**MultiShifter: software to generate structural models of extended two-dimensional defects in 3D and 2D crystals**

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Extended defects in crystals, such as dislocations, stacking faults and grain boundaries, play a crucial role in determining a wide variety of materials properties. Extended defects can also lead to novel electronic properties in two-dimensional materials, as demonstrated by recent discoveries of emergent electronic phenomena in twisted graphene bilayers. This paper describes several approaches to construct crystallographic models of two-dimensional extended defects in crystals for first-principles electronic structure calculations, including (i) crystallographic models to parameterize generalized cohesive zone models for fracture studies and meso-scale models of dislocations and (ii) crystallographic models of twisted bilayers. The approaches are implemented in an open source software package called MultiShifter.

I. INTRODUCTION

Extended defects play an important role in many materials applications, affecting electronic, functional and mechanical properties. The simplest two-dimensional extended defect is a surface created upon cleaving two halves of a crystal²–⁴. The two halves of a crystal can also be rigidly sheared relative to each other as occurs in many layered battery materials that undergo stacking sequence phase transformations upon intercalation⁵–⁷. Plastic deformation of crystals is mediated by the passage of dislocations that glide along a slip plane. While a dislocation is an extended one-dimensional defect, it often dissociates into a pair of partial dislocations that bound an extended two-dimensional stacking fault⁸.

Two-dimensional extended defects are also present in heterostructures of 2D materials⁹. 2D materials exhibit unique electronic properties that are absent in their three-dimensional counterparts. Furthermore, new electronic behavior can emerge when a pair of two-dimensional building blocks are twisted relative to each other¹⁰–¹², as was recently demonstrated for graphene¹³. The twisting of a pair of stacked two-dimensional materials by an angle $\theta$ around an axis that is perpendicular to the sheets breaks the underlying translational symmetry of the 2D building blocks. A new field of "twistronics" has emerged that seeks to understand and exploit the electronic properties that arise upon twisting a pair of two-dimensional materials¹⁴.

Here we describe a generalized framework and accompanying software package, called MultiShifter, to facilitate the study of the thermodynamic and electronic properties of periodic two-dimensional defects in crystalline solids from first principles. We introduce the concept of a generalized cohesive zone model that simultaneously encapsulates the energy of decohesion and rigid glide of two halves of a crystal. We then detail how super cells and crystallographic models can be constructed to accommodate a pair of two-dimensional slabs that have been twisted relative to each other by an angle $\theta$. MultiShifter automates the construction of crystallographic models of extended two-dimensional defects and of two-dimensional layered materials to enable (i) the study of surfaces, (ii) the calculation of cohesive zone models for fracture studies, and (iii) the construction of models of twist grain boundaries and twisted 2D materials. The crystallographic models generated by MultiShifter can then be fed into first-principles electronic structure codes to calculate a range of thermodynamic and electronic properties. MultiShifter consists of C++ and Python routines and draws on crystallographic libraries of the CASM software package."
to the minimum of $u_{cz}(\delta, \tau)$ as a function of $d$ at fixed $\tau$. The dependence of $u_{cz}(\delta, \tau)$ on $\tau$ at fixed $\delta$ or $d$ is periodic and has the same 2-dimensional periodicity as that of the crystallographic plane $P$. This means that the parameters that appear in Equation (1) are also periodic functions of $\tau$.

When all the $\alpha_n$ are set to zero, we recover the universal binding energy relation (UBER) that is able to describe the cohesive properties of metals with remarkable accuracy.\cite{UBER1, UBER2, UBER3, UBER4}

The additional parameters $\alpha_n$ are necessary to capture deviations from the UBER form due to deviations from purely metallic bonding.\cite{UBER5}

The coefficient $2\gamma$ is related to the energy of cleaving a crystal into two bicrystals

$$2\gamma(\tau) = \frac{E_{\text{cleaved}} - E_{\text{bulk}}(\tau)}{A_{\text{surface}}}$$

where $E_{\text{bulk}}(\tau)$ is the energy of the bicrystal at separation $d = d_0(\tau)$ and translation $\tau$, $E_{\text{cleaved}}$ is the energy of the bicrystals at infinite separation and $A_{\text{surface}}$ is the area of the exposed surfaces. $2\gamma(\tau)$ corresponds to the minimum of $u_{cz}(\delta, \tau)$ at each $\tau$ with respect to interslab separation $\delta$ and is referred to as the generalized stacking fault energy (GSFE), also known as the $\gamma$ surface. The GSFE is an essential ingredient of mesoscale simulation techniques such as phase field models\cite{phase_field1, phase_field2, phase_field3} and Peierls-Nabarro models of dislocations.

