Comparing calculated and measured grain boundary energies in nickel

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Abstract

Recent experimental and computational studies have produced two large grain boundary energy data sets for Ni. Using these results, we perform the first large-scale comparison between measured and computed grain boundary energies. While the overall correlation between experimental and computed energies is minimal, there is excellent agreement for the data in which we have the most confidence, particularly the experimentally prevalent Σ3 and Σ9 boundary types. Other CSL boundaries are infrequently observed in the experimental system and show little correlation with computed boundary energies. Because they do not depend on observation frequency, computed grain boundary energies are more reliable than the experimental energies for low population boundary types. Conversely, experiments can characterize high population boundaries that are not included in the computational study. Together the experimental and computational data
provide a comprehensive catalog of grain boundary energies in Ni that can be used with confidence by microstructural scientists.

1. Introduction

Grain boundary free energy provides the driving force for grain growth in polycrystalline materials, and as Cyril Smith recognized over sixty years ago, variations in grain boundary energy alter the kinetics and morphology of microstructural evolution [1]. Because of its importance, many investigators have attempted to survey grain boundary energies using various experimental techniques [2-16], but their data sets were limited due to the difficulty of measuring accurate interfacial energies in a bulk solid. Computational models have also been applied, c.f. reviews in [17-21], but computational restrictions limit data sets primarily to high symmetry boundaries.

Two recent studies have changed this situation. Researchers at Carnegie Mellon University (CMU) [22] and Sandia National Laboratories (SNL) [23] have utilized new, high-throughput methods to measure energy for large ensembles of grain boundaries. Taken together, these studies increase our database of grain boundary energy measurements by orders of magnitude.

The CMU and SNL studies differ in approach – experimental versus computational – and also in sources of uncertainty and error. In this paper, we will compare the CMU and SNL data sets for grain boundary energy in Ni. Our goal is to perform the first large-scale comparison between measured and computed grain boundary energies, in order to develop a comprehensive, validated, well-characterized database of grain boundary energies in Ni that can be used with confidence by microstructural scientists.
2. Method

2.1. Experimental grain boundary energy measurements

2.1.1. Methodology

An experimental method for measuring a large number of relative grain boundary energies in a polycrystalline microstructure has previously been described in detail [24] and applied to MgO [25] and Ni [22] polycrystals. This method presumes that all grain triple junctions are in equilibrium so that each satisfies the Herring condition [26] (i.e. balance of surface tensions and torques). If that is the case, then each triple junction may be characterized by an equation with four (vector) unknowns: three capillarity vectors and the triple junction line vector [27, 28]. Since the triple junction line vector may be measured directly, we are left with N equations (one for each triple junction) and 3N unknowns (one for each grain boundary).

In order to solve this system of linear equations, we must decrease the number of unknowns. To do this, we make the assumption, first articulated by Brandon [29], that grain boundaries that are close in crystallographic space are similar in energy. Thus, we can group boundaries in energetic bins. Presuming all boundaries in a given bin have the same energy, we reduce the 3N unknowns sufficiently to permit solution of the system of equations using an iterative optimization procedure [24]. The result is a set of capillary vectors that can be transformed into grain boundary energies if the grain boundary tangent and normal vectors are known. Because boundary energy is derived from triple junction geometry, the measurement yields boundary energies that are relative to one another, not absolute magnitudes.

Applying this method requires characterizing a large number of equilibrated triple junctions in three spatial dimensions and five crystallographic dimensions. While this would have been prohibitively difficult even a decade ago, new automated techniques now make it tractable. Four of the five macroscopic variables of grain boundary crystallography can be measured efficiently using EBSD and the fifth boundary
orientation parameter, which also gives the triple junction geometry, can be observed using FIB-based serial sectioning and reconstruction techniques. The synthesis of these two methods has yielded large data sets of fully-characterized grain boundary ensembles that were analyzed to give the grain boundary character distribution (GBCD) and relative grain boundary energy distribution (GBED) in MgO [25, 30] and Ni [22].

2.1.2. Sources of error and uncertainty

To compare these data with other studies, it is important to be aware of sources of non-systematic error (uncertainty) and systematic error (bias). If triple junctions are not in equilibrium, then the Herring condition will not be fulfilled, and calculated energies will be inaccurate for all boundary types. We ensure equilibrium by a careful anneal schedule. Non-systematic errors also arise during the grain boundary reconstruction process, as well-described in [30]; again, these errors should not vary with boundary type but should provide uniform error bars.

