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Atomistic Simulation and Virtual Diffraction Characterization of Alumina Interfaces: Evaluating Structure and Stability for Predictive Physical Vapor Deposition Models

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Atomistic Simulation and Virtual Diffraction Characterization of Alumina Interfaces: Evaluating Structure and Stability for Predictive Physical Vapor Deposition Models
Atomistic Simulation and Virtual Diffraction Characterization of Alumina Interfaces: Evaluating Structure and Stability for Predictive Physical Vapor Deposition Models

A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy in Mechanical Engineering

by

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Bachelor of Science in Chemical Engineering, 2010

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This dissertation is approved for recommendation to the Graduate Council.

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ABSTRACT

The objectives of this work are to investigate the structure and energetic stability of different alumina (Al$_2$O$_3$) phases using atomistic simulation and virtual diffraction characterization. To meet these objectives, this research performs molecular statics and molecular dynamics simulations employing the reactive force-field (ReaxFF) potential to model bulk, interface, and surface structures in the $\theta$-, $\gamma$-, $\kappa$-, and $\alpha$-Al$_2$O$_3$ system. Simulations throughout this study are characterized using a new virtual diffraction algorithm, developed and implemented for this work, that creates both selected area electron diffraction (SAED) and x-ray diffraction (XRD) line profiles without assuming prior knowledge of the crystal system. First, the transferability of the ReaxFF potential is evaluated by modelling different alumina bulk systems. ReaxFF is shown to correctly predict the energetic stability of $\alpha$-Al$_2$O$_3$ among the crystalline alumina phases, but incorrectly predicts an even lower energy amorphous phase. Virtual XRD patterns uniquely identify each phase and validate the minimum energy bulk structures through experimental comparison. Second, stable and metastable alumina surfaces are studied at 0, 300, 500, and 700 K. ReaxFF predicts minimum energy surface structures and energies in good agreement with prior studies at 0 K; however, select surface models at 500 and 700 K undergo significant reconstructions caused by the unnatural bias for a lower-energy amorphous phase. Virtual SAED analysis performed on alumina surfaces allow advanced characterization and direct experimental validation of select models. Third, ReaxFF is used to model homophase and heterophase alumina interfaces at 0 K. Predicted minimum energy structures of $\alpha$-Al$_2$O$_3$ interfaces show good agreement with prior works, which provides the foundation for the first atomistic study of metastable alumina grain boundaries and heterophase alumina interfaces. Virtual SAED patterns characterize select alumina interfaces and help guide the construction of
low-energy heterophase alumina interfaces by providing insight into crystallographic compatibilities. Combined, the energetic data extracted from bulk, surface, and interface simulations as well as insights gained through virtual diffraction will aid the development of mesoscale predictive models of polycrystalline alumina formation during physical vapor deposition.
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DEDICATION

Mathew Stewart Nye

&

Leigh Whitney Nye
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LIST OF ABBREVIATIONS AND VARIABLES

Applications

PVD  Physical vapor deposition
SZD  Structural zone diagram

Crystallography and Grain Boundaries

XRD  X-ray diffraction (pattern)
SAD  Selected area diffraction (pattern)
SAED Selected area electron diffraction (SAED)
FCC  Face-centered cubic crystal
BCC  Body-centered cubic crystal
HCP  Hexagonal closed-packed crystal
GB  Grain boundary
STGB Symmetric tilt grain boundary
CSL  Coincident site lattice
$\Sigma$ Reciprocal density of coincident lattice sites between crystal regions

Atomistic Simulations

MD  Molecular dynamics
MS  Molecular statics
NVE Microcanonical ensemble – number of atoms, volume and total energy constant
NVT Canonical ensemble – number of atoms, volume and temperature constant
NEB  Nudged elastic band
ReaxFF Reactive Force-Field
LAMMPS Large-scale Atomic/Molecular Massively Parallel Simulator
EAM  Embedded-atom method

$N$ Number of atoms
$V$ System volume
$T$ System temperature

$F_i$ Force acting on atom $i$
$U_i$ Interatomic potential function evaluated at atom $i$
$\nabla$ Spatial gradient operator
$r_i$ Initial position of atom $i$
$\dot{r}_i$ Velocity of atom $i$
$\ddot{r}_i$ Acceleration of atom $i$
$m_i$ Atomic mass of atom $i$
$k_B$ Boltzmann constant.

$t$ Time at step

$\Delta t$ Duration of a timestep

$\zeta$ Frictional term added for Nosé-Hoover style thermostats

$\nu_T$ Thermostat rate added for Nosé-Hoover style thermostats

$T_0$ Temperature of external thermal reservoir for Nosé-Hoover style thermostats

$r_N$ All atom positions

$U_i(r_N)$ Potential energy of the system evaluated with all atoms

$k$ Arbitrary state of the system during molecular statics

$\alpha^{(k)}$ Magnitude of the step used during molecular statics

$d^{(k)}$ Search direction used during molecular statics

$\beta^{(k)}$ Update parameter used during molecular statics

$r_{ij}$ Interatomic distance between atoms $i$ and $j$

RDF Radial distribution function

CNA Common neighbor-analysis

HF Hartree-Fock

DFT Density functional theory

LDA Local density approximation

GGA Generalized gradient approximations

**Interatomic Potentials**

$E_{\text{Pair}}$ Potential energy contribution due to pair-wise interactions

$A$ Magnitude of repulsive interaction in Buckingham potential

$\rho$ Radius of repulsive interaction in Buckingham potential

$C$ Optimization parameter for attractive interaction in Buckingham potential

$E_{\text{Coulomb}}$ Potential energy contribution due to Coulombic forces (electrostatic interactions)

$q_i$ Electrostatic charge associated with atom $i$

$C$ Energy-conversion and magnitude constant for Coulombic interactions

$\varepsilon$ Dielectric constant

$\gamma_{ij}$ Overlap parameter that adjusts for shielding effects

$E_{\text{Bond}}$ Potential energy contribution due to bonds (ReaxFF)

$BO_{ij}$ Continuous bond order parameter

$BO_{ij}$ Corrected bond order parameter
\( \sigma \) Single bond
\( \pi \) Double bond
\( \pi\pi \) Triple bond
\( p_{bo,n} \) Bond order parameters
\( D_e \) Bond energy parameter
\( p_{be} \) Bond energy parameter
\( \Delta_i' \) Corrected difference between the bond order and the expected from its valency
\( \text{Val}_i \) Valency of atom \( i \)
\( \Delta_{lp,i} \) Difference between the number of lone pair electrons and expected value
\( n_{lp,i} \) Number of lone pair electrons around atom \( i \)
\( \Delta_i^e \) Difference between the number of valence electrons in the outer shell and the sum of the bond orders
\( \lambda_n \) General parameters for ReaxFF \((n=1-33)\)

\( E_{vdWaals} \) Potential energy contribution due to van der Waals interactions (ReaxFF)
\( D_{ij} \) ReaxFF parameter describing the magnitude of the van der Waals energy
\( \alpha_{ij} \) ReaxFF parameter describes the width of the minima well
\( r_{vdW} \) ReaxFF parameter describing the ideal interatomic distance for the interaction
\( \lambda_W \) ReaxFF parameter optimizing shielding interactions

**Virtual Diffraction**

relp Reciprocal lattice point
relrod Reciprocal lattice rod

\( I_e(K) \) Electron diffraction intensity
\( I_x(K) \) X-ray diffraction intensity
\( F(K) \) Structure factor
\( F^*(K) \) Complex conjugate of structure factor
\( Lp(\theta) \) Lorentz-polarization factor
\( f_j \) Atomic scattering factor

\( \theta \) Diffraction (scattering) angle
\( \lambda \) Wavelength of monochromatic radiation
\( d_{hkl} \) Interplanar distance

\( K \) Reciprocal lattice vector
**$k_D$**  
Diffracted wave vector

**$k_i$**  
Incident wave vector

**$k$**  
A spherically averaged position in reciprocal space

**$B_n$**  
Reciprocal lattice axis

**$A_n$**  
Real space lattice axis

**$c_n$**  
Resolution parameter for virtual diffraction

**$a_i$**  
Length of a unit cell lattice in direction $i$

**$N_i$**  
Number of units cells in direction $i$

**Computer Science**

**MPI**  
Message passing interface

**OpenMP**  
Open multi-processing

**FFT**  
Fast Fourier transform

**MIC**  
Many integrated core

**VSMP**  
Virtual symmetric multiprocessing

**HPC**  
High performance computing

**AHPCC**  
Arkansas High Performance Computing Center

**TACC**  
Texas Advanced Computing Center

**SDSC**  
San Diego Supercomputing Center

**NCSA**  
National Center for Supercomputing Applications

**XSEDE**  
Extreme Science and Engineering Discovery Environment

**ECSS**  
Extended Collaborative Support Services
LIST OF PAPERS


Chapter 1: Introduction

1.1 Motivation

The motivation for this research is a desire to advance the understanding of microstructure formation and evolution during physical vapor deposition of polymorphic thin films. Despite vast experimentation and industrial use (cf. [1,2]), no quantitative, predictive model for microstructure formation and evolution has been established for the vapor deposition process of polycrystalline materials. This research uses atomistic simulations to elucidate important energetic and structural information pertaining to the surfaces and interfaces that influence phase nucleation and growth during physical vapor deposition. In addition, preliminary simulated ion bombardment on select alumina models are performed to understand the nanoscale mechanisms leading to phase nucleation and transformation. The data gathered from these atomistic simulations lay the framework for advanced phase-field simulations to provide a detailed understanding of the role of process conditions on microstructure morphology at the mesoscale.

Alumina is an ideal material system for this research due to its well-known polymorphism and vast industrial use in vapor-deposited coatings [3–6]. Vapor-deposited alumina thin films are used in electronic devices, optics, and as durable coatings in the cutting tool industry [7,8]. While these industries have different needs for alumina, the process by which the thin films are created is similar and the impact of microstructure on material properties is common to all applications.

Atomistic simulations provide an established route to compute the crucial energetic and structural data needed to create mesoscale predictive models of physical vapor deposition of polycrystalline alumina. Atomistic simulations provide the flexibility and control needed to systematically create, test, and characterize a variety of alumina interface and surface structures.
that could only be achievable through a much larger experimental study. However, the complex structures of the various alumina phases pose a unique challenge for atomistic simulation characterization. The traditional atomistic simulation characterization methods that work well with cubic based structures (such as radial distribution functions [9], centrosymmetry [10], common neighbor-analysis [11], etc.) do not clearly identify the subtle structure differences among the various alumina phases. This motivated the development and implementation of a characterization technique for atomistic simulations based on the phenomena of x-ray and electron diffraction.

### 1.2 Alumina

Alumina is an abundant ceramic material that exhibits extraordinary structural flexibility [3–6]. The different Al₂O₃ phases are characterized by a range of unique physical properties which make them useful in a variety of coating applications [7,8]. These properties stem from only subtle differences within the crystal structure of its phases. In general, the alumina phases are composed of a close-packed oxygen sublattice surrounded by aluminum interstitials partially filling the octahedral and tetrahedral sites to maintain stoichiometry [12], shown in Figure 1.1. The type of close-packed arrangement of the oxygen sublattice and the degree of symmetry of the aluminum interstitials within each alumina unit cell determines the phase and properties of the material.

The structure of alumina phases has been studied extensively in the literature using both simulation and experimental approaches (cf. [13–23]). These studies show that both the κ- and α-Al₂O₃ crystal structures contain a face-centered-cubic (FCC) oxygen sublattice, whereas θ- and γ-Al₂O₃ oxygen atoms form a the hexagonal closed pack (HCP) arrangement [17]. Aluminum atoms partially fill 2/3 of the oxygen interstitial sites with varying degrees of symmetry. The α-
Al$_2$O$_3$ phase, corundum, is the only thermodynamically stable alumina phase. The $\alpha$-Al$_2$O$_3$ crystal is formed from a five atom tetragonal unit cell that contains the most symmetric ordering of the aluminum interstitials which are present only in octahedral sites [3]. The symmetry of the aluminum interstitials increases the density of $\alpha$-Al$_2$O$_3$ and promotes directionality of the ionic
bonds between the atoms.

The metastable alumina phases have decreasing symmetry among the aluminum interstitials, which decreases their density and weakens bond strength by reducing bond directionality. The κ-Al₂O₃ metastable phase consists of an eight molecular unit (40 atom) orthogonal unit cell. In κ-Al₂O₃, the aluminum interstitials favor the octahedral sites as shown in a first-principles study by Yourdshahyan et al. [13–15]. The θ-Al₂O₃ phase has a ten atom unit cell with monoclinic symmetry as determined by Zhou et al. [22] and Borosy et al. [23] using experimental and simulation studies, respectively. Using a first-principles study, Borosy et al. showed that the aluminum atoms in θ-Al₂O₃ are evenly distributed in both the octahedral and tetrahedral sites [23]. The γ-Al₂O₃ phase is the least symmetric metastable phase examined in this research. It’s structure was studied in detail by Paglia et al. using both experimental and simulation approaches [20,21]. In their work, Paglia et al. showed that γ-Al₂O₃ forms both cubic and slightly tetragonal-distorted spinel-like structures with aluminum interstitials located in 60-75% of oxygen octahedral sites of a 160-atom unit cell.

Due to subtle structural differences among the phases as well as the large unit cells for some of the metastable phases, identification and characterization of alumina based solely on local atomic positions is difficult. However, experimental characterization techniques such as x-ray diffraction (XRD) and selected area electron diffraction (SAED) have proven successful in distinguishing between alumina phases [24–28].

Alumina surfaces, in particular those formed by the κ- and α-Al₂O₃ phases, are widely used as protective coatings due to their wear resistance, chemical inertness, resistance to thermal shock, and high hardness [29,30]. However, the alumina material system is known to have several other metastable states whose properties are less ideal for these purposes that can readily
form during deposition depending on the processing conditions [31]. For example, the less symmetric $\theta$- and $\gamma$-$\text{Al}_2\text{O}_3$ metastable phases form readily at lower processing temperatures, approximately 704 K (30% of the melting temperature) [31]. These metastable phases are characterized by having lower surface energies and therefore exhibit higher surface areas, which would be more appropriate for catalytic supports.

The alumina material system experiences a complex phase transition series that is dependent on the material precursors and processing conditions [32–34]. Schematics of the phase transition series, such as the one in Figure 1.2 modeled after the results of Levin et al. [32], can be used as guides in determining the necessary processing conditions required for phase transformation. However, these schematics alone cannot accurately predict the criteria for phase transformation during physical vapor deposition. Instead, researchers must continually characterize alumina surfaces to determine the state of transformation under their particular processes.

Figure 1.2: Transition sequence for alumina for different precursor materials and processing conditions [32].
1.3 Physical Vapor Deposition

Physical vapor deposition (PVD) is a process by which atoms condense from the vapor phase to form a solid thin film or coating on a substrate [35]. For example, in sputtering, atoms are removed from a target by high-energy ion bombardment. The removed atoms are ejected into the deposition chamber and upon reaching the substrate they condense to reform a solid material, resulting in energy reduction through bond formation. Deposition may be performed in either inert (Ar) or active atmospheres, i.e., prior to condensation atoms in the vapor phase may chemically react with gaseous species in the chamber to form compounds (e.g., Al$_2$O$_3$ [36,37]).

The microstructure of the solid deposit is largely determined by deposition conditions and the materials employed. For example, at low substrate temperatures or high deposition rates the adatoms that accumulate on the substrate surface may not have sufficient mobility to migrate to lower energy configurations prior to being covered up by the next deposited layer. Schematic maps, known as structure zone diagrams (SZDs) [38], have been proposed to correlate coating

![Image](image_url)

Figure 1.3: Example SZD showing the influence of deposition variables on thin film microstructure based on phenomenological observations [39].
morphology with deposition temperature and pressure, as shown in Figure 1.3 from Thornton [39]. The mechanical, electrical, and optical properties of the coating will change (often dramatically) within the deposition space in Figure 1.3. For evaporated or sputtered films, process zones are identified (1, 2, 3, and T in Figure 1.3) which differ by level of porosity and grain morphology.

While experimental observations of the type that lead to the SZD shown in Figure 1.3 are of tremendous importance, SZDs are heavily phenomenological and are only predictive if deposition is performed within the same process space. Consequently, different SZDs have been proposed in the literature for different PVD techniques, including sputtering [39], metal evaporation [40,41], RF sputtering [42], and magnetron sputtering [43,44]. Furthermore, SZDs do not account for the individual nanoscale mechanisms associated with microstructure formation and evolution (only processing conditions). Therefore, extensive experimentation would be required to extend SZDs to polymorphic thin films. Despite vast experimentation and industrial use, a quantitative model for microstructure formation and evolution during vapor deposition of polymorphic thin films has not been established.

1.4 Objectives

There are three main objectives of this work: (1) to develop and implement a virtual diffraction algorithm that generates x-ray and electron diffraction patterns directly from atomistic simulation data without a priori knowledge of the simulated crystal structure, (2) to utilize molecular statistics and molecular dynamics simulations to evaluate the energetic stability of different bulk alumina models as well as characterize metastable and stable alumina surfaces, and (3) to utilize molecular statistics and virtual diffraction to characterize homophase and heterophase alumina interfaces. Combined, the objectives of this work lay the framework for a
larger research study with the overarching goal to elucidate the fundamental nanoscale
mechanisms that drive microstructure formation and evolution during PVD of polymorphic thin
films.

1.5 **Dissertation Structure**

Because a portion of this research has been published previously in scholarly journals, this
dissertation is structured in the "Published Papers" format in line with the University of
Arkansas-Fayetteville Graduate School where published works are presented in their entirety as
unique chapters to the dissertation. Before presenting published works, Chapter 2 provides a
thorough background on the methods of atomistic simulations used throughout this research as
well as outlines the traditional characterization methods commonly used in atomistic simulations.
Chapter 2 also includes documentation outlining the search for an appropriate modeling method
(i.e., interatomic potential) to describe the complex interatomic interactions within Al₂O₃.

Chapter 3 contains work conducted in collaboration with Dr. Stephen Foiles at Sandia
National Laboratories that investigates grain boundary structures and mobility in nickel [45].
This work is included to highlight the capability of atomistic simulations to study interface
structures and dynamics. Specifically, this work emphasizes the construction of grain boundary
interfaces, techniques for analyzing grain boundary motion, and the capabilities of traditional
atomistic methods to characterize face-centered cubic materials.

The development and implementation of a novel computational algorithm that produces
virtual diffraction patterns are discussed in Chapters 4-6. Specifically, Chapter 4 includes a
detailed description of the virtual diffraction algorithm along with a thorough case study
examining select symmetric tilt grain boundaries in nickel, which was done in collaboration with
Dr. Laurent Capolungo from the Georgia Institute of Technology [46]. Chapter 5 includes an
additional description of the virtual diffraction algorithm as well as results highlighting its capability and versatility through findings from atomistic simulations of select symmetric tilt nickel grain boundaries, nanocrystalline copper models, and a heterogeneous interface formed between $\alpha$-Al$_2$O$_3$ (0001) and $\gamma$-Al$_2$O$_3$ (111) [47]. Note, analysis of copper nanocrystalline samples found in Chapter 5 are performed by fellow doctoral student Mehrdad M. Sichani.

Chapter 6 highlights the implementation and optimization of the virtual diffraction algorithm into the LAMMPS molecular dynamics simulator [48] as well as the creation of an automated workflow that facilitates high throughput computation and visualization. Work presented in Chapter 6 is done in collaboration with Extended Collaborative Support Services (ECSS) provided though the Extreme Science and Engineering Discovery Environment (XSEDE). Specifically, the optimization of the virtual diffraction code for multilevel parallelism and offloading to heterogeneous computing architecture discussed in Chapter 6 are done in collaboration with Dr. Yang Wang and Dr. Lars Koesterke. Workflow development and integrated visualization discussed in Chapter 6 are done in collaboration with and Dr. Sudhakar Pamidighantam and Dr. Mark Vanmoer, respectively.

Chapter 7 discusses atomistic simulations of bulk alumina and alumina surfaces structures modeled using the reactive force-field (ReaxFF) interatomic potential [49,50] at 0, 300, 500, and 700 K [51]. Here, virtual x-ray diffraction patterns uniquely identify each phase and validate minimum energy bulk structures through experimental comparison. In addition, virtual selected area electron diffraction patterns identify significant structural reconstructions affecting select alumina surface models at 500 and 700 K. Energetic data computed from these simulations are tabulated for use in the larger-scale predictive models of polymorphic thin film PVD.

Chapter 8 contains work discussing homophase and heterophase alumina interfaces modeled
with ReaxFF [52]. The contained study was specifically limited to molecular statics simulations performed at 0 K after observing limitations in the surface study in Chapter 7 that more greatly affect simulations at temperature when using the ReaxFF interatomic potential. ReaxFF shows good agreement to prior studies in its predicted interface structures and energies for most homophase \( \alpha \)-\( \text{Al}_2\text{O}_3 \) twins providing confidence to the novel study of metastable \( \text{Al}_2\text{O}_3 \) homophase boundaries and heterophase \( \text{Al}_2\text{O}_3 \) interfaces. Here, virtual diffraction is used to aid the construction of low energy heterophase interfaces as well as to provide further validation of select modeled interfaces through experimental comparison.

Lastly, Chapter 9 summarizes the major findings and makes recommendations for future research. Specifically, this includes recommendation for further analysis of nanoscale mechanisms leading to phase transformation though simulated ion bombardment on alumina surfaces as well as an in-depth study highlighting the use of virtual diffraction characterization to optimize interface misorientation. Preliminary results for such studies are included in this chapter. The conclusion includes discussions of the broader impact of the virtual diffraction algorithm addressing its ability to act as a bridge between materials simulation and experiments.
References


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[46] Coleman SP, Spearot DE, and Capolungo L. (2013) Virtual diffraction analysis of Ni [0 1 0] symmetric tilt grain boundaries, Modelling and Simulation in Materials Science and Engineering, 21, 055020.


Chapter 2: Background

2.1 Atomistic Simulations

This research utilizes molecular dynamics (MD) and molecular statics (MS) simulations to model alumina surfaces and interfaces as well as to identify the atomic mechanisms that drive phase formation and transformation during physical vapor deposition. In the MD and MS frameworks, each atom is represented by a single point mass in space that encompasses both the nucleus and the orbiting electrons. Interactions between atoms are governed by an interatomic potential that describes the conformational (potential) energy of the system. The accuracy of these atomic models depends on the choice of interatomic potential, which depends on the material system and phenomena of interest.

In the MD and MS frameworks, the forces acting on an atom, $F_i$, are computed from the specified interatomic potential function $U_i$, using the spatial gradient operator $\nabla$ via

$$ F_i = -\nabla U_i, $$

(2.1)

where $U_i$ is written as a function of the atom positions within the simulation. There are many types of interatomic potentials that are created and optimized to model specific materials or phenomena. In its simplest form, interatomic potentials will contain repulsion terms that increase in energy when atomic nuclei overlap and attraction terms that seek to bring atoms together due to the interactions between the negatively charged electron cloud of one atom and the positively charged atomic nuclei of another atom. Together, the repulsive and attractive terms will have energy minima that bring atoms to an ideal separation distance and confirmation. The atomic forces, $F_i$, computed through the interatomic potential are conservative and thus are not specific on the path taken by the atoms. More details concerning interatomic potentials and
the interatomic potential chosen to model the alumina phases in this work are included in Section 2.1.3 and Section 2.2 respectively.

Atomistic simulations commonly use periodic boundary conditions, introduced by Born and von Kármán in 1912 [1], to mimic infinite systems while modeling a finite collection of atoms. Periodic boundary conditions are constructed from 26 (in three-dimensions) replicated simulation cells surrounding a primary simulation model. The atom positions within each replica are identical after making appropriate adjustments for the periodic length defined by the dimensions of the primary simulation model. Atom velocities within each cell are also identical among the replicas and atom motion across replica boundaries results in symmetric motion of atoms across the mirroring boundaries.

This research utilizes the classical atomistic simulation package LAMMPS, which is distributed by Sandia National Laboratories under the terms of the GNU Public License [2]. LAMMPS runs in serial or in parallel using message-passing techniques (MPI) and a spatial-decomposition of the atomistic simulation domain. LAMMPS is easily extendable with “plug-and-play” functionality so that new interatomic potentials, new atomistic computations, or new integration schemes can be easily integrated without modifying the core of the atomistic simulation package. This "plug-and-play" functionality serves as the framework for an enhanced virtual diffraction computation, discussed in detail in Chapters 3-5, which computes diffraction intensity data concurrently while the atomistic simulation progresses.

2.1.1 Molecular Dynamics

In MD simulations, the forces acting on each atom $\mathbf{F}_i$, calculated with Eq. (2.1), are incorporated with a set of equations of motion to compute the atomic trajectories (position and momentum) over time. To solve the equations of motion, an integration algorithm requires
initial positions, \( \mathbf{r}_i \), velocities \( \dot{\mathbf{r}}_i \), and acceleration, \( \ddot{\mathbf{r}}_i \), for the \( N \) atoms contained within the simulation. Initial atom positions, \( \mathbf{r}_i \), are determined by the crystal structure of the material of interest. Initial atom velocities, \( \dot{\mathbf{r}}_i \), are typically randomized to achieve a predefined temperature, \( T \), computed through the equipartition formula [3],

\[
\frac{1}{2} \sum_{i=1}^{N} m_i \dot{\mathbf{r}}_i^2 = \frac{3}{2} N k_B T
\]  

(2.2)

where \( m_i \) is the atomic mass and \( k_B \) is the Boltzmann constant. The acceleration for each atom, \( \ddot{\mathbf{r}}_i \), is computed from the forces acting on an atom by coupling the forces \( \mathbf{F}_i \) computed through Eq. (2.1) with Newton’s 2\textsuperscript{nd} Law of Motion (for the microcanonical ensemble)

\[
\mathbf{F}_i = m_i \ddot{\mathbf{r}}_i .
\]  

(2.3)

Using initial values for atom positions \( \mathbf{r}_i \), velocities \( \dot{\mathbf{r}}_i \), and acceleration \( \ddot{\mathbf{r}}_i \), the integration algorithm solves the 6\( N \) coupled differential equations in an iterative manner to determine the 3\( N \) atom positions and 3\( N \) atom velocities over time. The microcanonical ensemble (NVE) conserves the number of atoms, the system volume, and total energy throughout the simulation.

A common integration algorithm for NVE dynamics used in LAMMPS is the velocity-Verlet method. The algorithm described by the velocity-Verlet method computes updated atom trajectories at time \( (t + \Delta t) \) from the initial trajectories at time \( t \) through the sequence of equations [4]:

\[
\dot{\mathbf{r}}_i \left( t + \frac{\Delta t}{2} \right) = \dot{\mathbf{r}}_i (t) + \frac{\Delta t}{2} \ddot{\mathbf{r}}_i (t) ,
\]  

(2.4)

\[
\mathbf{r}_i (t + \Delta t) = \mathbf{r}_i (t) + \dot{\mathbf{r}}_i (t) \Delta t .
\]  

(2.5)

\[
\mathbf{F}_i = -\nabla U_i \left[ \mathbf{r}_i^N (t + \Delta t) \right] , \text{ and}
\]  

(2.6)
where $\Delta t$ is the duration of the timestep. For MD simulations, the duration of the timestep is often on the order of 1 femtosecond ($10^{-15}$ s) such that the simulation accurately captures the atomic vibrations critical to represent temperature within a collection of atoms.

If desired, augmented versions of Newton's second law can be implemented into the integration algorithm to couple the system dynamics to the surrounding environment. The augmented versions can incorporate thermodynamic boundary conditions such as temperature and pressure control into the simulation [5–9]. For example, this work utilizes the canonical ensemble (NVT), which is designed to maintain a constant number of particles, volume, and system temperature within the simulation. LAMMPS implements the NVT ensemble using the equations developed by Shinoda et al. [10] which are based on a Nosé-Hoover [7] style thermostat. Nosé-Hoover [7] style thermostats augment the equations of motion by adding an additional frictional term, $\zeta$, that effectively connects the system to an eternal thermal reservoir of constant temperature, $T_0$, by using the equations

$$\mathbf{F}_i = m_i \ddot{\mathbf{r}}_i + \zeta m_i \dot{\mathbf{r}}_i \quad \text{and}$$

$$\zeta = v_T^2 \left( \frac{T}{T_0} - 1 \right)$$

where, $v_T$ is the thermostat rate. Application of the Nosé-Hoover thermostat dynamically adjusts the system temperature to the desired thermal reservoir temperature, $T_0$, by scaling the atomic velocities.

### 2.1.2 Molecular Statics

Molecular statics simulations seek to find the minimum energy conformation of a system at
In MS simulations, the forces acting on each atom $F_i$, calculated with Eq. (2.1), are incorporated into numerical algorithms that search for atom coordinates, $r_N$, that minimize a defined objective function. For the microcanonical ensemble, MS simulations seek to minimize the potential energy of the system as computed by the interatomic potential, $U_i(r_N)$.

Commonly, MS simulations use iterative methods to displace the atoms within the simulation along a specified search direction to find a lower energy state,

$$r_N^{(k+1)} = \begin{cases} r_N^{(0)} & \text{if } k = 0 \\ r_N^{(k)} + \alpha^{(k)} d^{(k)} & \text{if } k > 0 \end{cases}.$$

(2.10)

Here, $k$ is a particular iteration representing an arbitrary state of the system, $\alpha^{(k)}$ is the magnitude of the step, and $d^{(k)}$ is the search direction.

The choice of search direction depends on the type of minimization routine used. For the microcanonical ensemble, the search direction employs the forces acting on the atoms within the simulation, $F = -\nabla U_i(r_N)$, which represents the direction of greatest decrease of potential energy function as calculated through the gradient operator. At each iteration, the search directions are computed through,

$$d^{(k+1)} = \begin{cases} F^{(0)} & \text{if } k = 0 \\ F^{(k+1)} + \beta^{(k+1)} d^{(k)} & \text{if } k > 0 \end{cases}.$$

(2.11)

where $\beta^{(k)}$ is an update parameter that scales the input from the previous search direction. If $\beta^{(k)} = 0$, which is known as the method of steepest decent, no input from the previous search direction is included and step directions follow normal to the contour lines of the potential energy surface. For most atomistic simulations, the simplicity of the method of steepest decent is overwhelmed by the complexity of the potential energy surfaces, which slows the convergence.
of this method. To handle the complex potential energy surfaces found in atomistic simulations efficiently, MS simulations will commonly utilize a non-linear conjugate gradient method that incorporates values for the update parameter $\beta^{(k)}$ and thus utilizes information from the previous search direction. For example, the LAMMPS implementation of the non-linear conjugate gradient method uses the update parameter introduced by Polak and Ribière [11],

$$\beta^{(k+1)} = \frac{\mathbf{F}^{(k+1)T} \left( \mathbf{F}^{(k+1)} - \mathbf{F}^{(k)} \right)}{\mathbf{F}^{(k)T} \mathbf{F}^{(k)}}. \quad (2.12)$$

Using this update parameter, the successive search direction is conjugate [12] to the previous which the improves convergence to a minimum energy configuration when examining complex potential energy surfaces.

The magnitude of the step size $\alpha^{(k)}$ is chosen to minimize the objective function defined by the MS simulation, which for the microcanonical system is,

$$U_i \left( r_{N}^{(k)} + \alpha^{(k)} d^{(k)} \right). \quad (2.13)$$

In LAMMPS, the step size $\alpha^{(k)}$ is evaluated through an iterative line search algorithm. The line search algorithm moves the atoms along the search directions until an energy minimum is attained. More details about specific minimization routines can be found in [12].

### 2.1.3 Interatomic Potentials

As shown in the prior sections, the specified interatomic potential function, $U_i$, plays a key role in modeling the interactions between atoms and thus effects the prediction of material behavior. Interatomic potentials can take many forms in order to describe different material behavior, and are often decomposed as a summation of unique terms describing individual atomic interactions,
The most simple of interatomic potentials will consist of a single formula that describes the pairwise interaction of atoms, \( E_{\text{pair}} \), based solely on the interatomic distance \( r_{ij} \) between atom \( i \) and atom \( j \). An example pair potential is the Buckingham potential, described by \cite{13},

\[
E_{\text{Pair}} = A \exp \left( -\frac{r_{ij}}{\rho} \right) - \frac{C}{r_{ij}^6} \quad r_{ij} < r_c \ .
\]  

(2.15)

In the Buckingham potential, the first term describes the repulsive interaction of atoms as their interatomic distances decreases and atomic nuclei begin to overlap. This repulsion has spherical symmetry, with a magnitude based on the energy coefficient \( A \) and a radius related to the distance parameter \( \rho \). In Eq. (2.15), the second term represents the energy due to attractive van der Waals forces between atomic nuclei and electron clouds that are optimized for a particular material through the \( C \) parameter.

