Microstructural Effect on the Ductile-to-Brittle Transition in Body Centered Cubic Metals Investigation by Three Dimensional Dislocation Dynamics Simulations

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Doctor of Philosophy in Mechanical and Aerospace Engineering

by

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2004
To my wife Cynthia . . .

for her support, love, encouragement, and understanding

during the completion of this thesis
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Abstract of the Dissertation

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The Ductile-to-Brittle Transition (DBT) is a phenomenon that is widely observed in Body Centered Cubic (BCC) metals and in covalently-based materials. Below a critical temperature (DBTT), the material suddenly loses ductility. The controlling mechanism of this transition still remains unclear despite of large efforts made in experimental and theoretical investigation. Dislocation based theories have been used to explain this phenomena. These fall into two broad categories based on either nucleation of dislocation at crack-tips, or mobility become higher at higher temperature.

Due to the limitation on the basic dislocation dynamics tool, all the previous numerical work is based on two-dimensional dislocation theory. In this work, the Parametric Dislocation Dynamics (PDD) method is reviewed and its key features examined. They are: utilization of a small number of degree of freedom, high accuracy, and high convergence rates. Applications of the PDD are demonstrated to the determination of the flow stress in irradiated materials and to the inves-
tigation of the mechanism of Persistent Slip Band (PSB) formation under fatigue condition.

The interaction of dislocation and crack is a key part to understand the shielding effect of the dislocation to the crack tip. A three dimensional discrete crack dislocation distribution method based on PDD is proposed to solve for any kind of crack tip field. By way of simulating the equilibrium distribution of the crack dislocations under the applied load and their mutual interactions, crack tip stress field can be easily obtained according to these crack dislocations. When the stress field of the crystal dislocation is treated as external load, by modifying the distribution of the crack dislocation, the shielding effect can be easily obtained. Dislocations are assumed to nucleated from different site, by way of coalescing, generates longer dislocation to shield the whole crack front, and there is a transition zone of 3D to 2D simulation when the dislocation is far away from the crack. At lower temperature, lower dislocation mobility prohibit the dislocation leaving the crack tip region fast, and thus inhibit the instantaneous nucleation of further dislocations, thus determines the fracture toughness of the considered material, and as a result, the ductility or brittleness.
CHAPTER 1

Introduction

Large scale ferritic steel structures, are used in pressure vessels, fusion reactor structures, and in the construction of nuclear reactors. After a service period, the originally ductile material may abruptly become brittle, and break without pre-warning. This kind of abrupt transition in the level of ductility is unacceptable in engineering applications, because it may cause loss in economic. Thus understanding the triggering mechanism of such kind of transition is of great importance.

Materials are not intrinsically perfect, since during manufacturing and processing, micro voids and cracks are unavoidable. Under external loading, stresses around these micro-cracks tend to be higher, causing their propagation. In most metals, extensive plastic deformation is generated around the crack tip, absorbing most of the external loading energy. In some other materials, like ceramics, ice, Body Centered Cubic(BCC) metals at low temperatures, the applied external energy can only be absorbed by way of extending the crack surface, causing the propagation of crack and final failure of the material. Figure 1.1 shows two types of stress-strain relations. According to their ability to generate plastic deformation, materials can be classified as ductile or brittle.

The ductility of some materials can change with the environment. In some BCC metals (e.g. W, α-Fe) and some alkali halide, the ductility tends to be lower as the temperature decreases. This kind of behavior is not observed in
Figure 1.1: Illustration of stress-strain curve for ductile or brittle material.

FCC metals. Neutron irradiation is another embrittlement factor for some of the reactor pressure vessels made of BCC materials in existing commercial nuclear-power plants. Because of the importance of this problem, many researchers have investigated the critical temperature which cause the transition from ductile to fully brittle behavior.

After world war II, this area has drawn many researchers’ attention in U.S. and Britain. The consideration of the brittle fracture of steels received a major attention through the generalization of the Griffith criterion of brittle fracture by Irwin(1948) and Orowan(1949). The basic consideration starts from Armstrong [2], and it was further amplified by Kelly et al. [53]. Rice and Thomson [72] proposed the fundamental notion of intrinsic brittleness and intrinsic ductility of materials from a dislocation viewpoint. Recently, a number of theories have been developed regarding this phenomenon. These theories are based on dislocation behavior near cracks, and views it as a competition between the cleavage fracture and crack tip emission of dislocations. The transition of ductile to brittle behavior is attributed to a decrease in dislocation emission (Nuclear based theories), or
due to the obstruction of the motion of dislocations (mobility based theories). Thus dislocation dynamics becomes a useful tool to understand the behavior of dislocation activities and their effects on crack tip.

First introduced in the 1980’s [63, 37], Dislocation Dynamics has now become an attractive tool for investigations of the collective processes that constitute plastic deformation of crystalline materials. In its earlier stage, numerical descriptions of dislocation ensemble evolution have been examined in considerable detail. Dislocations are approximated as 2-D and straight, making rigorous calculation of close-range interactions rather difficult. Recently, a number of numerical simulation approaches have been developed which differ mainly in the representation of dislocation loop geometry, the manner of the calculation of the elastic field and self-energy, and the description of boundary conditions. According to these differences, these methods can be categorized as follows:

1. **The Lattice Method:**[58, 57, 18, 13, 56, 14, 20, 17, 19, 67] Straight segments, either pure screw or edge in the earliest versions, or of a mixed character in more recent versions, are allowed to jump on specific lattice sites and orientations.

2. **The Force Method:**[45, 95] Straight segments of mixed character are moved in a rigid body fashion along the normal to their mid-points. No information of the elastic field is necessary, since explicit equations of interaction forces, developed by Yoffe [94] are directly used.

3. **The Differential Stress Method:**[80, 78, 79] The stress field of a differential straight line element on the dislocation is computed and integrated numerically to give the necessary Peach-Koehler force. The Brown procedure [10] is then utilized to remove the singularities associated with the self force
calculation.

4. **The Parametric Method:** Dislocation loops are divided into contiguous segments represented by parametric space curves. The equations of motion for nodal attributes (e.g. position, tangent and normal vectors) are derived from a variational energy principle, and once determined, the entire dislocation loop can be geometrically represented as a continuous (to second derivative) composite space curve.

5. **The Phase Field Microelasticity Method:** Based on Khachaturyan - Shatalov (KS) reciprocal space theory of the strain in an arbitrary elastically homogeneous system of misfitting coherent inclusions embedded into the parent phase, a consideration of individual segments of all dislocation lines is not required. Instead, the temporal and spatial evolution of several density function profiles (fields) are dealt with.

Based on the pioneer work of Eshelby in the 1950’s, and later developed by Bibly, Cottrell and Swinden (BCS) [5], Bibly and Eshelby [4] and many others, a continuous distributed dislocation technique for the simulation of crack problems becomes an attractive alternative one. In this way, the elastic-plastic crack tip field can be viewed as not only a distribution of dislocations on the crack plane, but also a distribution of non-redundant dislocations within the plastic zones of the crack tip, which in general can be viewed as the interaction of different kinds of dislocations. For clarity, we name here the dislocation on crack plane as *crack dislocation*, and *real* dislocations in the plastic zone around the crack tip as *crystal dislocation*. So far, two dimensional crack has been extensively studied by continuous crack dislocation distribution method, which leads to solving the singular integral equations, but neither the numerical nor analytical solution for
the equations can be easily solved when dealing with real crack and dislocation interactions in three dimensional space. In this work, we proposed a discrete dislocation distribution (DDD) Method based on Parametric Dislocation Method (PDD) to avoid solving the hyper singular equations, and based on this, we will study the effect of the dislocation motion ahead of crack tip, as an aid to understand the material ductility changes.

In what follows, in Chapter 2 we will survey the experimental methods and data on the ductile-to-brittle transition behavior. In Chapter 3, current two dimensional BDT theories will be discussed. Our research progress on 3-D dislocation dynamics will be presented in Chapter 4. The details of the DDD method for the simulation of general three dimensional crack is given in Chapter 5. The interaction of crack and dislocations as well as the temperature effect on the motion of the dislocations and the changes of the fracture toughness ahead of crack tip is given in Chapter 6, and finally conclusion and future directions is given in Chapter 7.
CHAPTER 2
An Overview of Experiment Investigation of DBTT on Single Crystals

Temperature and strain rate sensitive transitions are generally determined by the competition between cleavage fracture mechanism and dislocation activity in the region of crack tip. Due to the limitation on mesoscopic theories on polycrystalline materials, most experiments are based on single crystals. It is found that the Ductile-to-Brittle transition(DBT) in single crystal materials can be classified into two categories, as shown in FIG. 2.1.

Figure 2.1: Two types of ductile brittle transition in single crystals.(a) soft transition (b) sharp transition[73]

1. Soft transition: The stress intensity factor for fracture rises over a wide temperature range. This rise is due to the increase in dislocation activity
at the crack tip with the temperature changes. This type of transition is displayed by germanium [82], MgO [7], zirconia[65], molybdenum[23], titanium aluminide[6], nickel aluminide[81, 3], tungsten[49].

2. Sharp transition: The transition occurs within a small range of temperatures, usually within 10°C. Below the transition temperature $T_c$, the stress intensity factor $K$ almost keeps as a constant value, around $K_{IC}$, but when temperature is enhanced above this critical value, there is a sharp increase in material ductility as shown in the figure the sharp increase in the applied load to causing fracture. This may due to the outburst of dislocation around the crack tip zone. This kind of phenomenon is observed in Si[51, 77, 8], Al[55].

It is also shown in FIG. 2.1 that the loading rate $dK/dt$ have strong effect on BDTT, higher $dK/dt$ may cause the increase in BDTT. It is shown that the transition type of DBT is strongly dependent on the initial distribution of dislocation sources [87], sharp transitions can only occur in initially dislocation free materials.

2.1 Silicon

Silicon is generally used as an experimental material, because of the availability of well-characterized dislocation-free crystals with known dopant levels. St. John[51] is the first to study crack tip dislocation activities by testing a tapered double cantilever beam specimen at different temperatures. In his experiments, he found that BDT occurs within a very narrow temperature range (i.e. sharp transition) and BDTT is rate dependent with an activation energy of 1.9 eV over the temperature range from 973 to 1223K. Michot et. al. (1980), Michot(1982),
Michot and George (1982, 1986) measured the influence of oxygen on the BDT in silicon. The BDT temperature was found to be about 40 K higher in Czochralski (CZ) material than in floating-zone (FZ) silicon. The activation energy of the BDT increases from 2.0 eV for FZ silicon to 2.4 eV for an oxygen-containing crystal. The threshold values of the stress intensity factor for the emission of dislocation at higher temperature are 0.24 and 0.31 MPa√m for FZ and CZ, respectively. Zhang and Haasen [93] in their dynamic fracture experiments with plasma hydrogenation found that there is no effect of hydrogen on the surface energy, but the BDT temperature is lowered by about 60 K and the BDT activation energy by 0.8 eV, and also the threshold K for dislocation emission from a crack tip is lowered by 0.2 MPa√m. Brede and Hassen[8] and Michot[66] conducted almost similar experiments on Si single crystals but with different impurity concentrations, and demonstrated the connection between DBTT and dislocation mobility as a function of the loading rate. Four-point bend specimens were used in the experiment of Samuels and Roberts[77], and they observed dislocation activities only at a few preferred sites along the crack surface. Both experiments[77, 8] show the evidences that BDTT is controlled mainly by dislocation motion. Chiao and Clarke[15] in their in situ TEM experiments clearly observed dislocation emission and dislocation loop expansion processes from a sharp crack tip.