When cleaving a single crystal across a plane $P$ (as opposed to a bicrystal consisting of two different materials), it is common to set the origin of translations $\tau$ at the shift coinciding with a perfect crystal (i.e. no stacking fault). In that case $\gamma_0 = \gamma(\tau = 0)$ becomes equal to the surface energy for the crystallographic plane $P$ in the absence of any surface reconstruction.\cite{surface_energy1, surface_energy2, surface_energy3} The generalized cohesive zone model can serve as a constitutive law to describe the response of a solid ahead of a transgranular crack tip\cite{crack_tip1, crack_tip2}. The elastic constant, $C$, along the direction of separation is a function of the parameters of the xUBER relation Eq. 1 according to\cite{elastic_constant1}

$$C = 2d_0 \frac{2\gamma}{\lambda^2} \left( \frac{1}{2} - \alpha_2 \right)$$

Since the parameters, $d_0$, $\gamma$, $\lambda$ and $\alpha_n$ of Equation (1) are periodic functions of $\tau$, they can each be expressed as a Fourier series. For example, the $\tau$ dependence of $\gamma$ can be expressed as

$$\gamma(\tau) = \sum_K \tilde{\gamma}_K e^{-iK\tau}$$

where the sum extends over $K$ vectors of the two-dimensional reciprocal lattice of the two-dimensional unit cell of the crystallographic plane $P$. The expansion coefficients, $\tilde{\gamma}_K$, are the Fourier transform of $\gamma$.

C. Extensions to account for twist

The twisting of two halves of a crystal or of a pair of two-dimensional materials will generally break any translational
symmetry that may have existed before. It is only for a subset of special twist angles \( \theta \) that a supercell translational symmetry is preserved\cite{20,22}. The energy of the bicrystal will not only depend on the relative separation \( d \) and translation \( \tau \), but also on the choice of rotation axis \( r \) and twist angle \( \theta \). This dependence can be formulated generally as

\[
u(d, \tau, r, \theta) = u_{cz}(d, \tau) + u_t(d, \tau, r, \theta) \tag{5}\]

where \( u_{cz} \) is the reference energy in the absence of a twist, and can be expressed using a form such as Eq \( 4 \). The function \( u_t \) accounts for the twist energy and is equal to zero when \( \theta = 0 \). While the \( \theta \) dependence of \( u_t \) could be represented as a Fourier series, as it is periodic in \( \theta \), it will generally exhibit cusps and will usually not be a smooth function of \( \theta \). The expansion coefficients of such a Fourier series would be a function of \( d, \tau \) and \( r \).

III. CREATING SLAB MODELS

Most electronic structure methods impose periodic boundary conditions on the crystallographic model. In this section, we describe how crystallographic models can be constructed to realize crystal separation, glide and twist within super cells that have periodic boundary conditions.

A. Constructing slab geometries to parameterize cohesive zone models

A cohesive zone model such as Equation \( 1 \) can be parameterized by fitting to training data as calculated with a first-principles electronic structure method. It is possible to accommodate the periodic boundary conditions that these methods impose using a slab geometry as illustrated in Figure 2a. The unit cell then consists of a slab of crystal with its periodic images separated by layers of vacuum parallel to the plane \( P \). For bulk crystals, the crystal slab must be sufficiently thick to avoid interactions between periodic images of the surfaces adjacent to the vacuum layer.

To construct a crystallographic model consisting of slabs parallel to a plane \( P \), it is necessary to first identify a supercell of the primitive cell vectors, \( a, b \) and \( c \), such that two supercell vectors, \( A \) and \( B \) span the plane \( P \). The vectors \( A \) and \( B \) can be determined by connecting the origin to the closest non-collinear lattice points that lie on a plane parallel to the plane \( P \) that also passes through the origin. A third vector, \( s \), can then be chosen as the shortest translation vector of the parent crystal structure that is not in the plane \( P \). The vector \( s \) determines the smallest possible thickness of the slab. The thickness of the slab can be adjusted by multiplying \( s \) by an integer, \( l \). It is usually desirable to translate the resulting vector \( l s \) parallel to \( P \) (by an integer linear combination of \( A \) and \( B \)) until its projection onto the plane \( P \) falls within the unit cell spanned by \( A \) and \( B \). This new vector, \( C \), is then the third super cell vector of the slab model.