For materials that have ionic bonding, additional terms maybe added to the interatomic potential to account for the energy due to Coulombic forces (electrostatic interactions), \( E_{\text{Coulomb}} \). The standard Coulombic interaction potential takes the form,

\[
E_{\text{Coulomb}} = \frac{C q_i q_j}{\varepsilon r_{ij}} \quad r_{ij} < r_c \ ,
\]  

(2.16)

where \( q_i \) and \( q_j \) are the charges associated with atom \( i \) and atom \( j \) respectively, \( C \) is an energy-conversion constant, and \( \varepsilon \) is the dielectric constant. Summed together, Eqs. (2.15) and (2.16) will describe an energy minima, like that shown in Figure 2.1, at the ideal interatomic separation between atom \( i \) and atom \( j \).

Interatomic potentials with greater complexity are derived from similar principles as the two-
body potential in order to model specific phenomena found in a material system. For example, many-body potentials will include additional angular terms that describe the energetics caused by bending and torsional interactions that can be calculated between groups of atoms. These types of interactions are important to describe the potential energy surface in hydrocarbon systems [14,15] as well as other complex material systems [16].

![Figure 2.1: Representative Buckingham-Coulombic interatomic potentials made with fictitious parameters.](image)

2.1.4 First-Principles Methods

Because of its important material properties and wide industrial use, the alumina material system has been studied extensively using atomistic simulations modeled using first-principles frameworks, as will be discussed in Chapter 7 and 8. First-principles methods are similar to molecular statics simulations in that they are used to study atomic structure and energetics at 0 K. However, unlike molecular statics simulations, first-principles methods rely on a quantum mechanical approach that models electron interactions explicitly (cf. [17–19]). Briefly, first-principles methods compute molecular energy and determine optimized atomic structures by
finding approximate solutions to the many-body Schrödinger equation for all electrons within the model. The approximations used in the prior alumina studies can be divided into two classes: (1) Hartree-Fock (HF) theory and (2) density functional theory (DFT). Methods based on HF theory attempt to find approximate solutions directly to the Schrödinger equation by posing a linear combination of functions that form a basis set approximating the interactions of all individual atoms. Whereas methods based on DFT attempt to find exact solutions to a set of modified equations which approximate the Schrödinger equation based on electronic density. The DFT methods also subdivide into two general classes that differ in how each represents the interactions between the different electrons (exchange-correlation). In DFT, the exchange-correlation can be modeled by functions relying on the local electron density alone, called the local density approximation (LDA), or by functions that also take into account non-local effects posed by gradients of the electronic density, called the generalized gradient approximations (GGA). The accuracy of first-principles models depends on the chosen method, basis set, and exchange-correlation; however, generally accuracy increases with DFT approaches compared to HF and in particular the greatest accuracy is achieved when using GGA exchange-correlation [20].

2.2 Alumina Interatomic Potentials

2.2.1 Pair-wise Potentials

Three pair-wise interatomic potentials are examined for use in this work developed by Matsui [21], Van Hoang [22], and Sun [23]. The Matsui potential was created to be transferable to several metal oxide materials and has been used to study amorphous alumina [24–26], melting $\alpha$-Al$_2$O$_3$ [27,28], liquid alumina [27–29], as well as atomic layer deposition of Al$_2$O$_3$ [30]. The Van Hoang potential is of the Born-Mayer type [31] and has been used to study amorphous
alumina [32], liquid alumina [33], and super cooled alumina [34,35]. Sun developed a modified Matsui potential that is transferable to other binary metal oxides. Sun’s modified Matsui potential has been used to study bulk $\alpha$-$\text{Al}_2\text{O}_3$ [23] and the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) surface [36,37].

Nonlinear conjugate gradient potential energy minimizations are performed on each phase at 0 K to determine the predicted structure and potential energy of each simulation as a function of the interatomic potential, shown in Table 2.1. Here, all three pair-wise potentials fail to represent the critical thermodynamic stability of $\alpha$-$\text{Al}_2\text{O}_3$. Instead, each potential predicts a lower potential energy $\gamma$-$\text{Al}_2\text{O}_3$ phase, which is inconsistent with experimental results.

Table 2.1: Minimized potential energy (kcal/mol) for various $\text{Al}_2\text{O}_3$ crystalline phases as a function of pair-wise interatomic potentials.

<table>
<thead>
<tr>
<th>Potential</th>
<th>$\alpha$-$\text{Al}_2\text{O}_3$</th>
<th>$\gamma$-$\text{Al}_2\text{O}_3$</th>
<th>$\kappa$-$\text{Al}_2\text{O}_3$</th>
<th>$\theta$-$\text{Al}_2\text{O}_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sun</td>
<td>-24.027</td>
<td>-25.911</td>
<td>-23.628</td>
<td>-25.605</td>
</tr>
<tr>
<td>Van Hoang</td>
<td>-30.349</td>
<td>-31.185</td>
<td>-29.392</td>
<td>-30.707</td>
</tr>
</tbody>
</table>

The inability to accurately predict the thermodynamic stability of $\alpha$-$\text{Al}_2\text{O}_3$ likely stems from the simplicity of these pair-wise models, specifically concerning the manner in which electrostatic interactions are described. The pair-wise interatomic potentials for alumina developed by Matsui, Van Hoang, and Sun all represent electrostatic interactions as simple point charges which cannot account for ion polarization effects. Wilson et al. [38] have noted that accurate modeling of oxygen polarization is a key element in the development of a valid transferable alumina interatomic potential. The results from the pair-wise interatomic potential study clearly show a more complex interatomic potential is needed to model the varying atomic environments within the different alumina phases.
2.2.2 Reactive Force Field Potential (ReaxFF)

In this work, the reactive force field (ReaxFF) potential is implemented to describe the interactions between the aluminum and oxygen atoms. ReaxFF was first developed by van Duin, Goddard, and coworkers [39] for use in hydrocarbon systems, but has since been expanded to many other material systems. To be transferable to a wide set of material systems, ReaxFF includes ten terms to represent various phenomena related to different atomic interactions [40],

$$U_j = E_{\text{Bond}} + E_{\text{Over}} + E_{\text{Under}} + E_{\text{val}} + E_{\text{lp}}$$
$$E_{\text{pen}} + E_{\text{tors}} + E_{\text{conj}} + E_{\text{vdWaals}} + E_{\text{Coulomb}}$$

(2.17)

However, when describing the aluminum and oxygen interactions, only four of these interactions are considered [41,42],

$$U_j^{\text{Al}_2\text{O}_3} = E_{\text{Bond}} + E_{\text{Over}} + E_{\text{vdWaals}} + E_{\text{Coulomb}}$$

(2.18)

where $E_{\text{Bond}}$, $E_{\text{Over}}$, $E_{\text{vdWaals}}$, and $E_{\text{Coulomb}}$ account for the energy contributions due to bonding energies, overcoordination penalties, non-bonded van der Waals interactions, and electrostatic interactions respectively. These four contributions allow ReaxFF to accurately model dynamic bond breaking and formation as well as the polarization of charged atomic species.

In ReaxFF, the bond energy term, $E_{\text{Bond}}$, gives the potential its ability to capture bond formation and breakage dynamically without explicitly determining connectivity. Dynamic bonding is made possible using a continuous bond order parameter, $BO_{ij}$, computed as a function of the interatomic distance, $r_{ij}$. The bond order parameter contains three terms to provide different functional forms for $\sigma$, $\pi$, and $\pi\pi$ (single, double, and triple bonds respectively) [40].
\[ \text{BO}_{ij} = \text{BO}_{ij}^\sigma + \text{BO}_{ij}^\pi + \text{BO}_{ij}^{\pi\pi} \]

\[ = \exp \left[ p_{\text{bo},1} \cdot \left( \frac{r_{ij}}{r_{\sigma}} \right)^{p_{\text{bo},2}} \right] + \exp \left[ p_{\text{bo},3} \cdot \left( \frac{r_{ij}}{r_{\pi}} \right)^{p_{\text{bo},4}} \right] + \exp \left[ p_{\text{bo},5} \cdot \left( \frac{r_{ij}}{r_{\pi\pi}} \right)^{p_{\text{bo},6}} \right], \quad (2.19) \]

where \( p_{\text{bo},n} \ (n = 1, 2, 3, \ldots) \) are parameters used to optimize the potential for the various atomic interactions. The bond order parameter is corrected for overcoordination, described in detail in the seminal publication [39], which accounts for atomic configurations exceeding their expected number of max bonds. This corrected bond order term, \( \text{BO}_{ij} \), is used to compute the bond energy contribution through [40],

\[ E_{\text{Bond}} = -D_{\sigma} \cdot \text{BO}_{ij}^\sigma \cdot \exp \left[ p_{\text{be},1} \left( 1 - \left( \text{BO}_{ij}^\sigma \right)^{p_{\text{be},2}} \right) \right] - D_{\pi} \cdot \text{BO}_{ij}^\pi - D_{\pi\pi} \cdot \text{BO}_{ij}^{\pi\pi}, \quad (2.20) \]

where the \( D \) and \( p_{\text{be}} \) are parameters that are optimized to describe the dissociation energy for the various bond types found in the material system.

Considerations for atom overcoordination are further addressed through an energy penalty, \( E_{\text{Over}} \), which is added to atoms exceeding their expected number of bonds based on their valency. The overcoordination energy is computed through [40],

\[ E_{\text{Over}} = \sum_{j=1}^{n_{\text{bond}}} p_{\text{be},3} \cdot \text{BO}_{ij} \cdot \left( \frac{1}{\Delta_i + \text{Val}_i} \right) \cdot \lambda_i \cdot \left( \frac{1}{1 + \exp(\lambda_6 \cdot \Delta_i)} \right), \quad (2.21) \]

where \( \Delta_i \) is the corrected difference between the bond order and the expected from its valency.
\( \text{Val}_i \), and the variables \( p_{\text{be},3} \) and \( \lambda_6 \) are bond energy and general parameters, respectively, optimized for a particular material system. For non-hydrocarbon systems, \( \Delta_i \) is influenced by the presence of lone pair electrons, as shown in its computation [40],

\[
\Delta_i' = -\Delta_{lp,j}' \frac{1}{1 + \lambda_{33} \exp(\lambda_{33} \text{SOV})} + \sum_{j=1}^{\text{nbond}} \text{BO}_{ij} - \text{Val}_i, \tag{2.22}
\]

where,

\[
\text{SOV} = \sum_{j=1}^{\text{neighbor}(j)} (\Delta_j - \Delta_{lp,j}') \text{BO}^\sigma_{ij}. \tag{2.23}
\]

Here, \( \Delta_{lp,j}' \) is the difference between the number of lone pair electrons, \( n_{lp,j} \), around an atom from the expected value based on its coordination. ReaxFF computes the number of lone pair electrons for each atom through [40],

\[
n_{lp,j} = \text{int} \left( \frac{\Delta_i}{2} \right) + \exp \left[ -\lambda_{16} \left( 2 + \Delta_i - 2 \cdot \text{int} \left( \frac{\Delta_i}{2} \right) \right)^2 \right] \tag{2.24}
\]

where \( \Delta_i \) is the difference between the number of valence electrons in the outer shell and the sum of the bond orders. In Eqs. (2.22) and (2.24), the \( \lambda \) variables are general parameters optimized for a particular material system. Note that in the alumina material system, overcoordination penalties are added to all atoms types; however, considerations for lone pair electrons primarily influences the expected valency of the oxygen atoms only.

In ReaxFF, non-bonded van der Waals interactions and Coulombic contributions are computed between all atom pairs within the simulation due to the consideration of dynamic bond breaking and formation through the bond energy. This differs from other bonded interatomic potentials that exclude bonded atoms when computing non-bonded van der Waals interactions.
and Coulombic contributions. The van der Waals interactions describe the repulsive forces that atoms experience as their atomic nuclei begin to overlap. In ReaxFF, the van der Waals interaction energy, $E_{\text{vdWaals}}$, is computed through [40],

$$E_{\text{vdWaals}} = D_{ij} \cdot \left\{ \exp \left[ \alpha_{ij} \cdot \left( 1 - \frac{f_{13}(r_{ij})}{r_{vdW}} \right) \right] - 2 \cdot \exp \left[ \frac{1}{2} \cdot \alpha_{ij} \cdot \left( 1 - \frac{f_{13}(r_{ij})}{r_{vdW}} \right) \right] \right\}, \quad (2.25)$$

where

$$f_{13}(r_{ij}) = \left[ r_{ij}^{\lambda_{29}} + \left( \frac{1}{\lambda_{W}} \right)^{\lambda_{28}/\lambda_{28}} \right]^{1/\lambda_{28}}. \quad (2.26)$$

Here the parameters $D_{ij}$ describes the magnitude of the van der Waals energy, $\alpha_{ij}$ describes the width of the minima well, and $r_{vdW}$ describes the ideal interatomic distance for the interaction. For the van der Waals energy, the interatomic distances are augmented through Eq. (2.26) to account for shielding interactions that decreases the repulsive interaction between bonded atoms. The shielding interactions are optimized using the $\lambda_{W}$ and general $\lambda$ parameters in Eq. (2.26).

Electrostatic interactions are modeled with the Coulomb energy contribution, computed in ReaxFF through [39],

$$E_{\text{Coulomb}} = C \cdot \frac{q_i \cdot q_j}{\left[ r_{ij}^3 + \left( \frac{1}{\gamma_{ij}} \right)^3 \right]^{1/3}}, \quad (2.27)$$

where $C$ relates the magnitude of the energy and $\gamma_{ij}$ is an overlap parameter that adjusts for shielding effects. An important feature of the ReaxFF interatomic potential is its incorporation of dynamic charge calculations. The atomic charges $q_i$ and $q_j$ in Eq. (2.27), are not fixed; rather, they are continuously optimized based on atomic geometry at each timestep using the electron equilibration method [43]. The dynamic charge calculation allows modeling of
polarization effects; however, to maintain numerical stability of this charge equilibrium scheme, MD simulations incorporating the ReaxFF potential generally require sub-femtosecond timesteps.

The ReaxFF parameters used in this study were first optimized by Zhang et al. in 2004 for Al and $\alpha$-$\text{Al}_2\text{O}_3$ using ab initio calculations on Al-O clusters [41] then further tailored by Sen et al. in 2013 to better incorporate varying Al oxidation states during oxidation simulations [42]. The ReaxFF interatomic potential is much more computationally demanding and memory intensive when compared to other standard pair-wise and many-body potentials [44]. However, these additional computational costs are necessary to accurately describe the structures and thermodynamics of multiple metastable and stable alumina phases.

Because of its important material properties and wide industrial use, the alumina material system has been studied extensively using atomistic simulations modeled from first-principles and Newtonian physics frameworks. However, none of these previous atomistic studies has attempted to catalog the structural and thermodynamic properties of interfaces and surface structures with more than a couple alumina phases using the same method. Instead, the majority of prior atomistic studies sought to identify the minimum energy structures of select bulk alumina phases, surfaces, and interfaces. By narrowing the focus on select alumina components, prior atomistic simulation studies have avoided the challenges of uniquely characterizing the subtle structural differences among the various alumina phases as well as the distortion created by atomic relaxations at the interfaces. In addition, because the various methods used previously to study alumina rely on different assumptions; direct comparisons between the results (especially quantitative thermodynamic values) across the different studies cannot be made.
2.3 Atomistic Simulation Characterization

2.3.1 Traditional Methods

The characterization techniques discussed in this section are described as "traditional methods" because they are well established and used extensively by researchers in the atomistic simulation community. A unique advantage of atomistic simulations is that they provide the coordinates and velocities (if performing dynamics) of every atom within the simulation. This allows researchers to study material phenomena by explicitly visualizing atomic structure and atomic velocities through time as well as enabling computation of spatially dependent properties.

For example, using the coordinates of the atoms, researchers can compute the radial distribution function (RDF) [45] to describe atom density as a function of separation distance from a reference atom. The radial distribution function is calculated through a binning procedure that counts the number of atoms that surround the reference atom within a thin shell region as shown in Figure 2.2. Binned amounts are computed for each atom and are normalized through spatial averaging to provide a unique description of the local atomic geometry up to the maximum separation distance examined.

Figure 2.2: Schematic of radial distribution function calculation [9].
To elucidate other material phenomenon, researchers will often employ various color-coding techniques to provide more insight into their visualizations. As a simple example, researchers may color-code atoms within a simulation by their per-atom potential energy to locate defect regions within the simulations where atoms possess elevated energy. Other per-atom quantities, such as centrosymmetry [46] and the common neighbor-analysis parameter [47], have been developed to provide further insights into material phenomena through similar color-coding visualization techniques.

Centrosymmetry is a per-atom quantity developed by Kelchner et al. [46] to uncover defect structures associated with dislocation nucleation in face-centered cubic Au during nanoindention. This technique takes advantage of a special centrosymmetric property defined by the bonding within face-centered cubic (FCC) and body centered cubic (BCC) materials. In centrosymmetric materials, the lattice is constructed from atoms with bond pairs that are equal and opposite in direction. The centrosymmetry parameter defined by Kelchner et al. measures the distortion of these bonds through,

\[ P = \sum_{i=1,6} \left| \mathbf{R}_i + \mathbf{R}_{i+6} \right|, \]  

(2.28)

where \( \mathbf{R}_i \) and \( \mathbf{R}_{i+6} \) are the vectors describing the bond pairs. Under homogenous elastic deformation, bond pairs in centrosymmetric materials within the bulk will remain equal and opposite in direction, thus showing no distortion in the centrosymmetry parameter \( (P = 0) \). However, atoms located at interfaces and near defects as well as bulk atoms experiencing non-homogenous elastic deformation will have positive centrosymmetry values \( (P > 0) \). A benefit of the centrosymmetry parameter is its ease of implementation and concept. However, the disadvantage of this method is that it is only useful and valid for materials with centrosymmetric
structures, which alumina is not as shown in Figure 1.1.

A more general per-atom characterization technique is the common neighbor-analysis (CNA) parameter, which identifies disorder found in arbitrary local environments. The common neighbor-analysis parameter was defined by Tsuzuki et al. [47] through,

$$Q = \frac{1}{n_i} \sum_{j=1}^{n_i} \left( \sum_{k=1}^{n_{ij}} R_{jk} + R_{jk} \right)^2,$$

where $n_i$ is the number of nearest neighbors of atom $i$, $n_{ij}$ is the number of common nearest neighbors between atom $i$ and atom $j$, and $R$ are vectors describing the bonds between the atoms identified through the subscript. The common neighbor-analysis parameter identifies deformation in many crystal systems and can distinguish between different types of defects such as stacking fault, dislocations, interface, etc. by relating these features to the common FCC, BCC, hexagonal closed packed (HCP), and “other” crystal systems.

![Graph](image)

**Figure 2.3:** Computed radial distribution functions (RDF) for select alumina phases at 0 K do not clearly identify the different crystal structures.

The traditional characterization methods discussed here have had great success in describing material structure and phenomena in metallic materials; however, they fall short to uniquely characterize the complex structures and phenomena found within the alumina material system.
due to the asymmetric placement of Al ions. The radial distribution functions computed for 0 K energy minimized structures using the ReaxFF potential [42], shown in Figure 2.3, show no clear distinction between the metastable phases. The complex, non-symmetric structures of the metastable phases have similar atomic densities within the 10 Å maximum cut-off distance explored; however, the specific placement of the aluminum interstitials within each phase is not clearly apparent.

Computed centrosymmetry and common neighbor analysis parameters are also unable to uniquely identify the various alumina phases. Figure 2.4 shows the unit cell for each alumina phase studied in this research colored by the centrosymmetry parameter. In general, the $\alpha$-$\text{Al}_2\text{O}_3$ and $\gamma$-$\text{Al}_2\text{O}_3$ phases are more centrosymmetric as seen by the lower magnitude of the centrosymmetry parameter for each atom; however, there is no clear differentiation among the phases. The common neighbor analysis conducted on the alumina unit cells did not identify any specific crystal system within the various phases; therefore, all atoms were declared as “other” using this method.
2.3.2 Kinematic vs. Dynamic Virtual Diffraction Methods

Because the traditional characterization methods used in atomistic simulations are unable to uniquely identify and differentiate the alumina phases, more advanced characterization methods based upon diffraction techniques are explored for his work. Diffraction is a common experimental method used to study the atomic scale structure of materials by examining the interaction between a beam of electromagnetic radiation (e.g. x-rays and electrons) and the periodic pattern of atoms within a material sample. Specifically, experimental diffraction patterns have been used to successfully identify and differentiate alumina phases [48–52]. Several previous virtual diffraction methods have been developed to analyze atomistic simulations; however, these prior methods have limited functionality and require a priori knowledge of the crystal system. Therefore, an advanced virtual diffraction technique is developed and implemented for this work that overcome these prior limitations, as will be discussed in Chapters 4-6.

Virtual diffraction methods utilize equations from diffraction theory in order to produce equivalent diffraction patterns in an effort to bridge atomistic simulations and experimental studies. More information on diffraction theory can be compiled from books by Warren [53] and Guinier [54]. Briefly, there are two classes of diffraction theory, (1) kinematic and (2) dynamic, that were developed starting in the early 1900s by W. H. Bragg, W. L. Bragg, Laue, Ewald, and Darwin soon after the discovery of coherent x-ray diffraction scattering in metals [55]. Kinematic diffraction theory models the interference patterns of diffracted radiation created by scattering off individual atoms. Similarly, dynamical diffraction theory models interference patterns; however, more rigorously incorporates the effects of reflected radiation. Figure 2.5 shows a schematic of diffracted versus reflected radiation related to the two classes of theory.
When studying large or perfect crystals, the dynamical theory is a more appropriate model for diffraction because the contributions from reflections become much more important. However, the computational cost to evaluate diffraction intensities via the dynamic diffraction theory is much larger as compared to kinematic diffraction theory due to the considerations of reflected radiation. When studying small volumes or imperfect crystal structures, such in the case for MD and MS simulations, reflected radiation becomes less important and a good approximation for diffraction is made by using kinematic diffraction theory. The virtual diffraction technique developed for this work utilizes equations based on kinematic diffraction theory. Details regarding its implantation are contained in Chapters 4-6 alongside various case studies to validate the method and showcase its versatility. Virtual diffraction methods are used to study alumina in Chapters 7 and 8 which focus on bulk and surface simulations as well as interface structures, respectively.
References


Chapter 3: The Effect of Synthetic Driving Force on the Atomic Mechanisms Associated with Grain Boundary Motion Below the Interface Roughening Temperature

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\textsuperscript{b}Computational Materials Science and Data Analysis Department, Sandia National Laboratories, Albuquerque, NM 87185

Abstract

The mechanisms associated with grain boundary motion induced by synthetic, crystal-orientation-dependent driving forces are investigated for a large-angle [001] Ni symmetric tilt grain boundary. The application of non-physical forces by this method brings legitimate concern that it could lead to non-physical results. This concern is especially relevant below the interface roughening transition temperature where there is a substantial drop in grain boundary mobility and large driving force dependence. Using slip-vector analysis and examining continuum metrics for microrotation and strain, this work shows that the application of synthetic-driving forces does not alter the fundamental mechanisms leading to grain boundary motion. Results in this work are compared directly to shear driven simulations which reveal that the path and deformation history of grain boundary motion is unbiased by the non-physical nature of the driving force acting on the boundary. Nudged elastic band calculations show that the transition path for grain boundary motion is independent of the driving force magnitude and the energy barriers for motion are not appreciably altered by the application of the synthetic driving force.
3.1 Introduction

A fundamental understanding of grain boundary motion and mobility is necessary to advance mesoscale predictive models of grain growth in polycrystalline materials [1]. Grain boundary motion has been simulated in recent years using synthetic driving force molecular dynamics simulations, first introduced by Janssens et al. in 2006 [2]. In these simulations [2], flat grain boundaries constructed from bicrystal models are driven by a crystal-orientation-dependent driving force superimposed on an embedded atom method (EAM) potential. This driving force stems from a continuous potential energy term that is added to all atoms which varies in magnitude based on the relative orientation of neighboring atoms as compared to a reference grain. Using this method, extra potential energy is added to the atoms of the reference grain resulting in a synthetic driving force that biases the motion of the grain boundary towards the higher energy grain in order to reduce the total system potential energy. The synthetic driving force stimulates motion of flat grain boundaries within the timescale of molecular dynamics simulations and allows any grain boundary misorientation to be studied independently. This facilitates high-throughput studies of grain boundary motion as a function of grain boundary crystallography and temperature such as the study by Olmsted, et al. [3].

Prior studies have applied the synthetic driving force method to a variety of grain boundaries spanning a wide misorientation range, but researchers have limited their analysis to the collective motion of the atoms at the grain boundary [2–7]. Here we include studies [4–6] which utilized the adapted interface-random-walk method introduced by Deng and Schuh in 2011 [4] as a hybrid method that combines synthetic driving force and statistical analysis of boundary fluctuations in order to analyze slow moving grain boundaries. By focusing the analysis on the collective motion of the atoms at the grain boundary, prior researchers [2–7] were able to
compute grain boundary velocity and mobility as well as compare their results to experimental values [3,5] and conventional molecular dynamics simulations with physically based driving forces such as shear and strain [6,7].

Studies of grain boundary motion have also reported an interface roughening transition which significantly impacts the motion of grain boundaries. Interfacial roughening is a thermodynamic phase transition where the spatial variation of the position of the interfacial plane is finite below the transition temperature and diverges with the interfacial area above the transition temperature. The transition temperature reflects the energy cost of a local fluctuation of the interfacial position normal to the interface. It has been shown that there is an abrupt increase in grain boundary mobility as the temperature is increased above the transition temperature for that interface [7]. Simulations that applied synthetic driving force methods revealed large driving force dependence for the smooth grain boundary configurations below the roughening transition temperature [3,7]. Since the synthetic driving force method applies non-physical forces, there is a legitimate concern that this approach might lead to non-physical results, especially below the roughening transition temperature.

While prior work [7] shows that the computed mobility agrees with physical driving force simulations, studies have not yet analyzed the effect of synthetic driving forces on the atomic mechanisms associated with the motion of smooth interfaces, nor has any study examined the effect of synthetic driving forces on the transition energy barriers for grain boundary motion. The purpose of the synthetic driving force method is to bias the system such that the grain boundary will move from its current position towards a region with artificially added energy without purposely raising the basins on the potential energy surface in an attempt to lower the transition energy barriers, as is done in the hyperdynamics method [8]. In this work, we
investigate the effect of synthetic driving forces on the atomic mechanisms associated with low-temperature grain boundary motion and compare these mechanisms to those observed in simulations driven by shear. In addition, we study the effect of applied synthetic driving forces on the potential energy surfaces revealing how transition energy barriers are affected by the synthetic driving force method.

3.2 Methods

The $\Sigma 37$ (570) [001] symmetric tilt grain boundary (STGB) modeled in this study was constructed originally by Olmsted et al. as part of their large survey of grain boundary energy and mobility [3,9]. The interface was chosen as a representative large-angle STGB constructed from sets of C type kite structural units [10] that zigzag at the interface. The initial studies by Olmsted et al. on this boundary provided evidence of grain boundary roughening at elevated temperatures and demonstrated motion-coupled shear when applying synthetic driving forces. Motion-coupled shear is the phenomenon whereby atoms in the neighboring grains collectively move parallel to the grain boundary plane in response to the synthetic driving force acting normal to the interface [11]. This phenomena is analogous to shear-coupled motion recently described in detail by Cahn et al. [12]. Boundaries experiencing motion-coupled shear are likely candidates for shear driven grain boundary motion, and would allow for direct mechanistic comparison.

The simulations conducted throughout this study use the LAMMPS molecular dynamics simulator [13] with the Foiles and Hoyt EAM potential for Ni [14]. To determine the roughening transition temperature, mobility calculations for the $\Sigma 37$ (570) [001] STGB are performed at 300-1200 K using synthetic driving forces of 0.001-0.025 eV/atom. In this study, we are able to extend into lower temperature and driving force regimes than originally studied by
Olmsted et al. [3] by increasing the effective statistical sampling size in order to reduce error in the computed mobility. Increased statistical sampling is achieved by using 25 independent simulations and offsetting initial conditions in a similar manner as described by Cheng and Schuh [4]. For each temperature and driving force, grain boundary displacement data are collected every 1 ps using an 2 fs offset. The combined 12,500 displacement data points collected are fit to a Gaussian distribution to determine the mean grain boundary displacement for each time interval. Using enhanced effective sampling, the minimum 10 Å grain boundary displacement criteria used by Olmsted et al. is relaxed in this study such that all simulations with computed grain boundary mobility greater than 1 GPa\(^{-1}\) m/s are analyzed.

The effects of the synthetic driving forces below the roughening transition temperature are explored through direct comparisons with shear driven grain boundary motion simulations. Simple shear simulations are constructed at 300 and 400 K such that the normal grain boundary velocities closely match those of the simulations using synthetic driving forces of 0.025 and 0.005 eV/atom. The shear simulations are designed similar to those by Tucker et al. [15], where grain boundary motion is induced by moving rigid edges on both sides of the bicrystal model at a constant (opposing) velocity parallel to the grain boundary plane. To avoid shock wave generation, a linearly ramped velocity profile is imposed on the atoms positioned between the rigid edges. Low-temperature simulations driven by shear and the synthetic driving force method are analyzed and compared using slip-vector analysis and continuum metrics of microrotation and strain as described by Zimmerman et al. [16] and Tucker et al. [17] respectively.

The energy barriers and transition states associated with the motion of the smooth \(\Sigma 37\) (570) \([001]\) Ni STGB are investigated using the nudged elastic band (NEB) method extracting data
from climbing images [18]. NEB calculations determine the 0 K minimum energy configurations along the path from an initial to a final state. In this study, a series of six NEB calculations are performed applying synthetic driving forces in the range of 0.000 - 0.025 eV/atom. It is important to note that the NEB calculations with no applied driving force are comparable to conditions observed for shear driven simulations; thus, providing a baseline for comparing the synthetically driven methods.

3.3 Results and Discussion

Figure 3.1 displays an Arrhenius plot of log(mobility) versus inverse temperature for the $\Sigma 37 \{570\} [001]$ STGB revealing an interface roughening transition between 400 and 500 K. From 500-1200 K, synthetic driving forces have minimal effect on the grain boundary mobility and log(mobility) remains nearly constant between 5.5-6.9 log(GPa$^{-1}$ m/s). However, from 300 to 400 K the mobility decreases significantly with application of lower driving forces, which is indicative of a transition into a smooth grain boundary configuration. Mobility data for the 300 K simulations driven by 0.0010 and 0.0025 eV/atom are not included as they fall below the minimum 1 GPa$^{-1}$ m/s criteria used in this study.

Figure 3.1: Arrhenius plot of log(mobility) versus inverse temperature for a $\Sigma 37 \{570\} [001]$ STGB computed with various driving forces. Grain boundary roughening occurs between 400 and 500 K as identified by a substantial drop in driving force dependence.
Shear-driven simulations are performed below the interface roughening transition temperature to directly compare the collective and individual atomic mechanisms associated with grain boundary motion. Table 3.1 shows the shear strain rates used to match the normal grain boundary velocity obtained using 0.005 and 0.025 eV/atom synthetic driving forces at 300 and 400 K. The effect of the different driving force types on the collective motion of the atoms near the grain boundary are presented in Figure 3.2, which displays the displacement of the grain boundaries normal ($x^*$) and parallel ($z^*$) to interface plane in dimensionless parameters. Here, the characteristic length is defined as 10 Å and the characteristic time, $\tau_{10}$, is equal to the time required for the grain boundary to move 10 Å in the direction normal to the grain boundary. Characteristic times used to generate the plots in Figure 3.2 are reported in Table 3.2.

Table 3.1: Shear strain rates used to match normal grain boundary velocity obtained through synthetic driving force molecular dynamics simulations.

<table>
<thead>
<tr>
<th>Driving Force</th>
<th>300 K (s$^{-1}$)</th>
<th>400 K (s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.005 eV/atom</td>
<td>1.107 $10^7$</td>
<td>2.527 $10^8$</td>
</tr>
<tr>
<td>0.025 eV/atom</td>
<td>3.159 $10^9$</td>
<td>3.790 $10^9$</td>
</tr>
</tbody>
</table>

Table 3.2: Characteristic time, $\tau_{10}$, required for the grain boundary to move 10 Å in the direction normal to the interface.

<table>
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<tr>
<th>Driving Force</th>
<th>300 K (ps)</th>
<th>400 K (ps)</th>
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</thead>
<tbody>
<tr>
<td>0.005 eV/atom</td>
<td>$1455^a$, $1685^b$</td>
<td>$100^a$, $98^b$</td>
</tr>
<tr>
<td>0.025 eV/atom</td>
<td>$10.2^a$, $10.8^b$</td>
<td>$8.5^a$, $8.7^b$</td>
</tr>
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</table>

$^a$ Synthetic driving force simulation
$^b$ Shear simulation

Simulations driven by 0.005 eV/atom and the corresponding shear strain rates are dominated by stick-slip behavior, which provides sufficient time for the shear-coupled motion and motion couple-shear to traverse through the bicrystal models. This stick-slip motion shows some
discontinuous jumps in Figure 3.2, but the normal and shear velocity remain relatively constant throughout the simulation. Differently, the shear and normal motion of the 0.025 eV/atom driven and corresponding shear simulations experience lag. Shear motion is delayed when applying 0.025 eV/atom synthetic diving forces because the driving force initiates motion at the grain boundary and requires time to propagate away from the boundary. Eventually, the shear motion in the 0.025 eV/atom driven simulations traverses through the simulation and recovers a similar velocity as the shear models. Likewise, motion normal to the grain boundary for shear simulations is delayed because the sheared regions are located a minimum of 75 Å away from the grain boundary at the edges of the bicrystal models. Thus, it takes time for the shear simulations to move the grain boundary, as compared to the synthetically driven simulations that act directly at the boundary and almost immediately initiate grain boundary motion.

