Grain boundary energies and cohesive strength as a function of geometry

Valerie R. Coffman and James P. Sethna
Laboratory of Atomic and Solid State Physics (LASSP), Clark Hall, Cornell University, Ithaca, New York 14853-2501, USA
(Received 6 July 2007; revised manuscript received 27 February 2008; published 23 April 2008)

Cohesive laws are stress-strain curves used in finite element calculations to describe the debonding of interfaces such as grain boundaries. It would be convenient to describe grain boundary cohesive laws as a function of the parameters needed to describe the grain boundary geometry; two parameters in two dimensions and five parameters in three dimensions. However, we find that the cohesive law is not a smooth function of these parameters. In fact, it is discontinuous at geometries for which the two grains have repeat distances that are rational with respect to one another. Using atomistic simulations, we extract grain boundary energies and cohesive laws of grain boundary fracture in two dimensions with a Lennard-Jones potential for all possible geometries which can be simulated within periodic boundary conditions with a maximum box size. We introduce a model where grain boundaries are represented as high symmetry boundaries decorated by extra impurities at the interface, and emission of dislocations to initiate at the site of these microscopic defects, the macroscopic fracture strength of the material is dependent on the microscopic structure of the grain boundaries. The debonding of an interface such as a grain boundary is described by a cohesive law, giving the displacement across the interface as a function of stress (Fig. 1).

Cohesive laws are used by finite element cohesive zone models (CZM), which simulate fracture initiation at interfaces. It has been shown that the shape and scale of the cohesive law has a large effect on the outcome of the finite element simulation. However, the CZM studies of grain boundary fracture have used cohesive laws that are guessed, chosen for numerical convergence, and do not take into consideration the effect of varying grain boundary geometries within the material—the same cohesive law is often used throughout the material despite the fact that in a real material, grain boundaries of varying geometries must occur.

It would be useful to find a formula for the cohesive laws of the grain boundaries of a given material as a function of geometry, for input into finite element simulations. The geometry of a three-dimensional (3D) grain boundary depends on five parameters that describe the orientations of the two grains. In addition, there are three different modes of fracture (normal to the crack plane, shear in a direction parallel to the crack line, or shear in a direction perpendicular to the crack line) to explore as well as dependencies on temperature, impurities at the interface, and emission of dislocations to consider.

Thus far, no systematic study of grain boundary cohesive laws as a function of geometry has been done with molecular dynamics or experiment. There is difficulty in measuring cohesive laws experimentally because it is difficult to isolate and measure the displacements on either side of the grain boundary.

Previous atomistic studies of the mechanical response of grain boundaries have concentrated on a small number of symmetric grain boundaries in three dimensions. Exploring the complete picture of 3D grain boundary cohesive laws involves exploring a five-dimensional space with three modes of fracture. Because of this difficulty, we have taken a step back. We seek to systematically explore the cohesive laws for mode I fracture for all possible grain boundary geometries in two dimensions that can be simulated in periodic boundary conditions for a particular size and strain. We will initially be focusing on symmetric grain boundaries and then expanding the picture to look at asymmetric 2D grain boundaries. One use of such data would be for finite element simu-

FIG. 1. An example of a cohesive law. The stress vs strain curve that describes the debonding of a 2D grain boundary with tilt angles 33.418° and 26.58°. Our measurements include the elastic response of the perfect crystal on either side of the grain boundary. In the results shown above, we have subtracted off the elastic response of the bulk in order to isolate the elastic response of the interface. This is described in detail in Ref. 7 where cohesive laws of grain boundaries are used in finite element simulations of polycrystals.

DOI: 10.1103/PhysRevB.77.144111
PACS number(s): 61.72.Mm, 62.20.M–, 31.15.xv
lations of polycrystals,\textsuperscript{7} as described above. For this purpose, it would be nice to find a functional form which describes the fracture strength of the grain boundary as a function of geometry.

We find, however, that the fracture strength as a function of tilt angles is discontinuous everywhere, with particularly large jumps at special, high symmetry grain boundaries composed of a simple arrangement of structural units with a low repeat distance. These special boundaries are also associated with cusps in the grain boundary energy. We will describe dependence of the energy and the fracture strength near these special boundaries by treating them as perfect, albeit complex, crystals with added dislocations to break the symmetry.