The next task is to sample different values of slab separations \( d \) and relative translations \( \tau \). Due to translational periodicity of the crystal, only translation vectors \( \tau \) within a two-dimensional unit cell spanned by the \( A \) and \( B \) vectors of the super cell need to be considered. One approach is to generate a uniform grid within the unit cell of possible translation vectors \( \tau \), as is illustrated in Figure 2b, for an fcc crystal in which one half is sheared relative to another half along a (111) plane. The two half crystals often have additional symmetries that make a subset of the translations \( \tau \) equivalent to each other. Symmetric equivalence between two different translation vectors \( \tau_1 \) and \( \tau_2 \) can be ascertained by mapping the resultant crystals onto each other with a robust crystal mapping algorithm\cite{26}. Figure 2b shows the orbits of equivalent translation vectors \( \tau \) for fcc by assigning the same color to all translation vectors that are equivalent by symmetry. The choice of supercell may break some symmetries of the

\[\text{FIG. 2.} \ \text{(a) Slab layer of fcc Al, where the exposed layer is the (1,1,1) plane. The unit cell of the slab has had 3Å of vacuum inserted above the exposed plane. The conventional unit cell of fcc is also shown for reference. (b) Schematic representation of the symmetry for different in plane shifts in Al. Spots with the same color are symmetrically equivalent structures, and will have the same DFT energy for equal cleavage values.}\]
underlying crystal, making symmetric equivalence of translation vectors $\tau$ dependent on the particular supercell. For each symmetrically distinct translation vector $\tau$ it is possible to generate a grid of separations $d$ over increments of $\Delta d$. This can be realized by adding $\Delta d\hat{n}$ to $C$ while keeping the Cartesian coordinates of the atoms within the unit cell unchanged. The vector $\hat{n}$ is a unit vector normal to the $A\cdot B$ plane.

The parameterization of a generalized cohesive zone model can occur in two steps. The first step is to calculate the energy of separation over a discrete set of separations $d$ for each symmetrically distinct translation vector $\tau$. The resultant energy versus separation $d$ relation can then be fit to an xUBER relation, Eq 1. The parameterization of an xUBER relation over all symmetrically distinct translation vectors $\tau$ will generate numerical values for the adjustable parameters, $d_0$, $\gamma$, $\lambda$ and $\alpha_n$, of the xUBER, Equation (1), over a uniform grid of translation vectors $\tau$. The dependence of the adjustable parameters on $\tau$ can then be represented with a Fourier series such as Eq. 4, thereby allowing for the accurate interpolation at any translation vector $\tau$.

B. Crystallographic models for twisted bicrystals

In addition to separation and glide, it is also possible to twist the two halves of a bicrystal by rotating one half relative to the other around a rotation axis, $r$, that is perpendicular to $P$. It is increasingly recognized that interesting electronic properties can emerge when a pair of two-dimensional materials are rotated relative to each other in this manner. In bulk materials, special grain boundaries, referred to as twist boundaries, can be generated by twisting the top half of a crystal around an axis that is perpendicular to the grain boundary. A challenge for electronic structure calculations is to identify a supercell that is able to accommodate the twisted half crystals.

A Moiré pattern emerges when a periodic two-dimensional lattice is rotated relative to a second periodic lattice. Figure 3a for example, shows the emergence of a Moiré pattern after a pair of two-dimensional triangular lattices (brown and green) have been rotated relative to each other by $5^\circ$. The Moiré pattern is itself periodic, but has lattice vectors that are usually much larger than those of the two-dimensional lattices that have been rotated. Furthermore, the lattice of the Moiré pattern is rarely commensurate with the lattices of the twisted half crystals. This is evident in Figure 3a where the lattice points of the Moiré pattern, shown as the intersections of the grey lines, do not exactly overlap with sites of the twisted triangular lattices shown in brown and green. Only a subset of particular twist angles $\theta$ produce Moiré patterns that are commensurate with the twisted two-dimensional lattice. The Moiré lattice can, nevertheless, serve as a guide to identify a supercell that can accommodate the twisted half crystals for first-principles electronic structure calculations. But since the Moiré lattices for most twist angles $\theta$ only approximately coincide with sites of the twisted lattices, both half crystals must usually

FIG. 3. (a) Superposition of a triangular lattice with itself after a 5 degree rotation. The resulting interference pattern is semi-periodic, forming the Moiré lattice. (b) Rotated triangular lattices in real (fig. 3b) and reciprocal (fig. 3c) space. (c) The reciprocal Moiré lattice is constructed by taking the difference of the lattice vectors in reciprocal space.
be deformed slightly such that they can be accommodated within a common supercell.