A dynamic DBTT test experiment was designed by Brede et al.[9], and later modified by Hsia and Argon [47]. In their experimental technique, the DBTT can be accurately measured for a propagating cleavage crack without explicitly differentiating the controlling crack tip mechanism of the phenomenon. Their results are somewhat different from that in static tests. In Hsia and Argon’s experiment [47], a constant $K$ field is exerted to the crack on $\{111\}$ and $\{110\}$ crack planes, a laser imaging technique was used to measure the crack velocity. They observed that due to the sluggish dislocation mobility, although disloca-
tion can be nucleated continuously from crack tips, they cannot move deep into the specimen to generate effective shielding. Furthermore, due to the free crack surface, image forces will draw these out of the surface in high temperature environments. Also in their experiment, it is shown that reaching the crack arrest temperature does not always cause general yielding. And if loading is continued, ductile to brittle transition can still happen. After crack arrest, continuous loading will cause a competition between the loading rate and the expansion rate of the crack tip plastic zone. When the rate of crack tip plastic zone (or motion of the dislocation) is lower than the loading rate, cleavage fracture was shown to still occur.

Figure 2.2: Depiction of a relatively idealized jerky crack advance scenario in which the stepwise advancing crack is systematically put into new environments at progressively increasing temperature until it is eventually arrested. After reference [30]

Recently, Gally and Argon [30] abandoned the constant displacement rate in Hsia and Argon’s experiment [47] due to its high sensitivity to minor perturbations. They used a double cantilever-beam (DCB) geometry specimen to test the DBTT since the crack is intrinsically stable due to the crack tip stress intensity
Figure 2.3: Dependence of the brittle-to-ductile transition temperature $T_{BD}$ on the average non-dimensional crack velocity $v_0$. After reference [30]

decreasing with increasing crack length. There is a temperature gradient on the specimen from low temperature at the initial crack tip to a critical temperature at the place where a travelling crack with a given velocity can arrest. The temperature was given by a series of up to ten thermocouples attached to the crack face of the specimen which can give excellent agreement with the numerical heat transfer solution, and the correction of the temperature was measured directly. In their experiment, shown in FIG. 2.2, a stepwise advancing crack is systematically put into new environment at higher temperatures, and during each step, dislocations will be possibly generated and followed by their multiplication and expansion, which leads to crack shielding and blunting. FIG. 2.3 shows the relationship between the normalized average crack velocity $v_0$ and $T_{BD}$. Here, $v_0$ is defined as:

$$v_0 = \left( \frac{\Delta a_f}{\Delta t_f} \right) \frac{1}{c} = \frac{\Delta a_f \dot{\delta}}{c \Lambda} \approx \exp \left( - \frac{\Delta U}{kT_{BD}} \right)$$ (2.1)

where, $\Delta a_f$ stands for final crack jump length, $c$ stands for shear wave velocity, $\Lambda$ is the characteristic DCB dimension, $\delta$ is the pin displacement, $\Delta U$ is the
Table 2.1: BDTT and fracture toughness K for tungsten on \{100\} and \{110\} cleavage planes with different crack front directories. Fracture toughness is in MPa$\sqrt{m}$, after reference [41]

<table>
<thead>
<tr>
<th>Crack System</th>
<th>BDTT(K)</th>
<th>K at room temperature</th>
<th>K at 77K</th>
</tr>
</thead>
<tbody>
<tr>
<td>{100} &lt; 010 &gt;</td>
<td>470</td>
<td>8.7</td>
<td>3.4</td>
</tr>
<tr>
<td>{100} &lt; 011 &gt;</td>
<td>370</td>
<td>6.2</td>
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</tr>
<tr>
<td>{110} &lt; 1\bar{1}0 &gt;</td>
<td>430</td>
<td>20.2</td>
<td>3.8</td>
</tr>
<tr>
<td>{100} &lt; 001 &gt;</td>
<td>370</td>
<td>12.9</td>
<td>2.8</td>
</tr>
</tbody>
</table>

activation energy. In their experiment, the activation energy is measured to be 1.82 eV which is considerably lower than most of the reported activation energies of 2.2 eV for dislocation glide. This may due to the inaccuracy in measuring the final period $\Delta t_f$. By way of combining etch pitting and Berg-Barrett imaging, they also find that there is an inclination of choosing only one slip activity on a set of two symmetrically placed vertical slip planes.

Argon and Gally [1] in their fracture experiments with dislocation-free Si single crystals finally pointed that, while the overall B-D fracture transition by crack arrest is indeed governed by the mobility of a very large group of dislocations moving away from the crack tip. These dislocations can be easily generated from the crack front cleavage ledges with lower energy barriers.

### 2.2 Tungsten

Gumbsch et al. [41] in their experiments examined the brittle-to-ductile transition behavior in tungsten single crystals. In their three point bend experiments, four crack systems: \{100\} < 010 >, \{100\} < 011 >, \{110\} < 1\bar{1}0 >, \{100\} < 001 >
Figure 2.4: Loading rate $\dot{K}$ dependence of fracture toughness $K$ of a specimen with $\{110\} < 110 >$ crack system. Triangles, $\dot{K}= 0.1 \text{ MPa m}^{1/2}/\text{s}$; squares, $\dot{K}= 0.4 \text{ MPa m}^{1/2}/\text{s}$; circles, $\dot{K}= 1.0 \text{ MPa m}^{1/2}/\text{s}$. After reference [41].

were carefully examined. The temperature range covered from liquid nitrogen temperature(77 K) to 650 K, and the test was performed at the constant loading rate $0.10 \pm 0.02 \text{ MPa} \sqrt{\text{m}}/\text{s}$. The corresponding BDTT of different crack systems is shown in Table 2.1. It is shown that for all the four systems, the BDTT falls into an interval of 100 K. It is also shown in FIG 2.4 that the high loading rate always lowers the fracture toughness, while increasing the BDTT, this is consistent with what Roberts et al’s estimation [73]. The activation energy is tungsten is measured as $Q_{BDT} \approx 0.2 \text{ eV}$. Chemical etching with an aqueous solution of $\text{K}_3\text{Fe}(\text{CN})_6$ and $\text{NaOH}$ makes the process of dislocation penetrating at fracture surface visible. In their experiment, motion of dislocations around the crack tip is clearly shown with this method. At higher temperature, there is an increase in the dislocation population, which suggest that additional sources are activated compared with prcracked specimens. Also, Gumbsch et al argue that
at low temperature, dislocation nucleation is limited because of the scarcity of active sources, while increasing the temperature sufficiently will activate a large number of sources. Dislocation mobility assumes to control the nucleation rate and thus the fracture toughness becomes rate dependent. And it is evident that any model for BDT cannot be based on nucleation while excluding dislocation mobility. In their whole temperature regime of investigation, it is shown that the main controlling effect of BDT is the dislocation mobility.

### 2.3 Experiments on Effect of Alloying

![Graph: Effect of Alloying](image)

Figure 2.5: The effect of various groups of alloy addition on the DBTT of Cr-based binary alloys. Group I: Ru, Al, Cu, Au, Ir, Nb, Mn, Mo, Pa, Pt, Rh, Ta, W, V; Group I*: Os, Re, Ag; Group II: Zr, U, Ti, Sn, Ni, Fe, Hf, Co; Group III: Y, Th, Sc, Ce, Gd, La. After reference [70].

Steel alloys have been investigated extensively during the past years due to its wide application in engineering. Recently, it was reported that substititional alloys,
interstitial alloys and alloys containing precipitates and second-phase inclusions can cause the so called solid-solution softening at low temperatures, hence the DBTT is somewhat different from pure metals, although that is not necessarily due to alloy softening (AS) [70]. FIG. 2.5 shows the change in DBTT with the solute concentration of Cr-based alloys with high purity. It is shown that most of the solutions result in the loss of ductility of alloyed Cr (i.e. increase in DBTT). For group II and III solutions, the ductility of Cr-based alloy was found to decrease with an increase in the solute concentration. The reason is still unclear.

Ductile crack growth initiated by the formation of disconnected ductile microcracks can be found in most of steel alloys. However, the mechanism of the formation of microvoids and their spacial distribution ahead of the crack tip maybe different. Recently, Ebrahimi and Seo [22] investigated crack initiations in a two specific type of steels: a ferritic-pearlitic and a bainitic structural steel. In the ferritic-pearlitic steel, relatively large inclusions, mainly manganese sulfide, were found to be close to the crack tip, where microvoids may form due to high shear stresses, Ductile cracks usually grow along the position of these particles. For bainitic steels, geometrical inhomogeneities was found to be the main reason for the formation of local microcracks.

As discussed in detail by Xu et al. [90, 91], \(\alpha - Fe\) has little or no energy barrier to kink motion along dislocation lines, thus the triggering effect of BDT is primarily through the formation of a dislocation embryo at the crack tip. This mechanism is different from crack behavior in silicon single crystals as discussed previously.
CHAPTER 3

Review of Current Theoretical Modelling

Many methods have been proposed to analyze the transition phenomenon, most of them show the intimate connection between the dislocation activity near the crack tip and fracture toughness. Among them, there are basically two groups. The first group is based on dislocation nucleation, and assume that dislocation nucleation at the crack tip is the controlling factor in the DBT. This is the characteristic of BCC transition metals. Another category is mobility-based models, which basically assume that dislocation nucleation is relatively easy, while temperature affects the motion of dislocations. The back stress due to these dislocations at the crack tip affects the fracture toughness. This behavior is typical of semiconductors and compounds. The distinguishing aspect between the two models is the mobility of kinks on dislocations. Computer simulations show that the kink motion along dislocations is hindered by very substantial energy barriers in Si. In some experiments, it is indicated that there is little resistance to the motion of kinks along dislocation lines in some BCC metals. These suggest that the transition is either controlled by nucleation (BCC materials) or is mobility controlled.
3.1 Nucleation-based Models

Rice and Thomson [72] were the first to consider the brittle versus ductile behavior of materials in terms of crack tip dislocation activity. As shown in FIG. 3.1, they proposed that the competition between dislocation emission and atomic decohesion at the crack tip is the controlling factor in the ductile versus brittle behavior of a material. In their model, they assumed that there is an embryonic dislocation nucleated on a gliding plane with inclination angle $\phi$. The distance between the embryonic dislocation and the crack tip is equal to the dislocation core radius $r_0$. Propagation of the crack will have two possibilities under external stress. The first is that the nucleation of the embryonic dislocation and its subsequent motion away from the tip, which makes the crack tip blunt. The other is that cleavage fracture proceed along the crack surface, and the crack tip remains sharp. The stress which causes the emission of the dislocation is mainly composed of an external K-field, the image stress due to the proximity to free crack surfaces, and any other stresses due to crack blunting. On the other hand, the energy release rate $G$ can cause the propagation of the crack, while the surface energy $\gamma_c$ acts as a retarding force. In the loading process, if dislocation
emission occurs earlier, then the material is assumed to be intrinsically ductile, otherwise it is brittle. In their model, the Griffith criterion is used for cleavage requirements. Rice and Thomson developed quantitative evaluations of the conditions for dislocation emission from the near-tip region. It was shown that the ratio $\mu b/\gamma_c$ (here $\gamma_c$ stands for surface energy, $\mu$ shear modulus, and $b$ the length of Burgers vector) was a good indicator of the ductile versus brittle response. Materials whose dislocations have wide core and with small value in $\mu b/\gamma_c$ tend to be ductile. The energy barrier to dislocation emission was calculated to be 19 eV for Fe, 329 eV for W, 111 eV for Si and 329 eV for Ge. Their values are much higher than the experimentally found activation energies for BDT.

In experimental observations, dislocations are not always emitted as straight long segments, as assumed in 2-D analysis shown in FIG. 3.1. Two dimensional models always underestimate the material ductility. Recently, researchers consider the initial dislocation loop configuration as rectangular or half elliptic, as shown in FIG. 3.2. Three dimensional loop models not only predict the ductility more precisely, but also decided the motion and configuration of dislocations after their nucleation. Gao and Rice [31] studied the half elliptical case, and obtained a lower $K$ needed for the emission, as compared with the 2-D case. Later, the Rice-Thomson model has evolved continuously, to account for elastic anisotropy, bimaterial interfaces, nonlinear dislocation core structures, and realistic slip system geometries. Xu, Argon and Otiz [91] in their paper studied different realistic slip system geometries. The activation configuration includes nucleation on inclined planes, on oblique planes and on cleavage ledges, all of them are treated within the framework of Peierls.