The atomic mechanisms associated with grain boundary motion below the interface
roughening transition temperature are analyzed using slip-vector analysis and continuum metrics of microrotation and strain as described by Zimmerman et al. [16] and Tucker et al. [17].

Application of these metrics shows that the path and deformation history created by the $\Sigma 37$ (570) [001] STGB motion is independent of the type of driving force and its magnitude. Example results for the 300 K simulation driven by 0.005 eV/atom are presented in Figure 3.3, but can be generalized for the eight cases studied. Slip vector analysis shown in Figure 3.3(a) reveals bands of inhomogeneous deformation within the region formerly associated with the structural units. The orientation of these slip bands clearly identifies the sheared path of the structural units; however, it does not clearly elucidate the mechanisms associated with grain

Figure 3.3: Metrics analyzing (a) slip-vector magnitude, (b) microrotation, (c) $E_{11}$ strain normal to the grain boundary plane, and (d) $E_{33}$ strain parallel to the grain boundary plane for the 300 K simulations induced by a 0.005 eV/atom synthetic driving force. The results are independent of the driving force type and magnitude and can be generalized for the eight, low-temperature boundary conditions studied
boundary motion.

Analysis of the microrotation results, shown in Figure 3.3(b), indicates that the motion of the $\Sigma 37 \ (570) \ [001]$ STGB is caused by a rotation of the atoms adjacent to the structural units. To accommodate the rotation, the region formerly associated with the structural units is strained both normal and parallel to the grain boundary plane. Figure 3.3(c) shows an atomistic estimation of the strain normal to grain boundary plane, which strongly affects the atoms along the deformation path as well as the current structural units within the grain boundary. At the start of the deformation path, atomic rotations also cause atoms associated with the regions containing the initial structural units to be compressed normal to the grain boundary plane. An estimation of strain parallel to the grain boundary is displayed in Figure 3.3(d) and is less severe compared to strain normal to the grain boundary plane. This lower relative impact is further evidence pointing to a rotational mechanism causing motion parallel to the grain boundary plane.

Nudged elastic band results are reported in Figure 3.4. In each NEB calculation, four energy barriers are computed from the saddle-energy associated with the individual reconstruction of the four C structural units from their ideal kite-shaped to a heart-shaped transition state, as shown in Figure 3.4(a). Analysis of the atom motion during NEB shows that the transition path and the structure of the transition phases are independent of the added synthetic energy in the range studied. The energy barriers associated with the reconstruction of the four C structural units are computed by first subtracting the extra energy associated with the synthetic driving force, then comparing the potential energy of the transition states to that of the initial configuration. These adjustments to the potential energy surface for simulations driven by synthetic driving forces produce a similar energy landscape that closely matches that of the sheared simulation with no added synthetic driving force, as shown in Figure 3.4(b). The transition energy barriers
associated with grain boundary motion are not significantly altered by the addition of a synthetic energy, as shown in Figure 3.4(c). The computed energy barriers range from 0.53-0.69 eV for each of the transition states with application of 0.000-0.0250 eV/atom synthetic driving force.

Figure 3.4: Nudged elastic band results showing (a) the four transition states to move the grain boundary, (b) NEB results for the 0.025 eV/atom driven simulation adjusted for the added synthetic energy compared to results from sheared driven simulations, and (c) the energy barriers relative to the energy of the initial grain boundary structure without the synthetic driving force energy.

3.4 Conclusion

In summary, this work identifies the grain boundary roughening temperature for the $\sum 37 (570) [001]$ Ni STGB and studies the mechanisms associated with motion in the low-temperature, smooth interface regime. At temperatures below the interface roughening transition, synthetic driving forces greatly affect the grain boundary mobility; however, the atomic mechanisms for grain boundary motion are not altered as shown through slip-vector
analysis and by examining continuum metrics of microrotation and strain. Additionally, comparison of these metrics with shear simulations shows that the synthetic driving force method induces the same mechanisms for motion as the physically based approach. NEB calculations confirm that synthetic driving forces do not alter the transition path for grain boundary motion and do not substantially change the transition energy barriers. The observation of consistent energy barriers, transition paths, and atomic mechanisms associated with grain boundary motion indicates the use of synthetic driving force molecular dynamics method does produce physically significant results.

**Acknowledgements**

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References


I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication. This work was done in collaboration with Dr. Stephen Foiles at Sandia National Laboratories, Albuquerque, NM.

Sincerely,

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Appendix 3.2  
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Chapter 4: Virtual Diffraction Analysis of Ni [010] Symmetric Tilt Grain Boundaries

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Abstract

Electron and x-ray diffraction are well-established experimental methods used to explore the atomic scale structure of materials. In this work, a computational method is implemented to produce virtual electron and x-ray diffraction patterns directly from atomistic simulations without \textit{a priori} knowledge of the unit cell. This method is applied to study the structure of [010] symmetric tilt low-angle and large-angle grain boundaries in Ni. Virtual electron diffraction patterns and x-ray diffraction 2θ line profiles show that this method can distinguish between low-angle grain boundaries with different misorientations and between low-angle boundaries with the same misorientation but different dislocation configurations. For large-angle Σ5 (210), Σ29 (520), and Σ5 (310) coincident site lattice [010] symmetric tilt grain boundaries, virtual diffraction methods can identify the misorientation of the grain boundary and show subtle differences between grain boundaries in the x-ray 2θ line profiles. A thorough analysis of the effects of simulation size on the relrod structure in the electron diffraction patterns is presented.
4.1 **Introduction**

Grain boundaries are planar defects accommodating the misorientation and elastic incompatibilities between two lattice regions of uniform crystallographic orientation [1]. Quantification and comprehension of the atomic scale structures within grain boundaries, such as full and partial dislocations, disclinations, etc., have greatly benefited from the use and interpretation of diffraction measures and simulations. Experimental work by Sass and coworkers [2–5] on grain boundaries used selected area diffraction (SAD) patterns from electron and x-ray sources to confirm that the atomic structure near grain boundaries is periodic. X-ray and electron diffraction patterns of low-angle twist grain boundaries in gold bicrystals showed weak subsidiary reflections caused by the regularly spaced screw dislocation grid within the grain boundary unit cell [5]. In studies of large-angle twist grain boundaries with low $\Sigma$ coincident site lattice (CSL) [6] designations, Tan et al. [7] observed subsidiary reflections and indexed these reflections in terms of the CSL unit cell. For [001] symmetric tilt boundaries, Guan and Sass [8] found extra reflections in the vicinity of the (200) relrod tails caused by the spacing of the edge dislocation array. Careful characterization of the extra reflections allowed researchers to identify the structure and misorientation of different grain boundaries [7–9].

To supplement these experimental studies, virtual diffraction models based on kinematic diffraction theory were developed to connect atomistic simulations of large-angle symmetric twist grain boundaries to the experimental SAD patterns [10–12]. In these models, the structure factor equation was computed over a region of reciprocal space constructed from a grain boundary unit cell known *a priori*. Work by Bristowe and Sass [10] showed that small displacements of the atoms within the unit cell can create identifiable changes to the diffraction pattern allowing researchers [11–14] to use the virtual SAD patterns to accurately predict grain
boundary structure validated by experimental comparisons. In these early virtual diffraction models [10–12], computational limitations restricted the resolution of the reciprocal space explored and the number of atoms included within each simulation. *A priori* knowledge of the symmetric twist grain boundary unit cell facilitated a significant reduction in computational demand during structural optimization by reducing the degrees of freedom through symmetry [11]. As computational capabilities increased, higher resolution scans of reciprocal space were performed by Bristowe and Balluffi [12] revealing more subtle diffraction reflections stemming from secondary relaxations that were more difficult to match with experimental SAD patterns.

To reconcile the effects of secondary relaxations, studies using larger simulations were performed on symmetric twist grain boundaries [13–15] examining the effects of multiple grain boundary unit cells within the simulation.

In addition to the SAD studies of grain boundary structure, x-ray diffraction line profile analysis has been used to study grain boundary structure in bicrystal and nanocrystalline samples. Several experimental x-ray diffraction studies have attempted to isolate the effects of grain boundaries on peak broadening [16–18]. Work by Eastman et al. [16,17] on nanocrystalline nickel found evidence of a reduction in line profile broadening due to relaxations in the grain boundary structure. Additionally, Gaudig et al. [18] showed that modulations in the lattice spacing near the grain boundary structure created subsidiary peaks primarily along the direction normal to the grain boundary plane.

Several researchers have applied virtual diffraction models based on the Debye scattering equation to study 2θ x-ray diffraction line profiles from atomistic models of nanocrystalline materials [19–22]. Derivation of the Debye scattering equation involves a spherical integration of the structure factor equation, which allows the diffraction intensity at a chosen scattering
angle to be related solely to the interatomic separation between pairs of atoms [23]. These researchers [19–22] verified the fidelity of the virtual diffraction line profiles using experimental comparisons. However, the complex structure of nanocrystalline materials and the assumptions inherent to the integration used to derive the Debye equation made isolating the influence of specific atomic scale defects on peak broadening difficult. Specifically, studies [19–21] have disputed the relative influence on peak broadening of the local displacement fields near grain boundaries versus the longer-range displacement fields associated with the nanocrystalline microstructure. Work by Stukowski et al. [19] found that the underlying cause for microstrain line broadening stems from longer-range coordinated displacements between grains. Whereas Alexsandrov et al. [20] and Derlet et al. [21] conclude that x-ray diffraction line profile broadening stems from the disorder around dislocations within the grain boundaries.

Together, experimental and virtual diffraction techniques have revealed important information regarding the atomic structure of grain boundaries and have enriched atomistic simulations by allowing a direct link between atomistic simulations and experimental results. This paper advances virtual diffraction methods by presenting an algorithm for performing virtual diffraction during an atomistic simulation via explicit evaluation of the structure factor equation without a priori knowledge of the grain boundary unit cell. The algorithm is sufficiently generic to be applicable to both electron and x-ray diffraction conditions and is integrated into the LAMMPS molecular dynamics simulator [24]. Within LAMMPS, the algorithm is capable of performing diffraction analyses either statically (after energy minimization) or dynamically during a molecular dynamics simulation to produce time averaged diffraction patterns at finite temperature. A visualization procedure is developed to create SAD patterns and 2θ x-ray diffraction line profiles from the intensities computed using the atomistic
simulation data. In this work, this algorithm is applied to study the structure of [010] symmetric tilt low-angle and large-angle CSL grain boundaries in face-centered cubic (FCC) Ni, differentiating from the previous literature on virtual diffraction which has primarily focused on the analysis of twist grain boundaries. Analogous to previous results in the literature, the virtual diffraction method is capable of extracting critical details related to the misorientation and structure of low-angle and large-angle CSL symmetric tilt grain boundaries.

4.2 Methodology

4.2.1 Virtual Diffraction Algorithm and Visualization

The diffraction algorithm implemented into LAMMPS generates a three-dimensional mesh of points filling a volume of reciprocal space constructed from the entire domain of the atomistic simulation cell. The mesh of reciprocal lattice points is built on a rectilinear grid with spacing \( c_n |A_n|^{-1} \) along each reciprocal lattice axis \( \mathbf{B}_n \). Each reciprocal lattice axis \( \mathbf{B}_n \) is determined from the associated vector \( \mathbf{A}_n \) corresponding to the \( n = 1, 2, \) or 3 edge of the simulation cell. The parameters \( c_n \) control the spacing of the reciprocal lattice points, as shown schematically in Figure 4.1, and determine the resolution within the reciprocal space volume explored. Specific values of \( c_n \) used in this work are provided in Section 4.2.2.

During electron or x-ray diffraction, each reciprocal lattice point is associated with a reciprocal lattice vector \( \mathbf{K} \) describing the deviation between the diffracted and incident wave vectors \( \mathbf{k}_d \) and \( \mathbf{k}_i \) [25],

\[
\mathbf{K} = \mathbf{k}_d - \mathbf{k}_i = \xi \mathbf{B}_1 + \eta \mathbf{B}_2 + \zeta \mathbf{B}_3 ,
\]

where \( \xi, \eta, \) and \( \zeta \) can be any real number. Assuming monochromatic incident radiation of wavelength \( \lambda \), the angle of diffraction \( \theta \) is computed utilizing the geometric relationship
between \( \mathbf{k}_i \), \( \mathbf{k}_b \), \( \mathbf{K} \), and \( \lambda \) through Bragg’s law [25],

\[
\frac{\sin(\theta)}{\lambda} = \frac{|\mathbf{K}|}{2} \quad . \tag{4.2}
\]

At certain reciprocal lattice points throughout the mesh, the Bragg condition is satisfied

\( (\mathbf{K} = \mathbf{K}_B) \) and the magnitude of the reciprocal lattice vectors is related to interplanar distances \( d_{hkl} \) of the associated \( \{hkl\} \) through [26],

\[
\frac{1}{d_{hkl}} = |\mathbf{K}_B| \quad . \tag{4.3}
\]

At these points, the atomic structure allows constructive interference of the radiation producing strong diffraction intensity.

![Diagram of the reciprocal space mesh illustrating the rectilinear spacing and criteria set on |K| to increase the computational efficiency of the algorithm by limiting the number or reciprocal lattice points explored.](image)

The diffraction intensity at each reciprocal lattice point is calculated differently for electron and x-ray radiation; however, both require evaluation of the structure factor \( F(\mathbf{K}) \). For each
type of radiation, the structure factor \( F(K) \) is computed utilizing the atomic positions \( r_j \) via [23],

\[
F(K) = \sum_{j=1}^{N} f_j \exp\left(2\pi i K \cdot r_j\right).
\]  (4.4)

Here, \( f_j \) are the atomic scattering factors which account for the reduction in diffracted intensity from an individual atom due to Compton scattering and vary by atom type, angle of diffraction \( \theta \), and type of radiation [23].

At each diffraction angle, the atomic scattering factors \( f_j \) are computed using analytical approximations parameterized for the specific atom type. For electron diffraction, the analytical approximation of the atomic scattering factor is the summation of five Gaussian functions of the form [27],

\[
f_j\left(\frac{\sin \theta}{\lambda}\right) = \sum_{i}^{5} a_i \exp\left(-b_i \frac{\sin^2 \theta}{\lambda^2}\right),
\]  (4.5)

which have been parameterized for the majority of neutral elements by Peng et al. [28]. For x-ray diffraction, the analytical approximation of the atomic scattering factor is found from a summation of four Gaussian functions plus a constant of the form [29],

\[
f_j\left(\frac{\sin \theta}{\lambda}\right) = \sum_{i}^{4} a_i \exp\left(-b_i \frac{\sin^2 \theta}{\lambda^2}\right) + c,
\]  (4.6)

and has been parameterized by Fox et al. for most atom types [30].

For electron diffraction, the diffraction intensity \( I_e(K) \) at each reciprocal lattice point is computed from the product of the structure factor and its complex conjugate \( F^*(K) \) normalized by the number of atoms in the simulation \( N \) via [23],

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To compute x-ray diffraction intensities \( I_x(K) \), an additional Lorentz-polarization factor \( L_p(\theta) \) is applied to account for the relative distribution of the reciprocal lattice points and the change in scatter intensity when using non-polarized incident radiation. The Lorentz-polarization factor is computed via (4.7),

\[
L_p(\theta) = \frac{1 + \cos^2(2\theta)}{\cos(\theta)\sin^2(\theta)},
\]

and thus the diffraction intensity from x-rays is computed through (4.9),

\[
I_x(K) = L_p(\theta) \frac{F(K)F^*(K)}{N}.
\]

In general, virtual selected area electron diffraction (SAED) patterns are created by examining the region in reciprocal space intersecting the Ewald sphere of radius \( \lambda^{-1} \). For a particular zone axis, the Ewald sphere is centered at the tail of the associated incident wave vector and intersects the origin of reciprocal space. To construct the electron diffraction pattern, a thin hemispherical slice of the reciprocal lattice mesh lying near the surface of the Ewald sphere is isolated and viewed parallel to the zone axis. The thickness of this slice is dependent on the resolution of the reciprocal space mesh and is chosen such that between one and five reciprocal lattice points are contained within the slice parallel to the zone axis at each point along the curvature of the Ewald sphere. To enhance features within the diffraction pattern, low intensity reciprocal lattice points are removed and the remaining points are coloured by intensity on a \( \log_{10} \) scale. X-ray diffraction line profiles are created by virtually rotating the Ewald sphere around the origin of reciprocal space to all possible orientations, mimicking powder diffraction conditions [31]. By making all diffraction orientations equally probable, all reciprocal lattice
points will intersect the surface of the Ewald sphere. Line profiles simulating x-ray powder
diffraction are constructed by collecting all reciprocal lattice points into bins corresponding to
their scattering angle $2\theta$ calculated using Eq. (4.2) and summing the intensity data. The
scattering angle bin size is optimized through trials to reduce the noise within the line profile
while maximizing peak features. Specific discussion of the parameters used to construct SAED
and $2\theta$ x-ray diffraction line profiles on simulated low-angle and large-angle CSL [010]
symmetric tilt grain boundaries is provided in Section 4.2.2.

**4.2.2 Application to Grain Boundaries**

The virtual diffraction algorithm is applied to six FCC Ni [010] symmetric tilt grain
boundaries (STGBs) modeled with the Foiles-Hoyt Ni embedded-atom method (EAM) potential
[32]. The grain boundaries are constructed utilizing the methods described by Spearot et al. [33]
with periodic boundary conditions applied in all directions forming bicrystal models with
specific misorientations. For each bicrystal, minimum energy grain boundary structures are
attained at 0 K by aligning the opposing lattice regions with the intended tilt and removing atoms
that are inside specified cutoff distances. This procedure effectively samples multiple starting
configurations with different relative displacements. Each grain boundary structure is optimized
to best minimize the potential energy using a nonlinear conjugate gradient method implemented
in LAMMPS [24]. Electron and x-ray virtual diffraction data are computed using the atomic
positions in the minimum energy structures.

In this study, three low-angle (dislocation) [010] STGBs and three large-angle CSL [010]
STGBs with low $\Sigma$ values are studied. The low-angle bicrystal models consist of one 5.00° [010]
STGB and two 10.39° [010] STGBs created by specifying different initial conditions during
energy minimization. These boundaries are chosen for this work to validate the ability of the
virtual diffraction algorithm to differentiate between different low-angle tilt boundaries and to
compare the results to prior experimental work [8]. The large-angle CSL bicrystal models
studied in this investigation are the \( \Sigma 5 \) (210), \( \Sigma 29 \) (520), and \( \Sigma 5 \) (310) [010] STGBs. In the CSL
notation, the \( \Sigma \) values correspond to the inverse density of the coincidence lattice sites between
the misoriented lattice regions and the Miller indices indicate the grain boundary plane [6].
These boundaries are chosen for this work because they exhibit the \(|B.B|\), \(|BC.BC|\), and \(|C|\) grain
boundary structures consistent with the structural unit model [34]. In the context of the structural
unit model, the \( \Sigma 5 \) boundaries used in this study correspond to special grain boundaries at
equilibrium while the \( \Sigma 29 \) boundary is non-special. From the point of view of continuum
mechanics, the transition between B and C structural units can be seen as a perturbation in an
otherwise uniform array of structural units. Such perturbations can be represented via either
secondary grain boundary dislocations or disclination dipoles [35]. For reference, descriptions of
all boundaries used in this study are listed in Table 4.1.

| Tilt Angle | \( \Sigma \) | Boundary Plane | \( |A_1| \) (Å) | \( |A_2| \) (Å) | \( |A_3| \) (Å) | \( N \) (atoms) | GB Energy (J/m²) |
|------------|-------|----------------|----------------|----------------|----------------|----------------|----------------|
| **Low-Angle** |       |                |                |                |                |                |                |
| 5.00°      | 265   | 23 1 0         | 161.63         | 28.16          | 81.01          | 33,760         | 0.789          |
| 10.39°     | 61    | 11 1 0         | 233.23         | 28.16          | 38.89          | 23,392         | 1.050¹         |
|            |       |                | 231.87         | 28.17          | 38.87          | 23,232         | 1.138²         |
| **Large-Angle** | 36.87° | 5 2 1 0        | 237.07         | 28.14          | 31.48          | 19,200         | 1.285          |
|            | 43.60°| 29 5 2 0       | 226.18         | 28.16          | 37.89          | 22,080         | 1.219          |
|            | 53.13°| 5 3 1 0        | 244.59         | 28.15          | 33.38          | 21,024         | 1.344          |

¹ Full dislocation grain boundary
² Partial dislocation grain boundary

Simulated 200 kV electron radiation (\( \lambda = 0.0251 \) Å [27]) and Cu K\( \alpha \) x-rays (\( \lambda = 1.54178 \) Å
[23]) are used to create SAED patterns and x-ray line profiles. Table 4.2 lists the parameters
used to compute analytical approximations of the Ni electron and x-ray atomic scattering factors.
For the SAED patterns, a very fine resolution of the reciprocal space lattice, approximately \(5 \times 10^7\) reciprocal lattice points per \(\text{Å}^{-3}\), is achieved by using \(c_n = 0.10\). This resolution is necessary to eliminate the mesh sensitivity in the analysis of relrods and subsidiary peaks. To increase the computational efficiency, the volume of reciprocal space that is explored is decreased by setting criteria on \(|\mathbf{K}|\), as illustrated in Figure 4.1. In this study, the SAED patterns are focused on regions where \(h^2 + k^2 + l^2 < 20\) limiting the explored volume of reciprocal space such that \(0 < |\mathbf{K}| < 1.70\ \text{Å}^{-1}\). SAED patterns aligned on the [010] misorientation axis are constructed by selecting reciprocal lattice points intersecting a 0.01 \(\text{Å}^{-1}\) thick Ewald sphere slice centered at (0 39.8406 0) \(\text{Å}^{-1}\) in reciprocal space. Initial SAED patterns are constructed by removing reciprocal lattice points with intensities less than 0.5% of the maximum intensity \(I_e^{\text{max}}\) outside the transmitted beam area. For more detailed patterns, this threshold is decreased to 0.2% of \(I_e^{\text{max}}\).

Relrod profiles are created by isolating a cylindrical region of radius 0.004 \(\text{Å}^{-1}\) centered on a specific (002) reflection with the average intensity of the reciprocal lattice points binned based on their position along the l-axis in reciprocal space.

For x-ray diffraction line profiles, the resolution of reciprocal space is set using \(c_n = 0.25\) resulting in approximately \(2 \times 10^7\) reciprocal lattice points per \(\text{Å}^{-3}\). Knowing that the ideal 2\(\theta\) scattering angles associated with \{111\}, \{002\}, and \{220\} planes of FCC Ni are located at 44.59°, 51.96°, and 76.55° respectively (assuming an equilibrium lattice parameter of 3.52 Å [32]) the range of \(|\mathbf{K}|\) is restricted such that \(35° < 2\theta < 80°\) as computed via Eq. (4.2). For each boundary, x-ray diffraction line profiles are created using an optimal 2\(\theta\) bin size of 0.067° determined through several trials.
Table 4.2: Parameters used to compute analytical approximations of the Ni atomic scattering factors for electron and x-ray diffraction as calculated via Eq. (4.5) and Eq. (4.6) respectively with $\sin(\theta)/\lambda$ (Å⁻¹).

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<td>12.1763</td>
<td>2.3800</td>
<td>66.3421</td>
<td>1.0341</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Peng et al. [28]

$^b$ Fox et al. [30]
4.3 Results and Discussion

4.3.1 Low-Angle Symmetric Tilt Grain Boundaries

The virtual diffraction methods are used initially to examine the structures of low-angle (dislocation) [010] STGBs to validate the algorithm implementation and to make comparison with the available literature that has previously discussed diffraction from low-angle tilt grain boundaries [8]. Figure 4.2(a) - (c) show the energy-minimized structure of each low-angle [010] STGB coloured by the centro-symmetry parameter [36]. The predicted structure of the 5.0° [010] STGB from energy minimization calculations includes periodic sets of partial dislocations along the interface plane. Energy minimization calculations predict two different structures for the 10.39° [010] STGB. In Figure 4.2(b) each dislocation in the periodic array along the grain boundary has dissociated into partial dislocations resulting in a boundary with slightly higher energy than that shown in Figure 4.2(c) where the edge dislocation array along the grain boundary plane remains intact. In Figure 4.2(b) the partial dislocations are spaced 7.74 Å apart within each grouping and separated from the neighboring group of partial dislocations by 11.69 Å, making the primary spacing of the dislocation array 19.43 Å. Similarly, the spacing of the primary array of intact edge dislocations in Figure 4.2(c) is 19.45 Å.

Selected area electron diffraction patterns with zone axis aligned along the [010] misorientation axis for each low-angle STGB are shown in Figure 4.2(d) - (e). Each SAED pattern explores approximately six million reciprocal lattice points and is indexed denoting the plane and the lattice region responsible for the diffraction reflection. Squares are drawn around the {220} reflections as guides to identify the misorientation between the two lattice regions in each low-angle grain boundary model. Figure 4.3 shows magnified portions of the SAED pattern in the vicinity of the (002) reflections for the two 10.39° [010] STGBs. In this figure, the
Figure 4.2: Low-angle [010] STGB structures for (a) \{23 1 0\} 5.0°, (b) \{11 1 0\} 10.39° with partial dislocations, and (c) \{11 1 0\} 10.39° with full dislocations. SAED patterns aligned with the [010] misorientation axis are shown in (d) - (f) for each corresponding grain boundary. The squares are included as a guide to distinguish the orientations of bicrystal regions.
lower intensity threshold is reduced to 0.2% of $I_{\text{max}}$ to better capture the presence of relrods and subsidiary reflections due to the grain boundary. The SAED patterns in Figure 4.3 show relrods orientated parallel to the grain boundary plane (vertical direction) and oriented normal to the grain boundary (horizontal direction) as well as subsidiary reflections in the vicinity of the FCC reflections.

![SAED patterns](image)

**Figure 4.3:** Magnified view of SAED patterns near the (002) reflections for (a) 10.39° [010] STGB with partial dislocations and (b) 10.39° [010] STGB with full dislocations. In (a) and (b) the zone axis is aligned with the [010] misorientation axis while in the insets the zone axis is aligned with the [100] normal to the grain boundary.

The subsidiary reflections in Figure 4.3 are the result of the periodic lattice strain created by the dislocations within the grain boundary analogous to those studied in detail in previous experimental works [2-14]. Specifically, studies on low-angle [001] STGBs indicated that the spacing between the extra reflections and the tails of the (002) reflections corresponds to the spacing of the primary dislocation array [8]. Rotating the zone axis to match the experimental results of Guan and Sass [8], the insets in Figure 4.3 correspond to SAED patterns aligned normal to the grain boundary plane and more clearly show the spacing between the extra
reflections and (002) tails. In the Figure 4.3(a) inset (corresponding to the 10.39° [010] STGB with partial dislocations) four extra peaks are observed due to the grain boundaries. The spacing between the reflections and the center of the (002) tails labeled 1-4 are 0.058, 0.055, 0.045, and 0.040 Å⁻¹ (17.28, 18.08, 22.22, and 25.08 Å) respectively. In the Figure 4.3(b) inset (corresponding to the 10.39° [010] STGB with intact dislocations) two extra reflections are observed and the associated spacing labeled 5 and 6 are 0.059 and 0.044 Å⁻¹ (16.90 and 22.87 Å) to the center of the (002) tails. As suggested by the experimental results of Guan and Sass [8], the average of these distances between the extra peaks and the (002) tails for each simulation is a close match to the primary spacing of the dislocation (19.43 and 19.45 Å) in the simulations.

In bicrystal models, relrods are expected to appear in the direction normal to each low-angle grain boundary; in this direction, the distortion field decays as a function of distance away from the interface providing a finite size effect. However, Figure 4.2 and Figure 4.3 show that the strongest relrods are those parallel to the grain boundary. Recall that the atomistic simulation model used in this work includes periodic boundary conditions in all directions meaning that atoms on or near the model surface interact via the interatomic potential through the periodic boundary with atoms on the opposing side of the model. However, the algorithm implemented to compute diffraction intensity from the structure factor equation does not account for an infinite size simulation model in the same way and thus the oscillating distortion field due to the dislocation structure at the low-angle grain boundary is truncated in the intensity calculation.

To illustrate the role of length scale on the relrod structure, traces of the (002) relrod along the l-axis are constructed, shown in Figure 4.4, using different simulation model sizes of the 10.39° partial dislocation [010] STGB. Here, only |A₃| (parallel to the grain boundary plane and perpendicular to the dislocation cores) is increased from 38.87 Å in the original study (Figure 4.2
and Figure 4.3) to 77.74 Å and 116.61 Å. The data computed from the relrod profiles are fit to [23],

\[
I = |F|^2 \left( \frac{\sin^2 (\pi N_1 \mathbf{K} \cdot \mathbf{a}_1)}{\sin^2 (\pi \mathbf{K} \cdot \mathbf{a}_1)} \right) \left( \frac{\sin^2 (\pi N_2 \mathbf{K} \cdot \mathbf{a}_2)}{\sin^2 (\pi \mathbf{K} \cdot \mathbf{a}_2)} \right) \left( \frac{\sin^2 (\pi N_3 \mathbf{K} \cdot \mathbf{a}_3)}{\sin^2 (\pi \mathbf{K} \cdot \mathbf{a}_3)} \right),
\]

(4.10)

which describes the predicted diffraction intensity observed from a small crystal constructed from many unit cells. For a single crystal \( \mathbf{a}_1 \), \( \mathbf{a}_2 \), and \( \mathbf{a}_3 \) in Eq. (4.10) are the lattice vectors describing the unit cell; however, because the bicrystal models are constructed using two unit cell orientations the vectors \( \mathbf{a}_1 \), \( \mathbf{a}_2 \), and \( \mathbf{a}_3 \) represents an averaged periodic repeating unit along each simulation cell axis. In Eq. (4.10), the variables \( N_1 \), \( N_2 \), and \( N_3 \) are the number of units cells in each direction within the grain boundary model. An excellent fit is achieved between the

Figure 4.4: Intensity profiles tracing the (002) relrods indicated in Figure 4.3(a) for the 10.39° [010] STGB containing partial dislocations with increasing \( |A_3| \) as labeled. The 0.5% minimum intensity cutoff is plotted to show the threshold used in the SAED pattern shown in Figure 4.2.
computed relrod profile and the fundamental equation describing the shape of relrods during electron diffraction from finite volumes. From the fit, the magnitudes of $a_3$ for the three simulations are 3.53, 3.53, and 3.54 Å with $N_3$ equal to 11, 22, and 33 respectively. These fitted parameters correspond to $|A_3|$ of length 38.86, 77.70, and 116.66 Å, which closely match $|A_3|$ in the simulations studied. In Figure 4.4, the 0.5% of $I^\text{max}$ threshold is illustrated showing the subsidiary maxima not visible in Figure 4.2. As $|A_3|$ in the simulations increases, the visible relrods become more concentrated around the (002) reflections.