Most studies use the coincidence site model\textsuperscript{13} to construct and classify special grain boundaries\textsuperscript{5,4–11} while a few mention the significance of structural units.\textsuperscript{8,10,14–16} The only 2D study we are aware of, the “bubble-raft” model, observes the structural units of several of the special, high angle grain boundaries for the triangular lattice and how patterns of structural units combine to create vicinal geometries.\textsuperscript{15} Other 3D studies discuss how grain boundaries with geometries close to those of special grain boundaries can be decomposed into the structural unit of the special grain boundary with added flaws.\textsuperscript{14,16} We develop a systematic way of finding high symmetry geometries and show that the combinations of patterns of structural units at vicinal grain boundaries are key to understanding the dependence of energy and fracture strength on geometry.

Sansoz and Molinari find the grain boundary energy by allowing the grains to relax together from an initial separation of a few Å.\textsuperscript{10} while others perform a conjugate gradients search.\textsuperscript{8} We use a systematic method for explicitly imposing a relative shift between the grain and using atomistic relaxation for finding the global energy minimum similar to that used in Refs. 9 and 16. It is well established that cusps exist in the grain boundary energy for special high angle grain boundaries.\textsuperscript{8,10,11,16–20} Recent studies of grain boundary constitutive properties focus on the response to shear\textsuperscript{8,10} or compare shear and tension.\textsuperscript{9,12} Warner \textit{et al.} claim that the tensile response does not depend on the geometry of the grain boundary.\textsuperscript{5} Others have seen jumps in the tensile fracture strength at high angle grain boundaries.\textsuperscript{8,11}

We will explain the cusps in energy and jumps in fracture strength by drawing an analogy between perfect crystals and high symmetry, high angle grain boundaries. The dislocation model of low angle grain boundaries gives a $\theta \ln \theta$ form for the grain boundary energy. Because vicinal grain boundaries can be thought of as high symmetry grain boundaries with added flaws, the energy cusps at special grain boundaries have the same $\theta \ln \theta$ form. A similar argument will apply to the peak fracture stress. Just as adding dislocations to the perfect crystal adds a nucleation point for fracture, and therefore a discontinuity (jump down) in fracture strength, adding a flaw to a high symmetry, high angle grain boundary abruptly changes the local fracture strength by adding a potential nucleation site for fracture.

II. PROCEDURE FOR CALCULATING GRAIN BOUNDARY ENERGY AND COHESIVE STRENGTH

We calculate energies and cohesive strengths for grain boundaries using the DigitalMaterial\textsuperscript{25} package to perform atomistic simulations. The potential we are using for 2D simulations is the Lennard-Jones potential with a smooth, fourth order cutoff between 2.41 and 2.7 Å. The ground state is a triangular lattice with a lattice constant of 1.11 Å.

We set up the grain boundary by initializing two rectangular grains with the given rotations that define the grain boundary geometry we wish to measure. An example of the configuration of atoms that make up a grain boundary simulation is shown in Fig. 2. We use periodic boundary conditions in the $y$ direction in order to avoid edge effects which could potentially cause cracks to nucleate at the intersection with the surface.\textsuperscript{22} This has the disadvantage of only allowing geometries that have finite repeat distances. A method for finding these geometries is discussed in detail in Sec. III B. If necessary, both grains are strained slightly by equal and opposite amounts in order to have both grains fit into a periodic box. We use a constrained layer of atoms to impose fixed boundary conditions in the $x$ direction. These are represented by the darker atoms in Fig. 2. The constrained layer of atoms has a width equal to twice the cutoff distance of the potential to eliminate surface effects from the free atoms.

Besides the tilt angles, there are other factors to consider in constructing the grain boundary geometries. For commensurate grain boundaries, there will be an ideal relative displacement in the direction parallel to the grain boundary (the $y$ direction in Fig. 2). This ideal displacement will correspond to the lowest energy and thus is the most natural configuration for a given pair of tilt angles.\textsuperscript{33}

We find the displacement along the boundary which gives the lowest energy by initializing the two grains with a small displacement in the $x$ direction and varying displacements in the $y$ direction, relaxing the atoms (with the boundary layers constrained to be rigid and nonrotating), and measuring the grain boundary energy per length. The range of displacements in the $y$ direction that we must search over is given by

\[
\Delta = \begin{cases} 
D_1 & D_2 \text{ mod } D_1 = 0 \\
\min(|D_2 - |D_2/D_1||D_1|) & D_2 \text{ mod } D_1 \neq 0 
\end{cases}
\]

(1)

where we assume $D_1 < D_2$. This is illustrated in Fig. 3. The grain boundary energy per length is defined as

FIG. 2. (Color online) An example of a grain boundary. The tilt angles are given by $\theta_1$ and $\theta_2$ and the repeat distances are $D_1 = |(a_1, b_1)|$ and $D_2 = |(a_2, b_2)|$. We have assumed the convention where $0^\circ$ indicates an orientation with Miller indices (2,1) for the vertical face of the grain boundary.
where $E_{\text{total}}$ is the total potential energy for the configuration of atoms (excluding the constrained atoms), $N_{\text{atoms}}$ is the number of unconstrained atoms, $E_{\text{bulk}}$ is the energy of a single atom in the bulk, and $L$ is the length of the grain boundary. An example of the results of such a search is shown in Fig. 4. The regions with the same final displacement and energy correspond to basins of attraction around the finite number of final configurations of atoms. The minimized energy per unit length [Eq. (2)] found by this method is what we record as the grain boundary energy.