The lattice of the Moiré pattern can be determined by working in reciprocal space. Consider a three dimensional unit cell with lattice vectors \( \mathbf{A}, \mathbf{B}, \mathbf{C} \). Assume that the vectors \( \mathbf{A} \) and \( \mathbf{B} \) form the two-dimensional lattice parallel to the plane \( \mathbf{P} \) (e.g. a two-dimensional triangular lattice) and that the third vector \( \mathbf{C} \) is perpendicular to \( \mathbf{P} \). The rotation axis \( r \) is therefore parallel to \( \mathbf{C} \). It is convenient to work with a \( 3 \times 3 \) matrix \( \mathbf{L} = [\mathbf{A}, \mathbf{B}, \mathbf{C}] \) where each lattice vector appears as a column of \( \mathbf{L} \). The reciprocal lattice vectors of the lattice \( \mathbf{L} \) are then the column vectors of the matrix \( \mathbf{K} \) defined by

\[
\mathbf{K} = 2\pi (\mathbf{L}^{-1})^T \tag{6}
\]

The application of a rotation \( \theta \) to a lattice \( \mathbf{L} \) around a rotation axis that is parallel to \( \mathbf{C} \) produces a lattice \( \mathbf{L}_\theta \). The lattice vectors of the two-dimensional Moiré pattern, \( \mathbf{M} \), will have a reciprocal lattice represented by the matrix \( \mathbf{K}_M \) in which the upper left \( 2 \times 2 \) block is equal to the corresponding difference between the reciprocal lattices of \( \mathbf{L} \) and \( \mathbf{L}_\theta \) (i.e. \( [\mathbf{K}_M]_{i,j} = [\mathbf{K}]_{i,j} - [\mathbf{K}]_{i,j} \) for \( i,j = 1,2 \)) and the third axis, which is unaffected by the rotation is the same as that of \( \mathbf{K} \) and \( \mathbf{K}_\theta \) (i.e. \( [\mathbf{K}_M]_{3,3} = [\mathbf{K}]_{3,3} = [\mathbf{K}_\theta]_{3,3} \)). The Moiré lattice is then

\[
\mathbf{M} = 2\pi (\mathbf{K}_M^{-1})^T \tag{7}
\]

This is illustrated in Figure 3b and Figure 3c for a pair of triangular lattices. The green triangular lattice of Figure 3b has been rotated by an angle \( \theta \) relative to the brown triangular lattice. The reciprocal lattice vectors of the green and brown triangular lattices are shown in Figure 3c. The two grey vectors in Figure 3c are the differences of the reciprocal lattice vectors of the rotated green lattice and those of the fixed brown lattice. The grey vectors, when translated to the origin of reciprocal space, span the unit cell of the reciprocal lattice of the Moiré pattern. This is illustrated by the thick black arrows in Figure 3c with the grid representing the reciprocal lattice points of the Moiré lattice. (For large rotation angles \( \theta \), it is possible that one of the reciprocal lattice vectors of \( \mathbf{K}_M \) falls outside of the Wigner-Seitz cell of either \( \mathbf{K} \) or \( \mathbf{K}_\theta \). In these situations, the offending reciprocal lattice vector of \( \mathbf{K}_M \) must be translated back into the Wigner-Seitz cell of either \( \mathbf{K} \) or \( \mathbf{K}_\theta \).)

The example illustrated by Figure 3b and Figure 3c is for a special angle \( \theta \) for which the Moiré lattice is commensurate with the two rotated lattices. For these special angles, the reciprocal lattice vectors \( \mathbf{K} \) and \( \mathbf{K}_\theta \) (brown and green vectors in Figure 3c) coincide with sites of the reciprocal lattice of the Moiré lattice \( \mathbf{K}_M \). Only a subset of twist angles produce Moiré lattices that are commensurate with the twisted lattices.

In general, the Moiré lattice points do not exactly coincide with sites of either the \( \mathbf{L} \) or \( \mathbf{L}_\theta \) lattices, as illustrated by the example of Figure 3a for a pair of triangular lattices that have been rotated by \( 5^\circ \). Nevertheless, the Moiré lattice can guide the search for a supercell that can simultaneously accommodate the twisted pair of two-dimensional lattices. The first task is to identify the super cells of \( \mathbf{L} \) and \( \mathbf{L}_\theta \) that are close to that of the Moiré lattice. A super cell of a lattice can be generated as an integer linear combination of the lattice vectors \( \mathbf{A}, \mathbf{B}, \mathbf{C} \) according to