Xu et al [90] suggested that the possible B-D transition temperature from the
activation energy as:

$$T_{BD} = \left( \frac{\ln \left( \frac{c}{v} \right)}{\alpha} + \eta \frac{T_0}{T_m} \right)^{-1} T_0$$  \hspace{1cm} (3.1)$$

Here, $T_0 = \mu b^3/k(1 - \nu) \approx 1.2 \times 10^5 K$, the melting temperature $T_m = 1809 K$ for $\alpha$-Fe, $\alpha = (1 - \nu) \Delta U_{activation}/\mu b^3$ is the normalized activation energy, $c$ is the speed of sound, $v \approx 1 \text{ cm/s}$ is the typical crack propagation velocity, giving $\ln(c/v) \approx 10$, $\eta \approx 0.5$ is a coefficient describing the temperature dependence of the shear modulus. Thus, BDTT is dependent on the activation energy of dislocation nucleation.

### 3.2 Mobility-based Models

In many materials, such as BCC metals, the dislocation velocity increases strongly with the temperature increase till the so-called athermal temperature. Above this temperature, the dislocation velocity is widely temperature independent. Below this temperature, the dislocation velocity is regarded as being thermally activated.

Figure 3.2: 3-D dislocation nucleation models from crack tip. (a) rectangular loop. (b) elliptical loop
Table 3.1: Material Properties of Tungsten single crystals and dislocation velocity. E is the Young’s modulus, \( \mu \) is the shear modulus, \( \nu \) is Poisson’s ratio, \( a_0 \) is the lattice constant, \( b \) is the length of Burgers vector, \( \alpha \) and \( \beta \) describes the temperature dependence of the stress exponent \( m \), and \( Q_{\text{dis}} \) (eV) is the apparent activation energy for dislocation motion.

<table>
<thead>
<tr>
<th>E(GPa)</th>
<th>( \mu )(GPa)</th>
<th>( \nu )</th>
<th>( a_0 )(nm)</th>
<th>( b )(nm)</th>
<th>( \alpha )(K)</th>
<th>( \beta )</th>
<th>A(m/s)</th>
<th>( Q_{\text{dis}} )(eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>388.7</td>
<td>152.7</td>
<td>0.29</td>
<td>0.3159</td>
<td>0.274</td>
<td>592</td>
<td>2.81</td>
<td>3.23 \times 10^{-9}</td>
<td>0.323</td>
</tr>
</tbody>
</table>

with some activation energy \( Q_{\text{dis}} \), and can be written in the Arrhenius-type form

\[
v = A \left( \frac{\tau}{\tau_0} \right)^{m(T)} \exp \left( -\frac{Q_{\text{dis}}}{k_B T} \right)
\]

(3.2)

Where \( A \) is a constant with the dimension of velocity, \( \tau \) is shear velocity exerted on the dislocation, \( \tau_0 \) is normalized stress, \( k_B \) is Boltzmann’s constant, \( T \) is the absolute temperature. \( m(T) \) is the temperature dependent constant, assumed to be of the form [71]

\[
m(T) = \frac{\alpha}{T} + \beta
\]

(3.3)

\( \alpha \) and \( \beta \) are the two fitted constants. Table 3.1 lists all the parameters for Tungsten dislocation velocity.

Let’s define dislocation mobility as

\[
M = A \exp \left( \frac{-U_v}{k_B T} \right)
\]

then EQN. 3.2 can be rewritten as:

\[
v = M \tau^m
\]

(3.4)

As can be seen from EQN. 3.4, the higher the total shear stress applied on the dislocation, the faster the dislocation moves. On the other hand, the higher
the temperature, the higher mobility of the dislocation, and thus the higher the dislocation velocity. At a given temperature, once the dislocation velocity is determined, and the external loading and internal interactions between dislocations are also set, then the position of the dislocation will be known. Thus, the shielding effect of the dislocations to the crack is known, and hence the local stress intensity factor is determined. Applying the Griffith criterion, the critical condition for crack propagation can be determined at any temperature. Numerical simulations show that the transition temperature is highly dependent on the loading rate[42].

The other category of models consider that the mobility is constant, while the total stress on the dislocation changes at different temperatures due to a change in the Peierls-Nabarro stress. The Peierls-Nabarro stress is defined as the applied resolved shear stress required to overcome the lattice resistance to movement by a dislocation loop. The Peierls stress is a consequence of the inter-atomic forces/displacement interaction between the dislocation loop and the surrounding crystals. And this resistance to dislocation movement is due to the periodic variation in the misfit energy of atomic half planes above and below the slip plane with the dislocation loop. For higher dislocation densities, the Peierls stress is comparable to long range interactions between dislocations. It is generally accepted that the Peierls stress is a dominant controlling factor in the plastic slip at low temperatures [84]. It decreases with an increase in temperature. Thus the effect of temperature on the Peierls stress can also help us understand the BDT phenomenon.
3.3 Current Computer Simulations and their limitations

With the fast development of the computer technique, more and more people tend to use numerical method to simulate the plastic deformation in mesoscopic scale. Dislocation Dynamics (DD) has now become an attractive tool for investigations of the ductile to brittle transition process, but most of them concentrated on the interaction on infinite straight dislocation with 2D cracks, which may underestimate or overestimate the BDT temperature. The Oxford group [42, 74, 45, 75] and related references, Xin and Hsia [89], Ferney and Hsia [26], carefully studied the BDTT on Si in 2D, and concluded that the BDT in silicon is mainly controlled by dislocation mobility, based both on peripheral evidence that the activation energy for the transition temperature is the same as that for dislocation mobility.

The basic procedure for the simulation is straightforward as shown in the following several steps:

1. **Calculate the total stress on the dislocation**

   The total stress applied on the $i$th dislocation $\tau_i$ can be written as follow:

   \[
   \tau_i = \tau_{iK} + \tau_{image}^i + \tau_{interaction}^i + \tau_{other}^i \tag{3.5}
   \]

   Here, $\tau_{iK}^i$ is the shear stress due to crack tip stress field, usually they are written as follow:

   \[
   \tau_{iK}^i = \frac{K_{applied}}{\sqrt{\pi r_i}} f(\theta) \tag{3.6}
   \]

   Here, $r_i$ is the distance toward the crack tip, $f(\theta)$ is the coefficient related to the inclination angle of the slip plane. Usually, 2D K-field is considered.

   For the image stress $\tau_{image}^i$, direct analytical results for infinite straight long
solution is used,

\[ \tau_{image}^i = \beta \frac{\mu b}{r_i} \]  

(3.7)

\( \beta \) is a correction coefficient.

The dislocation-dislocation interaction term \( \tau_{interaction}^i \) can be written as [89]:

\[ \tau_{interaction}^i = \frac{\mu b}{4\pi(1-\nu)} \sum_{j \neq i} \left[ \left( \frac{r_i}{r_j} \right)^{1/2} \frac{1}{r_i - r_j} + \frac{8r_i r_j}{(r_i + r_j)^3(r_i - r_j)} \right] \]  

(3.8)

Summing up all the stress term related to dislocation and crack, also stresses generated by grain boundary, or other impurities, we can get the net shear stress on the dislocation.

2. **Update the positions of each dislocation**

According to EQN 3.4, calculate the new position of each dislocation at a small time step by way of any implicit or explicit integration method.

3. **Update the locale stress intensity factor \( K \)**

\[ K_{total} = K_{applied} - K_D \]  

(3.9)

Here, \( K_D \) is the factor of the dislocations, and can be written as follows: [26, 43]

\[ K_D = \sum_i \frac{\mu b f(\theta)}{(2\pi r_i)^{1/2}(1-\nu)} \]  

(3.10)

Here, \( f(\theta) \) is a coefficient related with slip direction and different slip system, in different simulation conditions, this value may vary. In [89], this value is chosen as \( 3\sqrt{2}/4\alpha \) where, \( \alpha \) is the number of equivalent active slip systems, referring as the shielding coefficient, for instance, \( \alpha = 2 \) means that two symmetric slip systems are activated, and \( \alpha = 1 \) means that only one slip system is activated. In their results, it is shown that increasing
the number of active slip systems will lead to a sharp transition, and the
sharpness is strongly dependent on $\alpha$. When $\alpha = 4$, the transition may
become extremely sharp. It is argued that with lower $\alpha = 1$, the number
of dislocations at the moment of cleavage fracture increases gradually over
a wide temperature range, whereas for large values of $\alpha$, there is a sudden
increase in the number of dislocation accompanied by sharp transition.

4. **Check the dislocation emission condition**

Similar to the Rice-Thomson model\[72\], embryonic dislocations exist at
$r = r_c$ from the crack tip, where $r_c$ is the core radius, once the crack tip
stress intensity factor reaches a critical value, one dislocation is emitted.
Note here that, emission of new dislocation doesn’t depend on the presence
of prior ones.

5. **Apply the fracture criteria**

For brittle fracture, according to Griffith criterion, check the stress intensity
$K_{total}$ whether it exceeds the intrinsic fracture toughness $K_{IC}$, or for ductile
fracture\[89\], when the far applied field $K_{applied}$ exceed some certain limit,
$7K_{IC}$ is chosen in \[89\]. If both two conditions are not satisfied then go back
to step 1, and continue on the computational cycle.

FIG. 3.3 shows a typical simulation model. FIG. 3.3a shows a truly 2-D
problem with straight dislocations and 2-D crack tip field, and in FIG. 3.3b, a
pseudo 3D case are studied, where dislocations can be emitted from position near
crack fronts with rectangular shape, and during their motion, the 3 segment of
the dislocation still keeps as straight.

As previously discussed, the limitations are obvious. First, in reality, there is
no straight dislocations, so for the Dislocation Dynamics(DD) part, it is difficult
Figure 3.3: Modelling the plastic zone as a single inclining slip plane (a) uniform dislocation nucleation (b) dislocation nucleation at separated sources. refer to [43] to simulate real 3D configurations, and especially difficult to solve for the dislocation motion at earlier stage, since their shielding effect is stronger, due to their short distance to the crack tip. And for 2D case, it is difficult to estimate the effect of different slip systems as discussed in Xu and Argon’s work[91]. How is the mutual interactions between those dislocations in different slip planes? For 2D simulation, it is impossible to answer this question clearly. Second, is the calculation the applied $K$-field, since most of the problem cannot be simplified to two dimension.
CHAPTER 4

Computational Aspect of Parametric

Dislocation Dynamics

Dislocation Dynamics (DD) has now become an attractive tool for investigations of both fundamental and collective processes that constitute plastic deformation of crystalline materials. Since it is a foundational issue in studying the dislocation emission and motion ahead of crack tip, we here have a detailed review of the three dimensional parametric dislocation dynamics and its numerical accuracy and convergence.