The requirement that grain boundaries be binned into groups provides a source of systematic error. Because low-energy cusps are very localized in crystallographic space, low energy boundaries are inevitably binned with nearby, higher energy boundaries; thus, the experimental measurement method cannot resolve cusps and over estimates the energies of low energy boundaries. Adjusting the bin width alters, but cannot eliminate, this error because a finite bin width is necessary for the energy calculation to converge.

Finally, the energy measurements are not uniformly distributed among each grain boundary bin. For example, in the Ni specimen a single boundary type, the coherent twin, makes up almost 30% of the boundaries observed, and the Σ9 tilt boundary type contributes almost 10%. On the other hand, about 15% of the boundary bins contain fewer than five energy measurements. This nonuniform sampling has two effects: First, calculated energies will be less accurate for less prevalent boundary types simply due to poorer statistics. Second, in the optimization algorithm used to calculate energy, there will be a preference to optimize variables that occur often and thus affect many of the N equations at the expense of variables that affect few equations. The result is that the error
increases (perhaps more than predicted by statistical analysis) as the observation frequency decreases.

While these errors are difficult to quantify individually or collectively, it is certainly reasonable to expect error bars on the order of 10%, with larger error bars for less frequently observed boundaries.

2.2. Computational method

2.2.1. Methodology
The computational method for constructing a large catalog of grain boundaries and measuring their energy has been described in detail elsewhere [23, 31]. To avoid bias towards particular boundary types, the method examines all planar boundaries that can be constructed within a periodic cell of a specified maximum size. Having determined which pairs of grain orientations can fit inside the simulation cell, we construct a multiplicity of boundary structures for each orientation pair. We first translate one crystal relative to the other by several offset vectors. For each of the resulting structures, we place the boundary plane at each non-equivalent position. For each of these structures, we remove any overlapping atoms using several different criteria. Finally, we use molecular statics to minimize the T=0K free energy (i.e. the enthalpy) of each distinct structure. For typical boundaries we minimize several hundred to several thousand configurations; the lowest energy structure is presumed the equilibrium structure.

The material model for these calculations is an embedded atom method (EAM) empirical interatomic potential function [32], parameterized to represent Ni [33]. This particular potential function predicts the stable and unstable stacking fault energies quite accurately, which suggests it is reasonable to use to examine planar defects such as grain boundaries.

2.2.2. Sources of error
Errors in calculated energies can arise from the boundary construction or minimization processes. While the boundary construction algorithm produces a large number of candidate structures for each grain orientation pair, there is no guarantee that the global
minimum energy structure is among them. However, most boundaries evince a large number of nearly degenerate, nearly minimum energy configurations without evidence of low-energy outliers. This suggests that most minimum energy structures are at least reasonable approximations to the true minimum.

Because the boundaries are constructed in a relatively small periodic box, high symmetry boundaries (which often have low energy) are over-represented in the sampling; while this does not induce error in the energy measurements themselves, it does bias the grain boundary distributions toward high symmetry boundaries.

The molecular statics energy minimization calculates the excess enthalpy, rather than the free energy, of the boundaries. It is known that entropic contributions cause the boundary free energy to decrease with temperature by about a factor of three from 0K to near the melting temperature [34-36]. It has been presumed that this decrease is similar for most boundaries, and recent studies support this viewpoint, suggesting that free energy scales with the elastic constants up to about 0.75 of the melting temperature [37]. If most boundaries vary with temperature in a similar manner, then the measured enthalpies can be interpreted as relative (but not absolute) free energy values. However, there is the potential that some boundaries do not scale like their peers, perhaps due to structural transformations. The free energy of such boundaries might be poorly represented by their enthalpies. We have no way to determine which, if any, boundaries are so affected.

Finally, as with all empirical atomistic simulations, errors can arise from the physical model for atomic interactions, i.e. the EAM interatomic potential function. Different potential functions for the same material can give substantially different property values, depending on which parameters were used to fit the function. For calculating grain boundary energies, we select a potential well suited to planar defects. Even so, defect energy accuracy is typically limited to about 0.1 eV, and accuracy may vary with boundary type.
While these sources for error are difficult to quantify, typical error bars of 10% seem reasonable, given the agreement between simulated and measured stacking fault energies.