We have also tried thermal annealing and have found that for certain geometries, the grain boundary migrates to form a jagged interface with segments of different grain boundaries that collectively have a lower total energy than the geometry given by the original set of tilt angles. The tendency of certain grain boundaries to corrugate is also discussed by Ishida and Pumphrey.\textsuperscript{15,18} As a corrugated grain boundary is torn apart to measure the cohesive law, the corners form stress concentrations which weaken the grain boundary. Despite the fact that this configuration may be more natural, it is not what we intend to measure. We wish to measure the fracture toughness of all possible pairs of tilt angles, even if those pairs of tilt angles happen to be unstable. In a few cases, the minimization procedure described above produced a curved grain boundary. In these cases, we constrained the displacements in the $y$ direction such that a consistent pattern of flaws along a straight grain boundary is achieved.

After finding the ideal $y$ displacement, we increment the strain by displacing the constrained layers of atoms in the $x$ direction, away from the grain boundary. We relax the atoms and measure the force in the $x$ direction per unit length on the constrained layer of atoms. If the measurement of the stress drops abruptly during one strain step, the simulation restores the positions from a previous step, reduces the size of the strain increment, and proceeds. The result of such a simulation is shown in Fig. 1. The maximum stress in the stress strain curve is what we define as the peak stress.

### III. GRAIN BOUNDARY GEOMETRIES

#### A. Lattice symmetries and tilt angles

For the triangular lattice, it is clear that we only need to explore tilt angles between 0° and 60°, but we can further reduce the space of grain boundary geometries to consider. Figure 5 represents the space of tilt angle pairs. Reflecting a point in this space through the $\theta_2 = \theta_1$ line corresponds to swapping the two grain orientations which, as shown in

![Image](144111-3)
Fig. 5, is equivalent to flipping the grain boundary in both the vertical and horizontal directions, resulting in the same grain boundary. Reflecting through the (30,30) point takes \((\theta_1, \theta_2)\) to \((60-\theta_1, 60-\theta_2)\), reversing the sense of rotation of each grain. As shown in Fig. 5, this is equivalent to flipping the grain boundary in the vertical direction. Reflecting through the \(\theta_2=60-\theta_1\) line takes \((\theta_1, \theta_2)\) to \((60-\theta_1, 60-\theta_1)\), both switching the grains and reversing the sense of rotation. This is equivalent to flipping the grain boundary in the horizontal direction, also shown in Fig. 5. Thus we only need to consider the pairs of tilt angles in the triangle enclosed by the \(\theta_1=0\) line, the \(\theta_2=\theta_1\) line, and the \(\theta_2=60-\theta_1\) line—the shaded region in Fig. 5.

B. Finding all possible geometries for periodic boundary conditions

In order to better simulate the grain boundary in the bulk as described in Sec. II, we would like to use periodic boundary conditions along the direction of the grain boundary (y direction). In order to simulate 2D grain boundaries in periodic boundary conditions, we need not only for both grain orientations to have finite repeat distances, but also for the repeat distances to be commensurate with one another. Let \((a,b)\) be the lattice vector that is parallel to the edge of a 2D triangular lattice. Since the basis vectors are at a 60° angle to one another, the repeat distance of a particular orientation of a 2D triangular lattice is given by

\[
D = \sqrt{(a + \frac{1}{2}b)^2 + \left(\frac{\sqrt{3}}{2}b\right)^2} = \sqrt{a^2 + b^2 + ab}.
\]

(3)

The tilt angle is then given by

\[
\theta = \sin^{-1}\left(\frac{a + b/2}{\sqrt{a^2 + b^2 + ab}}\right).
\]

(4)

In order to expand the number of possible geometries, we have also considered geometries for which applying a small strain to each grain allows us to fit both grains inside the same periodic box. For any pair of orientations that have repeat distances that are not commensurate, we can find a continued fraction approximation to the ratios of their lengths, \(p/q = D_1/D_2\). We can strain each grain equally into a box of size \(L = (1/2)(pD_2 + qD_1)\), where the strain required is \(|L - pD_2|/L\). For a strain of 0.05%, the grain boundary would have to be 1000 Å long before the strain would alter the structure of the grain boundary. We find all pairs of tilt angles that correspond to commensurate or near commensurate grain boundaries by looping over pairs of surface vectors and comparing the repeat distances.