4.1 Formulation of Parametric Dislocation Dynamics (PDD)

The method of PDD is described in sufficient details in references [36, 39, 40, 38], and we will attempt here to give only a brief description. The first step is to calculate the stress field of curved parametric segments. Let the Cartesian orthonormal basis set be denoted by $\mathbf{1} \equiv \{\mathbf{1}_x, \mathbf{1}_y, \mathbf{1}_z\}$, $\mathbf{I} = \mathbf{1} \otimes \mathbf{1}$ as the second order unit tensor, and $\otimes$ denotes out tensor product. Now define the three vectors $(\mathbf{g}_1 = \mathbf{e}, \mathbf{g}_2 = \mathbf{t}, \mathbf{g}_3 = \mathbf{b}/|\mathbf{b}|)$ as a covariant basis set for the curvilinear segment, and their contravariant reciprocals as[46]: $g^i \cdot g_j = \delta_j^i$, where $\delta_j^i$ is the mixed Kronecker delta and $V = (\mathbf{g}_1 \times \mathbf{g}_2) \cdot \mathbf{g}_3$ the volume spanned by the vector basis, as shown in FIG. 4.1. The parametric representation of a general curved
Figure 4.1: Parametric representation of a general curved dislocation segment, with relevant vectors defined (after reference [38])

dislocation line segment, shown in the figure, can be described by a parameter $\omega$ that varies from 0 to 1 at end nodes. The segment is fully determined as an affine mapping on the scalar interval $\{\omega \in [0, 1]\}$, if we introduce the tangent vector $\mathbf{T}$, the unit tangent vector $\mathbf{t}$, and the unit radius vector $\mathbf{e}$ as follows:

$$\mathbf{T} = \frac{d\ell}{d\omega}, \quad \mathbf{t} = \frac{\mathbf{T}}{|\mathbf{T}|}, \quad \mathbf{e} = \frac{\mathbf{R}}{R}$$

Ghoniem, Huang and Wang [38] have shown that the elastic field of such a parametric segment can be obtained as an affine mapping transformation of the scalar parameter $\omega$, and that the stress field differential $d\sigma$ introduced by a parametric differential $d\omega$ are related as:

$$\frac{d\sigma}{d\omega} = \frac{\mu V |\mathbf{T}|}{4\pi(1-\nu)R^2} \left\{ \left( \mathbf{g}^1 \otimes \mathbf{g}_1 + \mathbf{g}_1 \otimes \mathbf{g}^1 \right) + (1-\nu) \left( \mathbf{g}^2 \otimes \mathbf{g}_2 + \mathbf{g}_2 \otimes \mathbf{g}^2 \right) - (3\mathbf{g}_1 \otimes \mathbf{g}_1 + \mathbf{I}) \right\}$$

(4.1)

The affine map described by EQN. 4.1 can also be given by the covariant,
contravariant and mixed vector and tensor functions [38]:

\[
S = \text{sym}[\text{tr}(A^i_j g_i \otimes g^j)] + A^{11}(3g_1 \otimes g_1 - 1 \otimes 1) \tag{4.2}
\]

The scalar metric coefficients \(A^i_j, A^{11}, B^{11}\) are obtained by direct reduction of EQN.4.1 into EQN.4.2. Once the parametric curve for the dislocation segment is mapped onto the scalar interval \(\{\omega \in [0, 1]\}\), the stress field everywhere is obtained as a fast numerical quadrature sum from EQN. 4.1 [39]. The self-force is obtained from knowledge of the local curvature at the point of interest.

To simplify the problem, let us define the following dimensionless parameters:

\[
\begin{align*}
  r^* &= \frac{r}{a}, & f^* &= \frac{F}{\mu a}, & t^* &= \frac{\mu t}{B}
\end{align*}
\]

Here, \(a\) is lattice constant, \(\mu\) the shear modulus, and \(t\) is time. Substitute these to the variational formula of the governing equation of motion of a single dislocation loop [40], we get the dimensionless matrix form as:

\[
\int_{\Gamma} \delta r^* \delta f^* \left( f^* - \frac{dr^*}{dt^*} \right) |ds^*| = 0 \tag{4.3}
\]

Here, \(f^* = [f_1^*, f_2^*, f_3^*]^T\), and \(r^* = [r_1^*, r_2^*, r_3^*]^T\), which are all dependent on the dimensionless time \(t^*\). Following reference [40], a closed dislocation loop can be divided into \(N_s\) segments. In each segment \(j\), we can choose a set of generalized coordinates \(q_m\) at the two ends, thus allowing parameterization of the form:

\[
r^* = CQ \tag{4.4}
\]

Here, \(C = [C_1(\omega), C_2(\omega), ..., C_m(\omega)]\), \(C_i(\omega), (i = 1, 2, ...m)\) are shape functions dependent on the parameter \((0 \leq \omega \leq 1)\), and \(Q = [q_1, q_2, ..., q_m]^T\), \(q_i\) are a set of generalized coordinates. Now substitute EQN.4.4 into EQN.4.3, we obtain:

\[
\sum_{j=1}^{N_s} \int_{\Gamma_j} \delta Q^T \left( C^T f^* - C^T C \frac{dQ}{dt^*} \right) |ds| = 0 \tag{4.5}
\]
Let,
\[ f_j = \int_{\Gamma_j} C^\top f^* |ds|, \quad k_j = \int_{\Gamma_j} C^\top C |ds| \]

Following a similar procedure to the FEM, we assemble the EOM for all contiguous segments in global matrices and vectors, as:
\[ F = \sum_{j=1}^{N_s} f_j, \quad K = \sum_{j=1}^{N_s} k_j \]

then, from EQN 4.5 we get,
\[ K \frac{dQ}{dt^*} = F \quad (4.6) \]

EQN. 4.6 represents a set of ordinary differential equations, which describe the motion of an ensemble of dislocation loops as an evolutionary dynamical system. Given the initial condition and boundary conditions, solving EQN. 4.6, the position and configuration of each dislocation is set, and hence at each time step, the total stress field can be known.

If we specifically choose cubic splines as shape functions, ignore the climbing effect and confine dislocation motion to be on its glide plane, we end up with only 8 DOFs for each segment with each node associated with 4 independent DOFs. These cubic spline shape functions are given by:
\[ C = [2\omega^3 - 3\omega^2 + 1, \omega^3 - 2\omega^2 + \omega, -2\omega^3 + 3\omega^2, \omega^3 - \omega^2] \]

\[ Q = [P_1, T_1, P_2, T_2]^\top \]

Here, \( P_i \) and \( T_i \) \((i = 1, 2)\) correspond to the position and tangent vectors, respectively.
4.2 Spatial and Temporal Resolution of Dislocation Mechanisms

4.3 Temporal resolution

Figure 4.2: The influence of the time integration scheme on the shape convergence of an F-R source. Here, Burgers vector is chosen as 1/2[\bar{1}01] with applied uniaxial stress \( \sigma_{11} = 80 \) MPa. (or \( \tau/\mu = 0.064\% \) )

As shown in previous section, after the initial conditions are set, the major problem is to solve EQN. 4.6. Two kinds of integration methods, implicit and explicit, are utilized. For the explicit integration, simple one step Euler forward method are used. Modified Gear’s implicit integration for stiffness equation [35] developed by Lawrence Livermore National Laboratory are used for implicit time integration. The comparisons between implicit and explicit integrations with different time steps are shown in FIG. 4.2, and error estimations at Table 4.1.

In the explicit scheme, it is noted that when the time step is larger than \( \approx 3000 \), there will be a numerical shape instability. For the parameters chosen
here, this corresponds to a physical time step of \( \approx 6 \) ps. The shape tends to diverge more along near screw segments of the F-R source. For a time step on the order of 1000 (i.e. \( \Delta t \approx 2 \) ps), the F-R shape is numerically stable, but not accurate. Finally, when the explicit time step is lowered to less than 500 (i.e. \( \Delta t \approx 1 \) ps), PDD tends to give a stable and accurate F-R source shape. Such small limit on the time step for high mobility crystals (e.g. FCC metals) can result in severe restrictions on the ability of current simulation for large scale plastic deformation. With the method designed by Gear for the numerical integration of ordinary differential stiff equations, a variable time step can be automatically determined based on the variation of any of the DOF. A level of relative accuracy of \( 10^{-6} \) is selected as a convergence constraint. Since the time step is automatically adjusted to capture the specified level of accuracy, the overall scheme is stable and convergent. It is shown in Table 4.1 that the overall running time in explicit integration is much less than that with explicit integration scheme at small time step. That is due to the ability of adjusting time step during implicit integration according to the stiffness of the equation, while explicit Euler method can’t, and only with very small time step can we get the same level of accuracy and convergence.

### 4.4 Spatial resolution limits on PDD

For large-scale computer simulations, there is an obvious need to reduce the computational burden without sacrificing the quality of the physical results. The smallest number of spline segments with the largest time step increment for integration is a desirable goal. However, one must clearly identify the limits of this approach. We study here the influence of the nodal density on the dislocation line, and the time integration scheme on the ability to satisfactorily resolve the
Table 4.1: Error Estimation for Different Integration Scheme. The implicit scheme is chosen as the reference configuration for error estimation.

<table>
<thead>
<tr>
<th>Integration Scheme</th>
<th>Absolute Error $\epsilon_a$</th>
<th>Relative Error $\epsilon_r$</th>
<th>Runtime (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Explicit Int. ($\Delta t^* = 3000$)</td>
<td>168.4</td>
<td>6.11%</td>
<td>0.92</td>
</tr>
<tr>
<td>Explicit Int. ($\Delta t^* = 1500$)</td>
<td>141.5</td>
<td>5.40%</td>
<td>1.82</td>
</tr>
<tr>
<td>Explicit Int. ($\Delta t^* = 1000$)</td>
<td>56.90</td>
<td>2.34%</td>
<td>2.76</td>
</tr>
<tr>
<td>Explicit Int. ($\Delta t^* = 500$)</td>
<td>0.06</td>
<td>0.003%</td>
<td>5.68</td>
</tr>
<tr>
<td>Implicit Integration</td>
<td>0</td>
<td>0</td>
<td>1.52</td>
</tr>
</tbody>
</table>

shape of a dynamic F-R source.

### 4.4.1 Single F-R source

FIG. 4.3 shows a stable (with applied $\sigma_{11} = 80$ MPa, $\tau/\mu = 0.064\%$) and an unstable ($\sigma_{11} = 200$ MPa, $\tau/\mu = 0.16\%$) F-R source configuration. The dislocation loop is divided into different number of segments, and its motion is gained by different numerical integrations. It is shown that one can achieve very high precision in describing the stable F-R shape with very small number of segments. The corresponding error is shown in Table 4.2. For comparisons, we choose here the result with 30 segments as the reference configuration (thus the relative and absolute error is set to zero). It is found that with the increasing number of segments, both the relative and absolute error are decreased sharply, but the running time is increased significantly. It is interesting to note here that with only 2 segments, one can achieve almost the same resolution as that with 30 segments, with the relative error less than 0.2%, 2% of the CPU time used in 30 segments. However, when the F-R source becomes unstable, the variation of curvature is considerable
Figure 4.3: The influence of number of segments on the shape convergence of an F-R source between its middle section and the sections close to the pinning points, 2 segments is not enough. FIG.4.3-b and Table 4.3 show the configuration and corresponding absolute, relative error and running time respectively in the unstable case. The reference configuration is chosen as that with 40 segments. It is found that only 2 segments is unable to achieve high accuracy, although it still converges. It is due to the complicate configuration compared to that at stable case. The curvature

<table>
<thead>
<tr>
<th>No. of Segments</th>
<th>Absolute Error $\epsilon_a$</th>
<th>Relative Error $\epsilon_r$</th>
<th>Runtime(sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>6.06</td>
<td>0.17%</td>
<td>0.12</td>
</tr>
<tr>
<td>6</td>
<td>6.01</td>
<td>0.15%</td>
<td>0.42</td>
</tr>
<tr>
<td>15</td>
<td>1.32</td>
<td>0.018%</td>
<td>1.53</td>
</tr>
<tr>
<td>30</td>
<td>0</td>
<td>0</td>
<td>5.77</td>
</tr>
</tbody>
</table>
Table 4.3: Error Estimation for Unstable Frank-Read Source at $t^* = 5 \times 10^6$.

<table>
<thead>
<tr>
<th>No. of Segments</th>
<th>Absolute Error $\epsilon_a$</th>
<th>Relative Error $\epsilon_r$</th>
<th>Runtime(sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>1408.8</td>
<td>20.15%</td>
<td>0.02</td>
</tr>
<tr>
<td>6</td>
<td>191.1</td>
<td>5.04%</td>
<td>0.20</td>
</tr>
<tr>
<td>15</td>
<td>133.8</td>
<td>3.24%</td>
<td>2.53</td>
</tr>
<tr>
<td>30</td>
<td>142.0</td>
<td>2.93%</td>
<td>24.14</td>
</tr>
<tr>
<td>40</td>
<td>0</td>
<td>0</td>
<td>27.57</td>
</tr>
</tbody>
</table>

is much higher at the zone near the fixed point. Only 2 segments is not enough to capture such a high curvature, the error is mainly from these two zones. In Table 4.3, it is shown that with the increasing number of segments, the accuracy increased greatly with the compensation of increasing CPU time.