Virtual 2θ x-ray diffraction line profiles for each low-angle [010] ST GB are shown in Figure 4.5. To the authors' knowledge, no prior studies report experimental or virtual 2θ x-ray diffraction line profiles created from isolated STGBs for comparison; however, insights into the profile features are gained via comparison with the virtual SAED patterns. Clearly, the {111} and {220} peaks show a broadening due to the presence of the low-angle grain boundaries. In the line profiles, smooth broadening of a peak is the result of gradual changes to the spacing between a particular set of crystallographic planes. However, discrete peaks (roughness) in the broadening and secondary peaks emerge as the result of non-continuous displacement fields corresponding to the dislocation cores and periodic secondary relaxations within the grain boundary region. In the virtual 2θ line profiles, the {002} peak shows less significant broadening and subsidiary peaks with higher intensity as compared to the {111} and {220} peak maxima. While some of these peaks are created from secondary periodic relaxations, analysis of the SAED patterns indicates the main source for these distinct peaks are the relrods associated with the finite size of the simulation cell and the implementation of the virtual diffraction algorithm.
Large-Angle Symmetric Tilt CSL Grain Boundaries

The virtual diffraction methods are applied to three large-angle CSL [010] STGBs to explore the capability of these methods to identify atomic scale structure, specifically the different structural units and their repeating pattern along the grain boundary. Previous experimental and virtual diffraction studies have not focused on isolated large-angle symmetric tilt CSL grain boundaries, thus no direct comparison can be made. Figure 4.6(a) - (c) show the energy-minimized structures of the $\Sigma 5$ (210), $\Sigma 29$ (520), and $\Sigma 5$ (310) [010] STGBs respectively, coloured by the centrosymmetry parameter. The energy minimization routine successfully converges on the $|B.B|$, $|BC.BC|$, and $|C|$ structural units as predicted by the structural unit model [34]. Selected area electron diffraction patterns with zone axis aligned along the [010] misorientation direction for each large-angle CSL grain boundary are shown in Figure 4.6(d) - (e). Each SAED pattern is indexed denoting the plane and the lattice region responsible for the
Figure 4.6: Large-angle [010] STGB structures for (a) $\Sigma 5 \{210\} 36.87^\circ$, (b) $\Sigma 29 \{520\} 43.60^\circ$, and (c) $\Sigma 5 \{310\} 53.13^\circ$ as well as corresponding virtual electron diffraction patterns (d) - (f) with beam aligned along the [010] misorientation axis. The squares are included as a guide to distinguish the orientations of bicrystal regions.
diffraction reflection and squares are drawn as a guide to successfully identify the misorientation
between the two lattice regions. The SAED patterns for the large-angle CSL [010] STGBs,
using a 0.5\% of $I_{\text{max}}$ threshold, do not contain obvious subsidiary reflections that can be linked
directly to dislocations or atomic structure. However, in Figure 4.6(d) and Figure 4.6(f), the
SAED patterns do show the alignment of the \{204\} reflections from both lattice regions
stemming from the special orientations of these \(\Sigma 5\) boundaries. Attempts at lowering the
minimum intensity threshold to reveal subsidiary reflections were unsuccessful due to more
pronounced vertical and horizontal relrods in the SAED patterns and extra reflections due to
relrods originating from FCC peaks off the reciprocal lattice plane viewed in Figure 4.6.

Figure 4.7 presents a size effect study, analogous to that performed on a low-angle [010]
STGB in Figure 4.4, for a (002) relrod profile created by the \(\Sigma 5\) (310) STGB with increasing
\(|A_3|\). In this study, \(|A_3|\) is extended from 33.38 Å in the original study (Figure 4.5) to 66.77 Å

![Diagram showing average intensity profile](image)

**Figure 4.7**: Average intensity profile of the \(\Sigma 5\) \{310\} [010] STGB (002) relrod from
simulations with increasing \(|A_3|\) as labeled.
and 100.13 Å. Figure 4.7 shows that the relrod profiles for the three simulation sizes exhibit an excellent fit to the predicted diffraction intensity computed from Eq. (4.10). The fitted magnitude of \( a_3 \) for all three simulations is 3.71 Å with \( N_3 \) equal to 9, 18, and 27 respectively. These fitted parameters correspond to \( |A_3| \) of length 33.38, 66.74, and 100.11 Å, which closely match the \( |A_3| \) used in the simulations. The larger magnitude of the fitted \( a_3 \) parameter compared to those determined from the 10.39° [010] STGB with partial dislocations compensates for the greater misorientation in the \( \Sigma 5 \) (310) [010] STGB.

Virtual x-ray diffraction line profiles of the three large-angle CSL [010] STGBs are presented in Figure 4.8. Similar to the diffraction patterns presented in this study for low-angle symmetric tilt grain boundaries, these patterns show broadening of the \{111\} and \{220\} peaks as well as distinct secondary peaks emerging near the \{002\} peak. For each large-angle CSL [010] STGB, the broadening surrounding the \{111\} peak contains more discrete roughness, compared to the line profiles in Figure 4.5, indicating the presence of a secondary periodic structure within the grain boundaries impacting the \{111\} planar spacing. Further, there are subtle differences in the relative intensities of the secondary peaks on either side of the primary \{111\} peak between the different grain boundaries. Similarly, differences in the relative intensities of the discrete secondary peaks surrounding the \{002\} peak could indicate the presence of secondary structure that is buried beneath the relrod structure observed in Figure 4.6 and analyzed in detail in Figure 4.7. These observations provide motivation for future work to (1) explore long-range displacement field corrections to the virtual diffraction algorithm to minimize the role of simulation cell size on relrod structure and (2) explore a wide variety of zone axes in the SAED data, such as around the \{111\} reflections, as the virtual diffraction algorithm is capable of generating the full three dimensional reciprocal space map of each grain boundary. In addition,
reduction of the relrod structures that stem from the simulation size effects will aid in future comparisons of the virtual diffraction results to experimental efforts.

Figure 4.8: X-ray diffraction patterns for large-angle CSL [001] STGBs constructed using a $2\theta$ bin size of 0.067°. The extra high intensity peaks near the {002} peak are the result of relrod structures due to the finite size effect of the simulations; however, the roughness of the {111} peak broadening stems from periodic secondary relaxations within the boundaries.

4.4 Conclusion

This work presents a general method to compute and visualize virtual diffraction patterns from atomistic simulation data; this method is applied to study select low-angle and large-angle CSL Ni [010] symmetric tilt grain boundaries. For each STGB, the virtual diffraction algorithm produces SAED patterns and $2\theta$ x-ray diffraction line profiles via explicit evaluation of the structure factor equation without a priori knowledge of the grain boundary unit cell. The virtual SAED patterns of the low-angle [010] STGBs differentiate the misorientation between the grain boundary regions and contain subsidiary peaks linked to the edge dislocation array within the simulations. Similarly, virtual SAED patterns of the large-angle CSL [010] STGBs, are successful in differentiating the misorientation between regions; however, subsidiary peaks are
not visible on these SAED patterns aligned with the misorientation axis due to prominent relrod structures. Thorough analysis on the effects of simulation size on computed relrod structures is performed for both low-angle and large-angle grain boundaries, which confirms their origin to be the finite size of the simulation cell. Virtual 2θ x-ray diffraction line profiles of the low-angle STGBs show smooth peak broadening around the {111} and {220} peaks due to gradual changes to their associated planar spacing, as well as discrete secondary peaks near the {002} linked to the relrods originating from the finite simulation cell size. For the large-angle CSL [010] STGBs, the {111} peak broadening contains small discrete peaks indicating the presence of a secondary periodic structure impacting this family of lattice planes.

Results from this work suggest that further virtual diffraction studies of bicrystal models can advance the understanding of non-local and non-homogeneous elasticity around grain boundaries which can be incorporated into constitutive models constructed for meso-scale simulations [37]. Non-locality and inhomogeneity in the elastic stiffness is typically associated with defects leading to incompatible strain and curvatures [38,39]. To treat this complex problem at the continuum scale, higher order and higher-grade constitutive laws have been proposed (cf. [38–43]). These necessarily introduce higher-order elastic-type stiffness tensors of rank fifth and higher, which are unknown. Their identification could be reached via fitting of virtual diffraction peaks between a continuum mechanics based representation of grain boundaries and those presented in this work.

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References


1953.


Appendix 4.1


I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication. This work was done in collaboration with Dr. Laurent Capolungo at the Georgia Institute of Technology.

Sincerely,

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Appendix 4.2

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Chapter 5: A Computational Algorithm to Produce Virtual X-Ray and Electron Diffraction Patterns From Atomistic Simulations

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Abstract

Electron and x-ray diffraction are well-established experimental methods used to explore the atomic scale structure of materials. In this work, a computational algorithm is developed to produce virtual electron and x-ray diffraction patterns directly from atomistic simulations. This algorithm advances beyond previous virtual diffraction methods by utilizing a high-resolution mesh of reciprocal space which eliminates the need for \textit{a priori} knowledge of the crystal structure being modeled or other assumptions concerning the diffraction conditions. At each point on the reciprocal space mesh, the diffraction intensity is computed via explicit computation of the structure factor equation. To construct virtual selected area electron diffraction patterns, a hemispherical slice of the reciprocal lattice mesh lying on the surface of the Ewald sphere is isolated and viewed along a specified zone axis. X-ray diffraction line profiles are created by binning the intensity of each reciprocal lattice point by its associated scattering angle, effectively mimicking powder diffraction conditions. The virtual diffraction algorithm is sufficiently generic to be applied to atomistic simulations of any atomic species. In this article, the capability and versatility of the virtual diffraction algorithm is exhibited by presenting findings from atomistic simulations of $\langle 100 \rangle$ symmetric tilt Ni grain boundaries, nanocrystalline Cu models, and a heterogeneous interface formed between $\alpha$-Al\textsubscript{2}O\textsubscript{3} (0001) and $\gamma$-Al\textsubscript{2}O\textsubscript{3} (111).
5.1 **Introduction**

Virtual diffraction is a simulation technique that generates experimentally comparable diffraction patterns from atomistic simulation data using equations from diffraction theory. For over thirty years, researchers have used virtual diffraction methods to make direct connections between atomistic simulations and experimentation [1–11]; however, computational limitations inherent to the time of these early studies restricted their capability and required researchers to make assumptions about the crystal structure or the diffraction conditions. The virtual diffraction algorithm discussed in this article advances these earlier studies by leveraging modern computational power in order to eliminate the need for assumptions related to the crystal structure of the material, allowing a single algorithm to produce both x-ray diffraction line profiles and selected area electron diffraction (SAED) patterns from atomistic simulation [12]. The virtual diffraction algorithm offers new routes to connect atomistic simulations directly to experimental studies in order to investigate the structure and deformation of nanoscale materials and offers new capability to characterize complex materials within the framework of atomistic simulations.

Virtual diffraction models based on kinematic diffraction theory were developed in the 1980s to characterize atomistic simulations of large-angle symmetric twist grain boundaries using area diffraction patterns [1–3]. In kinematic models, the diffraction intensity, $I$, is computed for $N$ atoms as the product of the structure factor, $F(K)$, with its complex conjugate, $F^*(K)$, as follows:

$$I(K) = F^*(K)F(K) \quad (5.1)$$

where

$$F(K) = \sum_{j=1}^{N} f_j \exp\left(2\pi i K \cdot r_j\right) \quad (5.2)$$
Here, $\mathbf{K}$ is the location of the diffraction peak in reciprocal space, $\mathbf{r}_j$ is the position of the atom in real space, and $f_j$ is the atomic scattering factor (often simplified as unity [1–5]). In the seminal studies [1–3], computational limitations restricted both the number of atoms used to compute diffraction intensities and the range of reciprocal space that could be explored. To improve computational efficiency, diffraction intensities were computed over a limited region of reciprocal space known to be important based on \textit{a priori} knowledge of the grain boundary unit cell. Using these techniques, work by Bristowe and Sass [1] showed that small displacements of the atoms within the grain boundary unit cell can create identifiable changes to the diffraction patterns. This breakthrough prompted several researchers [2–5] to incorporate virtual diffraction patterns in their subsequent studies to verify predicted grain boundary structures.

Following the early kinematic diffraction studies, a second method to generate virtual diffraction patterns became popular among researchers modeling nanocrystalline materials that utilized assumptions based on powder diffraction conditions [6–11]. These researchers used formulations of the Debye scattering equation to compute diffraction intensity based on the interatomic distance between atoms, $r_{ij}$, via [13],

$$I(k) = \sum_{i=1}^{N} \sum_{j=1}^{N} f_i f_j \frac{\sin(2\pi k r_{ij})}{2\pi k r_{ij}}.$$  \hspace{1cm} (5.3)

Here, $k = 2\sin(\theta)/\lambda$ represents a spherically averaged position in reciprocal space that is related to the diffraction angle, $\theta$, and monochromatic radiation of wavelength, $\lambda$. By spherical averaging the positions in reciprocal space, all orientations of the simulated crystals are mathematically represented mimicking the random distribution associated with powder diffraction conditions. Using this relationship, researchers [6–11] constructed x-ray diffraction line profiles to investigate peak shift and peak broadening in order to extract data on the mean

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grain size and microstrain in nanocrystalline models with different grain diameters.

The virtual diffraction algorithm discussed in this work advances beyond these previous methods to create both SAED and x-ray diffraction line profiles using the same algorithm without any *a priori* knowledge of the crystal structure. The algorithm is sufficiently generic for all atomic species and is integrated into the LAMMPS molecular dynamics simulator [14] as a user-defined compute; it can also be implemented into other atomistic simulation packages.

Following a concise discussion of the computational algorithm, this article presents three unique applications of the virtual diffraction method, displaying its versatility and its capability to connect atomistic simulations with experimental studies.

5.2 Computational Algorithm

The diffraction algorithm generates a high-resolution, three-dimensional mesh of points filling a volume of reciprocal space constructed from the entire domain of the atomistic simulation cell. Each point on the reciprocal lattice mesh is associated with a reciprocal lattice vector $\mathbf{k}$ describing the deviation between the diffracted and incident wave vectors $\mathbf{k}_D$ and $\mathbf{k}_I$ [13],

$$
\mathbf{K} = \mathbf{k}_D - \mathbf{k}_I = \xi \mathbf{B}_1 + \eta \mathbf{B}_2 + \zeta \mathbf{B}_3 ,
$$

(5.4)

where $\xi$, $\eta$, and $\zeta$ can be any real number. The mesh of reciprocal lattice points is built on a rectilinear grid with spacing, $c_n|A_n|^{-1}$, along each reciprocal lattice axis $\mathbf{B}_n$, as shown in Figure 5.1. Each reciprocal lattice axis, $\mathbf{B}_n$, is determined from its associated vector, $\mathbf{A}_n$, corresponding to the $n = 1, 2, \text{or } 3$ edge of the simulation cell. The parameters $c_n$ control the spacing of the reciprocal lattice points, and tune the resolution within the reciprocal space volume explored. To increase the computational efficiency, the volume of reciprocal space that is explored is reduced by setting criteria on $|\mathbf{K}|$ as shown in Figure 5.1. By constructing a high-
resolution reciprocal space mesh, strong intensity peaks associated with the constructive interference of Bragg reflections are intuitively captured without \textit{a priori} knowledge of the crystal structure. The interplanar distances $d_{hkl}$ associated with these Bragg reflections can be computed at each reciprocal lattice point utilizing the geometric relationships between $k_1$, $k_0$, $k$, and $\lambda$ described in Bragg's Law \cite{13},

$$\frac{2\sin(\theta)}{\lambda} = \frac{1}{d_{hkl}} = |K|.$$  \hspace{1cm} (5.5)

Diffraction intensity at each reciprocal lattice point is computed using the structure factor equation, Eq. (5.2), with the variations described below to differentiate between electron and x-ray diffraction. In the diffraction algorithm, atomic scattering factors $f_j$ are explicitly computed for each atomic species to account for the reduction in diffracted intensity from an individual atom due to Compton scattering and vary by angle of diffraction and type of radiation \cite{15}. At
each diffraction angle, the atomic scattering factors \( f_j \) are computed using analytical approximations parameterized for the specific atomic species. For electron diffraction, the analytical approximation of the atomic scattering factor is taken as the summation of five Gaussian functions of the form [16],

\[
f_j \left( \frac{\sin \theta}{\lambda} \right) = \sum_{i}^{5} a_i \exp \left( -b_i \frac{\sin^2 \theta}{\lambda^2} \right), \tag{5.6}
\]

which have been parameterized for the majority of neutral elements by Peng et al. [17]. For x-ray diffraction, the analytical approximation of the atomic scattering factor is found from a summation of four Gaussian functions plus a constant of the form [18],

\[
f_j \left( \frac{\sin \theta}{\lambda} \right) = \sum_{i}^{4} a_i \exp \left( -b_i \frac{\sin^2 \theta}{\lambda^2} \right) + c, \tag{5.7}
\]

and have been parameterized by Fox et al. for most atomic species [19].

For electron diffraction, the diffraction intensity, \( I_e(K) \), at each reciprocal lattice point is computed from the product of the structure factor and its complex conjugate normalized by the number of atoms being studied within the simulation via [15],

\[
I_e(K) = \frac{F(K)F^*(K)}{N}. \tag{5.8}
\]

To compute x-ray diffraction intensities, \( I_x(K) \), the Lorentz-polarization factor, \( L_p(\theta) \), is applied to account for the relative distribution of the reciprocal lattice points and the change in scatter intensity when using non-polarized incident radiation. The Lorentz-polarization factor is computed via [15],

\[
L_p(\theta) = \frac{1 + \cos^2(2\theta)}{\cos(\theta)\sin^2(\theta)}, \tag{5.9}
\]
and thus the diffraction intensity from x-rays is computed at each reciprocal lattice point through [15],

\[ I_s(K) = L_p(\theta) \frac{F(K) F^*(K)}{N} . \]  

(5.10)

Virtual SAED patterns are created by examining the region in reciprocal space intersecting the Ewald sphere of radius \( \lambda^{-1} \). For a particular zone axis, the Ewald sphere is centered at the tail of the associated incident wave vector and intersects the origin of reciprocal space. To construct the electron diffraction pattern, a hemispherical slice of the reciprocal lattice mesh lying on the surface of the Ewald sphere is isolated and viewed along the zone axis. Using visualization software, intensity values are interpolated between points on the reciprocal space mesh allowing the slice to take on no thickness while maintaining a continuous intensity field along the hemisphere. To enhance features within the diffraction pattern, reciprocal lattice points are colored by intensity on a \( \log_{10} \) scale. To achieve higher computational efficiency when creating SAED patterns for a user-specified zone axis, the diffraction algorithm can be augmented to limit the intensity calculation to only those reciprocal mesh points that lie near the surface of the Ewald sphere.

X-ray diffraction line profiles are created by virtually rotating the Ewald sphere around the origin of reciprocal space to all possible orientations, mimicking powder diffraction conditions [20]. By making all diffraction orientations equally probable, every reciprocal lattice point will intersect the surface of the Ewald sphere. Line profiles simulating x-ray powder diffraction are constructed by collecting reciprocal lattice points into bins corresponding to their scattering angle \( 2\theta \) calculated using Eq. (5.5) and summing the intensity data. The scattering angle bin size is optimized through trials to reduce the noise within the line profile while maximizing peak features. To achieve higher computational efficiency while computing x-ray diffraction line
profiles within known 2θ limits, additional criteria on |\mathbf{K}| can be assigned using Bragg’s law, Eq. (5.5), to associate 2θ with |\mathbf{K}|.

The new diffraction algorithm leverages parallelization techniques and modern computing hardware to compute diffraction intensities using a much larger number of atoms and a much larger number of reciprocal lattice points compared to any prior study. The diffraction algorithm is implemented into LAMMPS as a user-defined compute and uses message passing interface (MPI) parallelization of the atoms matching the standard spatial decomposition built within the LAMMPS framework. To further increase computational efficiency, the diffraction compute uses shared memory parallelization over the reciprocal space mesh via OpenMP and is capable of offloading computation to external multicore hardware.

5.3 Applications

The capability and versatility of the virtual diffraction algorithm is illustrated via three applications: (1) low-angle Ni <100> symmetric tilt grain boundaries, (2) Cu nanocrystalline models, and (3) a complex heterogeneous alumina interface. These studies highlight the flexibility of the virtual diffraction technique to successfully model different material systems. In the following applications, simulated 200 kV electron radiation (λ = 0.0251 Å [16]) and Cu Kα x-rays (λ = 1.54178 Å [15]) are used to create SAED patterns and x-ray line profiles, respectively. In the electron diffraction studies, the SAED patterns are produced limiting \mathbf{K} from 0 ≤ |\mathbf{K}| ≤ 0.85 Å⁻¹ and restricting mesh points to a 0.01 Å⁻¹ hemispherical slice from the associated Ewald sphere in the intensity calculation. X-ray diffraction line profiles utilize a 2θ bin size of 0.07 degrees.

5.3.1 Ni Bicrystals

Virtual diffraction SAED patterns are constructed to analyze two bicrystal samples
containing low-angle symmetric tilt grain boundaries (STGBs) with a [010] tilt axis modeled using the Foiles-Hoyt Ni embedded-atom method (EAM) potential [21]. Here, low-angle [010] STGBs with tilt angles of 10.39° and 12.68° are constructed utilizing the methods described by Spearot et al. [22] with periodic boundary conditions applied in all directions. For each bicrystal, minimum energy grain boundary structures are attained at 0 K by aligning the opposing lattice regions with the intended tilt and removing atoms that are inside specified cutoff distances. This procedure effectively samples multiple starting configurations with different relative displacements. Each grain boundary structure is optimized according to the minimum potential energy using a nonlinear conjugate gradient method implemented in LAMMPS. Electron and x-ray diffraction data are computed using the atomic positions of the minimum energy grain boundary structures.

The two energy minimized [010] Ni STGB structures contain a periodic array of edge dislocations separated by a distance, \( d_D \), of 19.45 Å and 15.94 Å for the 10.39° and 12.68° tilts, respectively, as shown in Figure 5.2(a-b). Computed SAED patterns with the zone axis aligned along the [010] tilt axis are shown in Figure 5.2(c-f). Figure 5.2(c-d) are viewed such that Bragg reflections in the range of \( h^2+k^2+l^2 \leq 8 \) are visible in order to capture the misorientation between the two grains. The figures also show the presence of relrods near the Bragg reflections orientated parallel to the grain boundary plane (vertical direction) and oriented normal to the grain boundary (horizontal direction). Relrods are a locus of non-negligible intensity peaks encompassing Bragg reflections that occur during diffraction within a finite volume [23]. The size and shape of the relrods are dependent on the size and shape of the finite volume, which correspond to the number of atoms included within the summation of the structure factor equation.
Figure 5.2(e-f) show magnified regions near the (002) reflections and highlight the presence of two sets of subsidiary reflections within each SAED pattern not attributed to the relrod structures. These subsidiary peaks were observed in previous experimental works and have been related to the periodic spacing of the dislocation cores within the grain boundary [24]. In this work, the average spacing between the subsidiary peaks and the (002) relrod tail is calculated as 1/19.89 Å⁻¹ and 1/15.94 Å⁻¹ which is in very good agreement with the spacing between the dislocations in the bicrystal models.

5.3.2 Cu Nanocrystals

X-ray diffraction line profiles and SAED patterns are created for nanocrystalline Cu samples with different grain diameters and number of grains. The different nanocrystalline samples are
constructed using the Voronoi method to create a random distribution of grain orientations within fully periodic, cubic simulation cells. The Voronoi method requires the user to specify a target mean grain size and uses this value to determine the number and distribution of grain centers within the simulation cell. There is no guarantee that the true mean grain diameter resulting from the Voroni construction algorithm will be exactly equal to the user-defined target value. Within each nanocrystalline sample, the atomic interactions are modeled using the Cu EAM potential parameterized by Mishin et al. [25]. Before computing the virtual diffraction patterns, the atomic structures are relaxed at 0 K using a non-linear conjugate gradient method.

X-ray diffraction line profiles are constructed for each nanocrystalline Cu sample using a mesh resolution of approximately $4.7 \times 10^7$ reciprocal lattice points per Å$^{-3}$, which is achieved by appropriately scaling the $c_n$ parameters for the different simulation dimensions. Figure 5.3 shows a representative x-ray diffraction line profile for a nanocrystalline sample containing 300 grains with a target mean grain diameter of 5 nm. Four peaks are observed at 20 locations $43.29^\circ$, $50.43^\circ$, $74.15^\circ$, and $89.96^\circ$ which correspond to the interplanar distances associated with

![Figure 5.3: XRD pattern for a nanocrystalline Cu samples with 300 grains and 5 nm mean grain diameter. The nanocrystalline sample is shown in the inset colored by grain number.](image)
{111}, {200}, {220}, and {311} planes, respectively. These peak locations are a close match to the predicted locations computed using Bragg’s law for single crystal Cu with a lattice parameter of 3.615 Å [25], indicating that there is no net tensile or compressive strain in the nanocrystalline samples after the Voronoi construction and energy minimization procedures.

Peak locations and broadening from the virtual x-ray diffraction line profiles are used to perform a Williamson-Hall analysis [26] to predict microstrain in the lattice due to the grain boundaries as well as the true mean grain diameter of the nanocrystalline samples. This analysis is performed for six samples containing 20 and 400 grains with target mean grain diameters of 5, 10, and 15 nm. Both Lorentzian and Lorentzian-Gaussian distributions are fit to the x-ray diffraction peaks using the Fityk software [27], which provides peak location and peak broadening information. It is found that the Lorentzian-Gaussian distribution provides a closer approximation to the peak maxima and shapes through minimization of the residuals between the computed diffraction data and each fitted distribution. Figure 5.4 shows a Williamson-Hall plot.
using the Lorentzian-Gaussian fit to the virtual diffraction data. The true mean grain diameter and the microstrain are extracted via a linear fit and are reported in Table 5.1 for the six different samples using both Lorentzian and Lorentzian-Gaussian fittings. For the 5 nm model, the true mean grain diameter is larger than the target grain diameter used during Voronoi construction and the microstrain within each nanocrystalline model is non-homogeneous as evident by the non-linearity of the data. On the other hand, for the 10 and 15 nm samples, the true mean grain diameter predicted with the Lorentzian-Gaussian distribution is smaller than the target grain diameter and data is linear, implying that the root-mean-squared microstrain is isotropic. These observations are consistent with work of Derlet et al. [10]. For simulations containing the same number of grains, the magnitude of the microstrain decreases in models built with increasing target grain diameter. This is attributed to the smaller fraction of atoms within the larger nanocrystalline samples whose lattice potions are distorted by the grain boundaries. Models constructed with the same target mean grain diameter show negligible dependence of the microstrain on the number of grains.

Table 5.1 True mean grain diameter (nm) and microstrain predicted from the Williamson-Hall analysis using different peak fitting functions

<table>
<thead>
<tr>
<th>Target Grain Diameter</th>
<th>20 grains Microstrain</th>
<th>True Diameter</th>
<th>50 grains Microstrain</th>
<th>True Diameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 nm</td>
<td>0.0188&lt;sup&gt;a&lt;/sup&gt;, 0.0192&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.64&lt;sup&gt;a&lt;/sup&gt;, 6.12&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.0186&lt;sup&gt;a&lt;/sup&gt;, 0.0183&lt;sup&gt;b&lt;/sup&gt;</td>
<td>7.79&lt;sup&gt;a&lt;/sup&gt;, 6.59&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>10 nm</td>
<td>0.0040&lt;sup&gt;a&lt;/sup&gt;, 0.0026&lt;sup&gt;b&lt;/sup&gt;</td>
<td>10.85&lt;sup&gt;a&lt;/sup&gt;, 8.11&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.0040&lt;sup&gt;a&lt;/sup&gt;, 0.0026&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8.43&lt;sup&gt;a&lt;/sup&gt;, 6.85&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>15 nm</td>
<td>0.0020&lt;sup&gt;a&lt;/sup&gt;, 0.0012&lt;sup&gt;b&lt;/sup&gt;</td>
<td>11.01&lt;sup&gt;a&lt;/sup&gt;, 9.23&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.0018&lt;sup&gt;a&lt;/sup&gt;, 0.0010&lt;sup&gt;b&lt;/sup&gt;</td>
<td>11.59&lt;sup&gt;a&lt;/sup&gt;, 9.70&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup>X-ray diffraction peaks fit to a Lorentzian distribution.

<sup>b</sup>X-ray diffraction peaks fit to a Lorentzian-Gaussian distribution.

Figure 5.5 shows SAED patterns generated with zone-axes oriented along the [100] direction for simulations containing 50 and 400 grains using target mean grain diameters of 5 and 10 nm. Each SAED pattern contains three rings associated with the {111}, {200}, and {220} planes, as
expected from experimental results which were performed on a thin copper film containing 45 nm grains [28]. In the 400 grain models the rings are more complete as compared with the 50 grain models implying that models with a larger number of grains are statistically more representative of nanocrystalline samples with random grain orientations. In addition, the rings in the 10 nm samples are thinner than the rings in the 5 nm samples because of microstrain effects, analogous to the role of microstrain on peak broadening in the x-ray diffraction line profiles.

![SAED pattern for nanocrystalline Cu models](image)

**Figure 5.5:** SAED pattern for nanocrystalline Cu models containing 50 grains having (a) 5 nm and (b) 10 nm grain diameter as well as 400 grain models with (c) 5 nm and (d) 10 nm grain diameter.

### 5.3.3 Heterogeneous Al₂O₃ interfaces

Finally, virtual diffraction methods are used to characterize a complex interface formed between α-Al₂O₃ (0001) and γ-Al₂O₃ (111). Here, both the individual components and combined
interface are constructed using fully periodic boundary conditions with a large vacuum region separating free surfaces normal to the grain boundary. The boundary dimensions along the grain boundary plane are chosen to minimize the strain associated with joining the $\alpha$-Al$_2$O$_3$ (0001) and $\gamma$-Al$_2$O$_3$ (111) components forming the interface. The Al$_2$O$_3$ phases are modeled using the ReaxFF potential [29] and the structures associated with the individual components and the combined interface are relaxed at 0 K to their minimum potential energy structures using a nonlinear conjugate gradient method implemented in LAMMPS.

![Figure 5.6: X-ray diffraction line profiles identifying an $\alpha$-Al$_2$O$_3$ slab (bottom), a $\gamma$-Al$_2$O$_3$ slab (middle), and a $\alpha$-Al$_2$O$_3$ (0001), and $\gamma$-Al$_2$O$_3$ (111) interface (top). The colored triangles are positioned at experimentally determined peak locations for bulk $\alpha$-Al$_2$O$_3$ (red) [54] and $\gamma$-Al$_2$O$_3$ (green) [55].](image)

Prior to this work, identification of different Al$_2$O$_3$ phases, much less interfaces, has not been possible in atomistic simulations using traditional computational methods (radial distribution functions, centrosymmetry [30], common neighbor-analysis [31], energy-filtration methods, etc.) due to their non-cubic nature and subtle differences between their atomic structures. However, virtual x-ray diffraction line profiles of the individual components and the combined interface, shown in Figure 5.6, uniquely identify the phases. These virtual x-ray
diffraction results are further validated through direct comparisons to experimental results (colored triangles, in Figure 5.6) [32,33]. Peak locations for the simulated alumina models are skewed relative to the experimental reference indicating a uniform expansion of the lattice structure. This expansion is related to the equilibrium lattice parameter predicted by the ReaxFF potential [29] as well as the relaxation of the lattice due to the free surfaces in the simulations being modeled.

The complex crystal structures associated with $\alpha$-$\text{Al}_2\text{O}_3$ and $\gamma$-$\text{Al}_2\text{O}_3$ makes optimization of the five macroscopic degrees of freedom required for the construction of their interface difficult [34]. During this study, virtual SAED patterns assisted in the search for an ideal interface structure by identifying orientations that achieve higher orders of crystallographic compatibility, as shown by the overlapping diffraction peaks in Figure 5.7. Interfaces constructed with more
overlapping peaks, as identified by analyzing patterns created by the individual components as well as comparisons to prior experimental work [35], were lower in energy compared to those with disregistry

5.4 Conclusions

The virtual diffraction algorithm introduced in this article computes diffraction intensities via the structure factor equation at each point within a high-resolution reciprocal space mesh. The three-dimensional intensity data are used to construct both SAED and x-ray diffraction line profiles from atomistic simulations without a priori knowledge of the crystal structure. The diffraction algorithm is implemented within the LAMMPS molecular dynamics simulator and is capable of being integrated into other atomistic simulation packages in order to take advantage of modern parallelization techniques and computing hardware.

Three unique applications discussed in this article show the versatility of this algorithm to produce diffraction patterns using different atomic species and model configurations as well as its capability to link virtual diffraction work directly to experimental findings. In addition, the study shows a new route virtual diffraction offers in the characterization and optimization of complex material structures within atomistic simulations.

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I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication. This work was an invited submission in collaboration with Ph.D. candidate Mehrdad M. Sichani at the University of Arkansas.

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Appendix 5.2

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Chapter 6: Performance Improvement and Workflow Development of Virtual Diffraction Calculations

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Abstract

Electron and x-ray diffraction are well-established experimental methods used to explore the atomic scale structure of materials. In this work, a computational algorithm is presented to produce electron and x-ray diffraction patterns directly from atomistic simulation data. This algorithm advances beyond previous virtual diffraction methods by utilizing an ultra high-resolution mesh of reciprocal space which eliminates the need for \textit{a priori} knowledge of the material structure. This paper focuses on (1) algorithmic advances necessary to improve performance, memory efficiency and scalability of the virtual diffraction calculation, and (2) the integration of the diffraction algorithm into a workflow across heterogeneous computing hardware for the purposes of integrating simulations, virtual diffraction calculations and visualization of electron and x-ray diffraction patterns.
1.1 Introduction

Diffraction is a common experimental method used to study the atomic scale structure of a material [1]. In x-ray diffraction, for example, a sample is exposed to a beam of monochromatic x-rays with a wavelength on the same order as the spacing between atomic planes in the sample. X-rays scatter upon interacting with the atoms in the sample and the constructive interference of the scattered x-rays is collected and analyzed to determine the crystal structure and lattice constants of the material. Similarly, a beam of electrons can be used within a transmission electron microscope (TEM) to produce a selected area electron diffraction (SAED) pattern. Both x-ray and electron diffraction can be used to study lattice distortion due to defects within crystalline solids (cf. [2,3]).