Figure 6 shows all possible geometries that can be simulated with a periodic length of 70 lattice constants or less, and a strain of 0.05% or less. The \(\theta_1=\theta_2\) line corresponds to perfect crystals, while the \(\theta_1=60-\theta_2\) line corresponds to symmetric grain boundaries. There is a gap near each of these lines because creating very small angle grain boundaries, or a geometry very close to symmetric grain boundaries, requires adding flaws that are separated by large distances. We see gaps near other high symmetry grain boundaries for the same reason. High symmetry grain boundaries are discussed in Sec. III C. The lines radiating out from high symmetry grain boundaries represent adding single flaws to those high symmetry grain boundaries at larger and larger distances, as you approach the high symmetry boundary. Each line represents a different type of flaw. The symmetry about the \(\theta_1=30\) line is due to the fact that grains with tilt angles \(\theta_1\) and \(60-\theta_1\) have the same repeat distance. Therefore the grain boundaries given by the tilt angles \((\theta_1, \theta_2)\) and \((60-\theta_1, \theta_2)\) have the same overall repeat distance, though these are different grain boundaries.

C. High symmetry grain boundaries

Certain grain boundary geometries have particularly low repeat distances. These grain boundaries have special properties. They mark the center of cusps in the grain boundary energy and discontinuous increases in the fracture strength as a function of tilt angle. Table I shows examples of high symmetry grain boundaries and Fig. 7 shows examples of geometries in between, which constitute adding a single flaw per supercell repeat distance to the high symmetry grain boundary. Note how the added flaws constitute a compromise between the two high symmetry geometries. Furthermore, a less high symmetry grain boundary can be repeated, and again have a single flaw added. Because of this hierarchical procedure for constructing lower-symmetry boundaries, the grain boundary energy and fracture strength as a function of tilt angle will have a self-similar nature.

The coincidence site lattice model (CSL) describes grain boundaries in terms of \(\Sigma\), the inverse density of lattice sites that are shared by the two grain orientations when rotated about a common lattice point. In two dimensions, CSL grain boundaries are necessarily commensurate and vice versa. For the triangular lattice, each common lattice point for a given pair of grain orientations has a repeat cell that is
an equilateral triangle with one edge defined by the surface vector with length \( D \). \( \Sigma \) is equal to the number of lattice points inside this cell and is given by the area of the cell divided by the area of one lattice triangle, \( \Sigma = D^2 \). While the CSL formulation gives a simple method for finding commensurate grain boundaries, it is misleading to suggest that the actual coincidence of sites plays a physical role. In fact, shifting the grains in the \( y \) direction so as to minimize the grain boundary energy (described in Sec. II) generally causes the atoms to no longer coincide. Without this shift, the two free surfaces will have atomic planes that meet at the same point. The elastic energy is lowered by staggering the dislocations.\(^{24}\) Wolf has pointed out that for three dimensions, the CSL formulation of grain boundaries involves a redundant number of parameters.\(^{19}\)

Bishop and Chalmers introduced the concept of “structural units”—polygonal structures of atoms along high angle grain boundaries.\(^{14}\) We find that the patterns of structural units are more relevant to the physical properties of the grain boundaries. When relaxed, the structural units of grain boundaries in two dimensions, triangular lattices are five atoms forming a regular pentagon or a pentagon that is slightly stretched. Each high symmetry grain boundary is comprised of a simple pattern of pentagonal structural units (Table I). When a high symmetry grain boundary is perturbed, an element of the pattern of structural units from the neighboring high symmetry grain boundary is introduced (Fig. 7). As the tilt angles move closer to the neighboring high symmetry grain boundary, the flaws become closer together until they outnumber the original structure. The roles of flaw and original structure are then reversed. In this manner, combinations of patterns of structural units can be used to build up any commensurate, high angle grain boundary.