4.4.2 Finite-Size Dipole Formation

FIG. 4.4 shows a 2-D projection on the (111)-plane of the dynamic process finite-size dipole formation. Two initially straight dislocation segments with the same Burgers vector $\frac{1}{2}[\bar{1}01]$, but of opposite line directions are allowed to glide on nearby parallel \{111\}-planes without the applying an external stress. The two lines attract one another, thus causing the two loop segments to move and finally reach an equilibrium state of a finite-size dipole. The two parallel dislocations are pined at both ends, the upper loop glides on the ”upper” plane, while the ”lower” one glides on the ”lower” one as shown in the figure. The mutual attraction between the two dislocations becomes significant enough to simultaneously reconfigure both of them only during the latter stages of the process. Because the two dislocations start with a mixed character, a straight and tilted middle section of the dipole forms. The length of this middle section, which we may
Figure 4.4: Two F-R source dislocations with the same Burgers vector \( \mathbf{b} = \frac{1}{2}[\overline{1}01] \) but opposite tangent vectors gliding on two parallel (111)-planes \( (h = 25\sqrt{3}a \text{ apart}) \) form a short dipole in an unstressed state. The view is projected on the (111)-plane. Time intervals are: (1) \( 2.5 \times 10^5 \), (2) \( 4.75 \times 10^5 \), (3) \( 5 \times 10^5 \), (4) Equilibrium state.
Table 4.4: Error Estimation for different nodal distribution of dipole formation. The configuration with 20 segments each dislocation is chosen as the reference configuration.

<table>
<thead>
<tr>
<th>No. of Segments</th>
<th>Absolute Error $\epsilon_a$</th>
<th>Relative Error $\epsilon_r$</th>
<th>Runtime(sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>26.5</td>
<td>3.16%</td>
<td>12.7</td>
</tr>
<tr>
<td>4</td>
<td>18.6</td>
<td>2.37%</td>
<td>43.3</td>
</tr>
<tr>
<td>5</td>
<td>7.8</td>
<td>1.13%</td>
<td>84.9</td>
</tr>
<tr>
<td>10</td>
<td>3.6</td>
<td>0.36%</td>
<td>438.2</td>
</tr>
<tr>
<td>20</td>
<td>0</td>
<td>0</td>
<td>1503.1</td>
</tr>
</tbody>
</table>

simply ascribe as the dipole length, is only determined by the balance between the attractive forces on the middle straight section, and the self-forces on the two end sections close to the pinning points. The separation of the two planes is $25\sqrt{3}$, which is approximately $60\ |\mathbf{b}|$.

The error estimation of different nodal distribution of same dipole formation is shown in Table 4.4. It is shown that with only 2 segments in each dislocation, high accuracy can be obtained, the relative error is less than 5%.

### 4.4.3 Dislocation Junctions

FIG.4.5 shows the dynamics of an attractive junction formation without any externally applied stress. Two initially straight dislocations $1/2[01\bar{1}](111)$ and $1/2[101](11\bar{1})$ are pinned at their ends, and allowed to move on the two intersected gliding plane respectively, approaching each other until they are locked at equilibrium. The length of the straight section (junction) that forms at the intersection of the two glide planes is approximately 200. FIG. 4.5-a shows a
2-D projection view of the successive motion of \( \frac{1}{2}[01\bar{1}](111) \), while the 3-D view of the junction structure is shown in FIG. 4.5-b. In order to calculate the error generated by different nodal distribution, the configuration with 12 nodes each dislocation is set as the reference one. The error estimation is shown in Table 4.5. It is shown that one can get good shape junction with less than 8 segments in each dislocation loop.

Figure 4.5: Dynamics of 2 unstressed F-R sources (\( \frac{1}{2}[01\bar{1}](111) \) and \( \frac{1}{2}[101](11\bar{1}) \)) forming a 3D junction along (\( \bar{1}10 \)), \( \mathbf{b} = \frac{1}{2}[110] \). (a) 2D view for the motion of the F-R source (\( \frac{1}{2}[01\bar{1}](111)\frac{1}{2}[101](1\bar{1}1) \)) on its glide plane(111). Time intervals are (1) initial configuration, (2) \( 1.5 \times 10^4 \), (3) \( 5.0 \times 10^4 \), (4) \( 1.3 \times 10^5 \), (5) Final configuration. (b) 3-D view of the junction

### 4.5 Adaptive Node Redistribution

One of the main feature of our PDD is the adaptive algorithm of space division. In order to capture details of small scale processes, such as the interaction between a
Table 4.5: Error Estimation for different nodal distribution of junction formation. The configuration with 12 segments each dislocation is chosen as the reference configuration.

<table>
<thead>
<tr>
<th>No. of Segments</th>
<th>Absolute Error $\epsilon_a$</th>
<th>Relative Error $\epsilon_r$</th>
<th>Runtime(sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>27.8</td>
<td>20.05%</td>
<td>406.2</td>
</tr>
<tr>
<td>4</td>
<td>19.7</td>
<td>14.25%</td>
<td>841.1</td>
</tr>
<tr>
<td>6</td>
<td>16.0</td>
<td>8.75%</td>
<td>2932.6</td>
</tr>
<tr>
<td>8</td>
<td>4.97</td>
<td>1.00%</td>
<td>4902.4</td>
</tr>
<tr>
<td>12</td>
<td>0</td>
<td>0</td>
<td>8320.2</td>
</tr>
</tbody>
</table>

dislocation and an atomic size defect cluster, or during the annihilation reaction between two dislocation segments of the same Burgers vector and of opposite tangent vector, large variations of the local dislocation line curvature would be expected. To effectively resolve these or similar mechanisms, we develop here a protocol for adaptively re-distributing of the nodal density on the dislocation line according to the variation in the local curvature. To show the level of resolution gained by this protocol, we study here the mechanism of dislocation segment annihilation in an expanding F-R source, and the subsequent generation of a fresh and closed dislocation loop.

In the annihilation event, the distance between any two nodes is tracked, and once the minimum distance between two segments of opposite tangent vectors is below a prescribed limit (e.g. 100), the two segment will annihilate, and generate two separate loops one closed and one open. The nodes are re-distributed in the immediate region, as can be seen in FIG. 4.6. The simulation conditions are the same as in previous figure, except that the applied stress $\sigma_{11} = 140$ MPa($\tau/\mu = 0.112\%$), and the Burgers vector of the loop is $\mathbf{b} = \frac{1}{2}[0\overline{1}1]$. 
Figure 4.6: Expansion of an initially mixed dislocation segment in an F-R source under the step function stress of $\sigma_{22} = 140$ MPa ($\tau/\mu = 0.112\%$). The F-R source is on the (1 1 1)-plane of a Cu crystal with Burgers vector $\mathbf{b} = \frac{1}{2}[0\bar{1}1]$. The time interval between different contours is $\Delta t^* = 5 \times 10^5$. 


As shown in the figure, after the annihilation event takes place, both new loops generate cusp regions, where the curvature is extremely high. In our computational strategy, we developed an adaptive scheme to resolve such an essential physical phenomena for sufficient accuracy without excessive computations.

In present algorithm, at prescribed time intervals, there will be a nodal redistribution process, it may not necessarily related to the integration time step. During the redistribution process, each cubic spline segment is divided into several sub-segments with equal arc length, and new ghost nodes are assigned. Thus, the entire loop is now filled with ghost nodes of equal nodal density per unit line length. Note that the total number of DOF for the loop is not changed up till this point. Also, the average loop curvature $\kappa_{avg}$ is determined simply as the mean value of the maximum $\kappa_{max}$, and minimum $\kappa_{min}$ curvatures of all ghost nodes. Now in order to distribute nodes evenly according to local curvature, we start from one end of the loop that has a current curvature $\kappa_C$, and skip a number of ghost nodes $N_{\text{skip}}$ determined by the relation:

$$N_{\text{skip}} = f(\kappa_C)$$  \hspace{1cm} (4.7)

Here, $f$ is a function related to current curvature. We can simply chosen as $f(x) = c\kappa_{avg}/x$, and $c$ is a constance which will be chosen to ensure desired nodes density are put in the dislocation after rearranging.

In our protocol, more nodes are distribute to the zone with high curvature while less in low curvature zone. FIG. 4.6 shows the node re-distribution in each closed loop and its recovering process from the cusp. The total number of nodes in each loop at a given time is generally kept under 25.
4.6 Interaction with SIA cluster atmosphere

FIG. 4.7 considers the interaction of F-R source with a row of 15 SIA clusters. During the F-R source expanding process under the external stress, it will subject to the elastic stress field of these clusters. Each segment of the dislocation line will subject to the attractive or repulsive force, together with its own self-force, external applied stress, the combined effect of all these forces will determine the dynamic process of the motion of the dislocation. As shown in the figure, the 15 SIA clusters each with diameter of 40 is located above the gliding plane at the distance of 50 with the same inter-cluster distance of 100 and normal direction [011]. The external stress $\sigma_{11}/\mu$ or CRSS($\tau/\mu$) is added gradually from 0 with an interval of $\Delta\tau/\mu = 0.008\%$. At each stage, it reaches a stable state as shown in FIG.4.7a. The clusters generate a repulsive stress field, and hence at low stress, the F-R source cannot go through the row of clusters, the dislocation line generated an unsymmetrical configuration. When CRSS is increased to 0.0984%, it starts to break away from one side, and once one side break through the effect of the cluster, due to the repulsive effect of the cluster, it will help other parts to break away quickly, it is a speed up process, and the whole breaking time is about a few ns as shown in FIG.4.7-b.

It is shown in FIG.4.7 that, when the dislocation approaches the cluster, the dislocation generated a wavy configuration, especially the part near the effect field of the cluster. The details are shown in FIG. 4.8. In FIG. 4.8a,b,c, the whole dislocation line is initially equally divided to 6, 18, 30 segments respectively. During the unlocking process, as shown in the figure, with more segments, the length of the segment is less than the inter-cluster distance, thus, the dislocation line can adjust itself more easily, and hence the dislocation is more flexible, configuration is more wavy. When the segment number is decreased to 6, the
Figure 4.7: Dynamics of dislocation *unsymmetrical unlocking* mechanism, from a cluster atmosphere of 15 equally distributed sessile interstitial clusters with diameter 40, stand-off distance 50 and inter-cluster distance 100. (a) Equilibrium state with equal shear stress interval 4 MPa $(\Delta \tau / \mu = 0.008\%)$. (b) Unlocking state at stress state $\sigma_{11} = 120$ MPa $(\tau / \mu = 0.0984\%)$ with equal time interval $\Delta t^* = 1 \times 10^5$. 
Figure 4.8: Comparisons of different nodal distribution of the details of unlocking mechanism. (a) 6 Segments. (b) 18 Segments. (c) 30 Segments.
segments are much longer than the inter-cluster distance, thus it can not sense the variation of the stress field of the clusters, and consequently, no local wavy configuration is generated, the precision is decreased. Compare all the previous results, it is interested to note that, for complicated shape (i.e. unstable F-R source, interaction with clusters, and so on), although less segment can also get convergence shape, their overall accuracy may not acceptable, more nodes are needed.

### 4.7 Formation of PSBs

The spontaneous formation of dislocation patterns is one of the most striking features of plastic properties of ductile crystalline solids at the micro scale. These patterns consist of alternating dislocation rich and dislocation poor regions usually in the $\mu$m range (e.g. dislocation cells, subgrains, bundles, veins, walls, channels). They are widely believed to be a result of collective interaction phenomena between complex dislocation configurations. Nonetheless, such collective phenomena play a prominent role in determining the general characteristics of plastic deformation, fatigue, and fracture properties of the ductile materials. The pattern formation during the cyclic deformation of metal single crystals has been aimed at explaining the occurrence of the so called matrix and persistent slip bands (PSB) structures. Regarding this, a sweeping mechanism [59] has been proposed. The basic model of the sweeping mechanism describes that during interaction between a glide dislocation segment and a dipolar loop (DL), the DLs are continuously swept away by the cyclic motion of the dislocation.

In the rigid body approximation, the equation of motion for the center of a
Figure 4.9: Interaction between a screw dislocation and a mobile dipolar loop under an external shear stress $\tau = 4$ MPa. Note that the scale on the axes is different. (a) Relative position of the dipolar loop and configuration of the dislocation at 0.88 ns and 2.83 ns, respectively. (b) Loop position and velocity as functions of time.