2.3. Comparing experimental and computational data

The experimental data set for polycrystalline Ni grain boundaries is described in [22]; 37,000 triple junctions were analyzed to provide approximately $10^5$ grain boundary free energy measurements. The grain boundaries were binned with $8.2^\circ$ resolution such that there are 17,894 discrete grain boundary types. Computational results indicate that the energies of boundaries within about $10^\circ$ of each other are well correlated [23, 31], so the $8.2^\circ$ bin size is reasonable. It is worth noting that the experimental data set includes both the distribution of grain boundary types in crystallographic space (i.e. the GBCD) and the extracted energy distribution (i.e. the GBED). The GBCD, which is measured directly, is considered to be more accurate than the GBED, which is a derived quantity.

The computational data set for Ni bicrystal boundaries is described in [23, 31] and is publicly available as online supplemental material to that paper; 388 distinct grain boundaries were constructed and characterized. To compare these boundaries to the experimental data, we extract the rotation angle and axis and boundary plane for each and apply all possible FCC symmetry operators. For each representation, we identify the appropriate grain boundary type (i.e. bin) in the experimental system and read the energy. The average energy for all the representations is taken as the relevant experimental energy, to be compared to the calculated value.

3. Results and discussion

3.1. All boundaries

Figure 1(a) compares the experimental and calculated grain boundary energy for all 388 boundaries in the computational data set. Taken in aggregate, the calculated and measured energies show little correlation (unweighted correlation coefficient $R_U = +0.18$) and the linear curve fit is unconvincing. For both sets, the lowest energy boundary is the coherent twin ($60^\circ/[111]$, pure twist) and, in general, boundaries with higher measured
energies have higher calculated energies. However, the distribution is broad and there are a significant number of outliers.

A challenge to interpreting this data arises from the unknown error bars. As discussed above, we expect the uncertainty in the experimental grain boundary energies to increase as the observation frequency decreases. The boundary population per bin, P, varies widely in the experimental data set, from 0.12 to 4500 times the population we would expect in a purely random system (termed multiples of a random distribution, MRD). If we weight the experimental energy measurements by P, we obtain a linear fit to the calculated energies with a much higher correlation coefficient (weighted correlation coefficient $R_W = +0.92$), as shown in Figure 1(b). The curve fit is intuitively more pleasing and fits the low energy boundaries, where we expect the most accurate data, especially well. Thus, the correlation between experimental and calculated grain boundary energy is very strong for the data in which we have the most confidence.

It is interesting to observe that the boundaries form three distinct and separate clusters in population space, as shown in Figure 1(b). The high population bins, with $P > 60$ MRD, each contain 300 or more measurements, and the curve fit is heavily weighted toward these boundary types; this group includes only the $\Sigma 3$ boundaries. The mid population bins, with $4 < P < 20$, contain 20 or more measurements per bin and include only $\Sigma 9$ boundaries. Intriguingly, they seem to suggest a different linear fit between experiment and simulation; the implications will be discussed below. The low population bins, with $P < 2.5$, contain fewer than 16 measurements each, and most of these bins contain fewer than 5 measurements. We would not expect the data quality to be high for these sparse bins, and indeed, there is substantial scatter about the linear fit.

Because high-energy boundaries are preferentially eliminated during grain growth (c.f. [38-40]), we expect that low-energy boundary bins will be more populated than high-energy bins in the experimental data. Figure 1(b) confirms this trend, although there are outliers.
3.2. **$\Sigma 3$ and $\Sigma 9$ boundaries**

Coherent and incoherent $\Sigma 3$ boundaries are the most frequently observed in the experimental system, comprising almost 40% of the total grain boundary length. Because of their high symmetry, they are also well represented in the computational data, where 41 of the 388 simulated boundaries are of $\Sigma 3$ type. Figure 2(a) shows experimental and computed energies for $\Sigma 3$ boundaries; as these were the most strongly weighted data in the overall fit, they unsurprisingly show even stronger correlation when considered separately ($R_U = 0.71$, $R_W = 0.95$). There are two apparent outliers, a (10 2 2)/(111) and an (811)/(544) boundary, circled in Figure 2(a). However, neither boundary corresponds to a local minimum in the relative boundary area distribution, as would be expected for a high-energy boundary. Combined with the fact that they are not consistent with the calculated energies, we consider these data questionable. With the two outliers excluded, the correlation improves slightly ($R_U = 0.80$, $R_W = 0.98$).