We find the structural units are a useful way of conceptualizing the different grain boundary geometries. We have not

<table>
<thead>
<tr>
<th>( \theta )</th>
<th>Miller Indices</th>
<th>Repeat Distance (lattice constants)</th>
<th>( \Sigma )</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>49.10</td>
<td>(1,4)</td>
<td>2.64</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>43.89</td>
<td>(2,5)</td>
<td>3.61</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>53.41</td>
<td>(1,7)</td>
<td>4.35</td>
<td>19</td>
<td></td>
</tr>
<tr>
<td>40.89</td>
<td>(1,2)</td>
<td>4.58</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>38.94</td>
<td>(4,7)</td>
<td>5.56</td>
<td>31</td>
<td></td>
</tr>
<tr>
<td>51.78</td>
<td>(2,11)</td>
<td>7.0</td>
<td>49</td>
<td></td>
</tr>
</tbody>
</table>

FIG. 7. (Color online) Finding the Burger’s vector for flaws along vicinal grain boundaries. Grain boundaries with tilt angles in the vicinity of high symmetry grain boundaries are comprised of the pattern of structural units of the high symmetry grain boundary with an element from the neighboring high symmetry geometry. When defining the Burger’s vector of a flaw added to a high symmetry grain boundary we are no longer comparing the flaw to the perfect lattice but to the repeating pattern of structural units. Therefore we must cross the grain boundary at equivalent portions of the structural unit in our reference loop and in our loop around the flaw. The loop on the bottom is the reference loop around the region of the grain boundary without the added flaw. The loop on the top surrounds the added flaw, which we model as a partial grain boundary dislocation. The arrows indicate the segments needed to make the top loop match the bottom loop; for example, the arrow on the upper right indicates that the path around the defect is one atomic length shorter on that leg than the corresponding leg of the reference loop. The arrows sum to the Burger’s vector of the added grain boundary dislocation. This grain boundary has a tilt angle of 44.3° and the added partial dislocation has a Burger’s vector with a length of 0.495 lattice constants and a direction along the negative \( x \) axis.
found the coincidence site lattice model useful in our investigations. In systematically constructing grain boundaries with small repeat distances (Sec. III B), we have found the surface lattice vectors of the two sides of the grain boundary to be a useful description. We suggest that the structure of high angle grain boundaries can best be described in terms of a dislocation model. The “extra flaws” added to create vicinal grain boundaries can be described as partial dislocations. The added flaws can also move, screen, or cancel the flaws that make up the high symmetry boundary.

IV. GRAIN BOUNDARY ENERGY

We have measured the grain boundary energy and peak stress for all symmetric grain boundaries with repeat distances under 20 lattice constants and asymmetric grain boundaries with repeat distances under 30 lattice constants. In order to explore the regions close to high symmetry grain boundaries, we have added a few geometries with longer repeat distances, close to the high symmetry boundaries. The energy associated with the series of flaws that make up the grain boundary is defined in Eq. (2). As stated earlier, and found by several earlier studies,24 cusp shapes occur at high symmetry boundaries. For small angle grain boundaries, the grain boundary energy has the form

$$E_{GB} = \frac{\mu b}{4\pi(1-n)}|\theta| \ln \left( \frac{e\alpha}{2\pi|\theta|} \right) \approx |\theta| \ln |\theta|,$$

where $\mu$ is the shear modulus, $b$ is the Burger’s vector, $n$ is the Poisson ratio, and $\alpha$ is a factor that includes the core energy. One can now imagine the same scenario applied to high symmetry boundaries. We can take a high symmetry grain boundary and add or subtract flaws a distance $d$ apart as shown in Fig. 8. By the same reasoning as used for the low angle grain boundaries, the energy near the high symmetry grain boundary will have the form

$$E_{GB} = E_0 + \frac{\mu b}{4\pi(1-n)}|\theta - \theta_0| \ln \left( \frac{e\alpha'}{2\pi|\theta - \theta_0|} \right),$$

where $E_0$ is the energy of the high symmetry grain boundary which occurs at the angle $\theta_0$ and $\alpha'$ incorporates the core energy of the flaw within the pattern of flaws.

The grain boundary energies for all of the symmetric grain boundaries that we have measured are shown in Fig. 9. Note that cusps occur at the angles listed in Table I. We are able to fit the data for symmetric grain boundaries to a function of the form

$$E_{GB}(\theta) = a_0|\sin 3\theta| \ln \left( \frac{b_0}{\sin 3\theta} \right) + a_{30}|\sin 3(\theta - 30)\theta| + \sum_{j=0}^{n} \left( a_i(\cos 6\theta + a_i(\cos 6\theta - \cos 6\theta) + \sum_{j=0}^{m} c_j \cos(6j\theta) + d. $$

The first two terms fit the cusps at $0^\circ$ and $30^\circ$, where the cusps are symmetrical about their respective center points. The next set of terms in the sum fit the cusps at high symmetry tilt angles. The function $|\cos 6\theta - \cos 6\theta| \ln |\cos 6\theta - \cos 6\theta|$ was chosen as a fitting function because it asymptotically gives a $\theta \ln(1/\theta)$ shaped cusp in the near vicinity of $\theta_i$ and because it has the correct symmetry: even mirror symmetry at $0^\circ$ and $30^\circ$ and an overall period of $60^\circ$. We use one term that is antisymmetric about $\theta_i$ and one term that is symmetric about $\theta_i$ so that we can fit the shape on either side of the cusp independently. We do not expect the slope of the curve on either side of the cusp to be the same since the Burger’s vectors of the additional flaws for the geometries on either side of the high symmetry geometry may differ (Fig. 7).