The loop can be written as:

$$\frac{dx^{DL}_i}{dt} = f^{dis} + \sum_{j\neq i} f^{DL}_j - f^{friction}$$  \hspace{1cm} (4.8)

Here, $x_i$ stands for the position of the $i$th dipolar loop center along the axis of its glide cylinder. $f^{dis}$, $f^{DL}_j$ and $f^{friction}$ are force components along the glide direction generated by the stress fields of the glide dislocation, other dipolar loops, and lattice friction [59], respectively.

Dipolar loop forces deform the dislocation line, and this in turn, produces stress gradients in the vicinity of the loop itself. Thus, the mere presence of a dipolar loop close to a straight dislocation generates forces on the loop tending to move it along. Once the effective total force on the dipolar loop exceeds the lattice friction force (the friction stress is assumed to be 8 MPa [59]), the loop moves along its glide cylinder, as governed by EQN. 4.8, and consequently changes
the force distribution on the dislocation again. FIG. 4.9 shows the simultaneous motion of a dipolar loop and the dislocation as obtained from the numerical integration of EQNs. 4.8 & 4.6. The projection of the configuration on the slip plane is shown in FIG. 4.9-a, while the position of the dipole center along its glide cylinder is given in FIG. 4.9-b.

The dipolar loop is swept away from its original location by the stress gradient generated from the curved dislocation. At the beginning of the process, the stress gradient is low, thus the velocity of the dipolar loop is low. After about 0.5 ns, the dipolar loop forces result in higher dislocation curvature. Thus, the loop is driven away faster with an increasing velocity. At about 2.5 ns, the loop reaches a terminal velocity of about 75 m/s, as shown in FIG. 4.9-b. During the simulation, the two ends of the dislocation are pinned. In order to remove boundary effects of the glide dislocation, we choose a long dislocation with an initial length of 4 µm.

Under cyclic loading conditions in FCC metals, the intimal vein structure is gradually replaced by the self-organized structure of PSBs. To shed some light on the possible cause of this transition, we perform here computer simulations for the dynamics of the collective motion of an initial random distribution of dipolar loops of the same Burgers vector. The dislocation is pinned at its ends, representing a Frank-Read source, and is cycled by the applied stress. The loops are initially randomly distributed in the vicinity of the dislocation slip plane, in the range of distances -300 nm ~ 300 nm, as shown in FIG. 4.10-a. The dislocation line is very long (4 µm), so as to remove the effects of pinning boundary conditions on dipolar loop dynamics. Successive configurations of this system are shown in FIG. 4.10-b through FIG. 4.10-d. After a critical number of cycles, dipolar loops separate into two clusters, and both groups move in the negative direction. The
Figure 4.10: The relative configuration of 20 dipolar loops at the end of different cycles: (a)-initial, (b)-5\textsuperscript{th} cycle, (c)-10\textsuperscript{th} cycle, (d)-15\textsuperscript{th} cycle
dynamics of the interaction process is displayed in a trajectory plot, shown in FIG. 4.11. Two well-separated loop clusters are formed after a number of cycles of the order of 20, as can be seen in FIG. 4.10-d and FIG. 4.11. Some loop pairs are strongly coupled together, retaining their relative positions throughout the entire simulation (see the trajectories of two loops at the lower end of FIG. 4.11), while most others execute complex trajectories that cannot be predicted a priori. Nevertheless, the final configuration is represented by dipolar loop clusters swept into two well-separated groups. The glide dislocation then moves in the loop-free channel.

4.8 PDD in Nonlinear Equation

Once the dislocation mobility changes at different temperature, and the velocity-stress law no longer obey the linear relation 4.3, the computational scheme of PDD should be slightly modified.
Similar as EQN. 4.3, the variational form of EQN. 3.2 in three dimensional form can be written as

$$\int_{\Gamma} \left[ \mathbf{v} - A \left( \frac{\tau}{\tau_0} \right)^{m(T)} \exp \left( -\frac{Q_{\text{dis}}}{k_B T} \right) \mathbf{n} \right] d\Gamma = 0 \quad (4.9)$$

Here, the integration is along the dislocation line $\Gamma$, $\mathbf{n}$ is the unit normal vector of the glide plane. The shear force $\tau$ is defined as $\tau = f_g/b$, $f_g$ is glide part of the resultant line force, and $b$ is the length of Burgers vector. With the same discretization method, similar governing equation as EQN. 4.6 can be obtained. The time integration scheme is the same as that of previous part.
CHAPTER 5

Three Dimensional Crack Simulation with Discrete Dislocation Representation Method

The distribution method originated in the investigation of the similarities of the crack and groups of pileup dislocation. Using classical orthogonal polynomials, Eshelby, et.al. studied the distribution of discrete dislocation pileups under external loading [25]. Under simple loading condition, they obtained the distribution function of the dislocations, and compared the stress field due to these dislocations with that of a crack with freely slipping surface, the same type of relationship of stress vs. distance to the tip as in the crack are obtained. Later, Leibfried [62] introduced a concept of continuous distribution dislocation to avoid solving complicate discrete distribution function. In his method, the slip plane is supposed to contain a smeared-out sheet of dislocation density, with an amount of \(bf(x)dx\) of Burgers vector between \(x\) and \(x + dx\), where \(x\) stands for the position in crack surface in 2-D. Thus the problem is converted to solve the following singular integral type equation:

\[
P \int_{-\infty}^{\infty} \frac{f(x')dx'}{x' - x} = F(x) \tag{5.1}
\]

Here, \(P\) and \(F(x)\) is determined by the boundary condition and applied load.

Friedel[28, 29] is the first one who introduced the continuous distribution of freely climbing crack dislocations to describe a crack with normal displacement
discontinuity. Bibly and Eshelby [4] pointed out that unlike the crystal dislocation, the crack dislocation is the natural discontinuities across the unwelded cuts when a body is stressed, while in the crystal dislocation, the cuts are being subsequently rewelded.

With the 2D continuous distribution method, the crack problems in elastic-plastic solid has been well solved analytically [61, 88], where the complicate situation of the image stress calculation have been avoided. But due to the complicate loading condition, the crack can not be treated in 2D. A 3D technique closely related to the eigenstrain method is developed by Mura [68], which itself based on earlier work of Eshelby [24] in solving ellipsoidal inclusion problem. In this method, the 3D crack plane are assumed to be filled with infinitesimal dislocation loops but with different densities everywhere. The stress field of this infinitesimal loop can be obtained analytically [12, 69], by solving the similar type equations as EQN. 5.1 but with hyper-singularity:

\[
\int_{-\infty}^{\infty} \frac{f(x', y')dS}{r^3} = \sigma^0(x, y)
\]

Here, \( r = \sqrt{(x - x')^2 + (y - y')^2} \) and, \( S \) is crack surface, \( \sigma^0(x, y) \) is a known function related to load and boundary condition.

5.1 Discrete Model in 3D cracks

In stead of solving EQN.5.2 which needs FEM method or other complicate technique [92], we here propose another discrete method based on Parametric Dislocation Dynamics.

According to Bueckner’s Principle [11], the stress state around a general 3-D crack in an elastic body subjected to general external loading is identical to
the summation of the following two parts:

1. The stress generated if the same external loading acting on the same solid but assuming the crack plane is welded

2. The stress from the same crack in the body subjects only to the internal pressure, or the the counterforce acting on the same plane generated in previous step.

Thus, it can be divided into two steps to solve a general crack problem in elastic solid. First, under the applied load $f_a$ and displacement constraint $u_a$ as shown in FIG. 5.1, and assume the original crack is welded, obtained the stress distribution $\sigma_1$, and then the force vector along the crack plane $f_c$. The second step is to apply $-f_c$ on the crack plane and solve for the stress distribution $\sigma_2$. The final stress distribution is thus obtained by summing up $\sigma_1$ and $\sigma_2$ together.
We here try to use the discrete dislocation distribution method in the second step.

Following Bibby and Eshely’s idea, the general 3D plain crack surface is assumed to be filled with layers of material strips then welded the crack and applied the load, thus produce a group of dislocation loops along the crack surface. The stress state in this solid $\sigma^{\text{combine}}$ is the combination of the applied load and stress from these dislocations. Note that the acting force along the original crack plane can be adjusted to zero by way of adjusting the position or thickness of these dislocation loops, then cut the welded surface, the overall stress state in the solid will not be changed since boundary condition is not changed. According to the Uniqueness theorem, the stress distribution of the crack system is the same as $\sigma^{\text{combine}}$ produced by previous steps. In this way, the problem is now to solve the distribution function of the dislocation loops.

By way of minimization of the total elastic energy of the whole system, which will lead to the variational form of the governing equation of motion of dislocation (EQN. 4.3), the distribution of the dislocation can be easily obtained. Thus by way of the dislocation dynamic process, one can obtain the final configuration when each dislocation loop reaches its equilibrium state, with the whole system reaching the minimum free energy. Due to the complexity of the stress field and the shape of the crack, an accurate method is needed to obtain reliable quantitative data on the process of the energy minimization. For that reason, the parametric dislocation dynamics (PDD) method is employed here for the simulation.

Note that either solving the length of the Burgers vector or its configuration can achieve a balanced state, to simplify the problem, we here set the value of
the Burgers vector $b$ of the moving dislocation as:

$$b = \alpha f_c$$

(5.3)

Here, $\alpha$ is a constant related to the selection of the length of the Burgers vector. $f_c$ is the force acting on the crack plane as defined in FIG. 5.1. It is interestingly to point out that $f_c$ is not uniform in the crack plane especially in general 3-D loading, hence the crack dislocations in generally is of Somigliana type.

we here provide a dynamic adding-deletion strategy for the simulation to obtain the final distribution of the dislocation, it is summarized as follows:

For a given crack, an initial distribution of 3 equally separated dislocation loops are put on the crack plane. The shape is chosen according to the crack plane shape with the outermost one chosen as the crack surface boundary and fixed during the whole simulation (Here we assume there is no crack propagation.).

Under the combined effect of their mutual interactions, stress from the applied load and their self-force, the crack dislocations will either move stretch toward the outermost one if the effect of the applied load is dominant, or shrink if its self-force is dominant, until reach equilibrium state. In the simulation process, new loops are continuously adding to the simulation, and old collapsed ones are taken out of the simulation, the strategy is:

**Adding:** In the adjusting process, once the innermost loop expands when the repulsive force from other loops is less than the applied load, its characteristic length $r$ will be larger than a critical value $r_0$, a fresh new loop is generated in the center. $r_0$ can be given by the equilibrium position of the loop under the applied load and self-force, or a very small set value. The new loop have the same property as the old one, except its Burgers vector is chosen, locally by EQN. 5.3. The total number of crack dislocations is increased by one.
Deletion: The repulsive force to the innermost loop is too large, making its characteristic length $r$ smaller than $r_0$. Notice that in this case its self-force makes the loop contract, or $r$ smaller, and finally to zero. This loop needs be taken out from calculation. Thus, the total number of crack dislocation in the crack plane is decreased by one.

By way of this dynamic adding-deletion process, a final equilibrium distribution of dislocation loops can be obtained easily.

5.2 Numerical Simulation of General 3-D Crack

We analyze here the stress field of different shape of cracks under either mode-I or mode-II load, and check the numerical precision of the distribution method.

5.2.1 Penny-shaped crack

Stress distribution in an infinite solid containing a penny-shaped crack under arbitrary loading has been extensively studied past. Sack [76] and Sneddon [83] first solve the problem of penny-shaped crack opened by pressure applied over its surface in an infinite elastic solid. Collins [16] later investigated further on the stress distribution of penny-shaped crack in thick plate. Kassir and Sih [52] studied elliptical crack under arbitrary loading, and compare the results of stress intensity factor under shear loading with different loading directions. A comparison between the dislocation distribution method and analytical method will be given below.
5.2.1.1 Pure shearing

Considering a penny-shaped crack with diameter of $2r_0$ as shown in FIG. 5.2a, under constant pure shear loading along $\sigma_{xz} = \tau$. We constructed the local coordinate system on the crack plane as shown in the figure. An initial distribution of dislocation is put inside the crack surface with 3 equally spaced circular dislocation loops, including the outermost static one which prevents all the others moving outside. The applied load generates a uniform shear stress field $\sigma_{xz}$. According to EQN.5.3, the Burgers vector can be chosen as:

$$b = bi$$

Here, $b$ is a constant number since $\tau$ is constant.