\[
\mathbf{S} = \mathbf{LT} \tag{8}
\]

where \( \mathbf{T} \) is a \( 3 \times 3 \) integer matrix and where the columns of \( \mathbf{S} \) contain the super cell lattice vectors. The sought after integer matrix \( \mathbf{T} \) is one that generates a super cell \( \mathbf{S} \) that is closest to that of the Moiré pattern \( \mathbf{M}_\theta \). This can be obtained by rounding each element of the matrix \( \mathbf{L}^{-1}\mathbf{M}_\theta \) to the nearest integer. A similar matrix \( \mathbf{T}_\theta \) must be determined by rounding the elements of \( \mathbf{L}_\theta^{-1}\mathbf{M}_\theta \) to the nearest integer. The resulting super cells, \( \mathbf{S} = \mathbf{LT} \) and \( \mathbf{S}_\theta = \mathbf{L}_\theta\mathbf{T}_\theta \), will usually not coincide exactly. However, they can both be strained and twisted slightly to a common supercell \( \mathbf{S} \) defined as the average of \( \mathbf{S} \) and \( \mathbf{S}_\theta \) according to

\[
\bar{\mathbf{S}} = \frac{1}{2} (\mathbf{S} + \mathbf{S}_\theta) \tag{9}
\]

This supercell can be used to accommodate the twisted bicrystal.

The amount of strain and twist needed to fit both bicrystals in \( \mathbf{S} \) can be calculated as follows. The dimensions of the bottom half of the bicrystal with super lattice \( \mathbf{S} \) will need to be deformed according to

\[
\mathbf{FS} = \bar{\mathbf{S}} \tag{10}
\]

where \( \mathbf{F} \), the deformation gradient, is a \( 3 \times 3 \) matrix. The deformation gradient \( \mathbf{F} \) can be factored into a product of a symmetric stretch matrix \( \mathbf{U} \) and a rotation matrix \( \mathbf{R} \) according to \( \mathbf{F} = \mathbf{RU} \). The stretch matrix \( \mathbf{U} \) describes the deformation of the crystal, while the rotation matrix \( \mathbf{R} \) in this situation corresponds to a rotation around \( \mathbf{C} \). A similar deformation gradient exists for the top half of the bicrystal with \( \mathbf{F}_\theta \mathbf{S}_\theta = \bar{\mathbf{S}} \) and \( \mathbf{F}_\theta = \mathbf{R}_\theta \mathbf{U}_\theta \). If the rotation angles of \( \mathbf{R} \) and \( \mathbf{R}_\theta \) are \( \phi_1 \) and \( \phi_2 \), respectively, then the two bicrystals need to undergo an additional relative twist of \( \Delta \theta = \phi_2 - \phi_1 \) to fit into \( \bar{\mathbf{S}} \). The actual rotation angle when describing the twisted bicrystal with a periodic supercell \( \bar{\mathbf{S}} \) is then \( \theta + \Delta \theta \). Hence, it is generally not possible to realize the target twist angle of \( \theta \) when using a periodic supercell to accommodate the twisted halves.

Information about the strains is embedded in the stretch matrices \( \mathbf{U} \) for the bottom half and \( \mathbf{U}_\theta \) for the top half. We use the Biot strain defined as \( \mathbf{E} = \mathbf{U} - \mathbf{I} \) where \( \mathbf{I} \) is the identity matrix. The strain is restricted to the two-dimensional space parallel to the twist plane. We assume that this plane is parallel to the \( x-y \) plane of the Cartesian coordinate system used to represent the lattice vectors. Conveni ent metrics of two-dimensional strain\footnote{\ref{footnote}} are

\[
\eta_1 = \frac{1}{\sqrt{2}} (E_{11} + E_{22}) \tag{11}
\]

\[
\eta_2 = \frac{1}{\sqrt{2}} (E_{11} - E_{22}) \tag{12}
\]
The first strain order parameter $\eta_1$ is a measure of symmetry preserving in-plane dilation. The two remaining strain order parameters, $\eta_2$ and $\eta_3$, measure deviatoric deformations within the plane. The strain energy within linear elasticity for a simple triangular or a simple square lattice takes the form $A\eta_1^2 + B(\eta_2^2 + \eta_3^2)$ where $A$ and $B$ are constants related to elastic moduli of the two-dimensional lattices. Two useful measures of the strain cost when using a particular supercell $S$ to achieve a target twist angle of $\theta$ are then $\eta_1$ for the dilation cost and $\sqrt{\eta_2^2 + \eta_3^2}$ for the deviatoric cost.