In polycrystals with many $\Sigma 3$ boundaries, there is a tendency for $\Sigma 3$ boundaries to form $\Sigma 3$ - $\Sigma 3$ - $\Sigma 9$ triple junctions (c.f. [41]), which was confirmed in this system [22]. $\Sigma 9$ boundaries are the second most prevalent boundaries in the experimental data set, and 23 of the 388 simulated boundaries are $\Sigma 9$ type. However, the computational boundary set included only four of the most experimentally prevalent $\Sigma 9$ boundaries, the low-energy $\Sigma 9$ tilt boundaries. Figure 2(b) shows the experimental and computed energies for $\Sigma 9$ boundaries. There is a modest correlation ($R_U = 0.43$, $R_W = 0.48$), which we would expect to greatly improve if the more populated $\Sigma 9$ boundary types were included in the comparison.

In twinned or grain boundary engineered materials, the $\Sigma 3$ and $\Sigma 9$ boundaries often form a network of boundaries and triple junctions that is relatively distinct from the high angle boundary network [41]. This implies that the triple junction equilibrium equations, which are minimized to calculate energy, may be divided into two disjoint types: $\Sigma 3$ - $\Sigma 3$ - $\Sigma 9$ triple junctions and other boundary junctions. The two groups have quite different computational characteristics, and it is possible that the optimization algorithm optimizes
each separately, assigning different relative energies to each set. This possibility requires further study.

3.3. **Other boundaries**

Other CSL boundary types, including $\Sigma 5$, $\Sigma 7$, $\Sigma 11$ and $\Sigma 15$, are well represented among the simulated boundaries. However, few of the simulated CSL boundaries are observed in the experimental data set at even the random frequency (i.e. $P < 1$ MRD for nearly all boundary types). As expected for low population boundaries, we see at best modest correlation between the experimental and computational energies for these boundaries. Furthermore, the slopes of the linear fits vary widely with boundary type, indicating that the correlation between data sets does not arise from proportionality between experimental and calculated energy.

It is interesting to consider how many additional experimental measurements would be required to resolve the energies of infrequently observed boundaries. Observation frequencies of high population boundaries are about two orders of magnitude higher than low population boundaries. Thus, to achieve confidence in low population boundary energies comparable to that of high population boundary energies would require about $10^2$ times more measurements than in the current data set. In this Ni system, that implies about $4 \times 10^6$ triple junction characterizations, which is currently an intractable size. In other materials, different boundary populations may require different system sizes. For example, because Al polycrystals lack the $\Sigma 3$ and $\Sigma 9$ boundary subnetwork seen in Ni, the boundary population distribution may be more uniform in Al, permitting smaller experimental system sizes.

Because the computed energies are not dependent on an observation frequency, they should be equally valid for low population boundaries. Furthermore, the good correlation between experimental and computed energies for high population boundaries suggests that the computed energies are physically realistic and representative. Thus, our conclusion is that the computed grain boundary energies are more reliable than the
experimental energies for low population boundary types and can be used to determine energies of boundaries that are experimentally inaccessible.

On the other hand, the experimental studies indicate that certain commonly simulated boundaries are not present in significant numbers in real microstructures and cannot be considered typical boundaries. For example, a number of computational studies have used the \( \Sigma 5 \) symmetric tilt boundary as a model high angle boundary (c.f. [42–44]). The GBCD data for Ni indicate that this boundary is rarely observed, with a population \( P < 0.4 \) MRD. Likewise, some frequently observed boundaries, such as most of the \( \Sigma 9 \) tilt boundaries, were not included in the computational boundary catalog. Clearly, experiments can guide computational boundary selection to ensure relevant and important boundary types are examined.

### 3.4. Boundary population correlations

While the grain boundary energy distribution is extracted from the grain boundary character distribution along with other geometric measurements, the GBCD is measured directly and has fewer sources for error. Both experimental [22, 25] and computational [38, 39] studies have indicated that the population of grain boundaries in a polycrystal is inversely proportional to their energies, or the GBCD scales with the reciprocal of the GBED. Because the GBCD is a potentially more reliable experimental measure, it is possible that the computed GBED correlates more strongly with the experimental GBCD than with the experimental GBED.