Figure 5.2: Distribution of crack dislocation loops of penny penny-shaped crack under mode-II loading with $\sigma_{xz} = 200$ MPa. (a) illustration of local coordinate system. (b) Final distribution of crack dislocations.

FIG. 5.2b illustrates the final equilibrium distribution of crack dislocations.
under the applied shear load $\tau = 200$ MPa. In the simulation, the constant $b$ is chosen as 0.4 $a$, the shear modulus $\mu = 50$ GPa, Poisson’s ratio $\nu = 0.31$, and crack radius $r_0 = 1000$ lattice constant $a$. As shown in the figure, totally 9 dislocations are fitted in the crack plane by using adding-deletion strategy. The energy of a screw segments is lower than that of a edge one per unit length, as a consequence, the self-force of screw component tends to be higher or stiffer than that of a edge one, thus the edge components tend to stretch more than screw ones as shown in the figure, and an oval configuration is generated for each loop. Due to the effect of the outmost circular loop, the ratio of the long to short axis of the oval loop decreases from inner to outer, as the effect field of the outer dislocation becomes stronger. It is shown in the figure that the minimum distance of the two outmost loops is within 10 $b$, in order to capture their mutual interaction in high precision, more Gaussian quadrature points are used in the line integral of stress calculation.

FIG. 5.3a shows the distribution of $\sigma_{xz}$ as a ratio of applied shear load $\tau$, when compared with the analytical solution obtained by Kassir and Sih [52], the relative error along $y$-direction is shown in FIG. 5.3b. Within a small range of distance to the tip ($\approx 2 \%$ of $R_0$), the stress drops quickly, but after that the stress goes to a saturated value which equals to the applied load. When the distance is far away from the crack tip, the stress is not proportional to $r^{-1/2}$ as that in two dimensional crack. When the distance is less than 0.5% of $r_0$, the relative error is larger than 10%. The main error comes from the the Fast Sum Method which is used in the stress calculation. As mentioned in [39], the relative error of stress tends to be higher when the field point is approaching the dislocation loop. More segments or Gaussian quadrature points are needed for higher precision. In our simulation, when the distance is extremely small (i.e. within a distance of 0.5 % of the radius to the crack tip), the error is inevitably high, since only 16 points
Figure 5.3: Comparisons of the stress component $\sigma_{xz}$ along $y$-direction from the final dislocation distribution, the same loading condition as that in FIG. 5.2 (a). The distribution of $\sigma_{xz}$. (b) The relative error of $\sigma_{xz}$.

are used for the integration for saving CPU time. When the distance increased to 1% of the radius, the relative error is decreased to 5% which is acceptable in numerical simulation.

5.2.1.2 Pure tension

In previous section, shear loading is applied to the penny-shaped crack, the only motion of the crack dislocation is glide along its glide plane. But when the same crack subjects to tensile loading, $\mathbf{b}$ is chosen uniformly according to EQN. 5.3 as follow:

$$\mathbf{b} = b \mathbf{k}$$

The direction of the Burgers vector is chosen normal to the glide plane, thus it is the climbing part of the force drives the motion of the dislocation. With
the same adding-deletion strategy, Fig. 5.4 shows the relative error of $\sigma_{zz}$ along both $z$ and $x$ direction, when compared with the analytical results [83]. With the increase of the length of Burgers vector, the mutual repulsive force is increased, hence, there is less number of dislocation in the final distribution, and as a result, there is a higher relative error in stress computation. When the length is 0.8 a (or 0.08% of radius $r_0$), a relative lower number of dislocation with 4 is enough for the representation of the crack, but the relative error is high, for the distance of 0.5% $r_0$ to the tip, it is 91.00%. When $b$ is decreased to 0.2 a (or or 0.08% of the radius), 18 dislocations are needed. As a result, the error is lowered to 7.65%. In this tendency, when the length is decreased, more dislocations enter into the computation, and when it approaches zero, a continuous distribution of dislocations can be obtained, and as a result, the error can be lowered to a neglected value, but the price for smaller $b$ is the sharp increase in the CPU time. As discussed in [48], CPU time of the interaction is proportional to $N^2$, where $N$ is the total number of segments in the dislocations.

Since each dislocation stands for a displacement jump which equals to the length of its associated Burgers vector, the byproduct of this simulation is that we can reconstruct the crack opening shape according to the distribution of the dislocation loops. The outermost loop stands for zero displacement, since its function is to confine the motion of the dislocation. When we count inwardly, the displacement increased one level for each crack dislocation, with a value $b$. Fig. 5.5b shows the final crack opening shape by way of reconstruction. The comparison of crack opening obtained by dynamic DDD with analytical results along the diameter direction is shown in Fig. 5.5a. When $b = 0.2$ a (or 0.02% of radius $r_0$), the number of dislocations is 18, the maximum relative error is 3.3% in the center. Since in the simulation scheme, the distribution is discrete, due to the self-force of each loop, no loop can exist in the center except
Figure 5.4: Comparisons of relative error of $\sigma_{zz}$ for penny-shaped crack under external load $\sigma_{zz} = 200$ MPa. (a) along radial direction, (b) along the vertical direction from the center

when the length of Burgers vector is chosen as zero, which means the continuous distribution. But with the decreasing of $b$, the maximum relative error for crack opening displacement (COD) can be decreased to an acceptable level, and the crack shape is more like an ellipsoid. That’s the why it is flat in the center shown in FIG. 5.5b.

5.2.2 Effect of Non-uniform Stress

When there are non-uniform applied stress directly acting on the surface of the same penny-shaped crack as that in FIG. 5.4, the shape can no longer preserve an ellipsoid configuration. Both FIG. 5.6 and FIG. 5.7 show the different crack shape recovered from the 3D DDD calculation. An decreasing pressure from the center is applied along radial direction as shown in FIG. 5.6a, the pressure is
Figure 5.5: Comparisons of crack opening displacement (COD) with the same condition as in FIG. 5.4 except the density of crack dislocation $n=18$. (a) The COD along diameter. (b) Recover of the crack opening shape in three dimension.

chosen as

$$\sigma = \frac{p}{r}$$

where $p$ is a constant value, $r$ is the distance to the center. $r_0$ is the crack radius. To avoid singularity in the center, a cut-off distance of $0.2 r_0$ is chosen here. And an increasing pressure

$$\sigma = \frac{p}{r}$$

is applied in FIG.5.7. In these two cases, the Burgers vector of each crack dislocation is different, and changes according to its position. Since a higher pressure is applied in the center, the center part of the crack penning shape is more obtrude as shown in FIG. 5.6, and more flat with a lower stress as shown in FIG. 5.7
Figure 5.6: The same penny-shaped crack as in FIG. 5.4 subjects to pressure inside crack surface with a decreasing gradient. (a) the applied stress. (b) crack opening shape

5.2.3 2-D straight crack

Two dimensional cracks has been extensively used in Ductile-to-Brittle transition simulation, hence it is more important to obtain the crack dislocation distribution under different loading condition. Notice that once the ratio of crack length to its width is very larger, stress field of its center part can be viewed as that of a 2-D crack, hence we will try to use the same method to simulate a 2D crack.

5.2.3.1 Simple loading

Considerer a straight crack with a length $2r_0$ in $x$-direction, but with finite width $20r_0$ in $y$-direction. The normal of the crack plane is chosen along $z$-direction as illustrated in FIG. 5.8a.
Figure 5.7: The same penny-shaped crack as in FIG. 5.4 subjects to pressure inside crack surface with an increasing gradient. (a) the applied stress. (b) crack opening shape

Unlike the penny-shaped crack, the boundary of the crack surface is a rectangular, and the two end in y-direction is open, the dislocation loops are assumed to be dipolar loops, the two long arms are F-R source type dislocations with opposite line directions, their 2 ends can move along x-direction, the Burgers vector of the two short arms assumed to be zero, since there are no displacement jump in the two end boundary.

When the crack subjected to the an applied tension $\sigma = 200$ MPa along $z$-direction, and choose $r_0 = 200 \ a$, the distribution of crack dislocation is shown in FIG. 5.8b. The middle part ($-2r_0 \leq y \leq 2r_0$) of the dislocation is perfect straight, and stress field in this region can be viewed as that in the 2-D crack. As a results of the distribution, the crack opening shape can also be recovered as shown in FIG. 5.8c. The relative error of $\sigma_{zz}$ in the center along $x$ direction from
Figure 5.8: Distribution of the crack dislocation of straight crack under mode-I loading with applied tension 200 MPa along $z$-direction. (a) the projection view of the crack. (b) Distribution of crack dislocation, the length of Burgers vector is chosen as 0.15 a. (c) Recovery of crack opening shape in three dimension
the crack tip is shown in FIG. 5.9. With the increase of number of dislocation pairs, the relative error drops significantly. With only 7 dipolar loop, the relative error is less than 2% when the distance is larger than 5 a (or 2.5 \% r_0).

Figure 5.9: Comparison of relative error of \( \sigma_{zz} \). The same condition as in FIG. 5.8

5.2.3.2 Mixed mode

When the same crack subjects to mixed mode I & II loading, the results is shown in FIG. 5.8a. The applied tension is chosen as 200 MPa, and shear stress 100 MPa, thus the stress tensor on crack surface is:

\[
\sigma = \begin{pmatrix}
0 & 0 & 100 \\
0 & 0 & 0 \\
100 & 0 & 200
\end{pmatrix} \text{ (MPa)}
\]
thus the Burgers vector of each crack dislocation is of the form:

\[ \mathbf{b} = \begin{pmatrix} 0.05 \\ 0 \\ 0.1 \end{pmatrix} \mathbf{a} \]

The Burgers vector has both in-plane and out-of-plane component, thus the motion of the dislocation is the mixture of climbing and gliding.

Figure 5.10: Straight crack under mixed model I & II loading. (a) Crack dislocation distribution. (b) Comparison of relative error of \( \sigma_{xz} \) with different inclination angle.

The final distribution of crack dislocation is shown in FIG. 5.10a, there are totally 11 crack dislocation pairs in the field. FIG. 5.10b shows the comparison of the relative error of \( \sigma_{xx} \) with different inclination angle to the crack plane. As shown in the figure, when the distance is larger than 10 a, the relative error is decreased within 5%.
CHAPTER 6

Dislocation Activity ahead of Crack Tip

Dislocation activity ahead of crack tip plays a dominant role in the investigating of Ductile-to-Brittle transition effect in BCC single crystal materials. Dislocations ahead of crack tip basically have two toughening effects, i.e., the shielding and blunting effects, which essentially increased the applied load necessarily to propagate the crack. In this chapter we will study the dynamically development of dislocation population in the vicinity of a loaded crack tip as a aid to understand the ductile-to-brittle transition.

6.1 Three Dimensional Elastic Dislocation Interaction with the Crack

The free crack surface result in the additional stress (or image stress), applied on the dislocation. On the other hand, this image stress decreases the crack tip fracture toughness. Many analytical models have been proposed to study the shielding of a pure edge or screw dislocation in two dimensional crack[64, 85], and its analytical estimation of shielding effect of single dislocation (i.e. EQN. 3.10) has been well utilized in previous 2-D simulation(e.g. [74]). Devincre and Roberts [21] provided an estimation of the local fracture toughness that is based purely on the crack opening or closing stress components of the dislocation stress field at the crack tip. In their estimation, if the crack extends into the z-direction, and
the crack plane is the $y=0$ plane, the $\sigma_{yy}$ component of the stress tensor causes crack opening. Then, the local stress intensity factor is written as

$$k^{\text{tip}}(z) = K^{\text{app}} + \sum_{i=1}^{N} \sqrt{\mu b} |\sigma_{yy}^i(zz)| \text{sign}(\sigma_{yy}^i(zz)) \quad (6.1)$$

where $N$ is the total number of dislocation segments and $\sigma_{yy}^i(zz)$ is the $yy$-component of the stress field of the $i$-th dislocation segment at position $z$ along crack tip. This formula gives a crude estimation for the shielding of general 3-D dislocation, but the effects of the crack surface changes due to the existence of the dislocation itself is not taken into consideration. Some other methods based on Boussinesq problem of point loading in a half space [27], or weight function method [33, 32, 34], has been attempted, but both are painstaking and not practical in the real computation.