A refinement of the above approach can be used to lower the error in the target angle $\Delta\theta$ and the incurred strains. However, the improvement comes at the cost of requiring larger super cells. Instead of identifying super cells $S$ and $S_\theta$ of the lattices $L$ and $L_\theta$ that are as close as possible to the Moiré lattice $M$, the super cells $S$ and $S_\theta$ can be matched to super cells of $M$. This is illustrated in Figure 4 for a pair of triangular lattices that are being rotated by a target angle of $\theta = 15^\circ$. When matching super cells $S$ and $S_\theta$ to the Moiré lattice $M$ itself, as described above, the actual twist angle is $\theta = 12.117^\circ$. In contrast, when matching the super cells $S$ and $S_\theta$ to a $\sqrt{3}a \times \sqrt{3}a$ supercell of the Moiré lattice $M$ the actual rotation angle becomes $14.911^\circ$, which is much closer to the target angle of $15^\circ$. However, the common super cell that accommodates the twisted bicrystal is three time larger.

There are special twist angles for which super cells of their Moiré lattice can be found that are commensurate with the under lying lattices of the bicrystals. For these special twist angles $\Delta\theta$ and the strain order parameters will be zero. Figure 5 plots the number of triangular lattice unit cells that are needed in the super cells of a subset of commensurate twist angles. It is clear that very large super cells are necessary for most commensurate twist angles.

IV. MULTISHIFTER

The MultiShifter software package constructs crystallographic models of extended two-dimensional defects that separate a pair of bicrystals as described in the previous section. This includes crystallographic models to parameterize generalized cohesive zone models such as Eq. 1 and crystallographic models of twisted bicrystals. Figure 6 illustrates a schematic flowchart of MultiShifter.

The first step of the MultiShifter workflow is to construct the slab building blocks. The user provides the primitive cell of a crystal (prim in Figure 6) and the Miller indices of the crystallographic plane $P$. In the slice step, MultiShifter constructs the thinnest supercell of prim with lattice vectors $A$ and $B$ parallel to the plane $P$ and the shortest vector $C$ that is not in the plane $P$. The resulting super cell contains the slab unit (Figure 5) that constitutes the fundamental building block of subsequent crystallographic models. The slab unit must next be made thicker when modeling three dimensional materials or stacked when modeling the twist of two-dimensional materials. The slab thickness is increased based on user input. This occurs with the stack step. Depending on the basis of the crystal, an additional translate may be required, where the basis atoms in the slab unit are rigidly shifted, changing the position through which the plane $P$ penetrates the crystal. For example, when exploring the glide of CoO$_2$ sheets in a layered battery material such as LiCoO$_2$, $P$ should extend between the oxide layers, not through them.

At this point, the MultiShifter workflow diverges into two tracks. The first (right arrows), generates crystallographic models of crystal cleavage and glide with respect to
a plane $P$. A list of separations $d$ is specified (cleave) and for each of these separations a second grid of glide vectors $\tau$ may be enumerated (shift). Symmetrically equivalent translation grid points are tagged as such. Directories with input files for a first-principles electronic structure software package are then generated. Upon completion of static electronic structure calculations, a list of first-principles energies are available to parameterize a generalized cohesive zone model.

An alternative to the imposed regular grid of translation vectors is to use the mutate step. With this approach, a single custom structure that has been shifted by an arbitrary vector $\tau$ and separated by a custom value $d$ from its periodic image is created.

A second track in the Multishifter workflow (left arrows) generates crystallographic models of twisted bicrystals. Here the user provides a target twist angle $\theta$ and a maximum number of two-dimensional unit cells for the super cell that will accommodate the twisted bicrystals. Multishifter next determines the Moiré lattice and then identifies the super cells of two bicrystals that best match a super cell of the Moiré lattice. A crystallographic model is output along with the actual twist angle $\theta + \Delta \theta$ and values of the strain order parameters $\eta_1$, $\eta_2$ and $\eta_3$. When twisting a pair of bicrystals within a unit cell constrained by periodic boundary conditions (including the $C$ axis) two interfaces are necessarily introduced. One is the twist interface of interest, while the other is usually separated by a large slab of vacuum. For two-dimensional materials, the vacuum is not necessarily a drawback. For twist grain boundaries, however, the thicknesses of the twisted slabs should be sufficiently large such that the free surfaces in contact with vacuum do not affect the energy and electronic structure of the twist boundary.