Calculations of diffraction patterns based on kinematic diffraction theory were first developed in the 1980s to characterize atomistic simulations of large-angle symmetric twist grain boundaries [4–6]. In kinematic models, the diffraction intensity, \( I \), is computed for \( N \) atoms as the product of the structure factor, \( F(K) \), with its complex conjugate, \( F^*(K) \),

\[
I(K) = F^*(K)F(K) \quad (6.1)
\]

where

\[
F(K) = \sum_{j=1}^{N} f_j \exp \left( 2\pi i K \cdot r_j \right) \quad (6.2)
\]

Here, \( K \) is the location of the diffraction peak in reciprocal space, \( r_j \) is the position of the atom in real space, and \( f_j \) is the atomic scattering factor. In [4–6], computational limitations inherent to the time restricted both the number of atoms in the calculation and the range of reciprocal space explored. To improve computational efficiency, diffraction intensities were computed over a limited region of reciprocal space known to be important based on a priori knowledge of
the grain boundary unit cell. Using these techniques, Bristowe and Sass [4] showed that small displacements of the atoms within the grain boundary unit cell can create identifiable changes to the diffraction patterns. This breakthrough motivated researchers [4–8] to incorporate virtual diffraction patterns in their subsequent studies to verify predicted grain boundary structures.

In the mid 2000s, a second method to compute diffraction patterns that utilized assumptions based on powder diffraction conditions became popular among researchers modeling nanocrystalline materials [9–14]. These researchers used formulations of the Debye scattering equation to compute diffraction intensity based on the interatomic distance between atoms, \( r_{ij} \), via [1],

\[
I(k) = \sum_{i=1}^{N} \sum_{j=1}^{N} f_i f_j \frac{\sin(2\pi k r_{ij})}{2\pi k r_{ij}}. \tag{6.3}
\]

Here, \( k = 2\sin(\theta)/\lambda \) represents a spherically averaged position in reciprocal space that is related to the diffraction angle, \( \theta \), and monochromatic radiation of wavelength, \( \lambda \). By spherically averaging the positions in reciprocal space, all orientations of the simulated crystals are mathematically represented mimicking the random distribution associated with powder diffraction. Using this relationship, researchers [9–14] constructed x-ray diffraction line profiles to investigate peak shift and peak broadening in order to extract data on the mean grain size and lattice strain in nanocrystalline models with different grain diameters.

The virtual diffraction algorithm in this work advances beyond these previous methods to create both SAED and x-ray diffraction line profiles using the same algorithm without any \textit{a priori} knowledge of the crystal structure. The algorithm is sufficiently generic for all atomic species and is integrated into the LAMMPS atomistic simulation package [15] as a user-defined compute; it can also be implemented into other atomistic simulation packages. Following a
concise discussion of the computational algorithm, this article presents advancements made to increase performance and scalability, and the integration of this algorithm into a workflow across heterogeneous computing hardware including visualization. A complete discussion of the virtual diffraction algorithm is provided in [16,17] along with application of this method to study grain boundary structure, nanocrystalline materials and interfaces between complex solids.

1.2 Virtual Diffraction Method

1.2.1 Diffraction Intensity

The diffraction algorithm generates a high-resolution, three-dimensional mesh of points filling a volume of reciprocal space constructed using the entire domain of the atomistic simulation cell. Each point on the reciprocal space mesh is associated with a reciprocal lattice vector \( \mathbf{K} \) describing the deviation between the diffracted and incident wave vectors \( \mathbf{k}_d \) and \( \mathbf{k}_i \) [1],

\[
\mathbf{K} = \mathbf{k}_d - \mathbf{k}_i .
\] (6.4)

The mesh of reciprocal space points is built on a rectilinear grid with resolution defined by the user [16,17]. By constructing a high-resolution reciprocal space mesh, strong intensity peaks associated with the constructive interference between the incoming x-ray or electron beam and the crystal lattice are intuitively captured without \textit{a priori} knowledge of the crystal structure of the material. The interplanar distances \( d_{hkl} \) associated with each reflection can be computed at each reciprocal lattice point utilizing the geometric relationships described by Bragg's Law [1],

\[
\frac{2 \sin(\theta)}{\lambda} = \frac{1}{d_{hkl}} = |\mathbf{K}| .
\] (6.5)
The diffraction intensity at each reciprocal lattice point is computed using the structure factor equation, Eq. (6.1), with the modifications described below to differentiate between electron and x-ray diffraction. In the diffraction algorithm, atomic scattering factors are explicitly computed for each atomic species to account for the reduction in diffracted intensity from an individual atom due to Compton scattering [18]. Specifically, at each diffraction angle, the atomic scattering factors are computed using analytical approximations parameterized for each atomic species. For electron diffraction, the analytical approximation of the atomic scattering factor is taken as the summation of five Gaussian functions of the form [19],

\[ f_j \left( \frac{\sin \theta}{\lambda} \right) = \sum_{i=1}^{5} a_i \exp \left( -b_i \frac{\sin^2 \theta}{\lambda^2} \right) \]  

which have been parameterized for the majority of neutral elements by Peng et al. [20]. For x-ray diffraction, the analytical approximation of the atomic scattering factor is the summation of four Gaussian functions plus a constant of the form [21],

\[ f_j \left( \frac{\sin \theta}{\lambda} \right) = \sum_{i=1}^{4} a_i \exp \left( -b_i \frac{\sin^2 \theta}{\lambda^2} \right) + c \]  

which have been parameterized by Fox et al. [22] for most atom species. In addition, to compute x-ray diffraction intensities, the Lorentz-polarization factor, \( L_p(\theta) \), is applied to account for the relative distribution of the reciprocal lattice points and the change in scatter intensity when using non-polarized incident radiation. The Lorentz-polarization factor is computed via [18],

\[ L_p(\theta) = \frac{1 + \cos^2(2\theta)}{\cos(\theta)\sin^2(\theta)} \]  

1.2.2 Diffraction Pattern Generation

X-ray diffraction line profiles are created by virtually rotating the Ewald sphere around the
origin of reciprocal space to all possible orientations, mimicking powder diffraction conditions [18]. By making all diffraction orientations equally probable, every reciprocal space point will intersect the surface of the Ewald sphere. Line profiles are constructed by collecting reciprocal space points into bins corresponding to their scattering angle, using Eq. (6.5), and summing the intensity data within each bin. The scattering angle bin size is optimized through trials to reduce the noise within the line profile while maximizing peak features. For example, Figure 6.1 shows the calculated powder diffraction line profile for bulk \( \alpha \)-alumina with comparison to experimental data [23].

Virtual SAED patterns are created by examining the region in reciprocal space intersecting the Ewald sphere of radius \( 1/\lambda \). For a particular zone axis, the Ewald sphere is centered at the tail of the associated incident wave vector and intersects the origin of reciprocal space. To construct the electron diffraction pattern, a hemispherical slice of the reciprocal space mesh lying on the surface of the Ewald sphere is isolated and viewed along the zone axis. Using VisIt [24], intensity values are interpolated between points on the reciprocal space mesh allowing the slice to take on no thickness while maintaining a continuous intensity field along the hemisphere. For

![Figure 6.1: Calculated selected area electron diffraction pattern for the \( \gamma \)-alumina surface with comparison to experiment [25].](image)
example, Figure 6.2 shows the calculated SAED pattern for a (001) γ-alumina surface with comparison to experimental data [25]. To enhance features within the diffraction pattern, the sampled reciprocal space nodes are colored by intensity on a $\log_{10}$ scale. To achieve higher computational efficiency when creating SAED patterns for a user-specified zone axis, the diffraction algorithm can be augmented to limit the intensity calculation to only those reciprocal space mesh points that lie near the surface of the Ewald sphere.

![Figure 6.2: Calculated powder diffraction pattern of α-alumina with comparison to experiment [23].](image)

1.3 **Algorithm Scalability**

1.3.1 **Initial Scalability**

The performance and scalability of the diffraction calculation was documented to provide benchmark data prior to the partnership with XSEDE Extended Collaborative User Services (ECSS). Benchmark diffraction computations were completed on the Texas Advance Computing Center (TACC) Stampede system, which is configured with compute nodes that host two 8-core Xeon E5-2680 processors and one Intel Xeon Phi SE10P coprocessor (MIC). The compute nodes are outfitted with 32 GB of memory (2GB/core) which is separate from the 8 GB of memory on the Xeon Phi coprocessor. The benchmark simulation contains 256,000 bulk Ni atoms and explores 9,006,316 reciprocal space nodes during a single x-ray diffraction computation of the static structure. The benchmark simulation is run within the LAMMPS
atomistic simulation package. The output of the compute is an x-ray diffraction line profile for Ni with \(2\theta\) ranging from 10° to 90°. Timing data and memory usage are extracted directly from primary x-ray diffraction compute to avoid any overlaying functionalities within LAMMPS.

Initially, the code was parallelized via the native message passing interface (MPI) parallelization within the LAMMPS atomistic simulation package, which performs a spatial decomposition of the atoms in a simulation, illustrated in Figure 6.3. The benchmark simulations were used to determine the speedup and efficiency of the diffraction calculation over 1, 2, 4, 8, and 16 nodes. Results of the initial scalability tests are shown in Table 6.1. The speedup values are determined by comparing the absolute time for the computation using multiple nodes, compared to the computation time run using one 16-core node. Efficiency is computed by comparing the computation core-time to the reference core-time using one 16-core node. The

![Figure 6.3: Schematic of the MPI parallelization technique in the initial diffraction code. The blue cube represents the atomistic simulation while the red sphere represents the reciprocal space nodes sampled in the diffraction calculation.](image-url)
results of the initial implementation of the code (prior to ECSS) show a 13.23 speedup with an
efficiency reduction to 83% when scaled to 256 cores. The reduction in efficiency is primarily
due to the finite number of atoms to parallelize over within the strong scaling study. An
identified weakness of the initial implementation of the virtual diffraction code is its memory
footprint. The initial code requires that both the locations of the atoms and the locations of the
reciprocal space nodes are kept in memory throughout the entire calculation. More importantly,
a copy of all these locations must be accessible to each MPI process. Therefore, as the
simulation size and resolution of reciprocal space increases, the memory required to complete the
calculation dramatically increases. For example, the modest size benchmark simulation requires
9.7 GB of the available 32 GB of memory per node to compute a single x-ray diffraction line
profile.

Table 6.1: Initial scalability testing showing speedup, efficiency, and total memory usage.

<table>
<thead>
<tr>
<th>Nodes</th>
<th>Cores</th>
<th>Speedup</th>
<th>Efficiency</th>
<th>Memory (GB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>16</td>
<td>1.00</td>
<td>100%</td>
<td>9.67</td>
</tr>
<tr>
<td>2</td>
<td>32</td>
<td>1.91</td>
<td>96%</td>
<td>19.34</td>
</tr>
<tr>
<td>4</td>
<td>64</td>
<td>3.65</td>
<td>91%</td>
<td>38.67</td>
</tr>
<tr>
<td>8</td>
<td>128</td>
<td>6.70</td>
<td>84%</td>
<td>77.35</td>
</tr>
<tr>
<td>16</td>
<td>256</td>
<td>13.23</td>
<td>83%</td>
<td>154.70</td>
</tr>
</tbody>
</table>

1.3.2 Scalability Improvements

Both speedup and efficiency of the diffraction calculation are improved through ECSS
collaboration, resulting in a second generation of the code which has improved memory
utilization and incorporated a second level of OpenMP parallelization shown schematically in
Figure 6.4. Further, a third generation of the diffraction code is developed via ECSS
collaboration to take advantage of offloading tasks to the available MIC coprocessors on
Stampede. Speedup values for each generation of the diffraction code are evaluated using the
same benchmark simulation as in Section 6.3.1. The speedup value is computed from the absolute time of computation using the new code on multiple nodes as compared to the computation time run using the initial code on one 16-core node. Thus, speedup values are directly comparable across the different generations of the diffraction code. Differently, efficiency values are computed for each specific code generation and utilization of MPI/OpenMP. Results from the new scalability tests are shown in Table 6.2.

![Figure 6.4: Schematic of the OpenMP parallelization technique of the reciprocal space nodes sampled in the diffraction calculation.](image)

In the second generation of the virtual diffraction code, the MPI and OpenMP-based parallelization show speedup of 1.69 and 1.78, respectively, due to a reorganization of data structures and incorporation of more efficient calculations. MPI-based parallelization shows a similar efficiency drop (82% using 256 cores) as the initial code due to the strong scaling effects. To mitigate this effect, OpenMP threads are added to parallelize over the reciprocal lattice.
points. As shown in Table 6.2, the efficiency of the OpenMP parallelized second generation code is less affected by the finite number of atoms due to its second level of parallelism over the reciprocal lattice nodes (95% using 256 cores). The MIC enabled third generation code shows approximately 2x speedup from the second generation code by offloading a section of computation to the MIC. The reported values are taken when 90% of the reciprocal lattices nodes are offloaded to the MIC to be used in the solution of the structure factor equation. The remaining 10% of the reciprocal lattice nodes are utilized within concurrent computations on the CPU.

Table 6.2: Scalability tests of the second and third generation diffraction code showing speedup compared to the original 16-core timing, efficiency relative to code generation and utilization of MPI/OpenMP, and total memory usage.

<table>
<thead>
<tr>
<th>Nodes - MPI/OpenMP/MIC</th>
<th>Speedup</th>
<th>Efficiency</th>
<th>CPU Memory (GB)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Second Generation</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 - 16/0/0</td>
<td>1.69</td>
<td>100%</td>
<td>8.1</td>
</tr>
<tr>
<td>2 - 32/0/0</td>
<td>3.21</td>
<td>95%</td>
<td>16.1</td>
</tr>
<tr>
<td>4 - 64/0/0</td>
<td>6.16</td>
<td>91%</td>
<td>32.2</td>
</tr>
<tr>
<td>8 - 128/0/0</td>
<td>11.71</td>
<td>86%</td>
<td>64.4</td>
</tr>
<tr>
<td>16 - 256/0/0</td>
<td>22.17</td>
<td>82%</td>
<td>128.9</td>
</tr>
<tr>
<td>1 - 1/16/0</td>
<td>1.78</td>
<td>100%</td>
<td>0.51</td>
</tr>
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<td>2 - 2/16/0</td>
<td>3.56</td>
<td>100%</td>
<td>1.01</td>
</tr>
<tr>
<td>4 - 4/16/0</td>
<td>7.12</td>
<td>100%</td>
<td>2.02</td>
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<td>8 - 8/16/0</td>
<td>14.16</td>
<td>99%</td>
<td>4.05</td>
</tr>
<tr>
<td>16 - 16/16/0</td>
<td>26.94</td>
<td>95%</td>
<td>8.10</td>
</tr>
<tr>
<td><strong>Third Generation (MIC – Enabled)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 - 1/16/240</td>
<td>4.61</td>
<td>100%</td>
<td>0.51</td>
</tr>
<tr>
<td>2 - 2/16/240</td>
<td>9.19</td>
<td>100%</td>
<td>1.01</td>
</tr>
<tr>
<td>4 - 4/16/240</td>
<td>17.43</td>
<td>95%</td>
<td>2.02</td>
</tr>
<tr>
<td>8 - 8/16/240</td>
<td>34.65</td>
<td>94%</td>
<td>4.05</td>
</tr>
<tr>
<td>16 - 16/16/240</td>
<td>60.93</td>
<td>83%</td>
<td>8.10</td>
</tr>
<tr>
<td>1 - 16/0/240</td>
<td>4.37</td>
<td>100%</td>
<td>8.1</td>
</tr>
<tr>
<td>2 - 32/0/240</td>
<td>8.38</td>
<td>96%</td>
<td>16.1</td>
</tr>
<tr>
<td>4 - 64/0/240</td>
<td>15.94</td>
<td>91%</td>
<td>32.2</td>
</tr>
<tr>
<td>8 - 128/0/240</td>
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<td>84%</td>
<td>64.4</td>
</tr>
<tr>
<td>16 - 256/0/240</td>
<td>49.32</td>
<td>70%</td>
<td>128.9</td>
</tr>
</tbody>
</table>
1.4 Workflow Implementation and Visualization

1.4.1 SEAGrid Lammps_DS Workflow

The coupled execution of LAMMPS molecular dynamics followed by the simulation of x-ray diffraction or selected area electron diffraction calculations of the appropriately averaged system of interest and consequent visualization of the diffraction patterns required setting up of a workflow. The need for a workflow is particularly significant when multiple platforms for computing and visualization are to be used to address the large memory VSMP implementation of the visualization software required in the final step. The workflow problem is tackled by using an existing XSEDE SEAGrid science gateway, that supported the LAMMPS application and the corresponding tools to achieve the remote job submission, but in this case using a remote workflow submission and managing the data such that a single jobID handle can provide all the data at the end of the multi-resource computations. Additionally, a script-based high-throughput submission of multiple jobs is implemented in SEAGrid client, as shown in Figure 6.5. This uses an XML script with tags to specify job requirements and input files as needed and the system is equipped to execute each of the jobs specified as independent jobs. This is particularly useful for parameter sweep type runs and may benefit other communities that use the SEAGrid gateway as well.

The workflow implemented consists of coupled execution of three main tasks: Lammps_MD followed by Lammps_XRD, Lammps_SAED and a set of parallel VisIt executions as depicted in the schematic in the Figure 6.6. Initial implementations of the workflow used the same compute resource or a closely coupled resource at the same host site (such as Stampede and Ranch). A complete implementation involves execution of initial Lammps_MD at TACC’s Stampede system and the latter two tasks in SDSC’s Gordon system. The large memory
available for the Gordon system was critical for the second stage of the workflow and it is useful in distributing the computing and visualization tasks. This also facilitates interactive access to the VisIt service deployed on Gordon in the future.

![Figure 6.5: Job setup panel in DESSERT client with multi-job submission script.](image)

The workflow execution required additional inputs to drive the latter tasks and the original LAMMPS input was modified with a remark line for this purpose. The initial LAMMPS execution ignores this line in the input while the same is parsed and the queue instructions are set using this data for the subsequent steps. SEAGrid uses GSISSH based execution of a local script for the execution of a job task at a remote HPC site (for details of SEAGrid operation please refer to [26] and [27]). The LAMMPS specific portion of the script is modified to include the
execution of the workflow transparently during this ECSS project. SEAGrid supported publicly released LAMMPS already, but a special module named DS (Doug Spearot’s version) is implemented and currently restricted to the developer group to accommodate the special workflow version. The initial job is launched with the modified input on the Stampede system as a standard SEAGrid job. The Stampede job in turn prepares the inputs and provides the data required for the Lammps_XRD (and Lammps_SAED) and the VisIt components for the second stage. The script verifies the normal termination of the first stage, moves all the data and inputs for the subsequent jobs to the SDSC Gordon system and launches the jobs in Gordon. The path hierarchy is consistently managed and all the results are archived at the end of each stage into the mass storage device in SEAGrid organization. The entire workflow is tracked as a single job and all the corresponding result files can be retrieved based on the job handle through MyCCG job monitoring function in the client. The files retrieved can be further processed using the post processing tools available in the GridChem client. Though SEAGrid integrated the Apache Airavata based XBaya workflow execution system [28] which is described in detail in [27] for Paramchem project, this explicit implementation is chosen as the Gordon system did not support the Gram based orchestration of workflow available in XBaya. Currently, we are exploring the
Unicore [29] based orchestration of tasks on Gordon system through Airavata/Xbaya system.

1.4.2 Visualization

Visualization uses the VisIt program and utilizes either a session-less protocol or a set of session files supplied along with the LAMMPS input by the user. The session files are staged to the appropriate computer resource and used in the final task. A rendering python script drives the VisIt computation in a parallel batch job for each of the vtk files generated during the processing of the diffraction patterns that are provided as LAMMPS dump files in the Lammps_XRD compute step. The VisIt runs result in the images outputted in .png format. The visualization of the images from the simulation can be launched from the GridChem client automatically and an example set for an alumina surface is shown in Figure 6.7. Alternatively if the visualization programs are locally installed, such as Ovito used to generate Figure 6.7(a), appropriate files can be exported to these applications from the GridChem post processing tool.

1.5 Summary

In this work, a computational algorithm is presented to produce electron and x-ray diffraction
patterns directly from atomistic simulation data. First, through XSEDE ECSS support, the scalability and performance of the virtual diffraction algorithm has been considerably improved. The diffraction algorithm parallelization has been enhanced to support offloading to the MIC coprocessor on Stampede. Second, through XSEDE ECSS support, the virtual diffraction algorithm has been integrated into a workflow platform that allows for job submission across heterogeneous computing hardware at different sites within the XSEDE network. Specifically, atomistic simulations using LAMMPS, diffraction calculations using LAMMPS and visualization using VisIt are coupled in the workflow using the SEAGrid Science gateway. Ultimately, the diffraction algorithm provides a novel bridge between experiments and computation and will be of significant use to both communities to study the atomic-level structure of materials.

**Acknowledgements**

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Appendix 6.1


I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication. This work was an invited submission in collaboration with Drs. Sudhakar Pamidighantam and Mark Van Moer at the National Center for Supercomputing Applications (NCSA), Dr. Yang Wang at the Pittsburg Supercomputing Center (PSC), and Dr. Lars Koesterke at the Texas Advanced Computing Center (TACC), as part of a grant received from the XSEDE supercomputing network.

Sincerely,

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Appendix 6.2

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Chapter 7: Atomistic simulation and virtual diffraction characterization of stable and metastable Al₂O₃ surfaces

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Abstract

The structures of select alumina surfaces are studied using molecular statics and molecular dynamics simulations and are characterized using virtual diffraction methods. First, bulk alumina simulations are performed to validate the transferability of the ReaxFF potential to model different alumina phases. Bulk alumina simulations accurately predict α-Al₂O₃ as the lowest energy crystalline phase; however, they unexpectedly predict an even lower energy amorphous phase. At 0 K, virtual x-ray diffraction patterns of the bulk crystalline phases and select alumina surfaces are validated by experimental studies. Molecular statics simulations of select alumina surfaces are consistent with prior first-principles studies. However, molecular dynamics simulations show that many surfaces experience significant reconstructions at temperatures below what is expected from experiments. It is believed that premature surface reconstructions are biased by the predicted lower energy amorphous phase and occur due to the extra degrees of freedom allowed by the free surfaces as well as the available thermal energy during dynamics. Discrete peaks appearing in virtual selected area electron diffraction patterns indicate that the reconstructions are not fully amorphous due to lattice constraints imposed by the internal bulk structure. Bulk and surface energies are tabulated for each simulation to be used in future predictive mesoscale models of polymorphic alumina.
7.1 Introduction

Alumina (Al₂O₃) is an abundant ceramic material that exhibits extraordinary structural flexibility [1–5]. The different Al₂O₃ phases display a range of unique physical properties which make them useful in a variety of coating applications [6,7]. These properties stem from only subtle differences within the crystal structure of its phases. In general, the alumina phases are composed of a close-packed O sublattice surrounded by Al interstitials filling 2/3 of the octahedral and tetrahedral sites to maintain stoichiometry [8]. The type of close-pack arrangement of the O sublattice and the degree of symmetry of the Al interstitials within each alumina unit cell determines the phase and properties of the material. Corundum, α-Al₂O₃, is the only thermodynamically stable alumina phase, which contains the most symmetric ordering of the Al interstitials [1]. High symmetry leads to the high density of α-Al₂O₃ compared to the other phases and promotes directionality and ionic bonding between the atoms leading to high hardness. The metastable alumina phases have decreasing symmetry of the Al interstitials, which decreases their density and weakens their bond strength by reducing bond directionality. Due to subtle structural differences, phase identification within atomistic simulations based solely on local atomic positions is difficult; however, experimental characterization techniques such as x-ray diffraction (XRD) and selected area electron diffraction (SAED) have proven successful in distinguishing between alumina phases [9–13].

Selective vapor deposition of single-phase alumina thin films is often desired in order to take advantage of specific material properties, but is difficult due to the complexity of the alumina material system. For example, κ- and α-Al₂O₃ surfaces are widely regarded as ideal protective coatings due to their wear resistance, chemical inertness, resistance to thermal shock, and high hardness [14,15]. However, several metastable alumina phases demonstrate properties less ideal
for these purposes and can form during deposition depending on the processing conditions [16]. For example, \( \theta \)- and \( \gamma \)-Al\(_2\)O\(_3\) metastable phases may form at lower processing temperatures \( T_m \sim 0.3 \) [16]. These metastable phases have lower surface energies and therefore exhibit higher surface areas making them more appropriate as catalytic supports. In addition, selective vapor deposition of single-phase alumina proves difficult due to alumina's complex phase transition series that is dependent on the material precursors and processing conditions [17–19]. To determine the current state under a particular synthesis process, researchers and manufacturers must continually characterize the surface structure.

Predictive mesoscale material simulations can aid in the search for the ideal processing conditions that produce tailored alumina coatings, similar to solidification studies of multicomponent systems [20–22]. By considering the mechanisms governing phase formation and evolution, predictive models can computationally explore the unique processing conditions that achieve single-phase coatings in polycrystalline materials. Analogous experimental approaches, such as those taken to develop structure zone diagrams [23,24], rely on phenomenological observations requiring an exhaustive experimental study for a polymorphic material system and are only predictive within the same processing space. Mesoscale models, such as phase-field methods [25], for physical vapor deposition of polymorphic materials will require energetic data (i.e., bulk, surface, and interface energies) that can be easily computed from atomistic simulations. However, to quantitatively compare these data the relevant energies need to be computed using the same computational model, as is done in this work.

Specifically, this work investigates select bulk and surface alumina structures using atomistic simulations modeled with the reactive force-field (ReaxFF) potential [26]. This article begins with a detailed discussion of previous atomistic simulations performed on alumina bulk and
surface structures highlighting the necessity for a consistent computational model to provide quantitatively comparable data. This is followed by a description of the current simulation methods using ReaxFF as well as the virtual diffraction method [27,28] used to characterize the nanoscale structure of the alumina simulations. Next, computed surface energies as well as characterization results are reported and analyzed. This article concludes with a summary and discussion of the future work needed to develop predictive mesoscale models of vapor deposition for polymorphic alumina.

7.2 Previous atomistic studies of alumina

Because of its important material properties and wide industrial use, the alumina system has been studied extensively using molecular dynamics (MD), molecular statics (MS), and first-principles computational models. However, none of these previous atomistic studies has attempted to catalog structural and energetic properties of bulk and surface structures across more than two alumina phases using the same computational model. By narrowing the focus on select alumina components, prior atomistic simulation studies have avoided the challenges of uniquely characterizing the subtle structural differences among the various alumina phases as well as the distortion created by atomic relaxations near surfaces. More importantly, because prior alumina studies used different computational models, direct comparisons between computed energetic values cannot be made.

First-principles simulations have studied alumina bulk and surface structures at 0 K using models based on quantum mechanics which rely on approximations to describe the electron interactions explicitly (cf. [29–31]). These approximations can be divided into three classes: (i) Hartree-Fock (HF) theory, (ii) density functional theory with local density approximations of the electron exchange-correlation (DFT-LDA), and (iii) density functional theory with generalized
gradient approximations of the electron exchange-correlation (DFT-GGA). Differently, MS and MD simulations use models based on Newtonian physics which represent atoms as point masses in space that encompass both the nucleus and the orbiting electrons. Interactions between atoms are governed by an interatomic potential to describe the potential energy of the system. Prior MS and MD simulations employed a variety of interatomic potentials with different approximations for electrostatic and non-electrostatic interactions (i.e., the styles describing pair, many-body, and bonded interactions). The approximations used to describe the electrostatic interactions in alumina can be divided into three different classes: (i) fixed point charges [32–43], (ii) charged shell models [44–48], or (iii) dynamic geometry dependent charges [49–52]. Each of these increases in complexity in an attempt to better represent the polarization of the O ions.

7.2.1 Bulk Alumina Studies

Prior atomistic simulations of alumina modeled bulk systems in order to assess the transferability of the computational model to multiple phases. Because \( \alpha\)-Al\(_2\)O\(_3\) is the only thermodynamically stable phase, transferable computational models should predict \( \alpha\)-Al\(_2\)O\(_3\) as the lowest potential energy structure per Al\(_2\)O\(_3\) unit. Table 7.1 lists a sample of previous atomistic simulation studies that compared the predicted \( \alpha\)-Al\(_2\)O\(_3\) potential energy to a selection of metastable alumina phases as well as the bixbyite structure. Here, relative energies are measured as the difference between the predicted potential energy of the specified phase and \( \alpha\)-Al\(_2\)O\(_3\) such that positive values indicate lower energy \( \alpha\)-Al\(_2\)O\(_3\).

Table 7.1 shows different computational models predict a different ordering of the energetic stability among alumina phases and shows that some models fail to predict the stability of \( \alpha\)-Al\(_2\)O\(_3\). Wilson et al. [53] commented that some of the discrepancy among the various models
Table 7.1: Sample values of computed energy deviations from $\kappa$-, $\theta$-, $\gamma$-, and bixbyite (B) $\text{Al}_2\text{O}_3$ compared to $\alpha$-$\text{Al}_2\text{O}_3$. Values are reported in eV per formula unit.

<table>
<thead>
<tr>
<th>Year</th>
<th>Comp. Model</th>
<th>$\Delta E(\kappa-\alpha)$</th>
<th>$\Delta E(\theta-\alpha)$</th>
<th>$\Delta E(\gamma-\alpha)$</th>
<th>$\Delta E(B-\alpha)$</th>
<th>Ref.</th>
</tr>
</thead>
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<tr>
<td>2007</td>
<td>DFT-LDA</td>
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<td>[54]</td>
</tr>
<tr>
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<td>MS$^a$</td>
<td>---</td>
<td>0.19</td>
<td>---</td>
<td>0.66</td>
<td>[52]</td>
</tr>
<tr>
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<td>---</td>
<td>0.18</td>
<td>---</td>
<td>[55]</td>
</tr>
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<td>DFT-GGA</td>
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<td>0.03</td>
<td>---</td>
<td>---</td>
<td>[56]</td>
</tr>
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<td>-0.01</td>
<td>---</td>
<td>0.08</td>
<td>[57]</td>
</tr>
<tr>
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<td>DFT-GGA</td>
<td>0.09</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>[58]</td>
</tr>
<tr>
<td>2001</td>
<td>DFT-LDA</td>
<td>0.15</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>[58]</td>
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<tr>
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<td>0.04</td>
<td>---</td>
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<td>[59]</td>
</tr>
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<td>0.25</td>
<td>---</td>
<td>0.21</td>
<td>[59]</td>
</tr>
<tr>
<td>1998</td>
<td>DFT-LDA</td>
<td>---</td>
<td>0.38</td>
<td>---</td>
<td>0.97</td>
<td>[60]</td>
</tr>
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<tr>
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<td>---</td>
<td>0.44</td>
<td>---</td>
<td>---</td>
<td>[61]</td>
</tr>
</tbody>
</table>

$^a$ Bond Style – Geometry Dependent  
$^b$ Pair Style – Charged Shell

stems from difficulties representing the polarization of the O ions. Another difficulty stems from uncertainty in the exact crystal structure of several of the metastable phases. In particular, the structure of the $\kappa$- and $\gamma$-$\text{Al}_2\text{O}_3$ phases has been the subject of much debate regarding the placement of the Al ions in the O interstitial sites. Using similar first-principles approaches, Yourdshahyan et al. [62,63] and Paglia et al. [64,65] studied the minimum energy structures of bulk $\kappa$-$\text{Al}_2\text{O}_3$ and $\gamma$-$\text{Al}_2\text{O}_3$, respectively. In their studies, candidate structures for the bulk phases were constructed using a chosen O sublattice and Al ions located in combinations of every possible interstitial site. The 40-atom $\kappa$-$\text{Al}_2\text{O}_3$ unit cell, studied by Yourdshahyan et al. [62,63], produced 169 candidate structures whose bulk energies could be computed directly from first-principles. Differently, the 160-atom $\gamma$-$\text{Al}_2\text{O}_3$ unit cell studied by Paglia et al. [64] produced $\sim$1.47 billion candidate structures that could not be studied directly from first-principles due to computational costs. Instead, Paglia et al. used selection criteria to reduce the number of $\gamma$-$\text{Al}_2\text{O}_3$
candidates to 56,064 that were then modeled using less computationally expensive MS simulations. The MS simulations produced a subset of 1,161 lower-energy structures with more-optimal atomic configurations. Paglia et al. [64] computed bulk energies of select low-energy structures using first-principles methods and observed the same energy trends as the MS simulation results.

Low-energy bulk $\kappa$- and $\gamma$-$\text{Al}_2\text{O}_3$ structures found by Paglia and Yourdshahyan were further analyzed using virtual diffraction methods. In their work, Yourdshahyan et al. showed that the virtual XRD line profile generated from the lowest-energy unit cell was the best match to the experimentally obtained pattern [2]. Differently, Paglia et al. showed that simulated neutron diffraction line profiles from low-energy $\gamma$-$\text{Al}_2\text{O}_3$ structures did not match the experimental results; instead, a better experimental match was found from unit cells slightly altered from the ideal spinel structure [64].