The $\theta_i$ can be any angles that have the shortest repeat distances, such as those given in Table I. The curve in Fig. 9 is the result of fitting Eq. (7) to the data shown in the same
TABLE II. Coefficients for fitting grain boundary energy to Eq. (7).

<table>
<thead>
<tr>
<th>i</th>
<th>$a_i^{(a)}$</th>
<th>$a_i^{(d)}$</th>
<th>$b_i^{(a)}$</th>
<th>$b_i^{(d)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.76</td>
<td>$7.4 \times 10^{-4}$</td>
<td>0.196</td>
<td>13.6</td>
</tr>
<tr>
<td>2</td>
<td>0.68</td>
<td>$0.023$</td>
<td>0.292</td>
<td>$9.19 \times 10^{-7}$</td>
</tr>
<tr>
<td>3</td>
<td>0.22</td>
<td>$-8.9 \times 10^{-3}$</td>
<td>0.104</td>
<td>4.76</td>
</tr>
<tr>
<td>4</td>
<td>0.44</td>
<td>0.039</td>
<td>0.154</td>
<td>$3.93 \times 10^{-3}$</td>
</tr>
</tbody>
</table>

FIG. 10. (Color online) Grain boundary energies for asymmetric geometries. The surface and contour represents a smooth interpolation between the particular geometries that have been simulated, represented by the black dots in the $\theta_1, \theta_2$ plane. Perfect crystals have zero grain boundary energy, shown by the groove along the $\theta_1 = \theta_2$ line.

FIG. 11. (Color online) Asymmetric grain boundary energies in neighborhood of high symmetry geometry. The grain boundary geometries considered in the above plot each have a misorientation of $21.79^\circ$ and are in the vicinity of the high symmetry grain boundary with tilt angles $24.79^\circ$ and $3.00^\circ$. The lines are fit to $\theta \ln \theta$ for the points on either side of the cusp.

V. FRACTURE STRENGTH

A. Low angle symmetric grain boundaries

The Frank conditions state that the total Burger’s vector for the dislocations making up a low angle grain boundary is equal to the difference of the surface vectors that define the orientation of each grain. In order to guarantee that the grain boundary will have only one dislocation per repeat distance after it is relaxed, we must choose surface vectors that have a difference equal to a basis vector and have the same repeat distance. One pair of surface vectors is $(2n+1,-n)$ and $(2n+1,n-1)$. This gives a symmetric grain boundary with a single dislocation with tilt angles close to $0^\circ$, a Burger’s vector equal to $(0,1)$ and a repeat distance of $\sqrt{3n^2+3n+1}$. Our simulations show that such grain boundaries fail via intragranular fracture rather than intergranular fracture as shown in Fig. 12.

The apex of the triangle $\theta_1 = \theta_2 = 30^\circ$ in Fig. 6 is a perfect crystal oriented at $30^\circ$, which is the end point of a series of symmetric tilt boundaries. At exactly $\theta_1 = \theta_2 = 30^\circ$, there is an abrupt jump up in fracture strength since the perfect crystal has no “extra dislocations” at which to nucleate fracture. Our
simulations find a peak stress of 4.31 (Lennard-Jones units) for the perfect crystal.

The low angle grain boundaries near the 30° lattice orientation have surface vectors \((1, n)\) and \((n, 1)\), repeat distances \(\sqrt{1+n+n^2}\), and total Burger's vector \((-1,1)\), which splits into two flaws with Burger's vectors \((0,1)\) and \((-1,0)\) shown in Fig. 13. For wide enough simulations, these dislocations glide in opposite directions until they are restricted by the constrained zones on either side. For narrower simulations, the dislocations do not glide but form nucleation points for grain boundary fracture causing an abrupt jump down in the peak stress compared to the peak stress of the perfect crystal.