Figure 3 compares the experimental GBCD with the computational GBED. Note that because boundary populations vary over a wide range, the GBCD is plotted as \( \ln(P) \). The three population groups described above are indicated in the figure; they are clearly distinct groups. Although a linear fit to the complete data set shows modest correlation \( (R_U = 0.45) \), the actual line is placed between the high \( P \) and low \( P \) boundary clusters and bears little resemblance to the data set. However, the linear fit to the high \( P \) \( \Sigma 3 \) boundary data gives a good fit and a much higher unweighted correlation coefficient \( (R_U = 0.96) \) than observed in the GBED comparison. (Note that in these fits, we do not weight by
population, since population is the dependent variable.) In addition, the outliers that were present in the GBED data are absent here, again indicating that those energy measurements are spurious. The Σ9 fit is also improved relative to the GBED comparison, with $R_U = 0.69$. Although the low P boundary population data is more tightly grouped and seems to show a clearer trend with energy, the GBCD remains poorly correlated with calculated GBED ($R_U = 0.24$), reemphasizing the dependence of data quality on observation frequency.

Overall, the experimental GBCD data provides a better correlation to the computed GBED. Since the GBCD is more straightforward to characterize experimentally than the GBED, it offers a more accessible metric for comparison to computed boundary energies. However, the improved correlation cannot compensate for the lack of observation data in low P boundary bins. For those boundaries, the computed energy remains the more reliable than the experimental data.

4. Conclusions

Recent independent experimental and computational studies have produced two large sets of grain boundary energy data for Ni [22, 23, 31]. The data were obtained by different approaches with distinct sources of error and uncertainty. The experimental data set contains a large number of relative grain boundary free energies binned into 17,894 boundary types. The computational data set includes only 388 boundaries, and the enthalpy of each boundary is characterized individually. Our comparisons between the experimental and computed boundary energies suggest several conclusions:

1. The unweighted correlation between experimental and computed grain boundary energies is minimal.
2. There is a strong correlation between the experimental and calculated energies when the fit is weighted by the experimental observation frequency. Because the quality of the experimental data scales with the observation frequency, we conclude that there is excellent agreement between experiment and computation for the data in which we have the most confidence.
3. For Σ3 boundary types, which are the most frequently observed boundaries in the experimental polycrystal, the correlation between experiment and computation is even stronger.

4. The next most populous boundaries, the Σ9 boundaries, show a modest correlation between experiment and computation. Other CSL boundaries are infrequently observed in the experimental system and show little correlation with computed boundary energies.

5. Boundary population can be used to determine the experimental system size required to achieve a desired accuracy in energy measurements. Different materials systems will require different system sizes.

6. Because they do not depend on observation frequency, computed grain boundary energies are more reliable than the experimental energies for low population boundary types and can be used to determine energies of boundaries that are experimentally inaccessible.

7. The computational data set excludes some frequently observed boundaries and includes some infrequently observed boundaries. Experiments should guide future computational studies to ensure relevant and important boundary types are examined.

8. The grain boundary character distribution, which can be measured more easily than the energy distribution, offers a strong inverse correlation to computed grain boundary energies and can be used in place of the grain boundary energy distribution in comparisons with computed energies.

Overall, the experimental and computational results validate each other for boundaries that are appropriately represented in both data sets. By understanding the limitations of each method, we can combine experimental and computational data to achieve a more comprehensive catalog of grain boundary energies in Ni that can be used with confidence by microstructural scientists.
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6. Figures

Figure 1. The relationship between experimental and calculated grain boundary energy in Ni. (a) An unweighted linear fit to all the data (solid line) shows little correlation and is not a good representation of the data. (b) A linear fit weighted by boundary population, P, shows much higher correlation and is a better representation of the data. Note that the data fall into three population groups: P > 60 (red diamonds), 4 < P < 20 (blue squares), and P < 2 (green circles).
Figure 2. The relationship between experimental and calculated grain boundary energy for CSL grain boundaries in Ni. (a) $\Sigma 3$ boundaries show a strong correlation between experimental and calculated energy. The two circled points are outliers in the experimental energy distribution that do not correspond to minima in the experimental population distribution, so are considered questionable. (b) $\Sigma 9$ boundaries show a weaker correlation between experimental and calculated energy.
Figure 3. The relationship between experimental GBCD and calculated GBED in Ni. For high population $\Sigma 3$ and mid population $\Sigma 9$ boundaries, the inverse correlation between GBCD and GBED (solid lines) is stronger than the direct correlation between experimental and calculated GBEDs. However, the low population boundaries remain poorly correlated, due to high experimental uncertainty.
7. References

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