. Figure 9 shows that as the size of the ensemble changes to include higher-energy free surfaces, the average site behavior can vary dramatically. In Fig. 9 the size of the ensemble is computed as the sum of free surface weights, $\sum w_j$, providing an “effective” number of free surfaces. Note, the prediction for sites 2, 5, and 8 in the $\Sigma 7 \{213\}$ STGB and site 5 in the $\Sigma 21 \{415\}$ STGB can be either embrittling or strengthening depending on the number of free surfaces included in the ensemble. As $k \rightarrow +\infty$ and the ensemble reduces to only the minimum energy free surface, we obtain the embrittling potency that would be reported in traditional segregation studies which rely only on the assumption of a single free surface. However, including higher-energy free surfaces in the ensemble, represented by lower k values, shows a broader spectrum of embrittling behavior. In general, how the weighted averages change based on the size of the ensemble is unique to each site type, but it is clear from Fig. 9 that the inclusion of data from non-minimum energy free surfaces is important to the evaluation of embrittlement behavior. Note the choice $k = 250$ corresponds to 73 effective free surfaces considered for the $\Sigma 7 \{213\}$ STGB and 6 effective free surfaces considered for the $\Sigma 21 \{415\}$ STGB.

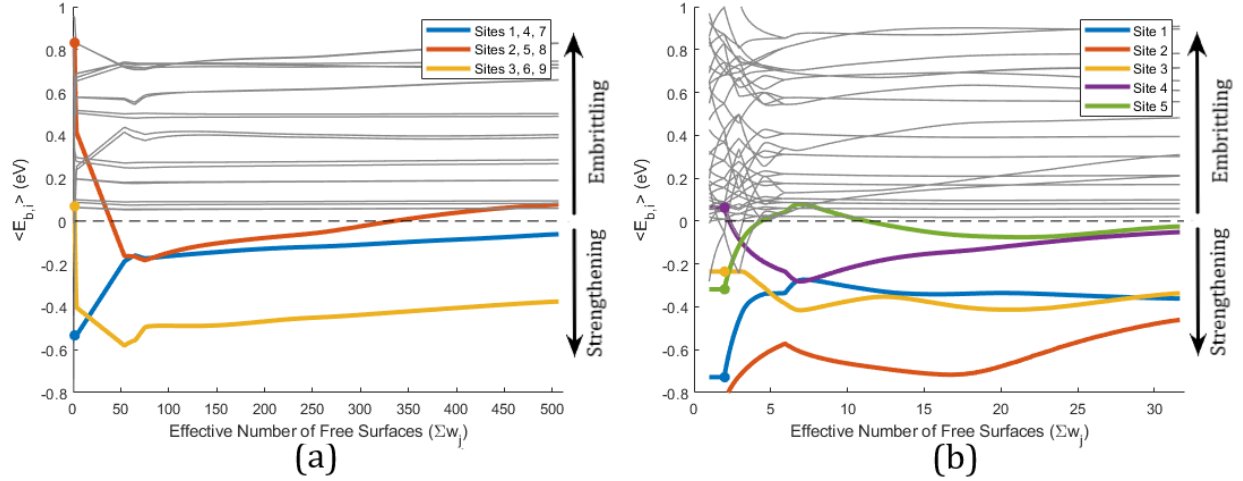


Figure 9: Ensemble average embrittling potencies for (a) sites 1-9 in the $\Sigma 7 \{213\}$ STGB and (b) sites 1-5 in the $\Sigma 21 \{415\}$ STGB calculated via Eq. (6) plotted against effective number of free surface topographies. Note that for $k = 250$, the $\Sigma 7 \{213\}$ STGB has 73 effective surfaces, while the $\Sigma 21 \{415\}$ STGB has 6. Grey lines mark the trajectories of sites off the GB, which are generally embrittling as corroborated by Fig. 8.

4.3. Site Occupancy Probability

To average the $\langle \Delta E_{b,i} \rangle$ into a singular term for each GB, we need to consider occupancy probability for each GB site. Though the at.% occupancy X_i^{GB} provides insight into what sites are occupied, it does not represent the overall probability of a site being occupied. We can compute the site occupancy probability P_i^{GB} as

$$P_i^{GB} = F_i^{GB} X_i^{GB} \quad (7)$$

The density of states for interface sites F_i^{GB} can be computed if the number of interface sites of type i , denoted N_i^{GB} , is known. Interface site types are distinguished only by their segregation energy $\Delta E_{seg,i}^{GB}$ [21], thus,

$$F_i^{GB} = \frac{N_i^{GB}}{\sum N_i^{GB}} \quad (8)$$

Usually, site occupancy is computed for a wide variety of GBs to emulate a polycrystal [16,21,53]. When using many GB sites, the density of states becomes a nearly continuous function of the segregation energy. However, when considering only two GBs in isolation, as is done in this work, we must treat the density of states discretely. To account for this, we simply consider the density of states as a constant value $F_i = \frac{1}{N}$ where N is the number of unique GB sites analyzed. The width of the GB, and hence N , is defined quantitatively using the segregation energy-based criterion detailed in Section 3.1. This is advantageous over a common neighbor analysis (CNA) approach [54], as CNA does not reveal all sites which impact the properties of the GB.