For narrow simulations of low angle grain boundaries in this region, we find that the peak stress has a parabolic dependence on angle for low angle grain boundaries that are constrained by width to fracture in qualitatively similar ways. Fracture nucleates exactly at the dislocation and the first nonvanishing term in the stress at this point due to neighboring dislocations goes as \(1/d^2\), where \(d\) is the distance between dislocations.

\[
\sigma_{\text{vol}}(x,y) = \frac{\mu b}{2\pi(1-\nu)} \frac{x(x^2-y^2)}{(x^2+y^2)^2},
\]

where the \(x\) direction is the direction of the Burger's vector. Since the Volterra solution is odd, the \(n=1\) term of the stress at each dislocation vanishes as we sum over the neighboring dislocations on either side. The first nonvanishing term in Eq. (7) is the \(n=2\) term which has three contributions. (i) The \(n>1\) terms are the multipole expansions of the stress field as well as nonlinear terms. (ii) The nonlinear term in strain field has the form \(du/dx \neq du/dx\), giving a power law of \(1/r^2\). (iii) Geometrical restrictions cause some grain boundaries to have flaws unequally spaced in the \(y\) direction (though for the results given in Fig. 14 we have only explored geometries with equally spaced flaws). The grain boundary geometries used in Fig. 14 do have flaws that are not aligned perfectly in the \(x\) direction. In each of these cases, shifting the dislocation constitutes adding a dislocation dipole (adding one positive and one negative, canceling a dislocation and adding a new one), and therefore is another contribution to the \(1/r^2\) term. The external stress needed to produce a stress equal to \(\sigma_c\) at each flaw is then

\[
\sigma_{\text{peak}} = \sigma_c - \frac{a_2}{d^2} = \sigma_c - A(\theta - \theta_0)^2,
\]

where \(a_2\) combines the three contributions described above. This parabolic dependence on \(\theta - 30°\) is depicted on the left-hand side of Fig. 15.
at the grain boundary vary from those of the perfect crystal. The fracture nucleates along the boundary in the region where the stress due to the added flaw is positive. Because fracture does not nucleate at the center of the added flaw, the Volterra solution as summed over the neighboring, added flaws does not cancel at the nucleation site. This leads to a linear law for fracture strength as a function of tilt angle, for grain boundaries near high symmetry geometries.

Consider the grain boundaries with tilt angles ranging from 49.39° to 53.41°, which are close to the high symmetry grain boundary at 49.11°. The additional flaws that characterize these grain boundaries have Burger’s vectors equal to \((-\sqrt{3}\sin\theta, -\sin\theta)\) and \((-\sqrt{3}\sin\theta, \sin\theta)\), where \(\theta\) is the symmetric tilt angle of the lattice. The norms are \(2|\sin\theta|\) and the angles are 210° and 150°. For these geometries, the \(xx\) component of the stress field [due to two dislocations a distance \(D = b/2(\theta_0 - \theta)\)] along the \(y\) axis is

\[
\sigma_{xx}(y) = \frac{\mu[2\sqrt{3}y(\theta_0 - \theta) - 3\sin(\theta_0 - \theta)]\sin(\theta_0 - \theta)}{2\pi(1 - \nu)[y(\theta_0 - \theta) - \sqrt{3}\sin(\theta_0 - \theta)]},
\]

where \(\theta_0\) is the tilt angle of the high symmetry grain boundary. We need to look at simulations in the fixed displacement (narrow width) regime in order to observe where fracture nucleates. We find that for geometries with this pattern of flaws, fracture nucleates at the same distance above the added flaw. The external stress needed to nucleate fracture at a distance \(y\) from the added flaw, along the grain boundary is then approximately

\[
\sigma_{xx}(y) = \frac{\mu(3-2\sqrt{3}y)(\theta_0 - \theta)}{2\pi y(1 - \nu)(y - \sqrt{3})} + O(\theta_0 - \theta)^3,
\]

for high angle grain boundaries, the added flaw is no longer the sole nucleation site for fracture and fracture does not necessarily nucleate in the core of the added flaw. The added dislocation creates a stress field given roughly by the Volterra solution [Eq. (10)] with a positive stress on one side, negative stress on the other, and a singularity at the center shown in Fig. 16. The stress field differs slightly from the Volterra solution because the elastic constants of the material...
Stress (Lennar along the diagonal. Not included these data points which would otherwise show a ridge grain boundaries. In order to use a smaller vertical scale, we have cone. The peak stress of perfect crystals is much higher than that of the interpolation, each high stress peak is rendered as a finite-width sent a smooth interpolation between particular geometries. Due to repeat distance of 9.53 lattice constants. Figure 19 shows the grain boundary with tilt angles 24.79° and 3.00° which has a repeat stress of this grain boundary and grain boundaries in its neighborhood along a line of constant misorientation.