V. EXAMPLES

A. Cohesive zone models of simple metals

As an illustration of a generalized cohesive zone model, we consider cleavage and glide with respect to a (111) plane of fcc Al. Figure 7a shows the energy of an Al crystal as it is cleaved along a pair of neighboring (111) planes for three different translational shifts $\tau$. The points were calculated with a plane-wave energy cutoff of 520 eV. K-points were generated using a fully automatic mesh with a length parameter $R_k = 50 \, \text{Å}^{-1}$. The UBER form of Eq. 1 was fit through the DFT and is shown as dashed lines in Figure 7a. As is clear from Figure 7a, the UBER curve is able to fit the DFT data very well. The blue points in Figure 7a reside on the energy versus separation curve for perfect fcc corresponding to $\tau = 0$. The difference in energy at $d_0$, where the curve has a minimum, and the energy for large separations corresponds to the surface energy $2\gamma$. The energy versus separation curve when separating pairs of (111) planes that form a stacking fault corresponding to $\tau = 1/3A + 1/3B$ is very similar, as is evident by the orange points in Figure 7a, although the minimum is not as deep and the equilibrium spacing $d_0$ is slightly shifted to a larger distance. The energies of separation for a translation that puts atoms of adjacent (111) planes directly on top of each other (green points in Figure 7a) is also very well described with an UBER curve. The minimum is at a much higher energy compared to other translations.

By performing similar DFT calculations over a uniform grid of symmetrically distinct translation vectors, it becomes possible to express the $\tau$ dependence of the adjustable parameters of the UBER curve as a Fourier series. Figure 7b for example shows the dependence of $2\gamma$ on the translation $\tau$. It exhibits the periodicity of the (111) plane of fcc Al, with the global minima corresponding to perfect fcc and the other minima corresponding to stacking faults. Figure 7c shows the dependence of the equilibrium separation, $d_0$, between two half crystals of fcc Al as a function of $\tau$. This plot looks very similar to the $2\gamma$ surface of Figure 7b with the minimum in $d_0$ coinciding with the fcc stacking and the maximum in $d_0$ coinciding with a stacking for which a pair of adjacent (111) planes are directly on top of each other.

While slip in fcc predominantly occurs on (111) planes, there are often multiple slip planes in more complex crystal structures such as hcp. Figure 8a schematically shows two slip planes in hcp: basal slip, prismatic slip and pyramidal slip. Each plane has a different periodicity. Figure 8b and fig. 8c shows the energy at fixed spacing $d$ for the pyramidal and prismatic slip planes of hcp Mg. The DFT method to calculate these energy surfaces was the same as that used for Al in Figure 7 except that a plane wave cutoff of 650 eV was used.
FIG. 7. Figure 7a: Calculated binding energy versus separation as two halves of an fcc crystal are separated between a pair of adjacent (111) planes. The dashed lines represent UBER fits through the first-principles (DFT) data points. Separation curve for perfect fcc, a stacking fault, and a translation that places a pair of adjacent (111) planes directly on top of each other are shown in blue, orange, and green respectively. Figure 7b: Surface energy for shifts at $d_0$ relative to equilibrium Al. Figure 7c: Lowest energy separation values $d_0$ relative to equilibrium Al.

FIG. 8. Figure 8a Schematic of different slip planes of an hcp system. Figure 8b Surface energy of the pyramidal plane for shifts at the minimum energy separation $d$. Figure 8c Surface energy of the prismatic plane for shifts at the minimum energy separation $d$. 
B. Crystallographic models of twisted two-dimensional materials: triangular lattices and honeycombs

**MultiShifter** facilitates the construction of crystallographic models of twisted two-dimensional materials. As described in section [III B], most twist angles \( \theta \) do not produce structures that have periodic boundary conditions. The imposition of periodic boundary conditions, therefore, requires an adjustment of the target twist angle \( \theta \) by \( \Delta \theta \) and some degree of strain within the twisted two-dimensional building blocks. We explore the variation in the error \( \Delta \theta \) in the target twist angle and the strain within the twisted building blocks due to the imposition of periodic boundary conditions for a pair of triangular lattices. This is of relevance for two-dimensional materials such as MoS\(_2\) and graphene since their two-dimensional lattices are triangular. Of interest is the variation of \( \Delta \theta \) and strain with twist angle \( \theta \) and maximum super cell size.

**MultiShifter** determines the best super cell for a twisted bicrystal with target twist angle \( \theta \) that is smaller than a user specified maximum. The “best” super cell is defined as the super cell that minimizes \( \sqrt{\eta_1^2 + \eta_2^2 + \eta_3^2} \). This strain metric is equal to zero for super cells that are commensurate with the lattices of the twisted bicrystal. The size of a super cell is measured in terms of the number of primitive two-dimensional unit cells of the fundamental slab building blocks.