Importantly, note that Eq. 7 provides $X_i^{GB} \cong P_i^{GB} \cong 0$ for sites where $\Delta E_{i,seg}^{GB} \geq 0$, i.e., the probability of an anti-segregation site being occupied is zero. Hence, when mean GB embrittlement is determined in Section 4.4, anti-segregation sites do not contribute to the mean, regardless of the value of $\langle \Delta E_{b,i} \rangle$.

4.4. Mean Grain Boundary Embrittlement

The ensemble average embrittling potency $\langle \Delta E_{b,i} \rangle$ provides a weighted average embrittling potency for each GB site. To develop a metric that describes the embrittling condition of the whole GB, we perform a sum weighted by site occupancy and normalized by the area of the unit cell A_{GB} ,

$$\langle \Delta E_b^A \rangle = \frac{\sum_i \langle \Delta E_{b,i} \rangle P_i^{GB}}{A_{GB}} \quad (9)$$

Here, the A superscript indicates normalization by area. $\langle \Delta E_b^A \rangle$ has units of energy per unit area and accounts for free surface energetics, site occupancy, and structural properties

of the GB. In this work, we refer to this term as the *Mean Grain Boundary Embrittlement*. The inspiration for this term comes from the Rice-Wang equation [55], which considers coverage as the number of solutes per unit area. To explain the development of Eq. (9), consider other possible formulations of $\langle \Delta E_b \rangle$. One formulation considered was a weighted sum normalized by the site occupancy,

$$\langle \Delta E_b^P \rangle = \frac{\sum_i \langle \Delta E_{b,i} \rangle P_i^{GB}}{\sum_i P_i^{GB}} \quad (10)$$

Here, the P superscript indicates normalization by the site occupancy. In this approach, the mean GB embrittlement has units of energy. To assess Eq. (10), consider a hypothetical GB with a single unique site type. Suppose we have two different alloying cases, such that case 1 has a low P_i^{GB} and positive, high magnitude $\langle \Delta E_{b,i} \rangle$, and case 2 has a high P_i^{GB} and positive, low magnitude $\langle \Delta E_{b,i} \rangle$. For the sake of this example, suppose that $\langle \Delta E_b^A \rangle^{(1)} = \langle \Delta E_b^A \rangle^{(2)}$, or otherwise stated, $\sum_i \langle \Delta E_{b,i} \rangle^{(1)} P_i^{GB(1)} = \sum_i \langle \Delta E_{b,i} \rangle^{(2)} P_i^{GB(2)}$ with $i = 1$. Eq. (9) suggests that these two cases result in identical embrittlement of the GB, which agrees with the Rice-Wang equation. However, since $P_i^{GB(1)} < P_i^{GB(2)}$, normalizing by the sum of P_i^{GB} as in Eq. (10) would result in case 1 being more embrittling than case 2. While it is true that case 1 has a higher embrittling potency for its site, case 2 overcomes the lower embrittling potency with higher occupancy. Hence, energetically, these two cases should be represented identically if a metric is to be used as a comparative term between GBs and clearly Eq. (10) cannot be used for this purpose.

To justify the area normalization in Eq. (9), consider the weighted sum with no normalization,

$$\langle \Delta E_b \rangle = \sum_i \langle \Delta E_{b,i} \rangle P_i^{GB} \quad (11)$$

To test Eq. (11), consider an example with two GBs which differ in size and structure. In this case, assume that $\sum \langle E_{b,i}^{(1)} \rangle P_i^{GB(1)} < \sum \langle E_{b,i}^{(2)} \rangle P_i^{GB(2)}$ and $A_{GB}^{(1)} < A_{GB}^{(2)}$, and also that $\frac{\sum \langle E_{b,i}^{(1)} \rangle P_i^{GB(1)}}{A_{GB}^{(1)}} = \frac{\sum \langle E_{b,i}^{(2)} \rangle P_i^{GB(2)}}{A_{GB}^{(2)}}$. Since the embrittlement effect in the Rice-Wang equation requires both embrittling potency and segregant coverage per unit area, we expect both cases to be equally embrittling over the same area. However, if we compare these GBs using Eq. (11), then case 2 would be more embrittling than case 1, which contradicts the motivation the Rice-Wang equation provides.

According to these hypothetical examples that consider both chemically and geometrically variate cases, if the objective is for $\langle \Delta E_b \rangle$ to be a term that characterizes the effect of a solute on a particular GB and provides fair comparison between two GBs, then it is necessary use Eq. (9) with area normalization. While Eq. (10) more closely resembles the original definition of embrittling potency (as a term with units of energy), the normalization deleteriously changes the way occupancy impacts the average. Similarly, Eq. (11) provides a sum of the embrittling potencies over a GB unit cell, but GBs with larger unit cells would appear to have larger embrittlement magnitude when compared against GBs with smaller unit cells if not area normalized.