### 7.2.2 Alumina Surface Studies

Many prior studies have independently examined select surface structures of several alumina phases. In particular, many studies focused on the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) surface due to its important industrial use in catalytic and electronic supports. Table 7.2 lists a sample of previous atomistic studies that explored the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) surface structure along with computed surface energy values. From Table 7.2, it is clear that the computational model can significantly influence the computed values for surface energy. Analysis of these studies shows that structural relaxations and size effects can further affect the results. Tasker [66] was the first to incorporate an energy minimization routine into his study, which revealed substantial structural relaxations of the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) surface. Verdozzi et al. [67] showed that the degree of structural relaxation is impacted by the number of layers contained in the surface model (i.e., the thickness
Table 7.2: Sample history of α-Al₂O₃ (0001) surface energy values computed using various atomistic simulation methods. Values in parenthesis are for non-relaxed surface structures.

<table>
<thead>
<tr>
<th>Year</th>
<th>Comp. Model</th>
<th>E_{surf} (0001) [J/m²]</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
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<td>DFT-GGA</td>
<td>(3.51) 1.54</td>
<td>[68]</td>
</tr>
<tr>
<td>2007</td>
<td>HF</td>
<td>(4.82) 1.85</td>
<td>[69]</td>
</tr>
<tr>
<td>2004</td>
<td>DFT-GGA</td>
<td>1.98</td>
<td>[70]</td>
</tr>
<tr>
<td>2004</td>
<td>MS⁻</td>
<td>1.00</td>
<td>[52]</td>
</tr>
<tr>
<td>2004</td>
<td>DFT-GGA</td>
<td>(3.15) 1.54</td>
<td>[55]</td>
</tr>
<tr>
<td>2004</td>
<td>DFT-GGA</td>
<td>(3.52) 1.55/1.57</td>
<td>[71]</td>
</tr>
<tr>
<td>2003</td>
<td>DFT-GGA</td>
<td>(3.58) 1.60</td>
<td>[72]</td>
</tr>
<tr>
<td>2003</td>
<td>DFT-LDA</td>
<td>(3.97) 1.94</td>
<td>[72]</td>
</tr>
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<td>2001</td>
<td>MS⁻</td>
<td>(6.5) 3.15 at 300K</td>
<td>[73]</td>
</tr>
<tr>
<td>1999</td>
<td>DFT-GGA</td>
<td>(3.5) 1.95</td>
<td>[74]</td>
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<tr>
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<td>DFT-LDA</td>
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<td>[75]</td>
</tr>
<tr>
<td>1998</td>
<td>MS⁻</td>
<td>2.06 at 300K</td>
<td>[76]</td>
</tr>
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<td>1995</td>
<td>MS⁻</td>
<td>(4.48) 2.00</td>
<td>[77]</td>
</tr>
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<td>MS⁻</td>
<td>(3.86) 1.70</td>
<td>[77]</td>
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<td>MS⁻</td>
<td>(3.67) 2.67</td>
<td>[49]</td>
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<td>1994</td>
<td>DFT-LDA</td>
<td>(3.77) 1.76</td>
<td>[78]</td>
</tr>
<tr>
<td>1993</td>
<td>MS⁻</td>
<td>(5.04) 2.04 at 300K</td>
<td>[36]</td>
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<td>(3.7) ---</td>
<td>[79]</td>
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<td>1992</td>
<td>HF</td>
<td>(3.30) ---</td>
<td>[80]</td>
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<tr>
<td>1989</td>
<td>HF</td>
<td>(6.53) 5.32</td>
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<td>1987</td>
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<td>[66]</td>
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<tr>
<td>1980</td>
<td>MS⁻</td>
<td>(4.82) ---</td>
<td>[83]</td>
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⁻ Bond Style – Geometry Dependent  
⁻ Pair Style – Charged Shell  
⁻ Pair Style – Point Charge  
⁻ Many-body Style – Geometry Dependent  
⁻ Many-body Style – Point Charge

of the simulation). In Table 7.2, size effects were not considered in earlier simulations as they focused on constructing a minimum size model to reduce the computational expense.

Considering all the findings from the α-Al₂O₃ (0001) studies, it is clear that direct comparisons of computed energetic values cannot be easily made unless the same computational models are implemented and similar considerations for relaxations and size effects are taken into account, as
is done in this work.

Table 7.3 lists a sample of prior atomistic studies on select $\alpha$-$\text{Al}_2\text{O}_3$ surfaces along with computed surface energies. For qualitative comparisons among the studies, relative surface energies are reported as a percentage value based on the ratio of the computed surface energy versus the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) surface. While the relative stability of the various $\alpha$-$\text{Al}_2\text{O}_3$ surfaces deviates in early studies, the later HF results by Marmier and Parker [70] and DFT-GGA results by Sun et al. [84] agree that the ordering of surface energies for $\alpha$-$\text{Al}_2\text{O}_3$ follows: $(0001)_{\alpha} < (10\overline{1}2)_{\alpha} < (11\overline{2}0)_{\alpha} < (10\overline{1}0)_{\alpha} < (10\overline{1}1)_{\alpha}$, indicating the basal surface as most stable.

Only a few prior atomistic studies have examined the structure and energetics of surfaces in metastable alumina phases. Table 7.4 lists a sample of these studies along with computed surface energies. Of these, only one study by Blonski and Garofalni [36] used the same computational model to examine multiple surfaces in both $\alpha$-$\text{Al}_2\text{O}_3$ and a metastable phase. Blonski and Garofalni utilized a many-body potential to study select $\alpha$- and $\gamma$-$\text{Al}_2\text{O}_3$ surfaces [36]. The many-body potential modeled pair interactions with a Born-Mayer Huggins form [85], three-body interactions for Al-O-Al and O-Al-O triplets, and electrostatic interactions using fixed point charges. In their work, Blonski and Garofalni showed that the potential successfully predicted minimum energy structures for bulk $\alpha$- and $\gamma$-$\text{Al}_2\text{O}_3$ when compared to experimental radial distribution functions. Additionally, the $\alpha$-$\text{Al}_2\text{O}_3$ surface energy calculations predicted the ordering of surface energies to be $(0001)_{\alpha} < (11\overline{2}0)_{\alpha} < (10\overline{1}0)_{\alpha}$ similar to the more recent first-principles results [70,84]. Differently, the predicted ordering of $\gamma$-$\text{Al}_2\text{O}_3$ surface energies by Blonski and Garofalni [36] contradicts more recent DFT-GGA values computed by Pinto and Elliot [55]. The DFT-GGA study [55] predicted the ordering of $\gamma$-$\text{Al}_2\text{O}_3$ surface energies as $(111)_{\gamma} < (001)_{\gamma} < (110)_{\gamma} < (150)_{\gamma}$, indicating the $(111)_{\gamma}$ surface as most stable.
Table 7.3: Sample of previous $\alpha$-$\text{Al}_2\text{O}_3$ surface energies computed using atomistic simulation methods. All values are in J/m$^2$. The values in parenthesis are for non-relaxed surface structures. For relaxed values, relative energy compared to the $(0001)_\alpha$ surface energy are reported.

<table>
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<tr>
<th>Year</th>
<th>Comp. Model</th>
<th>$(0001)_\alpha$</th>
<th>$(10\bar{1}2)_\alpha$</th>
<th>$(1\bar{1}20)_\alpha$</th>
<th>$(10\bar{1}1)_\alpha$</th>
<th>$(1\bar{1}0\bar{1})_\alpha$</th>
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<td>(2.94) 2.14</td>
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<td>129%</td>
<td>137%</td>
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<td>(4.45) 2.65</td>
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<td>(2.77) 1.82</td>
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<td>145%</td>
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<tr>
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<td>(3.49) 2.27</td>
<td>---</td>
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<td>HF</td>
<td>6.72 (5.32)</td>
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<td>---</td>
<td>---</td>
<td>(5.65) 5.59</td>
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<td>(3.63) 2.29</td>
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<td>1980</td>
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<td>4.83</td>
<td>4.8</td>
<td>3.8</td>
<td>2.55</td>
<td>---</td>
<td>[88]</td>
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$^a$ Pair Style – Charged Shell  
$^b$ Pair Style – Point Charge  
$^c$ Many-body Style – Geometry Dependent  
$^d$ Many-body Style – Point Charge
Table 7.4: Sample of computed surface energies from previous atomistic simulations of metastable alumina. Subscripts indicate the alumina phase and unrelaxed surface energy values are in denoted by parenthesis.

<table>
<thead>
<tr>
<th>Year</th>
<th>Comp. Model</th>
<th>Metastable Alumina Surface Energies [J/m²]</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2004</td>
<td>DFT-GGA</td>
<td>(2.97) 1.05 (3.43) 1.53 (1.62) 0.95 (2.79) 1.91</td>
<td>[55]</td>
</tr>
<tr>
<td>2002</td>
<td>DFT-GGA</td>
<td>0.97 1.54 --- ---</td>
<td>[89]</td>
</tr>
<tr>
<td>1993</td>
<td>MS&lt;sup&gt;a&lt;/sup&gt;</td>
<td>(3.37) 0.79 (4.62) 1.21 (9.45) 0.87</td>
<td>[90]</td>
</tr>
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<td>2004</td>
<td>DFT-GGA</td>
<td>(0.70) (1.65) 1.00 ---</td>
<td>[91]</td>
</tr>
<tr>
<td>1994</td>
<td>HF</td>
<td>3.88 --- --- 4.41</td>
<td>[61]</td>
</tr>
<tr>
<td>2003</td>
<td>DFT-GGA</td>
<td>(7.00) 5.20 (7.00) 5.80</td>
<td>[72]</td>
</tr>
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</table>

<sup>a</sup> Many-body Style – Point Charge

Table 7.4 also lists previously computed surface energy values for a selection of θ- and κ-Al<sub>2</sub>O<sub>3</sub> surfaces. While similar DFT-GGA models were used in these calculations, the authors did not match basis sets or use similar boundary conditions necessary for quantitative comparison with the other studies. Thus, direct comparison of the computed surface energy values among the various studies is not possible. However, the reported data do indicate the lowest energy surfaces for θ- and κ-Al<sub>2</sub>O<sub>3</sub> are the (100)<sub>θ</sub> and (001)<sub>κ</sub> surfaces, respectively.

It is important to note that the previously discussed studies were conducted for clean alumina surfaces. In experimental settings, alumina surfaces exposed to water can become hydroxylated. Several atomistic studies have examined hydroxylated alumina surfaces [91–96] and reported that hydroxylation increased the stability of alumina surfaces. For example, Łodziana et al. [91] computed negative surface energies for the (110)<sub>θ</sub> surface in θ-Al<sub>2</sub>O<sub>3</sub> caused by the generation of a new, more porous equilibrium surface structure that exhibits a lower energy than bulk θ-Al<sub>2</sub>O<sub>3</sub>. 

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7.3 Methods

7.3.1 Simulation Details

Atomistic simulations of alumina are performed with LAMMPS [97] using the ReaxFF potential [26]. ReaxFF is chosen for this work because of its hypothesized transferability to many alumina phases due to its use of a dynamic bond order term and geometry dependent charge optimization. Specifically, ReaxFF computes van der Waals and Coulombic contributions to the potential energy between all atom pairs and implements a continuous bond order term to allow dynamic bond formation and breaking. Additionally, ReaxFF uses an electron equilibration method [98] to optimize atomic charges based on the atomic geometry at each time step. Dynamic charge optimization has the potential to increase the transferability of ReaxFF by modelling different oxidation states within different alumina phases. The ReaxFF parameters used in this study were first optimized by Zhang et al. for Al and $\alpha$-$\text{Al}_2\text{O}_3$ using first-principles simulations of Al-O clusters [52] then further tailored by Sen et al. to better incorporate varying Al oxidation states during oxidation simulations [51].

Slab models are created for the 11 surfaces described in Table 7.5. Each slab model is constructed using fully-periodic boundary conditions with a vacuum region normal to the surface greater than 50 Å, as shown in Figure 7.1. For each model, the simulation dimensions and terminating planes are chosen such that the correct stoichiometry of alumina is maintained. The simulation dimensions along the surface plane are chosen to minimize strain stemming from small misalignments of the periodic images created by the oriented crystals.

For each slab model, a series of initial structures are constructed by varying the terminating planes on each opposing surface. The initial structures are relaxed at 0 K using a non-linear conjugate gradient method. Surface energies for each structure are computed using two
Table 7.5: Description of the simulation size for alumina slabs modeled in this study. Here each subscript indicates a particular alumina phase associated with the surface plane.

<table>
<thead>
<tr>
<th>Surface</th>
<th>$L_x$ [Å]</th>
<th>$L_y$ [Å]</th>
<th>$L_z$ [Å]</th>
<th>N [atoms]</th>
</tr>
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<tbody>
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<td>$(0001)_\alpha$</td>
<td>201.2721</td>
<td>203.6963</td>
<td>194.9318</td>
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<td>$(1120)_\alpha$</td>
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<td>195.8093</td>
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<td>$(1\overline{1}00)_\alpha$</td>
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</tr>
<tr>
<td>$(001)_\gamma$</td>
<td>198.9950</td>
<td>199.0289</td>
<td>189.7136</td>
<td>800,000</td>
</tr>
<tr>
<td>$(110)_\gamma$</td>
<td>203.6746</td>
<td>190.2056</td>
<td>202.9786</td>
<td>829,440</td>
</tr>
<tr>
<td>$(111)_\gamma$</td>
<td>202.1025</td>
<td>175.9054</td>
<td>207.5978</td>
<td>777,600</td>
</tr>
<tr>
<td>$(001)_\kappa$</td>
<td>197.3578</td>
<td>201.6267</td>
<td>198.2140</td>
<td>844,800</td>
</tr>
<tr>
<td>$(010)_\kappa$</td>
<td>361.5524</td>
<td>118.2935</td>
<td>184.9249</td>
<td>844,800</td>
</tr>
<tr>
<td>$(100)_\kappa$</td>
<td>336.2477</td>
<td>217.3695</td>
<td>107.7260</td>
<td>844,800</td>
</tr>
<tr>
<td>$(001)_\theta$</td>
<td>201.3005</td>
<td>201.0150</td>
<td>194.7824</td>
<td>844,560</td>
</tr>
<tr>
<td>$(110)_\theta$</td>
<td>158.9278</td>
<td>191.0272</td>
<td>161.4515</td>
<td>512,085</td>
</tr>
</tbody>
</table>

Figure 7.1: Schematic of slab model used in atomistic simulations of alumina surfaces showing the thermostat regions and placement of Gibb's dividing surfaces.
methods. The first method considers the entire slab and computes an average surface energy, \( \gamma^\text{ave} \), accounting for both the upper and lower surfaces via,

\[
\gamma^\text{ave} = \frac{E_{\text{Slab}} - nE_{\text{Bulk}}}{2A}.
\]  

(7.1)

Here, \( E_{\text{Slab}} \) is the minimized potential energy for a particular slab, \( E_{\text{Bulk}} \) is the minimized potential energy for the bulk alumina unit cell of the matching phase, \( n \) is the number of alumina unit cells contained within the slab, and \( A \) is the area of the slab surface. Note, the average surface energy \( \gamma^\text{ave} \) for a slab model with identical surface structures will be equivalent to the surface energy of the upper and lower surfaces. However, for complex non-symmetric alumina phases there is no guarantee the surface structures on opposing sides of the slab will be identical. Therefore, a second method of computing surface energies is employed that uses Gibbs dividing surfaces to divide the slab into upper and lower stoichiometric regions [99]. The Gibbs dividing surfaces are aligned parallel to the slab surfaces and are positioned within the slab through an iterative process that identifies stoichiometric regions exposed to the vacuum. The iterative process places an initial Gibbs dividing surface in the middle of the slab and steps normal to the desired surface until the atoms within the region exposed to the vacuum are stoichiometric. Surface energies for the upper \( \gamma^U_s \) and lower \( \gamma^L_s \) regions are computed via,

\[
\gamma^U/L_s = \frac{E_{\text{Region}} - mE_{\text{Bulk}}}{A}
\]  

(7.2)

where \( E_{\text{Region}} \) and \( m \) are the potential energy and equivalent number of unit cells contained within the Gibbs region, respectively. Surface energies are computed using both methods for each series of initial structures to determine the lowest energy configuration for each slab model.
For low-energy degenerate structures, slab models with the smallest deviation between upper and lower surface energies calculated from Eq. (7.2) are selected for the MD study as this indicates the presence of similar surface structures.

Molecular dynamics simulations are performed on the lowest-energy slab models for each surface using the canonical (NVT) ensemble at 300, 500, and 700 K. The MD simulations use a 0.5 fs time step with the Hoover damping parameter set to 100 fs. The NVT ensemble is chosen in order to decrease the time for equilibration and is sufficient for alumina models because of their low coefficient of thermal expansion [42]. To avoid localized temperature fluctuations near the surfaces caused by extra degrees of freedom experienced by atoms exposed to the vacuum, the slab models are divided into three NVT thermostat regions. Two thermostat regions are defined for the upper and lower surface regions by grouping all atoms exposed to the surfaces that are located within 25% of the direction normal to the surface (Lz) while the remaining atoms are grouped into the third thermostat region. Each region is coupled to a unique external thermal reservoir held at the desired temperature.

The MD simulations for each slab are performed at each temperature for 25 ps and equilibration is monitored by reporting the total potential energy every 0.05 ps. Energetic data are extracted over the last 5 ps of the simulation when the maximum deviation of the potential energy is less 0.1% of the average value. Surface energies are computed for each slab at each temperature via Eqs. (7.1) and (7.2) using the equilibrated potential energy values for the slab and from bulk alumina simulations equilibrated using the same procedures as discussed above.

**7.3.2 Virtual Diffraction Analysis**

Characterization of alumina atomistic simulations proves difficult due to the subtle structural differences observed among the phases. While traditional characterization methods used for
atomistic simulation results (i.e., radial distribution functions [100], centrosymmetry [101], common neighbor-analysis [102]) are successful in analyzing cubic or other high-symmetry crystal structures, none of these methods are capable of uniquely identifying alumina phases due to the subtle differences in the Al ion placement within the non-cubic crystal structures. Therefore, in this work, virtual diffraction methods developed by Coleman et al. [27,28] are chosen to identify the alumina models and characterize the stable and metastable alumina surfaces.

The virtual diffraction method developed by Coleman et al. [27,28] uses kinematic diffraction theory to create both SAED patterns and XRD line profiles from atomistic simulation data. Previously, other virtual diffraction methods based on kinematic diffraction theory have studied atomistic simulations of (i) grain boundary structures in bicrystals [103–108], (ii) grain size and microstrain in nanocrystals [109–114], (iii) defect formation and evolution during shock of nanocrystals [115–118], and (iv) bulk alumina simulations [2,64]. The virtual diffraction method developed by Coleman et al. [27,28] advances beyond these prior studies because it generates both SAED patterns and XRD line profiles without requiring a priori knowledge of the crystal structure. Additionally, the method by Coleman et al. takes advantage of modern computer hardware and external accelerators in order to directly compute the Fourier components in the structure factor equation [119]. The algorithm is generic for all atom species and is integrated into LAMMPS [97] which leverages multi-level parallelization techniques to rapidly compute diffraction intensities for large atomistic simulations [119].

Briefly, the method introduced by Coleman et al. [27,28] computes diffraction intensities, \( I(\mathbf{K}) \), for N atoms as the product of the structure factor, \( F(\mathbf{K}) \), with its complex conjugate, \( F^*(\mathbf{K}) \), via
\[ I(K) = \frac{F^*(K)F(K)}{N} \]  

where \( F(K) \) is computed by the structure factor equation [120],

\[ F(K) = \sum_{j=1}^{N} f_j \exp(2\pi i \mathbf{K} \cdot \mathbf{r}_j). \]  

Here, \( \mathbf{K} \) is the location of the diffraction peak in reciprocal space, \( \mathbf{r}_j \), is the position of the atom in real space, and \( f_j \) is the atomic scattering factor computed for each atom species.

Diffraction intensities are explored across a high-resolution, three-dimensional reciprocal space mesh that is constructed either based on the dimensions of the entire simulation or as a user defined value. The resolution of the reciprocal space mesh is tuned in order to ensure important peak locations are sampled. In this work, all atoms within the simulation box (except for select surface analyses as denoted) are used to compute the diffraction intensities. Reciprocal lattice nodes are spaced 0.00475 Å\(^{-1}\) and 0.02 Å\(^{-1}\) in each direction to generate SAED and XRD patterns, respectively. SAED patterns and XRD line profiles are constructed from the three-dimensional intensity data using simulated 200 keV electron radiation (\( \lambda = 0.0251 \ \text{Å} \) [121]) and Cu K\( \alpha \) x-rays (\( \lambda = 1.54178 \ \text{Å} \) [120]) by applying unique data analysis and visualization methods developed for each technique [27].

### 7.4 Results and Discussion

#### 7.4.1 Bulk Alumina

The transferability of the ReaxFF potential is evaluated by modeling bulk \( \gamma-, \kappa-, \theta-, \) and \( \alpha- \) \( \text{Al}_2\text{O}_3 \) systems using MS and MD simulations. Potential energy values per \( \text{Al}_2\text{O}_3 \) unit extracted from these simulations are reported in Table 7.6. At all temperatures explored, the ReaxFF interatomic potential successfully predicts \( \alpha-\text{Al}_2\text{O}_3 \) as more energetically stable (i.e., lower potential energy) than any other crystalline phase. Table 7.6 also shows the difference in
potential energy per Al₂O₃ unit at 0 K for the metastable phases compared to α-Al₂O₃ computed using ReaxFF and for the most recent DFT-LDA study by Lee et al. [54]. ReaxFF predicts the metastable structures to be nearly energetically degenerate having potential energies each approximately 0.20 eV per Al₂O₃ unit greater than α-Al₂O₃. Lee et al. predict a similar magnitude for the energy deviation; however, their DFT-LDA results show the deviation for γ-Al₂O₃ to be nearly twice that as κ-Al₂O₃. The larger energy deviation in the prior DFT-LDA study could be the result of a different γ-Al₂O₃ initial structure as compared to that used in the current study [64,65].

A series of MS simulations are performed exploring an isotropic expansion of the α-Al₂O₃ lattice parameters to investigate the minimum energy lattice structure and to compute the predicted bulk elastic modulus. To calculate the bulk modulus, the minimized potential energies are fitted to the Birch-Murnaghan equation of state [122]. As seen in Table 7.7, the resulting minimum energy lattice parameters and the predicted bulk modulus closely match both experimental and first-principles computed values. Specifically, the a and c lattice constants calculated by ReaxFF for this work deviate from the reported experimental values by only 0.1% while the bulk modulus deviates by 2.8%.
Due to the complexity of the metastable alumina phases, structural properties for these phases are unable to be computed using the same procedure. However, XRD patterns computed for each energy-minimized structure are shown in Figure 7.2. The virtual XRD line profiles

![XRD Profiles](image)

**Figure 7.2:** Computed XRD profiles (colored) compared to experimental references [4,11,129,130] (black) for various energy minimized phases using the ReaxFF potential.

**Table 7.7:** Computed properties for $\alpha$-Al$_2$O$_3$. Prior experimental data were observed at high pressure and 300 K.

<table>
<thead>
<tr>
<th>$\alpha$-Al$_2$O$_3$</th>
<th>Expt.</th>
<th>DFT-GGA</th>
<th>ReaxFF</th>
<th>This Work</th>
</tr>
</thead>
<tbody>
<tr>
<td>a [Å]</td>
<td>4.758$^a$</td>
<td>4.783$^c$</td>
<td>4.810$^c$</td>
<td>4.81</td>
</tr>
<tr>
<td>c [Å]</td>
<td>12.995$^a$</td>
<td>13.252$^c$</td>
<td>13.100$^c$</td>
<td>13.13</td>
</tr>
<tr>
<td>B [Gpa]</td>
<td>253$^b$</td>
<td>250$^b$</td>
<td>248$^c$</td>
<td>244</td>
</tr>
</tbody>
</table>

$^a$ Reference [123]

$^b$ Reference [124]

$^c$ Reference [52]
uniquely identify each alumina phase. In addition, the virtual XRD line profiles (colored) show excellent agreement with experimental powder diffraction data (black). The ability of ReaxFF to predict the expected energetics and correct structures suggests good transferability of this potential to different alumina phases at 0 K.

### 7.4.2 Alumina Surfaces

Average surface energies evaluated at 0, 300, 500, and 700 K for each of the 11 alumina slab models are listed in Table 7.8. Among the $\alpha$-Al$_2$O$_3$ surfaces, ReaxFF predicts the (0001)$_\alpha$ surface to be the lowest energy surface at all temperatures, which is consistent with previous HF [70] and DFT-GGA results [84]. Among the $\gamma$-Al$_2$O$_3$ surfaces, ReaxFF predicts the (001)$_\gamma$ surface to be the lowest surface energy at 0 and 700 K; however, at 300 and 500 K, the ReaxFF simulations predict an even lower energy for the (110)$_\gamma$ surface. The 0 K results for $\gamma$-Al$_2$O$_3$ are consistent with previous MD simulation results [90] but disagree with results found by DFT-GGA [55], which found the lowest surface to be the (111)$_\gamma$ surface. Among the $\kappa$-Al$_2$O$_3$ surfaces, ReaxFF predicts the (010)$_\kappa$ surface to be the lowest surface energy at 0, 500, and 700 K; however, at 300 K, the ReaxFF simulations predict an even lower surface energy for the (001)$_\kappa$ surface. Interestingly, a negative average surface energy is found for the (010)$_\kappa$ surface at 700 K indicating the presence of a lower-energy surface structure within the slab model as compared to the bulk. Negative values for the surface energies are also found for both $\theta$-Al$_2$O$_3$.

<table>
<thead>
<tr>
<th>Temp.</th>
<th>$\alpha$-Al$_2$O$_3$</th>
<th>$\gamma$-Al$_2$O$_3$</th>
<th>$\kappa$-Al$_2$O$_3$</th>
<th>$\theta$-Al$_2$O$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(0001) (1100) (1120)</td>
<td>(001) (110) (111)</td>
<td>(001) (010) (100)</td>
<td>(001) (110)</td>
</tr>
<tr>
<td>0 K</td>
<td>0.749 1.756 1.767</td>
<td>2.166 2.897 2.826</td>
<td>0.434 0.189 0.773</td>
<td>-0.187 -1.948</td>
</tr>
<tr>
<td>300 K</td>
<td>0.768 1.505 1.524</td>
<td>1.853 1.131 1.456</td>
<td>0.238 0.294 0.604</td>
<td>-0.101 -1.012</td>
</tr>
<tr>
<td>500 K</td>
<td>0.805 1.174 1.158</td>
<td>1.751 1.254 1.574</td>
<td>0.786 0.254 0.620</td>
<td>-1.174 -0.812</td>
</tr>
<tr>
<td>700 K</td>
<td>0.913 0.948 0.961</td>
<td>0.507 0.536 1.025</td>
<td>0.574 -0.638 0.849</td>
<td>0.078 -1.946</td>
</tr>
</tbody>
</table>

Table 7.8: Computed average surface energy values, Eq. (7.1), for select alumina surfaces using the ReaxFF potential [60]. Values are reported in J/m$^2$. 

156
surface structures at 0, 300, and 500 K and for the (110)₀ surface at 700 K. Between the θ-Al₂O₃ surfaces, ReaxFF predicts lower surface energy for the (110)₀ at 0, 300, and 700 K. The upper and lower surface energy values computed using Eq. (7.2) are tabulated in the supplementary materials. Of the 11 alumina slab model studied, only the (001)γ, (111)γ, and (001)κ surfaces showed significant deviation between the computed upper and lower surface energy values due to different terminating structures.

Figure 7.3 shows virtual SAED patterns observed normal to the α-Al₂O₃ (0001) and γ-Al₂O₃ (110) surface after the 0 K energy minimization as well as comparisons to prior experimental results obtained by Heffelfinger et al. [125] and Morrissey et al. [8], respectively. For visualization, low diffraction intensity data in the virtual SAED patterns are removed so that peak locations become more evident. The virtual SAED results show distinct diffraction peaks as well as regular vertical and horizontal relrod structures due to the finite volume of the simulation [27]. For the α-Al₂O₃ (0001) surface, the virtual SAED results clearly show the same hexagonal pattern as observed in the experimental results. For the γ-Al₂O₃ (110) surface, the underlying shape of the diffraction peaks is consistent with the experimental results; however many low-intensity diffraction peaks are evident in the virtual SAED pattern which are not observed experimentally. These low-intensity peaks arise from a regular placement of Al interstitials within the slab created by repeating the γ-Al₂O₃ unit cell when constructing the model. This periodicity of the Al interstitials is especially pronounced in the static structure. In general, the one-to-one comparison between experimental and virtual SAED patterns, shown in Figure 7.3, validate the computational methods used and provide confidence in the virtual SAED results when no experimental comparisons are available.
The nanoscale structure of the lowest average surface energy models (at 0 K) for each phase are characterized using virtual SAED patterns as well as by analyzing snapshots of the atomic positions within an approximate 2 x 2 x 1.25 nm$^3$ region near the upper and lower surfaces. Characterization results from all models are contained in the supplementary materials. At temperature, time averaged virtual SAED patterns are constructed by averaging diffraction data taken every 2.5 ps over the entire 25 ps equilibration period.

Figure 7.4 shows the characterization results of the (0001) surface in α-Al$_2$O$_3$ at 0, 300, 500, and 700 K. Here, both the SAED and atomic snapshots are viewed along the [1012] direction. The upper and lower (0001)$_a$ surfaces at 0, 300, and 500 K are cleaved with Al terminations that have displaced inwardly from their bulk configurations into the slab. However, at 700 K the (0001)$_a$ surface experiences a reconstruction in which clusters of Al and O ions protruding from
the surface. At 0, 300, and 500 K, smearing of the diffraction peaks is minimal which is consistent with the lack of surface reconstructions at these temperatures; however, at 700 K, peak smearing occurs at the higher (hkl) peak locations due to the surface reconstruction.

Characterization results for the (001) surface in γ-Al₂O₃ at 0, 300, 500, and 700 K are shown in Figure 7.5 viewed along the [010] direction. At 0, 300, and 500 K, O ions terminate a nearly cleaved upper surface while the Al ions terminating the lower surface extend slightly into the vacuum region. At these temperatures, the Al terminated surface energy is approximately 1.4

<table>
<thead>
<tr>
<th>0 K</th>
<th>300 K</th>
<th>500 K</th>
<th>700 K</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="0 K" /></td>
<td><img src="image2" alt="300 K" /></td>
<td><img src="image3" alt="500 K" /></td>
<td><img src="image4" alt="700 K" /></td>
</tr>
<tr>
<td><img src="image5" alt="0 K" /></td>
<td><img src="image6" alt="300 K" /></td>
<td><img src="image7" alt="500 K" /></td>
<td><img src="image8" alt="700 K" /></td>
</tr>
<tr>
<td><img src="image9" alt="0 K" /></td>
<td><img src="image10" alt="300 K" /></td>
<td><img src="image11" alt="500 K" /></td>
<td><img src="image12" alt="700 K" /></td>
</tr>
<tr>
<td><img src="image13" alt="0 K" /></td>
<td><img src="image14" alt="300 K" /></td>
<td><img src="image15" alt="500 K" /></td>
<td><img src="image16" alt="700 K" /></td>
</tr>
<tr>
<td><img src="image17" alt="0 K" /></td>
<td><img src="image18" alt="300 K" /></td>
<td><img src="image19" alt="500 K" /></td>
<td><img src="image20" alt="700 K" /></td>
</tr>
</tbody>
</table>

Figure 7.4: Characterization results for the α-Al₂O₃ (0001) surface viewed along the [10\bar{1}2] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
than the O terminated surface. The 500 K surfaces show minimal surface relaxations in the outer layers. However, at 700 K both upper and lower surfaces experience significant reconstructions extending substantially into the slab models, which reduces the deviation between the upper and lower surface energies to 0.8 J/m$^2$. Virtual SAED patterns of the cleaved surfaces at 0 and 300 K show minimal peak smearing. However, relaxations and reconstructions at 500 and 700 K result in increased smearing of diffraction peaks as compared with the 700 K (0001)$_\alpha$ surface in Figure 7.4.

Figure 7.5: Characterization results for the $\gamma$-Al$_2$O$_3$ (001) surface viewed along the [010] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.6 shows characterization results for the $\kappa$-Al$_2$O$_3$ (010) surface viewed along the [100] direction. At 0 and 300 K both upper and lower surfaces form a regular concave structure with terminating O ions. Similar to the (001)$_\gamma$ surfaces, at 500 K the outermost layers of the (010)$_\kappa$ surfaces are relaxed, whereas substantial surface reconstruction is seen at 700 K. At 700 K, the surface reconstruction extends far beneath the surface of the slab creating a lower-energy configuration as compared to the bulk, which results in the negative average surface energies reported in Table 7.8. The virtual SAED patterns for (010)$_\kappa$ surfaces at
Characterization results for the (110) surface in $\theta$-Al$_2$O$_3$ at 0, 300, 500, and 700 K are shown in Figure 7.7 viewed along the [001] direction. At 0 and 300 K, the outer layers of the surfaces are relaxed with a mixture of O and Al terminating ions. Substantial surface reconstructions are observed at 500 and 700 K appearing almost non-crystalline in nature. However, distinct diffraction peaks within the virtual SAED patterns indicate that some crystallinity is preserved at 0, 300, and 500 K show no significant smearing of the diffraction peaks; however at 700 K, substantial peak smearing is evident.