Figure 9a and Figure 10 shows \( \Delta \theta \) and \( \sqrt{\eta_1^2 + \eta_2^2 + \eta_3^2} \) as a function of \( \theta \) for two scenarios. The error \( \Delta \theta \) versus \( \theta \) of Figure 9a is for super cells that were generated using the primitive Moiré lattice only. For small angles close to zero, the absolute error is small, however, for larger angles, the error \( \Delta \theta \) can be very large. The black circles correspond to angles for which there is a commensurate super cell that can accommodate the twisted pair of bicrystals. As is evident in Figure 9a, the primitive Moiré lattice is not sufficiently large to identify the commensurate super cell for those special angles, as large fraction of them have large errors \( \Delta \theta \). Similarly, the strain for super cells generated using the primitive Moiré lattice is also large as shown in Figure 10a. Figure 2b plots \( \Delta \theta \) versus \( \theta \) as determined by considering super cells of the Moiré lattice to identify the optimal super cell of the twisted bicrystals. A cap of 1000 primitive unit cells was used for Figure 9b and Figure 10b. The error in the target angle \( \Delta \theta \) is dramatically reduced and almost all special angles for which commensurate super cells exist now have an error \( \Delta \theta \) equal to zero, indicating that all commensurate super cells have been found.

VI. DISCUSSION AND CONCLUSION

Two-dimensional defects in bulk crystals play an out-sized role in determining the properties of many crystalline materials. The energy of two-dimensional extended defects are a crucial ingredient to a variety of meso-scale models of fracture and dislocations.\(^{[25-32,35,39]}\) The information of relevance for such models can be encapsulated in a generalized cohesive zone model that describes the energy of a pair of bicrystals as a function of the perpendicular separation and parallel glide of a pair of crystallographic planes. In this paper, we have described how such a generalized cohesive zone model can be formulated and how crystallographic models can be constructed for the first-principles electronic structure calculations that are needed to parameterize the cohesive zone model. The **MultiShifter** code automates the construction of periodic crystallographic models of decohesion and glide. It can also be used to generate crystallographic models for surface calculations. However, with the exception of the simplest materials, most surfaces undergo surface reconstructions to eliminate dangling bonds. This requires an additional enumeration step.\(^{[25]}\)

Twisted two-dimensional materials are currently attracting much attention due to the promise of emergent electronic properties in such structures.\(^{[33]}\) We have described an approach to construct crystallographic models of twisted bilay-
FIG. 10. Resultant error from approximating the moiré lattice. Allowing for super cells that have more than one moiré can reduce the error, as shown by allowing the minimum, and up to 1000 lattice sites to occupy the periodic unit.

ers within periodic unit cells. Only a subset of twist angles generate bilayer structures that are periodic. For all other angles, the imposition of periodic boundary conditions results in an error in the target twist angle and some degree of strain within the constituent bilayers. MultiShifter constructs these crystallographic models and quantifies the twist angle error and the strain.

In the construction of cohesive zone models, a question often arises as to whether to allow for relaxations or not. When including relaxations, it is important to carefully extract the energy of homogeneous elastic relaxations of the adjacent slabs since the cohesive zone model should only describe the energy between the cleaved planes. These subtleties are discussed in great detail in [18]. For simple metals, empirical evidence suggests that relaxations do not need to be explicitly taken into considerations when parameterizing a cohesive zone model for decohesion [18].

Many applications require a generalized cohesive zone model for a multi-component solid. The cohesive zone model will then not only depend on the local concentration, but also on the local arrangement of different chemical species. This dependence can be accounted for with the cluster expansion approach [43], as has been done in the context of hydrogen embrittlement [44] and fracture in Li-ion battery electrode material [45]. Often cohesive zones must be treated as open systems to which mobile species can segregate and thereby alter cohesive properties. In these circumstances it is convenient to formulate cohesive zone models at constant chemical potential as opposed to constant concentration [46,47]. A cluster expansion approach has also been used to describe the dependence of the \( \gamma \) surface on composition and short-range ordering in a refractory alloy [48].

In summary, MultiShifter is a code that automates the construction of crystallographic models within periodic unit cells to enable the construction of cohesive zone models and the study of twisted bilayers of two-dimensional materials with first-principles electronic structure methods. MultiShifter can also be used to construct models with highly distorted local environments that are representative of dislocations, grain boundaries and surfaces for the purposes of training machine learned inter-atomic potentials.

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