Thus, using the segregation and FS energies generated by atomistic calculations in this study, we find that $\langle \Delta E_b \rangle = 1.018 \times 10^{-3} \text{ eV}/\text{\AA}^2$ for the $\Sigma 7 \{213\}$ STGB and $\langle \Delta E_b \rangle = 1.482 \times 10^{-3} \text{ eV}/\text{\AA}^2$ for the $\Sigma 21 \{415\}$ STGB. Thus, the $\Sigma 21 \{415\}$ STGB is predicted to be

45.7% more embrittled by the presence of Cr solutes than the $\Sigma 7 \{213\}$ STGB when averaged over the ensemble of many free surfaces and considering differences in GB area.

Recall from Fig. 5 that the black dots represent the embrittling potency of sites considering only the lowest-energy free surface pair, where $k \rightarrow +\infty$. Previous studies have only considered this data point and have ignored the influence of the intergranular fracture surface topography. Suppose we calculate a mean GB embrittlement using only values for the minimum energy free surface for the $\Sigma 7 \{213\}$ STGB. This results in $\Delta E_b^A = 1.778 \times 10^{-3} \text{ eV}/\text{\AA}^2$. This value overpredicts the $k = 250$ ensemble average value of $\langle \Delta E_b^A \rangle = 1.018 \times 10^{-3} \text{ eV}/\text{\AA}^2$ by 75%. Similarly, if we calculated the mean GB embrittlement for the $\Sigma 21 \{415\}$ STGB using only the minimum energy free surface, this results in $\Delta E_b^A = 1.878 \times 10^{-3} \text{ eV}/\text{\AA}^2$, which overshoots the $k = 250$ ensemble average value by 27%. Although it still predicts the same relative relationship, namely that the $\Sigma 21 \{415\}$ STGB is more embrittled by Cr solutes than the $\Sigma 7 \{213\}$ STGB, using the minimum energy fracture surface only predicts a less significant discrepancy of 5.6%, whereas the discrepancy is 45.7% in the ensemble case for the selected value of k . Note that the relative difference is not preserved for different k , and may change depending on the ensemble.

These results highlight the significance of considering a statistical perspective when studying solute induced intergranular embrittlement. The value for k here is selected as an example considering the free surface energy range, but for any ensemble where the $k \neq \infty$, as shown in Fig. 9, there will be a difference between the embrittlement for the minimum energy free surface and the embrittlement for an ensemble of free surfaces. This ensemble

perspective is more inclusive of different intergranular fracture surface topographies that are observed during experiments and atomistic simulations.

5. Conclusions

In this work, we propose a statistical perspective on solute induced GB embrittlement by considering an ensemble of free surfaces in the analysis. Specifically, atomistic simulations of Cr segregation at Ni $\Sigma 7 \{213\}$ and $\Sigma 21 \{415\} \langle 111 \rangle$ STGBs are performed to compute distributions of embrittling potencies for each GB site generated by sampling different free surface topographies. Statistical mechanics principles are used to condense the resulting distributions of embrittling potencies into a singular value for each GB site, called the ensemble average embrittling potency. These embrittling potencies are then weighted by the site occupancy and averaged over the unit cell area to provide a measure of mean GB embrittlement. Ultimately, the ensemble perspective provides a new term to compare the embrittling effects of solutes between different GBs (Eq. (9)), which is then used to assess the average embrittlement effect of Cr segregation at Ni $\Sigma 7 \{213\}$ and $\Sigma 21 \{415\} \langle 111 \rangle$ STGBs. All calculations are performed within the dilute limit, ignoring solute-solute interactions.

Previously, studies of embrittling potencies have generally ignored the impact of free surface topography formed during fracture and have treated embrittling potency as a singular value for a selected GB. This work demonstrates that an ensemble of free surfaces is necessary to completely understand embrittlement; there is a spectrum of embrittlement behavior exhibited by each GB site. The distributions of embrittling potency demonstrate cases where a site can be either strengthening or weakening depending on the free surface

topography. In addition, the magnitude of the embrittlement or strengthening effect of a solute depends on the free surface considered.

Acknowledgements

M.F. and D.E.S. acknowledge support provided by Sandia National Laboratories and the University of Florida. R.D. is supported by the United States (U.S.) Department of Energy (DOE) Office of Basic Energy Sciences (BES), Department of Materials Science and Engineering. The authors acknowledge University of Florida Research Computing for providing computational resources and support that have contributed to the research results reported in this publication. The authors thank Dr. Chongze Hu for providing useful comments on a draft of this manuscript. This work was performed, in part, at the Center for Integrated Nanotechnologies, an Office of Science User Facility operated for the U.S. Department of Energy. Sandia National Laboratories is a multi-mission laboratory managed and operated by National Technology and Engineering Solutions of Sandia, LLC., a wholly owned subsidiary of Honeywell International, Inc., for the U.S. Department of Energy's National Nuclear Security Administration under Contract No. DE-NA0003525. This paper describes objective technical results and analysis. Any subjective views or opinions that might be expressed in the paper do not necessarily represent the views of the U.S. Department of Energy or the United States Government.

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