Figure 7.7: Characterization results for the $\theta$-Al$_2$O$_3$ (110) surface viewed along the [001] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
all temperatures. With increasing temperature, certain diffraction peaks disappear and new peaks emerge while smearing of existing peaks becomes more substantial. To further investigate the negative surface energy computed for (110)$_{θ}$ surfaces, slab models are analyzed using the virtual XRD patterns, as shown in Figure 7.8. XRD patterns produced from 0 K energy minimized slabs as well as at each temperature do not match the predicted or experimental XRD patterns for bulk $θ$-$Al_2O_3$, thus showing ReaxFF predicts a substantially different structure for (110)$_{θ}$ slab models. XRD profiles for the (110)$_{θ}$ slab models at 0, 500, and 700 K show the strongest reflection at $2θ = 31.65^\circ$ corresponding to an interplanar spacing of 2.83 Å. Additionally, diminishing peaks at larger $2θ$ values suggest a transformation towards an amorphous configuration.

![XRD Patterns](image_url)

**Figure 7.8:** Virtual XRD patterns computed for bulk $θ$-$Al_2O_3$ and slab models of the (110)$_{θ}$ surface at varying temperatures which show the prediction of a new low-energy phase in the slab modeled by the ReaxFF potential.
ReaxFF predicts substantial reconstructions of several alumina surfaces at temperatures below what is expected from experimental studies (c.f. [126–128]). To investigate the cause for the premature reconstructions, amorphous bulk alumina is modeled using ReaxFF. Samples of bulk amorphous alumina are constructed by heating models of each crystalline phase to 3,000 K using a NPT ensemble for 25 ps then cooling the amorphous melt to 30 K over an additional 25 ps. This is followed by a non-linear conjugate gradient minimization of the solidified amorphous system. Virtual SAED diffraction analysis performed on each relaxed amorphous structure shows similar diffuse ring patterns, as expected for an amorphous material, which indicates the final structures are independent of the original phase selected. Similarly, virtual XRD profiles show a single strong reflection at $2\theta = 25.57^\circ$ corresponding to an interplanar spacing of 3.48 Å. A representative amorphous structure is equilibrated to each of the three temperatures via the NVT equilibration procedure described in Section 3.

Unexpectedly, at all temperatures explored ReaxFF predicts the potential energy of the amorphous alumina phase to be approximately 0.10 eV per $\text{Al}_2\text{O}_3$ unit lower than $\alpha$-$\text{Al}_2\text{O}_3$. Therefore, it is believed that ReaxFF unnaturally biases surface reconstruction within slab models to a lower energy amorphous-like configuration, especially due to the extra degrees of freedom available to the exposed surfaces. Note, virtual SAED patterns constructed only using the reconstructed surface regions reveal distinct (but smeared) peaks signifying that these regions are not fully amorphous. It is believed the lattice constraints imposed by the bulk interior of the slab models prevents the formation of fully amorphous reconstructed surface structures. In addition, it is believed that the bias for reconstruction primarily affects surfaces modeled at temperatures when the available thermal energy increases the ability for transformation into a different surface configuration. Therefore, 0 K results from this study are believed to be more
appropriate for use in future predictive mesoscale simulations.

7.5 Conclusions

In this work, select bulk and surface alumina structures are modeled using the ReaxFF interatomic potential in order to characterize their nanoscale structure and to compute bulk and surface energies. In comparison to prior efforts, this work provides the widest study to date on alumina bulk and surface structures within a consistent computational model. ReaxFF accurately predicts bulk $\alpha$-$\text{Al}_2\text{O}_3$ as the lowest energy crystalline phase; however, it also predicts an even lower energy amorphous phase. Virtual XRD patterns computed for each bulk crystalline structure uniquely identify alumina phases and are validated by experimental data.

Surface energies computed using ReaxFF of many alumina phases are consistent with prior atomistic simulation results. However, results from certain surfaces significantly deviate from prior experimental and atomistic results. For example, ReaxFF predicts negative surface energies for the $\theta$-$\text{Al}_2\text{O}_3$ surfaces due to the creation of a new lower-energy $\text{Al}_2\text{O}_3$ surface structure within the slab models as determined by virtual XRD patterns. Several alumina surfaces also experience significant surface reconstructions at temperatures below what is expected from experiments. It is believed the premature surface reconstruction is biased by the lower energy amorphous phase predicted by ReaxFF. Free surfaces enable the premature surface reconstructions due to the extra degrees of freedom available, which is especially true during MD simulations when thermal energy increases the ability for the transformation to occur.

Because a consistent computational model is employed throughout this study, the computed energies can be quantitatively compared and used in future predicted mesoscale models of vapor deposition for polymorphic alumina. It is believed that the limitations found for the ReaxFF potential primarily affect simulations of surface properties at temperature. Therefore, data
extracted at 0 K is the most appropriate for use in predictive mesoscale models. A complete mesoscale model of vapor deposition for polymorphic alumina will require future work to investigate the structures and energetics of homogenous and heterogeneous (solid/solid) alumina interfaces. For quantitative comparison with data in this study, it is essential that this future work be conducted using the same computational model employed here.

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Appendix 7.1


I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication.

Sincerely,

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Dissertation Director
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University of Arkansas
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Office: 479-575-3040
E-mail: dspearot@uark.edu
Appendix 7.2

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Appendix 7.3: Supplementary Materials:

7.3.1 Local Surface Energies

Computed surface energies for the upper $\gamma_{s}^{U}$ and lower $\gamma_{s}^{L}$ regions are evaluated through

$$\gamma_{s}^{U/L} = \frac{E_{\text{Region}} - mE_{\text{Bulk}}}{A}$$

in the manner discussed in the text are reported in Table 7.9.

<table>
<thead>
<tr>
<th></th>
<th>$\alpha$-Al$_2$O$_3$</th>
<th>$\gamma$-Al$_2$O$_3$</th>
<th>$\kappa$-Al$_2$O$_3$</th>
<th>$\theta$-Al$_2$O$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>0 K</strong></td>
<td>(0001) (11\overline{1}0) (11\overline{2}0)</td>
<td>(001) (110) (111)</td>
<td>(001) (010) (100)</td>
<td>(001) (110)</td>
</tr>
<tr>
<td>Upper</td>
<td>0.749 1.754 1.721</td>
<td>2.881 2.800 1.629</td>
<td>0.199 0.189 0.773</td>
<td>-0.181 -1.936</td>
</tr>
<tr>
<td>Lower</td>
<td>0.749 1.756 1.794</td>
<td>1.450 2.936 3.730</td>
<td>0.701 0.189 0.773</td>
<td>-0.194 -1.942</td>
</tr>
<tr>
<td><strong>300 K</strong></td>
<td>(0001) (1\overline{1}00) (11\overline{2}0)</td>
<td>(001) (110) (111)</td>
<td>(001) (010) (100)</td>
<td>(001) (110)</td>
</tr>
<tr>
<td>Upper</td>
<td>0.771 1.496 1.482</td>
<td>2.479 0.940 0.361</td>
<td>-0.471 0.293 0.600</td>
<td>-0.101 -0.997</td>
</tr>
<tr>
<td>Lower</td>
<td>0.765 1.510 1.548</td>
<td>1.226 1.337 2.453</td>
<td>0.969 0.295 0.607</td>
<td>-0.101 -1.101</td>
</tr>
<tr>
<td><strong>500 K</strong></td>
<td>(0001) (1\overline{1}00) (11\overline{2}0)</td>
<td>(001) (110) (111)</td>
<td>(001) (010) (100)</td>
<td>(001) (110)</td>
</tr>
<tr>
<td>Upper</td>
<td>0.806 1.169 1.116</td>
<td>2.487 1.033 0.307</td>
<td>-0.002 0.243 0.615</td>
<td>-1.176 -0.810</td>
</tr>
<tr>
<td>Lower</td>
<td>0.803 1.176 1.180</td>
<td>1.015 1.529 2.662</td>
<td>1.567 0.264 0.625</td>
<td>-1.172 -0.803</td>
</tr>
<tr>
<td><strong>700 K</strong></td>
<td>(0001) (1\overline{1}00) (11\overline{2}0)</td>
<td>(001) (110) (111)</td>
<td>(001) (010) (100)</td>
<td>(001) (110)</td>
</tr>
<tr>
<td>Upper</td>
<td>0.908 0.935 0.915</td>
<td>0.900 0.333 -0.408</td>
<td>0.135 -0.626 0.863</td>
<td>0.121 -1.957</td>
</tr>
<tr>
<td>Lower</td>
<td>0.918 0.959 0.984</td>
<td>0.114 0.800 2.342</td>
<td>0.996 -0.650 0.835</td>
<td>0.034 -1.918</td>
</tr>
</tbody>
</table>
7.3.1 Characterization Results

The characterization results for alumina surfaces studied but not reported in the text are shown here in the supplementary materials.

![Characterization results for the α-Al₂O₃ (1100) surface viewed along the [0001] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.](image)
Figure 7.10: Characterization results for the $\alpha$-Al$_2$O$_3$ (1120) surface viewed along the [0001] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.11: Characterization results for the $\gamma$-Al$_2$O$_3$ (110) surface viewed along the $\{110\}$ direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.12: Characterization results for the $\gamma$-Al$_2$O$_3$ (111) surface viewed along the [011] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.13: Characterization results for the $\kappa$-Al$_2$O$_3$ (001) surface viewed along the [100] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.14: Characterization results for the κ-Al₂O₃ (100) surface viewed along the [010] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns.
Figure 7.15: Characterization results for the $\theta$-Al$_2$O$_3$ (001) surface viewed along the [010] direction showing snapshots of the (a-d) upper and (e-h) lower surfaces as well as (i-l) virtual SAED patterns. (Color online)
Chapter 8: Atomistic Simulation and Virtual Diffraction Characterization of Homophase and Heterophase Alumina Interfaces

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\textsuperscript{a}Department of Mechanical Engineering, University of Arkansas, Fayetteville, AR 72701

Abstract

The minimum energy structures of twelve homophase and heterophase alumina interfaces are studied using the ReaxFF potential. First, the computational methods are validated by exploring a set of five $\alpha$-$\text{Al}_2\text{O}_3$ symmetric tilt twin interfaces that have been studied in depth through prior simulations. The interface structures and energies for most homophase $\alpha$-$\text{Al}_2\text{O}_3$ twins are in good agreement with prior atomistic studies; however, small deviations from prior works occur in select $\alpha$-$\text{Al}_2\text{O}_3$ interfaces due to the larger interface areas explored in this work. Next, select experimentally observed $\kappa$-$\text{Al}_2\text{O}_3$, $\gamma$-$\text{Al}_2\text{O}_3$, and $\theta$-$\text{Al}_2\text{O}_3$ homophase interfaces as well as heterophase $\alpha$-$\text{Al}_2\text{O}_3$ // $\gamma$-$\text{Al}_2\text{O}_3$ and $\theta$-$\text{Al}_2\text{O}_3$ // $\gamma$-$\text{Al}_2\text{O}_3$ interfaces are investigated for the first time using atomistic simulations. The computed interface energies for the $\gamma$-$\text{Al}_2\text{O}_3$ \{111\} twin and the $\theta$-$\text{Al}_2\text{O}_3$ \{200\} twin interfaces are on the same order as that found for the lowest energy $\alpha$-$\text{Al}_2\text{O}_3$ prismatic twin boundary. ReaxFF predicts the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) // $\gamma$-$\text{Al}_2\text{O}_3$ (111) interface to have the lowest energy of the structures studied. Lastly, virtual selected area diffraction patterns of select interfaces are used to experimentally validate the modeled interface structures. Because a consistent computational method is implemented throughout this work, the tabulated interface energies can be incorporated in future predictive mesoscale simulations of polymorphic alumina.
8.1 Introduction

Alumina (Al$_2$O$_3$) films are used in a variety of applications, ranging from electronic and catalytic supports [1,2] to thermal barrier and protective coatings [3,4], due to the wide range of material properties exhibited by its various polymorphs. Deposition methods and processing conditions influence the microstructure and morphology of the alumina films, which influence the overall performance of the coatings. Both physical vapor deposition (PVD) and chemical vapor deposition (CVD) processes commonly result in the formation of mixtures of $\gamma$-Al$_2$O$_3$, $\theta$-Al$_2$O$_3$, $\kappa$-Al$_2$O$_3$, and $\alpha$-Al$_2$O$_3$ phases [5–13]. Generally, higher percentages of $\gamma$-Al$_2$O$_3$ and $\theta$-Al$_2$O$_3$ metastable phases form using PVD methods or at lower processing temperatures [5–8], whereas higher percentages of $\kappa$-Al$_2$O$_3$ and $\alpha$-Al$_2$O$_3$ phases form using CVD methods or at elevated temperatures [9–12]. The $\gamma$-Al$_2$O$_3$ and $\theta$-Al$_2$O$_3$ metastable phases are desired in catalytic support applications due to their high surface area and acid-basic properties [14,15]. Differently, $\kappa$-Al$_2$O$_3$ and $\alpha$-Al$_2$O$_3$ phases are desired in protective coating applications due to their wear resistance, chemical inertness, resistance to thermal shock, and high hardness [16,17]. Because $\alpha$-Al$_2$O$_3$ is the only thermodynamically stable alumina phase, researchers and manufacturers seeking protective coatings will often employ thermal treatments to induce the necessary phase transformations to form $\alpha$-Al$_2$O$_3$ [18,19]. Of course, the morphology of heat-treated films is dependent on the microstructure of the as-deposited polymorphic coatings [18,20].

As-deposited alumina films can be fully amorphous, nanocrystalline, or coarse-grained depending on the processing conditions used to create the coating [21]. Internal interfaces within polycrystalline alumina films can affect the properties and performance of the coating [22–26] as well as influence microstructure evolution during thermal treatment [27–29]. It is
known that abnormal grain growth in alumina occurs due to the presence of impurities and pores at the interfaces [30,31]. However, the effects of interface misorientation on grain growth in polycrystalline alumina films is not fully known [32–34], in particular for heterophase interfaces constructed from different adjoining alumina polymorphs. Despite not fully understanding the role of interfaces, researchers and manufacturers commonly create interfaces through deposition of multilayers which have been shown to be more wear resistant and easier to manufacture than single phase coatings [35,36].

Predictive mesoscale simulations of alumina vapor deposition can help determine the processing conditions needed to produce tailored film morphology and properties. To perform mesoscale simulations, such as phase-field modelling [37–39], extensive knowledge of alumina energetic data (i.e., bulk, surface, and interface energies) is needed. Atomistic simulations can easily compute the relevant energies required, but care must be taken to utilize the same computational method in order for these values to be used quantitatively. The current work utilizes the reactive force-field (ReaxFF) potential [40] to study homophase and heterophase alumina interfaces. An identical computational method has been used in a previous atomistic simulation study by the authors on bulk and surface alumina structures [41]. Together, these works tabulate the energetic data necessary to enable predictive mesoscale models to explore the unique processing conditions that achieve a desired tailored morphology and properties in polycrystalline and polymorphic alumina coatings.

This article begins with a detailed discussion of previous studies performed on alumina interfaces, focusing on prior atomistic simulations and the need for a consistent computational method to provide quantitatively comparable data. This is followed by a description of the current simulation and analysis methods imposed to extract interface energies and characterize
the atomic geometry near the alumina interfaces. Here, virtual diffraction methods [42,43] are implemented for the first time to aid in the construction of complex, heterophase alumina interfaces. Next, the computed interface energies are tabulated and reported alongside characterization results. Select alumina interfaces in this work are characterized using virtual selected area electron diffraction (SAED) patterns [42,43] that are directly comparable to experimental studies.

8.2 Previous alumina interface studies

Alumina interfaces have been studied using a variety of experimental and computational methods. The majority of previous atomistic simulation studies focused on five twin interfaces formed within \( \alpha\)-Al\(_2\)O\(_3\), as described in Table 8.1, using (i) molecular dynamics (MD), (ii) molecular statics (MS), (iii) density functional theory with local density approximations of the electron exchange-correlation (DFT-LDA), and (iv) density functional theory with generalized gradient approximations of the electron exchange-correlation (DFT-GGA). Each of these interfaces is created by a rotation of the basal plane about the \([1\bar{2}0]\) axis as shown schematically in Figure 8.1. Computed \( \alpha\)-Al\(_2\)O\(_3\) interface energies collected from previous atomistic simulations are listed in Table 8.2. Four previous research studies examined multiple \( \alpha\)-Al\(_2\)O\(_3\) interface structures using a consistent computational method within each study. These results

<table>
<thead>
<tr>
<th>ID</th>
<th>Interface Planes</th>
<th>Rotation</th>
<th>( \Sigma )</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(0001) // (0001)</td>
<td>0°</td>
<td>3</td>
<td>Basal Twin</td>
</tr>
<tr>
<td>B</td>
<td>(10\bar{1}0) // (10\bar{1}0)</td>
<td>90°</td>
<td>3</td>
<td>Prismatic Twin</td>
</tr>
<tr>
<td>C</td>
<td>(\bar{1}02) // (\bar{1}02)</td>
<td>56.9°</td>
<td>7</td>
<td>Rhombohedral Twin</td>
</tr>
<tr>
<td>D</td>
<td>(1\bar{1}4) // (1\bar{1}4)</td>
<td>142.5°</td>
<td>13</td>
<td>Pyramidal Twin</td>
</tr>
<tr>
<td>E</td>
<td>(10\bar{1}1) // (10\bar{1}1)</td>
<td>79.1°</td>
<td>11</td>
<td>General Twin</td>
</tr>
</tbody>
</table>

Table 8.1: Description of \( \alpha\)-Al\(_2\)O\(_3\) twin interfaces studied using atomistic simulation. Rotation is about the [1\(\bar{2}\)0]tilt axis and measured from the [10\(\bar{1}\)0] direction in the basal plane.
showed that the ordering of interface energy can depend on several factors, including the computational method. For example, molecular statistics simulations performed by Galmarini et al. predicted the lowest interface energy in the basal twin structure (A) whereas the first-principles study performed by Elsässer et al. [44–51] and the molecular dynamics studies performed by Suzuki et al. [52], predicted the prismatic twin structure (B) to have the lowest energy.

Deviations among the computed interface energies result partially from the different approximations used to model electrostatic interactions. The different approximations for electrostatic interactions affect the ability of the computational method to capture the correct polarization of the O ions, which directly affects the accuracy of the energy calculation [62]. In addition, some of the deviations among the computed interface energies result from the construction of different energy-minimized interface structures created when different periodic

Figure 8.1: Diagram of five $\alpha$-Al$_2$O$_3$ twin interface planes explored, viewed along the [1210] direction. In all figures within this work, the O and Al ions are colored red and gray, respectively.
simulation cell dimensions are used. Different periodic simulation cell dimensions impose different image forces that affect the interface structure. Both the computational method and available resources can limit the simulation cell dimensions, which restricts the number of interface structures capable of being modeled. To the authors' knowledge, no prior atomistic study has examined homophase interfaces constructed from a metastable alumina phase nor has any atomistic study examined heterophase interfaces constructed from multiple alumina polymorphs, as is done in this work.

A much wider scope of alumina interface structures has been studied using experimental methods, such as high-resolution transmission electron microscopy and electron diffraction. Several research groups have characterized the structure [63–67], analyzed impurities [68,69], and determined interface energies [70,71] in diffusion bonded \( \alpha\text{-Al}_2\text{O}_3 \) bicrystals, while others have performed similar analyses using polycrystalline \( \alpha\text{-Al}_2\text{O}_3 \) [72–74]. Experimentally
measured interface energies for $\alpha$-$\text{Al}_2\text{O}_3$ are reflective of the type of samples considered. For example, the 0.85 J/m$^2$ interface energy measured by Shin et al. [74] using polycrystalline $\alpha$-$\text{Al}_2\text{O}_3$ represented an averaged value for all interfaces sampled. Differently, analysis of $\alpha$-$\text{Al}_2\text{O}_3$ [0001] symmetric tilt bicrystals conducted by Sakuma and coworkers [70,71] revealed three classifications of interfaces with different energy values: (i) special $\Sigma 7$ (\{2310\} // (3210)) and $\Sigma 3$ prismatic twin boundaries with low-energy interfaces approximately 0.054 J/m$^2$, (ii) low-angle, $\Sigma 21$ (\{4510\} // (5410)) and $\Sigma 13$ (\{3410\} // (4310)) boundaries with interface energies approximately 0.4 J/m$^2$, and (iii) large-angle and $\Sigma 19$ (\{3520\} // (5320)) boundaries with interface energies approximately 0.7 J/m$^2$.

Table 8.3: Sample of experimentally observed homophase and heterophase alumina interfaces containing metastable $\text{Al}_2\text{O}_3$.

<table>
<thead>
<tr>
<th>ID</th>
<th>GB Planes</th>
<th>Ref.</th>
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</thead>
<tbody>
<tr>
<td>F</td>
<td>$\gamma$-$\text{Al}_2\text{O}_3$ {111} Twins</td>
<td>[75,76]</td>
</tr>
<tr>
<td>G</td>
<td>$\kappa$-$\text{Al}_2\text{O}_3$ {001} 120° Twins</td>
<td>[77]</td>
</tr>
<tr>
<td>H</td>
<td>$\theta$-$\text{Al}_2\text{O}_3$ {200} Twins</td>
<td>[78]</td>
</tr>
<tr>
<td>I</td>
<td>$\theta$-$\text{Al}_2\text{O}_3$ {110} Twins</td>
<td>[79]</td>
</tr>
<tr>
<td>J</td>
<td>(110)$<em>\gamma$ // (100)$</em>\gamma$</td>
<td>[80]</td>
</tr>
<tr>
<td>K</td>
<td>(010)$<em>\theta$ // (011)$</em>\gamma$</td>
<td>[82,84]</td>
</tr>
<tr>
<td>L</td>
<td>(0001)$<em>\alpha$ // (111)$</em>\gamma$</td>
<td>[80,81]</td>
</tr>
<tr>
<td>M</td>
<td>(100)$<em>\alpha$ // (110)$</em>\gamma$</td>
<td>[80]</td>
</tr>
<tr>
<td>N</td>
<td>(1120)$<em>\alpha$ // (112)$</em>\gamma$</td>
<td>[82,84]</td>
</tr>
</tbody>
</table>

Homophase interfaces constructed from a metastable alumina phase and heterophase interfaces constructed from multiple alumina phases have also been studied experimentally, as described Table 8.3. Multiple \{111\} $\gamma$-$\text{Al}_2\text{O}_3$ twins (F) were observed during CVD, which promoted texture within the films grown [75,76]. An analysis of pure $\kappa$-$\text{Al}_2\text{O}_3$ observed platelet like crystal formations oriented along \{001\} containing 120° twinning (G) of three crystal
regions, which mimicked pseudo-hexagonal structures [77]. Twinning also occurred in \( \theta \)-Al\(_2\)O\(_3\) on the \{200\} [78] and the \{110\} [79] mirror planes (H and I, respectively). Five heterophase alumina interfaces were identified from studies exploring the phase transition series of alumina which exhibit good crystallographic compatibility [75–83]. These commensurate interphases included two \( \theta \)-Al\(_2\)O\(_3\) // \( \gamma \)-Al\(_2\)O\(_3\) interfaces (J-K) and three \( \alpha \)-Al\(_2\)O\(_3\) // \( \gamma \)-Al\(_2\)O\(_3\) interfaces (L-N).

### 8.3 Methods

Atomistic simulations of alumina are performed with LAMMPS [85] using the ReaxFF potential [40]. ReaxFF is chosen for this work because of its transferability to many alumina phases, as shown in a previous study by the authors on bulk and surface alumina [41]. The ReaxFF parameters used in the previous [41] and the current study were first optimized by Zhang et al. [86] for Al and \( \alpha \)-Al\(_2\)O\(_3\) using first-principles simulations of Al-O clusters, then further tailored by Sen et al. [87] to better incorporate varying Al oxidation states during oxidation simulations. In the previous study [41], the authors showed that ReaxFF [87] correctly predicts \( \alpha \)-Al\(_2\)O\(_3\) as the lowest energy crystalline alumina phase; however, it unexpectedly predicts an even lower energy amorphous phase. The lower energy amorphous phase biased substantial surface reconstructions during MD simulations at temperatures well below what is expected from experiments. However, because no substantial surface reconstructions were observed at 0 K, it was concluded that the added thermal energy during MD helped promote the surface transformations. Thus, to avoid unnatural reconstructions, the current study on alumina interface structures is restricted to MS simulations performed at 0 K. Further, interface structures in this work are validated by experiment, when possible, using virtual SAED to ensure ReaxFF provides accurate results.

Alumina interfaces are studied using stoichiometric fully periodic bicrystal models and
bicrystal slabs, as shown schematically in Figure 8.2. For most homophase interfaces, fully periodic bicrystal models are constructed with dimensions normal to the boundary chosen such that two identical interface structures are created with a minimum 40 Å separation, without the vacuum region. For other interfaces, asymmetric lattice terminations prohibit the modeling of two identical interface structures; therefore, these select interfaces are studied using bicrystal slab models created by the adjoining lattice regions and separated by a minimum 100 Å vacuum region. For all models, the dimensions parallel to the interface plane are chosen to minimize the strain experienced by both lattices due to the application of periodic boundary conditions. When possible, the adjoining planes of each opposing lattice region are terminated with Al and O ions, respectively, in order to satisfy the electrostatic interactions within the interface structure. For all models, the minimum energy interface structure is found through an iterative process that (i)
systematically translates one of the orientated crystal regions and (ii) performs a non-linear conjugate gradient energy minimization at each translation.

The minimum interface energy found through translations and energy minimizations is associated with a selected interface orientation relationship. When constructing heterophase interface models that are described solely by boundary planes, one must also consider the relative rotation in the interface plane between the lattice regions to find an optimal orientation relationship. In this work, the orientation relationship between lattice regions is chosen with the aid of virtual diffraction. First, three-dimensional diffraction intensities are computed for each lattice region using the methods described by Coleman et al. [42]. Next, the diffraction intensity data for each lattice are superimposed using the VisIt visualization software [88]. An optimal orientation relationship between the lattice regions is chosen by virtually rotating the diffraction data for one lattice until a maximum overlap of the diffraction intensities is achieved. This method proved successful in identifying the ideal minimum energy orientation relationship for the $\alpha$-$\text{Al}_2\text{O}_3$ (0001) // $\gamma$-$\text{Al}_2\text{O}_3$ (111) interface [43] and is uniquely enabled by the virtual diffraction method [42].

Interface energies are calculated using two methods. For fully periodic bicrystals containing two identical interface structures, an averaged interface energy, $E_{\text{Int}}^{\text{ave}}$, for the entire model is computed via,

$$E_{\text{Int}}^{\text{ave}} = \frac{E_{12}^{\text{tot}} - nE_{1}^{\text{Unit}} - mE_{2}^{\text{Unit}}}{2A}. \quad (8.1)$$

Here, $E_{12}^{\text{tot}}$ is the total potential energy of the fully periodic bicrystal model, $n$ and $E_{1}^{\text{Unit}}$ are the number of unit cells and minimized potential energy for the bulk alumina phase in Grain 1, $m$ and $E_{2}^{\text{Unit}}$ are the number of unit cells and minimized potential energy for the bulk alumina phase.
in Grain 2, and $A$ is the area of the interface. Interface energies are also computed for all models using a method employing Gibbs dividing surfaces [89] to isolate a stoichiometric region containing the interface. Interface energies computed using Gibbs dividing surfaces, $E_{\text{Gibbs}}^{\text{Int}}$, are computed via,

$$E_{\text{Gibbs}}^{\text{Int}} = \frac{E_{12}^{\text{Gibbs}} - hE_{\text{Unit}}^{\text{Gibbs}} - kE_{\text{Unit}}^{\text{Gibbs}}}{A}. \quad (8.2)$$

Here, $E_{12}^{\text{Gibbs}}$ is the total potential energy of the atoms within the Gibbs region containing the interface, $h$ is the number of unit cells associated with Grain 1 in the Gibbs region, and $k$ is number of unit cells associated with Grain 2 in the Gibbs region. The Gibbs dividing surfaces are placed through an iterative process that takes equal steps in both directions normal to the interface while careful consideration is taken to maintain stoichiometry of the enclosed region and for each lattice. For complex orientations and phases, planar slices normal to the interfaces are not sufficient to enclose a stoichiometric region. In these cases, the necessary number of Al or O ions nearest to the Gibbs dividing surfaces are added to the Gibbs region in order to achieve stoichiometry. Because ReaxFF implements a variable charge equilibration scheme [90], the charge neutrality of the selected Gibbs region is not guaranteed at every placement of the dividing surfaces. Interface energies computed using Gibbs dividing surfaces are reported for Gibbs dividing surfaces located the smallest distance from the interface structure, $L_{\text{Gibbs}}^{\text{Gibbs}}$, where the averaged charge of the ions within the Gibbs region is below 0.005 e/atom.

Additionally, the work of adhesion for select bicrystal slab models is defined as the force needed to separate the adjoined lattice regions at the interface, creating two additional surfaces. Work of adhesion, $W_{\text{ad}}$, is computed via,
Here, $E_i^{\text{Tot}}$ ($i = 1,2$) is the total potential energy computed for slab models of each lattice region and $E_{12}^{\text{Tot}}$ is the total energy of the bicrystal slab containing the interface. When computing work of adhesion, slab models for each region maintain the same terminating structure as found in the interface model, but are relaxed via a non-linear conjugate gradient method before computing $E_i^{\text{Tot}}$.

8.4 Results and Discussion

8.4.1 $\alpha$-Al$_2$O$_3$ homophase interfaces

In order to validate the methods for computing interface energies, this work first reexamines five previously studied [44–61] twins formed by $\alpha$-Al$_2$O$_3$ using both fully periodic bicrystal and bicrystal slab models. Table 8.4 lists the computed interface energies and work of adhesion for each of the $\alpha$-Al$_2$O$_3$ interfaces. The averaged interface energy for each fully periodic bicrystal model is in good agreement with the interface energy computed using Gibbs dividing surfaces. However, computed work of adhesion do not show a clear correlation with the interface energies. ReaxFF predicts the ordering of $E_{\text{Int}}^{\text{ave}}$ and $E_{\text{Int}}^{\text{Gibbs}}$ as $B < C < D < A < E$ indicating the prismatic twin (B) is the most energetically stable $\alpha$-Al$_2$O$_3$ interface, whereas the ordering for $W_{\text{ad}}$ follows $A > B > D > C > E$ indicating the basal interface (A) requires the largest force for separation. The lack of correlation between the work of adhesion and interface energies stems from the creation of non-minimum energy alumina surfaces upon the separation of select bicrystal slab models. For example, the opposing lattice regions forming the minimum energy basal interface (A) are terminated by Al and O ions, respectively, which upon separation form high-energy surfaces ($2.87 \text{ J/m}^2$) as compared to the stable $\alpha$-Al$_2$O$_3$ (0001) surface ($0.75 \text{ J/m}^2$) [41]. In Eq. 8.3, the work of adhesion $W_{\text{ad}}$ is given by:

$$W_{\text{ad}} = \left( E_i^{\text{Tot}} + E_2^{\text{Tot}} - E_{12}^{\text{Tot}} \right) / A.$$  

(8.3)
(8.3), the creation of two additional high-energy surfaces increases the values for $E_{i}^{\text{tot}}$ as compared to $E_{i2}^{\text{tot}}$, resulting in a larger $W_{ad}$ for the basal interface. Because the work of adhesion are found to be dependent on the surface structures created upon separation, energetic data for the remainder of this work is restricted to the interface energies computed using Gibbs dividing surfaces.

With a few exceptions, the magnitude and relative order of the $\alpha$-$\text{Al}_2\text{O}_3$ interface energies predicted by ReaxFF are consistent with those predicted by Elsässer et al. [44–51] who used DFT-LDA computational methods. Both methods predict the prismatic twin (B) to be the most energetically stable $\alpha$-$\text{Al}_2\text{O}_3$ boundary with interface energies of 0.35 and 0.30 J/m$^2$ predicted by ReaxFF and DFT-LDA [51], respectively. ReaxFF and DFT-LDA also predict the next energetically stable $\alpha$-$\text{Al}_2\text{O}_3$ boundary as the rhombohedral twin (C) with interface energies of 0.60 and 0.63 J/m$^2$ [45]. Different from the results predicted by Elsässer et al. [45–51], ReaxFF predicts the pyramidal twin (D) to be more energetically stable than the basal twin (A). The interface energies for the pyramidal twin and basal twin modeled in this study by ReaxFF are 0.66 and 0.81 J/m$^2$, respectively, versus 1.88 and 0.73 J/m$^2$, respectively, predicted using DFT-LDA [44,46].