VI. CONCLUSION

We have systematically explored the space of 2D grain boundaries, the patterns of structural units that make up short repeat distance grain boundaries, and the flaws in the patterns of structural units that make up vicinal grain boundaries. We have shown that the patterns of structural units are key to understanding the singularities in energy and peak stress at special grain boundaries by drawing an analogy to perfect crystals. We have used this insight to find a functional form for the energies of 2D symmetric grain boundaries and

\[ \sigma_{\text{peak}} = \sigma_c - \frac{\mu(3 - 2\sqrt{3})}{2\pi\nu(1 - \nu)(\nu - \sqrt{3})} \theta_1 - \theta \] (14)

which explains the linear dependence on angle shown in Figs. 16 and 17.

The results for the peak stress of all grain boundaries with repeat lengths less than 30 lattice constants are shown in Fig. 18. We also note spikes in peak stress for grain boundaries with particularly low repeat distances. Consider again the grain boundary with tilt angles 24.79° and 3.00° which has a repeat distance of 9.53 lattice constants. Figure 19 shows the peak stress of this grain boundary and grain boundaries in its neighborhood along a line of constant misorientation.

FIG. 18. (Color online) Grain boundary peak stresses for asymmetric geometries. As with Fig. 10, the surface and contours represent a smooth interpolation between particular geometries. Due to the interpolation, each high stress peak is rendered as a finite-width cone. The peak stress of perfect crystals is much higher than that of grain boundaries. In order to use a smaller vertical scale, we have not included these data points which would otherwise show a ridge along the diagonal.

FIG. 19. (Color online) Asymmetric grain boundary peak stresses in neighborhood of high symmetry geometry. The grain boundary geometries plotted above are the same as those in Fig. 11, along a line of constant misorientation in the vicinity of a high symmetry geometry. The lines show a possible linear fit to the points on either side of the low repeat distance grain boundary, to understand the tilt angle dependence of peak stress near special grain boundaries.

In principle, with enough computer time it would be possible to conduct a similar study for the 5D space of idealized 3D grain boundaries. We suspect that the perfect crystal analogy could also explain the cusps in energy and discontinuous spikes in peak stress for special 3D grain boundaries. The geometry dependence of energy and peak stress surrounding these special grain boundaries will be more complicated since the grain boundaries can have a pure tilt, pure twist, or a mixed type of geometry. In real polycrystals, the fracture strength and energy are further complicated by impurities at the interface, emission of dislocations during fracture, and more complex geometries such as triple junctions of grains, which are often the nucleation site for fracture. Systematic studies of such systems are infeasible and better suited by on the fly simulations of local regions of interest.

ACKNOWLEDGMENTS

This work was supported by NSF Grants No. ITR/ASP ACI0085969 and No. DMR-0218475. We also wish to thank Nicholas Bailey, Drew Dolgert, Gerd Heber, Anthony Ingraffea, Surachute Limkumnerd, Chris Myers, and Paul Wawrzynek.
simulation of the same geometry, but with periodic boundary conditions in the y direction. We see that if we use rolled boundary conditions, we need a simulation that is 48 times longer in the y direction in order to converge to within 2% of the results found by a simulation with periodic boundary conditions. Hence for computational efficiency, we choose to use periodic boundary conditions in the y direction.

23 The Frenkel-Kontorova model (Ref. 31) describes a one-dimensional chain of atoms, connected by springs, subject to an external, sinusoidal potential. If the relaxed length of the spring and the width of the potential wells are commensurate a minimum energy state exists. For the incommensurate case, there will be a pinned phase with many local minima and an unpinned phase, depending on the depth of the wells in relation to the stiffness of the springs.


25 Since this Burger’s vector may be a sum of lattice vectors from either side of the grain boundary, the total may not be a full translation vector of the triangular lattice.

26 We use “energy” for “energy per unit length” throughout.


30 For simulations of a few Å wide, the crack opens up along the grain boundary with a slow unzipping mechanism, starting at particular flaws. For wider simulations, the crack snaps open, soon after a peak stress is reached. The slow unzipping of the narrow simulations is characteristic of fixed displacement boundary conditions—since the displacements of a region close to the interface is fixed, the opening of the crack is controlled, allowing us to see the details of how the crack opens. The snapping open, seen with a wider simulation, is characteristic of fixed force boundary conditions. In the wider simulations, the material on either side of the grain boundary acts as a spring, which effectively softens the boundary conditions, approximating fixed-force boundaries for long lengths. When the constrained region is far away, the interface snaps open once it reaches the maximum stress that it can sustain.