The differences between the current work and that predicted using DFT-LDA stems partly
from the different approximations within the computational methods as well as deviations between the energy minimized structures. Figure 3 shows the structures of special low \( \Sigma \) \( \alpha \)-Al\(_2\)O\(_3\) twin interfaces, which are identified by their corresponding IDs in Table 1. Similar to prior studies [44,61], ReaxFF predicts the minimum energy basal interface structure (A) to be
centered about a mirror-symmetric O plane. Relaxations of the atomic structure normal to the basal interface are minimal; however, ReaxFF does predict more substantial relaxation within the central O plane than previously reported [44,61]. The minimum energy interface structures found using ReaxFF for the α-Al₂O₃ prismatic twin (B) and rhombohedral twin (C) boundaries are consistent with those found in the prior DFT-LDA work. Figure 4 shows the minimum energy structures predicted for the higher Σ α-Al₂O₃ interfaces studied. ReaxFF predicts an Al ion terminated α-Al₂O₃ pyramidal twin (D) structure, which is consistent with the prior DFT-LDA results [46].

Figure 8.4: Minimum energy structures viewed along the <i210> directions of higher Σ α-Al₂O₃ interfaces: (D) Σ13 pyramidal twin and (E) Σ11 (1011) // (1011) twin.
In Figure 8.4, the minimum energy $\alpha$-Al$_2$O$_3$ $\Sigma 11$ (10\overline{1}I) // (10\overline{1}I) interface (E) was not explored by Elsässer et al. [45–51], but contains local structures similar to those predicted by Milas et al. [56] and Kenway [61] using DFT-GGA and MS computational methods, respectively. Due to computational limitations, the simulation cells used in the prior works were much smaller than those used in the current study, with interface plane dimensions of approximately 4.8 x 15.5 Å$^2$ versus 24.46 x 75.05 Å$^2$ used currently. The larger simulation cell used in this work allows a proper 79.1° rotation about the [12\overline{1}0] axis of O ions oriented within the basal plane versus the ~73° rotation used previously to aid the construction of a smaller simulation cell. In addition, the larger simulation cell enables a more complete representation of the structures found within the interface plane. ReaxFF predicts large voids at local regions near the interface of the adjoining lattice regions which are similar to those found in low-energy interfaces predicted by both Milas et al. [56] and Kenway [61]. In addition, ReaxFF also predicts local regions of ions having more uniform spacing across the interface which was observed as the global minimum energy structure created by Milas et al. [56].

8.4.2 Metastable Al$_2$O$_3$ homophase interfaces

Three metastable homophase Al$_2$O$_3$ interface structures are studied in this work: a $\gamma$-Al$_2$O$_3$ {111} twin (F), a $\kappa$-Al$_2$O$_3$ {001} 120° twin (G), and a $\theta$-Al$_2$O$_3$ {200} twin (H) boundary. Figure 8.5 shows snapshots of the energy minimized interface structures as well as describes the specific lattice orientations used in the study. The initial structures for each oriented lattice are derived from bulk $\kappa$-, $\gamma$-, and $\theta$-Al$_2$O$_3$ structures determined by Yourdshahyan et al. [91], Paglia et al. [92] and Zhou et al. [93], respectively. As shown, the minimum energy $\gamma$-Al$_2$O$_3$ {111} twin interface (F) contains a central plane of O ions that relax to optimize the Al-O interactions.
Note, determination of the atomic relaxation within each region is difficult due to the complex structure of $\gamma$-Al$_2$O$_3$, which appears slightly non-crystalline even in bulk models due the distribution Al ions in O interstitial sites [92]. In Figure 8.5, the $\kappa$-Al$_2$O$_3$ {001} 120° twin (G) structure is formed with Al and O ion terminations at the interface. Similar to the $\alpha$-Al$_2$O$_3$ $\Sigma$11
(10\bar{1}l) // (10\bar{1}l) interface (E), local regions of the \(\kappa\)-Al\(_2\)O\(_3\) \{001\} 120° twin contain voids near the interface of the adjoining lattices while other local regions are filled with more uniformly spaced ions across the interface. Lastly, ReaxFF predicts that the \(\theta\)-Al\(_2\)O\(_3\) \{200\} twin (H) is centered about an O plane. The minimum energy \(\theta\)-Al\(_2\)O\(_3\) \{200\} twin shows asymmetric relaxations of the ions on either side of the central O plane. Specifically, small relaxations of the Al and O ions occur in the first layer of the upper \(\theta\)-Al\(_2\)O\(_3\) \{200\} lattice, whereas no relaxations are evident in the bottom region.

Table 8.5: Computed metastable Al\(_2\)O\(_3\) interface energies modeled with ReaxFF.

<table>
<thead>
<tr>
<th>ID</th>
<th>(E_{int}^{Gibbs}) (J/m(^2))</th>
<th>(2L_{Gibbs}) (Å)</th>
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<tbody>
<tr>
<td>F</td>
<td>0.29</td>
<td>34.4</td>
</tr>
<tr>
<td>G</td>
<td>1.17</td>
<td>10.31</td>
</tr>
<tr>
<td>H</td>
<td>0.37</td>
<td>35.15</td>
</tr>
</tbody>
</table>

Table 8.5 lists the computed interface energies for each of the metastable homophase Al\(_2\)O\(_3\) interface structures. ReaxFF predicts that both the \(\gamma\)-Al\(_2\)O\(_3\) \{111\} twin (F) and the \(\theta\)-Al\(_2\)O\(_3\) \{200\} twin (H) have energies on the same order as that found for the lowest energy \(\alpha\)-Al\(_2\)O\(_3\) prismatic twin boundary (0.29 and 0.37 J/m\(^2\), respectively). Differently, the computed interface energy for the \(\kappa\)-Al\(_2\)O\(_3\) \{001\} 120° twin (G) is 1.17 J/m\(^2\), which is on the order of the highest energy \(\alpha\)-Al\(_2\)O\(_3\) \(\Sigma\)11 (10\bar{1}l) // (10\bar{1}l) interface. It is interesting to note that the atomic charge distribution within the higher energy \(\kappa\)-Al\(_2\)O\(_3\) \{001\} 120° twin requires a smaller \(L_{Gibbs}\) to achieve an effective neutral averaged charge within the Gibbs region than the other homophase interfaces constructed from metastable alumina phases.

### 8.4.3 Heterophase Al\(_2\)O\(_3\) interfaces

The heterophase Al\(_2\)O\(_3\) interfaces studied in this work include the (J-M) boundaries listed
in Table 8.3. Figure 8.6 shows snapshots of each energy minimized heterophase interface, oriented such that $\gamma$-Al$_2$O$_3$ occupies the lower lattice region. In the (110)$_\theta$ // (100)$_\gamma$ interface (J), both lattices are terminated with a mixture of Al and O ions that relax into the interfacial region. The magnitude of the relaxations experienced by the Al and O ions located in the lower (100)$_\gamma$ lattice is greater than that found in the upper (110)$_\theta$ lattice and continues further into the bulk, upwards of 10 Å. Differently, in the (010)$_\theta$ // (0$\bar{1}$1)$_\gamma$ interface (K), the magnitude of the relaxations in the lower (0$\bar{1}$1)$_\gamma$ lattice is less than the relaxations in the upper (010)$_\theta$ lattice. These findings suggest that interface relaxations could be correlated to the $<100>$ direction for both lattices in $\theta$-Al$_2$O$_3$ // $\gamma$-Al$_2$O$_3$ interfaces.
In Figure 8.6, the (0001)\(_\alpha\) // (111)\(_\gamma\) interface (L) is constructed with an upper (0001)\(_\alpha\) lattice region terminated by a plane of Al ions and a lower (111)\(_\gamma\) lattice region terminated by a mixture of Al and O ions. Both Al and O ions in the (0001)\(_\alpha\) lattice relax away from the interface towards the bulk whereas Al and O ions in the (111)\(_\gamma\) lattice relax asymmetrically toward both the interface and bulk. The (10\(\bar{1}\)0)\(_\alpha\) // (110)\(_\gamma\) interface (M) appears at a nearly cleaved plane with isolated O ion clusters extending across the interface. In the (10\(\bar{1}\)0)\(_\alpha\) // (110)\(_\gamma\) interface, the behavior of relaxation is similar to the (0001)\(_\alpha\) // (111)\(_\gamma\) interface. However, the magnitude of the relaxations found in both \(\alpha\)-\(\text{Al}_2\text{O}_3\) // \(\gamma\)-\(\text{Al}_2\text{O}_3\) interfaces examined is much less than that found in the \(\theta\)-\(\text{Al}_2\text{O}_3\) // \(\gamma\)-\(\text{Al}_2\text{O}_3\) interfaces discussed previously.

Table 8.6 provides a list of the computed interface energies for the heterophase \(\text{Al}_2\text{O}_3\) boundaries. Of all boundaries studied in this work, the heterophase (0001)\(_\alpha\) // (111)\(_\gamma\) interface (L) has the lowest interface energy, computed at 0.25 J/m\(^2\); while, the heterophase (010)\(_\theta\) // (0\(\bar{1}\)1)\(_\gamma\) interface has the highest interface energy, computed at 2.61 J/m\(^2\). Both the (110)\(_\theta\) // (100)\(_\gamma\) and the (10\(\bar{1}\)0)\(_\alpha\) // (110)\(_\gamma\) interfaces, have similar energies computed at 1.35 and 1.47 J/m\(^2\), respectively, which are on the order of the highest interface energies calculated for the homophase boundaries.

Table 8.6: Computed heterophase alumina interface energies modeled with ReaxFF

<table>
<thead>
<tr>
<th>ID</th>
<th>(E^{\text{Gibbs}}_{\text{Int}}) (J/m(^2))</th>
<th>(2L^{\text{Gibbs}}) (Å)</th>
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<tr>
<td>J</td>
<td>1.35</td>
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<tr>
<td>K</td>
<td>2.61</td>
<td>25.2</td>
</tr>
<tr>
<td>L</td>
<td>0.25</td>
<td>14.0</td>
</tr>
<tr>
<td>M</td>
<td>1.47</td>
<td>18.5</td>
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</tbody>
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8.4.4 Virtual diffraction of select \(\text{Al}_2\text{O}_3\) interfaces

To provide further validation of the interface structures sampled in this work, virtual SAED
is used to make direct comparisons of select interface models with available experimental patterns. Virtual SAED patterns are created for select interfaces using the methods described by Coleman et al. [42,43], which only require the wavelength of radiation $\lambda$, atomic positions, and atom types; no a priori knowledge of the crystal structure is assumed. The virtual diffraction algorithm generates an ultrahigh resolution, three-dimensional reciprocal lattice mesh using a $0.001 \text{ Å}^{-1}$ spacing in each direction. At each reciprocal lattice point, the algorithm computes the

![Figure 8.7: Comparison between virtual and experimental SAED patterns produced by (a-c) simulated and experimentally observed $\alpha$-Al$_2$O$_3$ basal twin [63], (d-e) simulated $\alpha$-Al$_2$O$_3$ $\Sigma$11 (10\(\overline{1}\)1) // (10\(\overline{1}\)\(\overline{1}\)) interface and experimentally observed $\alpha$-Al$_2$O$_3$ near $\Sigma$11 (\(0\overline{1}\)\(\overline{1}\)\(\overline{1}\)) // (0\(\overline{1}\)\(\overline{1}\)\(\overline{1}\)) interface [67], and (f-g) simulated and experimental (0001)$_a$ // (111)$_\gamma$ interface [81].](image)
diffraction intensities using the structure factor equation derived from kinematic diffraction theory (cf. [94,95]). Virtual SAED patterns are constructed by viewing a spherical slice of the diffraction intensity data parallel to the incident radiation, which represents the intersection of an appropriate Ewald sphere for the chosen radiation wavelength.

Figure 8.7(a-c) show virtual and experimental [63] SAED patterns for the $\alpha$-Al$_2$O$_3$ basal twin interface (A) produced along the [1210] direction using 125 keV ($\lambda=0.03275$ Å) electron radiation. In Figure 8.7(a), the virtual SAED pattern has been rotated to match that of the experimental orientation. Similar to previous works [42,43], vertical and horizontal relrods emerge from the diffraction peaks in the virtual SAED pattern due to the finite volume of the simulation. The virtual SAED pattern computed for the $\alpha$-Al$_2$O$_3$ basal twin interface closely matches the experimental pattern; however, it misses a single set of \{2202\} peaks as evident from the schematic of the diffraction pattern [63] in Figure 8.7(c). Analysis of the three-dimensional diffraction data shows that the \{2202\} peaks lie slightly off the Ewald sphere used to construct the virtual SAED pattern and would be visible through small tilts of the electron beam (or sample).

Figure 8.7(d-e) show a virtual SAED patterns for the $\alpha$-Al$_2$O$_3$ $\Sigma$11 (10$\bar{1}$)$\bar{1}$ // (10$\bar{1}$)$\bar{1}$ interface (E) alongside an experimental SAED pattern taken for a near $\Sigma$11 (0$\bar{1}$11) // (0$\bar{1}$11) boundary created by diffusion bonding [67]. Both patterns are aligned parallel to their respective $<1\bar{2}10>$ tilt axis and use 400 keV ($\lambda=0.016439$ Å) electron radiation. The SAED patterns clearly identifies the differences between the two interfaces, showing that the modeled $\alpha$-Al$_2$O$_3$ $\Sigma$11 (10$\bar{1}$1) // (10$\bar{1}$1) interface has a $<1\bar{2}10>$ tilt misorientation of $43.8^\circ$ while the near $\Sigma$11 (0$\bar{1}$11) // (01$\bar{1}$1) experimental sample contains a $35.2^\circ$ misorientation. However, the SAED patterns does show similar symmetry of the diffraction peaks within the virtual and experimental SAED
patterns. Note, the modeled interface structure with the larger misorientation angle predicts a 0.29 J/m² lower interface energy from that reported in [67].

Figure 8.7(f-g) show virtual and experimental [81] SAED patterns for the (0001)ₓ//(111)ᵧ interface (L) taken along the [10̅10]ₓ and [01̅1]ᵧ directions using 1200 keV (λ=0.00736 Å) electron radiation. The major diffraction peaks found in the virtual SAED pattern closely match those found in the experimental pattern. However, additional peaks appear in the experimental patterns due to the presence of a second γ-Al₂O₃ orientation contained within the experimental sample [81] but not represented in the simulation.

## 8.5 Conclusions

In this work, alumina interfaces are modeled using the ReaxFF interatomic potential in order to characterize their atomic structure and to extract interface energy data. This study includes a selection of homophase α-Al₂O₃, γ-Al₂O₃, θ-Al₂O₃, and κ-Al₂O₃ interfaces as well as heterophase α-Al₂O₃ // γ-Al₂O₃ and θ-Al₂O₃ // γ-Al₂O₃ interfaces. The computational methods are validated using five α-Al₂O₃ homophase interfaces, which have been studied in depth by prior atomistic simulations. The current computational methods are then applied to study homophase interfaces constructed with metastable alumina lattices and heterophase alumina interfaces constructed from multiple alumina phases, which have never before been modeled. Virtual SAED patterns are constructed for select alumina interfaces in order to obtain direct experimental validation of the simulation results.

Interface energies computed using ReaxFF for α-Al₂O₃ homophase boundaries show good agreement with those found in prior atomistic studies. However, the minimum energy structures found for the Al₂O₃ basal interface structure and the α-Al₂O₃ Σ11 (10̅1) // (10̅1) interface models deviate slightly from prior results. ReaxFF predicts the homophase γ-Al₂O₃ {111} twin and the
θ-Al₂O₃ twin to have energies on the same order as that found for the lowest energy α-Al₂O₃ prismatic twin boundary, whereas the predicted interface energy for the κ-Al₂O₃ {001} 120° twin is on the order of the highest energy α-Al₂O₃ Σ11 (101̅1) // (101̅1) interface.

The lowest energy boundary modeled in this study is the (0001)ₐ // (111)₇ interface. As mentioned, virtual diffraction is used to help optimize the orientation relationship of the (0001)ₐ // (111)₇ interface and other heterophase interfaces defined solely by their interface planes. While results for the (0001)ₐ // (111)₇ interface show that greater overlap of three-dimensional diffraction intensities from each lattice region correlate with lower energy interfaces, further studies should be performed to prove if this correlation holds true for more general interfaces and other material systems.

The interface energies tabulated in this work cover the greatest number and widest span of alumina boundaries explored using a consistent computational model, ReaxFF. These data add to recent alumina bulk and surface energies computed using ReaxFF [41] such that future predicted mesoscale models polymorphic alumina may be performed.

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Appendix 8.1


I certify that Mr. Shawn Coleman is the first author of the paper and completed greater than 51% of the work in this publication.

Sincerely,

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Appendix 8.2

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Chapter 9: Conclusions

9.1 Summary of Major Findings

Recall, the three main objectives of this work are (1) to develop and implement a virtual diffraction algorithm that generates x-ray and electron diffraction patterns directly from atomistic simulation data without a priori knowledge of the simulated crystal structure, (2) to utilize molecular statistics and molecular dynamics simulations to evaluate the energetic stability of different bulk alumina models as well as characterize metastable and stable alumina surfaces, and (3) to utilize molecular statistics and virtual diffraction to characterize homophase and heterophase alumina interfaces. The major findings of this work towards meeting these objectives are highlighted below.

- In preparation for the development and implementation of an advanced virtual diffraction algorithm, traditional methods of characterizing atomistic simulations are explored in a study that identifies the mechanisms of motion associated with a large-angle Ni grain boundary induced by synthetic, crystal-orientation-dependent driving forces [1]. The face-centered cubic Ni structure enables both a slip-vector analysis [2] and exploration of continuum metrics for microrotation and strain [3] in the Ni $\Sigma 37$ (570) [001] symmetric tilt grain boundary. The study focuses on the motion of this large-angle Ni grain boundary below the interface roughening temperature, where there is greater concern that the non-physical nature of the driving force could promote non-physical results [4,5]. Comparison between synthetically driven and shear driven simulations using traditional atomistic simulation characterization methods reveals that the synthetic-driving forces do not alter the mechanism of grain boundary motion. In addition, nudged elastic band
calculations show that the transition path and energy barriers for motion are not appreciably altered by the application of the synthetic driving force. It is important to emphasize that although great insights are gained from the use of a slip-vector analysis and the exploration of continuum metrics of microrotation and strain within this work, the same characterization techniques are not possible in the alumina material system due to the complexity of their crystal structures.

- The traditional methods of characterizing atomistic simulations are unable to uniquely identify each alumina phase considered; therefore, a novel virtual diffraction algorithm is developed and implemented [6–8]. Virtual diffraction methods are analogous to their experimental counterparts and model the scattering of electromagnetic radiation from individual atoms in order to identify the structure of a material system. The new virtual diffraction algorithm, implemented into LAMMPS [9], creates an ultra-high resolution three-dimensional mesh of reciprocal space. Diffraction intensities are computed at each reciprocal space mesh point using the structure factor equation [10], which is evaluated via the summation of inverse Fourier transform functions incorporating each atomic position. Unlike prior virtual diffraction algorithms [11–18], the current method does not require a priori knowledge of the crystal system and does not map the atomic positions onto a regular grid in order to utilize fast Fourier transforms. Instead, the algorithm requires only knowledge of the atom type and position as well as the wavelength of the incident radiation. Additionally, the new virtual diffraction method uses the same mesh algorithm to create both x-ray diffraction (XRD) line profiles and selected area diffraction (SAED) patterns, which are then constructed by a unique analysis and
The virtual diffraction algorithm is validated by studying low-angle and large-angle Ni [010] symmetric tilt boundaries, such that direct comparisons can be made to prior experimental and atomistic results [6,7]. Virtual SAED patterns viewed along the tilt axis clearly reveal the misorientation angle created by each bicrystal model. Similar to prior experimental work, subsidiary peaks appear near the {002} reflections in low-angle grain boundaries, which correspond to the dislocation spacing found in the grain boundary structure. Vertical and horizontal relrods aligned with the simulation domains also appear from the main diffraction peaks due to the finite volume explored [19]. A relrod analysis for varying sized simulations show that the predicted intensity profiles match well to the analytical expression derived from fundamental diffraction theory [19]. Relrods from the finite volume simulations are also shown to influence the predicted XRD line profiles. The predicted XRD line profiles for each grain boundary simulation show diffraction peaks at the expected Bragg angles; however, secondary peaks emerge near the {002} reflection due to relrods. At other Bragg angles, the predicted XRD line profiles show broadening due the local strain associated with the grain boundary structure.

The computational costs to perform virtual diffraction in its initial implementation could have limited the scope of this study. Therefore, the initial implementation of the virtual diffraction algorithm within LAMMPS is upgraded to take advantage of multilevel parallelism and heterogeneous computing [8]. In addition, the virtual diffraction
algorithm is also incorporated into an automated workflow to allow high throughput studies of alumina bulk, surfaces, and interfaces [8]. By incorporating OpenMP parallelization and enabling offloading to many integrated core (MIC) hardware, the virtual diffraction algorithm gains a 4.6x speed up over the original MPI based implementation. The improved algorithm is then linked to the SEAGrid science gateway [20,21] to automate the molecular dynamics simulation, virtual diffraction characterization, and visualization desired for this study through the submission of a single input deck.

• Molecular dynamics and molecular statics simulations of bulk α-, γ-, θ-, and κ-Al₂O₃ at 0, 300, 500, and 700 K show that ReaxFF [22] correctly predicts α-Al₂O₃ as the lowest energy crystalline phase [23]. Virtual SAED patterns and XRD line profiles taken of each phase are unique, which enables a distinct method for identification of each phase. Additionally, XRD line profiles of 0 K energy minimized bulk alumina are in good agreement with their experimental diffraction patterns [24–27]. Despite these successes, however, bulk simulations using ReaxFF predict a lower energy amorphous alumina phase. Virtual SAED patterns are used to characterize the amorphous Al₂O₃ systems and confirm that similar amorphous structures arise from the same thermal treatment imposed on each alumina phase.

• Atomistic simulations of 11 α-, γ-, θ-, and κ-Al₂O₃ surface structures are performed at 0, 300, 500, and 700 K using the ReaxFF potential [22] in order to extract surface energies and characterize the surface structures [23]. At 0 K, the predicted minimum energy
surface structures and energies are in good agreement with previous atomistic studies [28–30]. However, ReaxFF predicts significant surface reconstructions within select structures at temperatures below what is to be expected from experiments [31–33]. It is believed that these reconstructions are driven by the unnatural bias for amorphization within ReaxFF potential, which are promoted by the extra degrees of freedom associated with the free surface and the thermal energy available at temperatures. Analysis of several reconstructed surfaces reveal negative surface energies resulting from the creation of lower energy structures as compared to the bulk phases.

- Virtual SAED patterns are used to help identify the orientation and structures of each surface model as well as to gain experimental validation through direct comparisons of select results to prior work [23]. At 0 K, virtual SAED patterns show mostly discrete reflection peaks at the expected orientations; however, virtual SAED patterns taken at temperature reveal peak smearing within select patterns due to surface relaxations and reconstructions. Because the diffraction peaks associated with the reconstructed surfaces remain intact, complete amorphization of these surface structures is not reached due to the constraints imposed by the internal structure. Virtual XRD line profiles of select negative energy θ-Al₂O₃ surfaces are also performed and reveal a distinct pattern that is different from bulk θ-Al₂O₃ indicating the transformation of the system into a lower energy crystalline phase.

- Molecular statics simulations of 12 homophase and heterophase alumina interfaces are performed at 0 K using the ReaxFF potential [22] in order to characterize the interface
structure and extract interface energies [34]. The structures and interface energies of five $\alpha$-$\text{Al}_2\text{O}_3$ twin interfaces show good agreement with prior atomistic studies [35–41] providing validation for the current analysis methods. Homophase interfaces constructed from adjoining regions of a single metastable alumina phase as well as heterophase interfaces constructed from multiple alumina phases are investigated for the first time through atomistic simulations. For heterophase interfaces defined solely by the interface planes, virtual diffraction is used to help achieve the optimal orientation relationship of the adjoining lattice regions. In addition, virtual SAED patterns computed for select interfaces show good agreement to prior experimental results [42–44], which provide further validation for the atomistic simulations.

In summary, virtual diffraction methods are developed, implemented, and improved throughout this work in order to be used as an advanced characterization tool for atomistic simulations. ReaxFF [22] is used to model alumina bulk, surface, and interface structures and to extract pertinent energetic data. Combined, the tabulated data provides the widest scope ever performed on the alumina material system using a consistent computational model. The consistent computational model will allow quantitative comparisons to be made in future phase-field simulations (cf. [45–47]) of polymorphic alumina vapor deposition, as is discussed in Section 9.2. Because this work identifies an incorrect bias within the ReaxFF potential for constructing lower energy amorphous structures, a detailed study of the effects of ion bombardment could not be performed. However, preliminary results from ion bombardment simulations as well as new techniques of using virtual diffraction as an advance analysis tool are discussed in Section 9.2.2.
9.2 **Recommendations for Future Works**

The data and insights gained from the atomistic simulations performed on the alumina system as well as the virtual diffraction methods developed for this study lay foundations for a multitude of future works. Atomistic simulations performed in the current study highlight the structures and energetics of bulk, surface, and interface alumina systems that influence the microstructure evolution and properties of polymorphic alumina under different processing conditions. However, these data alone cannot predict the necessary processing conditions to achieve tailored alumina properties because of their limited spatiotemporal resolution. Instead, future mesoscale simulations, such as phase-field methods, are needed to extend the materials simulations into a larger spatiotemporal regime such that accurate predictions of the developing microstructure can be made varying processing conditions. The energetic data collected throughout this study will be key inputs into these predictive mesoscale models of polymorphic and polycrystalline alumina physical vapor deposition. Specifically, these energies can aid in the development of an accurate free-energy functional to evolve the solid-state alumina transformations explored by phase-field simulations.

The virtual diffraction algorithm developed for this work shows tremendous promise in advancing future materials simulations by providing additional insights and characterization capabilities, especially in connection with similar experimental studies. In its current state, the virtual diffraction algorithm can be easily applied to any atomistic simulation with corresponding experimental comparison to provide both validation of the modeled system and offer greater understanding to the experimentally observed phenomena. For example, virtual diffraction methods implemented into combined atomistic simulation and experimental studies can provide
insights into the evolution of materials properties in systems at extreme conditions, such as shock or severe plastic deformation. In these cases, virtual SAED patterns and XRD line profiles would be useful to help provide connections between the underlying microstructure evolution, such as grain growth and rotation, to the effects on the material properties.

Future works can also take advantage of the extensive three-dimensional intensity data computed by the virtual diffraction algorithm to help construct more accurate simulations and provide insights to damage and deformation processes. For example, advance analysis of diffraction data using visualization software may help optimize the necessary interface orientation relationships to achieve low-energy boundaries as discussed in [34] and in a further preliminary study detailed in Section 9.2.1. Similarly, visualization software can be used to provide greater insights to damage and deformation by tracking the deviation of diffraction intensities throughout a simulation. Examples of this analysis technique are shown in the preliminary results for ion bombardment simulations that are detailed in Section 9.2.2.

Lastly, the current virtual diffraction algorithm can be augmented to model other experimental characterization techniques based on diffraction, such as electron backscatter diffraction (EBSD). For example, preliminary work conducted in collaboration with Professor Eric Homer at Brigham Young University shows that new analysis techniques applied to the three-dimensional diffraction intensity data can simulate EBSD and produce Kikuchi patterns from atomistic simulations [48]. The analysis techniques implemented for virtual EBSD integrates different regions of the three-dimensional diffraction intensity data, which could help increase the resolution capabilities of the resulting patterns.

9.2.1 Optimization of Interface Orientation Relationships

Advanced analysis techniques using virtual diffraction data suggest a promising method for
optimizing interface orientation relationships in complex systems. Preliminary work conducted on heterogeneous alumina interfaces [34] and for the Al (111) / α-Al₂O₃ (0001) interface, has led to a method of optimizing the relative rotation of grains by using virtual SAED patterns. Figure 9.1 shows a depiction of the method whereby computed diffraction patterns created by the individual lattice regions are superimposed to create a prediction for the diffraction pattern created by the interface. In Figure 9.1, the crystal regions are rotated to achieve a maximum overlap of the virtual diffraction pattern peaks, which corresponds to higher coincidence of atoms at the interface.

![Figure 9.1](image)

**Figure 9.1:** Schematic showing the method used to obtain an optimal orientation relationship for the Al (111) / α-Al₂O₃ (0001) interface by superimposing diffraction peaks from the adjoining lattice regions.

The work of adhesion, \( W_{\text{ad}} \), is computed for both the optimized misorientation and an initial misorientation constructed with the α-Al₂O₃ region rotated 90° counterclockwise from optimal. Work of adhesion represents the force needed to separate the two regions at the interface and is computed using the equation,
\[ W_{ad} = \left( E_{1}^{\text{Tot}} + E_{2}^{\text{Tot}} - E_{12}^{\text{Tot}} \right)/A, \]  
(9.1)

where \( E_{i}^{\text{Tot}} \) \((i = 1, 2)\) is the total energy of the slab model for each region, \( E_{12}^{\text{Tot}} \) is the total energy of the interface slab, and \( A \) is the area of the grain boundary. The work of adhesion of the optimal misorientation is higher than the initial model (0.8 J/m\(^2\) versus 0.4 J/m\(^2\)). Because the same terminating planes are used in both models, the higher work of adhesion indicates the presence of a more energetically stable interface. A selected area diffraction pattern oriented along the grain boundary plane for the stable interface is shown in Figure 9.2 and compared to a pattern obtained by Medlin et al. in their experimental study [49]. The virtual diffraction pattern matches well to the experimental pattern and provides further validation of the optimal orientation relationship created by the analysis of virtual diffraction patterns.

Future work is needed to investigate the relationship between the overlapping diffraction intensities of opposing lattice regions and their corresponding interface energies in order to prove if this correlation holds true for more general interface and other material systems. Ideally, this
work will be conducted initially on a simpler FCC metal such that greater analysis can be conducted in order to connect the diffraction patterns directly to previously studied interface relationships.

9.2.2 Ion Bombardment

Preliminary simulations of a single aluminum ion bombarding onto alumina surfaces using the ReaxFF potential [22] have incorporated advanced virtual diffraction analysis techniques to measure the deviations in SAED patterns throughout the simulation. In this study, slab models of the $\alpha$-Al$_2$O$_3$ (0001) and $\gamma$-Al$_2$O$_3$ (111) surfaces as well as a slab model containing a $\gamma$-Al$_2$O$_3$ (111) // $\alpha$-Al$_2$O$_3$ (0001) interface are constructed approximately 5 nm thick and equilibrated to

![Figure 9.3: Simulated 1 keV Al ion bombardment captured at 2.000 ps impacting (a-b) $\alpha$-Al$_2$O$_3$ (0001), (c-d) $\gamma$-Al$_2$O$_3$ (111), and (e-f) $\gamma$-Al$_2$O$_3$ (111) // $\alpha$-Al$_2$O$_3$ (0001) slab models. Atoms are filtered for displacements greater than 1.5 Å or having 0.5 eV/atom kinetic energy. Colored here by kinetic energy 0.5-1.0 eV.](image)
300 K. Note, the interface slab model is oriented such that the $\alpha$-Al$_2$O$_3$ (0001) lattice is located on the top region. After equilibration, the centermost Al ion on the upper regions of each slab model is provided an additional 1 keV velocity normal to the surface in order to simulate ion bombardment while keeping the correct stoichiometry of the system. Snapshots of the ion bombardment simulations at 2.0 ps show no appreciable phase transformation as shown in Figure 9.3. However, measurements of the deviation of between the current and equilibrated SAED patterns, shown at 2.0 ps in Figure 9.4, are able to identify damage correlated to small changes in the crystal structure.

In Figure 9.4, the deviation between the SAED patterns taken at equilibration and after 2.0 ps from the initial ion bombardment reveal emerging peaks (red) and decaying peaks (blue) at different locations for each slab model. Here, emerging peaks closer to the origin of the SAED patterns indicate an expansion of the corresponding lattice direction while peaks emerging further from the origin indicate a contraction in the lattice direction. Figure 9.4 shows that expansion and compression of the lattice are isolated to specific directions within the each model, which provides insight into possible mechanisms for the initiation of phase transformations. However, because ReaxFF has been shown to unnaturally bias amorphization of the alumina structure [23], the accuracy of these ion bombardment simulation is questionable. To further characterize the nanoscale mechanisms leading to nucleation and phase transformation during physical vapor deposition using ion bombardment future work is needed that incorporates a more appropriate interatomic potential.
Figure 9.4: Deviation in the computed SAED patterns 2.0 ps after 1 keV Al ion bombardment captured at 2.000 ps impacting (a-b) $\alpha$-$\text{Al}_2\text{O}_3$ (0001), (c-d) $\gamma$-$\text{Al}_2\text{O}_3$ (111), and (e-f) $\gamma$-$\text{Al}_2\text{O}_3$ (111) // $\alpha$-$\text{Al}_2\text{O}_3$ (0001) slab models.
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