A new method based on the 3D dislocation distribution method is proposed here. Recall in CHAP. 5, when we solve the 3D crack problem, the external load is the only applied stress, but when there is crystal dislocation loops inside the material, since the nature of the stress field and the external load are the same, it can be treated the same as the external load. The Burgers vector of crack dislocation is thus modified as:

$$b = \alpha (f_a + f_d) = b_a + b_d \quad (6.2)$$

Here, $f_a$ and $f_d$ are the counterforces of force vectors generated by the applied load and crystal dislocation respectively as described in FIG. 5.1. $b_a = \alpha f_a$ is the contribution to the Burgers vector by the applied load, usually it is constant on the crack surface, $b_d = \alpha f_d$ is the part contributed by the crystal dislocation, it is usually not uniform on the crack surface, and it decreases with the increase of the distance to the crystal dislocation. Since the stress field of the dislocation decreases quickly, the effective area to the Burgers vector of the crack dislocation is very small.
With the same procedure of obtaining the final distribution as that in CHAP. 5, one can obtain the overall stress field under both the external load and the crystal dislocations. The stress field from such a distribution can be also divided into two parts, one is the original stress from crack under external loading, and the other is the extra part due to the crack free surface, and this part is the so-called image stress.

The local fracture toughness at each point along the crack tip can be easily obtained from this stress distribution by:

$$K = \lim_{r \to 0} \sigma(r) \sqrt{2\pi r}$$

(6.3)

Thus the shielding effect of the dislocation $K^d$ can be obtained by

$$K^d = K^a - K^t$$

(6.4)

Here $K^t$ is the total toughness obtained by the crack dislocation distribution, and $K^a$ is the applied fracture toughness. When $K^d$ is larger than zero, the overall toughness is decreased, hence the dislocation has shielding effect on the crack, on the contrary, when $K^d < 0$, it has anti-shielding effect to the crack.

As a comparison, we here use the 3-D model to simulate the shielding effect of an 2D straight edge dislocations ahead of a finite slit crack, and compare that with the analytical results obtained by Wang and Lee[85]. Suppose a finite crack with length $2r_0 = 1000$ a, the width is sufficiently long of $20r_0$, its tip along [010]-direction and (001)-plane as its crack plane, a straight edge dislocation with $b = \frac{1}{2}[101](\bar{1}0\bar{1})$ is introduced to the calculation. It sits on the slip plane (101) with a distance $d$ to the crack tip. Before the introduction of the edge dislocation, the crack plane is already filled with equilibrium state crack dislocations at the applied stress $\sigma = 200$ MPa. With the existence of the dislocation, the equilibrium state is broken, the distributed dislocation will re-adjust their config-
uration till the new equilibrium state. From this new equilibrium distribution of crack dislocation, the image stress as well as its shielding effect can be obtained.

Figure 6.1: Interaction of an edge dislocation and a straight crack in 2D. (a) relative position of crack and the edge dislocation, \( r_0 \) is the half width of the crack. (b) comparison of fracture toughness with the analytical solution.

It is shown in FIG. 6.1 that with the increase of the the distance to the tip, the shielding effect decreases, and the numerical results is about 2 times of analytical results. This is due to the consideration of crack surface roughness generated by the existence of the crystal dislocation, and hence the numerical results have a higher shielding effects.

Under the same condition as in FIG. 6.1, when the straight dislocation is replaced by a circular dislocation loop with \( b = \frac{1}{2}[101] \), the shielding effect is shown in FIG. 6.2. The distance of the center of the loop to the crack tip is chosen as 100 \( a \). The the stress intensity factor of the loop with the change of the radius \( R_0 \) is shown in FIG. 6.2b. The contour plot of the \( \{33\} \) component of
the stress tensor of the shear loop $\sigma_{33}$ is shown in FIG. 6.2a when $R_0$ is chosen as 50. $\sigma_{33}$ is always negative, having a tendency to close the crack, thus it has a shielding effect to the crack. When the radius is increased, the shear loop is more close to the tip, thus it has a higher shielding effect. And since the stress field of the loop can only affect a small area, the shielding effect is within a small zone. The effective shielding area is within $-500a \sim 500a$ as shown in the figure.

![Image](image.png)

Figure 6.2: Shielding effect of a shear dislocation loop ahead of crack tip. (a) contour of the $\{33\}$ component of the stress tensor by the shear dislocation loop. the radius $R_0$ is 50 $a$. (b) comparisons of the shielding at different radius.

### 6.2 Motion of Dislocation ahead of Crack Tip

Dislocation motion ahead of crack tip is quite complicate, and it is a key issue to understand the change of fracture toughness.

Suppose a straight dislocation emanated from a finite crack tip and starts to move away from the tip, FIG.6.3 shows its configuration and corresponding
Figure 6.3: Motion of dislocation half loop ahead of crack tip. The initial size of the loop is chosen 400 a. (a) configuration of the dislocation at different time. (b) the corresponding shielding effect of the dislocation to the crack.
shielding effect to the crack. The size of the crack is chosen as \(2r_0 = 1000a\), and (100) is its crack plane. The uniaxial applied stress is \(\sigma_{11} = 3.0\) GPa (or \(K_a = \sigma_{11}\sqrt{\pi r_0} = 2.11\) MPa\(\sqrt{m}\)). The Burgers vector of the crystal loop is chosen along (111)-direction with length 0.274 nm and its glide plane is (101) with an angle of 45° to the crack plane. The length of the loop is initially chosen as 400 a with its two ends at (-200 a, 15 b), (200 a, 15 b) respectively. The two ends are confined to slide along the crack tip direction. b is chosen as 0.274 nm. The material constant \(m\) is chosen as 2, the temperature is chosen as 450 K, so the force-velocity relation EQN. 3.2 can be rewritten as:

\[
v = M \left(\frac{\tau}{\tau_0}\right)^2
\]

(6.5)

Here, M is effective mobility and \(M = 2.47 \times 10^{-3}\) sec/a, \(v\) is the nodal velocity.

The effect of the free surface of the crack dislocation, or the image stress tends to pull the dislocation back to the crack, thus it has a shielding effect. In order to avoid being totally pulled back, the initial distance of the dislocation to the crack tip is chosen as 15 b. Under the driving force of the crack, the dislocation starts to move away from the tip. At its early stage, when \(t < 3.5\) ms, due to the image stress, the total stress applied on the dislocation in the center part is much lower than that at the two ends, thus its speed is much lower, and a concave configuration is generated. With the dislocation continually moving away from the tip, the effect of the image stress decreases, and the concave configuration gradually disappeared as shown in FIG. 6.3a when \(t = 3.88\) ms. After that, due to the fast decaying of the crack stress, the speed of the nodes in the center part is still lower than that at the two side, thus a flat roof is generated and the flat roof becomes more longer during the motion. The shielding effect \(K_d\) of the dislocation is shown in FIG. 6.3b. It initially has a maximum shielding effect of around 50 % of \(K_a\) at the center, but after 0.59 ms, the shielding of the center
Figure 6.4: Contour of the \{11\} component of the stress tensor from crack tip due to the applied stress and image stress. (a) t=0, (b) t=0.59 ms, (c) t=1.59 ms, (d) t=2.50 ms, (e) t=3.42 ms, (f) t=4.35 ms
is only 27% of $K_a$. Due to the stretch of the dislocation, the dislocation can cover more part of the crack tip field, thus more area of the crack tip fracture toughness is lowered. FIG. 6.4 shows the contour of $\{11\}$ component of the total stress (original crack tip stress plus the image part) from the crack at $t=0$, 0.59 ms, 1.59 ms, 2.50 ms, 3.42 ms, 4.35 ms respectively. The images stress only affect a small area near the dislocation, and it decreases the crack tip stress field. With the increase of the distance of the dislocation to the crack tip, this effect becomes more and more weak.

As a comparison, when the initial size of the straight dislocation is a small value of 40 a, FIG. 6.5 shows the motion of the dislocation loop and its corresponding shielding effect with a different shape. The initial effect field of the dislocation is very narrow as can be seen from FIG.6.5b when $t<0.01$ ms, and due to its two ends are confined sliding along crack tip, the concave configuration is very small which is different from FIG. 6.3, it is very easily recovered at around $t=0.07$ ms. At the time of recovery, the difference between the maximum distance at the center part and minimum distance at two sides part is only around 5 a. Thus when compared with 83 a in FIG. 6.3, variance of the nodal speed in the dislocation is not very large, thus there is no flat roof in its configuration as shown in FIG.6.5a, the center part starts to protrude out during the motion.

### 6.3 Dislocation Nucleation

In this section, we will focus on the way of the dislocation nucleation around the three dimensional crack. Xu et. al. [90, 91] has studied the saddle-point configurations of embryonic dislocation loops near crack tip and their associated energy with a variational boundary integration method based on the continuous distribution dislocation method. Here, we will directly assume that the nucle-
Figure 6.5: Motion of dislocation half loop ahead of crack tip. The initial size of the loop is chosen as 40 a. (a) configuration of the dislocation at different time. (b) the corresponding shielding effect of the dislocation to the crack.
ation of the embryonic dislocation is easy and can be nucleated immediately at
different site along the crack tip without any cost. The initial configuration of
the embryonic dislocation is assumed to be short straight dislocation segment,
and can only bow out with its two ends sliding along the direction of crack tip,
forming half loops. When the neighboring dislocation half loops meet, they will
coalesce to form a longer one and continue bowing out.

Suppose a semi-infinite crack extends along (010) direction, the crack plane
is chosen as (100). A slip plane is chosen as (101) with an angle of 45° to the
crack surface. The simulation temperature is chosen as 200 K, the initial applied
load is set to 1.3 MPa√m, and the loading rate $\dot{K}$ is chosen as 0.05 MPa√m/s.
The constant $m$ is chosen as 2. Thus the dislocation has the same stress-velocity
relation as EQN.6.5, but the constant $M = 5.14 \times 10^{-6}$ a/sec.

The detailed dislocation nucleation process on the slip plane from the crack
tip and the corresponding loading history is shown in FIG. 6.6 & 6.7 respectively.
The length of the straight dislocation embryonic is chosen as 20 a. In order to
avoid the complicate boundary effect, a periodic boundary condition along the
direction of the crack tip is used during the simulation, the width of the crack is
chosen as 3000 a. Assume that there are five equally separated nucleation sites
with a distance of 600 a between the neighbors. Under the crack tip stress field,
the dislocation embryo begins bowing out with their two ends confined to slide
along the crack tip direction. At each time step, we will check these five sites,
one there is enough space to put a new embryo loop, a new loop is assumed
nucleated, but after this time step, due to the repulsive force of earlier nucleated
dislocations, if the new dislocation is pushed back to the tip, it will be taken
outside the simulation. FIG. 6.6 shows the successive loop nucleation process.
The loops starts to nucleate at $t = 0$, and after around 2.5 sec, new embryo
Figure 6.6: Stages of dislocation nucleation and the coalescing process.
Figure 6.7: The applied and the effective fracture toughness as a function of loading time.

loops generated at the middle three nucleation site as shown in FIG. 6.6a. After another 5 sec, the outer dislocation loops begin meet and merge to form a longer one as shown in FIG.6.6 (b), (c) and (d). The two ends of the longer dislocation are still confined to slide along the crack tip direction. Once the two ends of the longer dislocation loop reaches the crack boundary, it covers the whole crack front, we assume they can move along the slip plane boundary, thus it moves away from the crack as shown in FIG.6.6e when $t > 8.5$ sec.

As discussed before, the short segments can only have shielding effects at its nearby location, the part that not covered by the loops tends to have highest fracture toughness, it has no contribution to the maximum fracture toughness along the crack tip, the crack always first propagates at the site with maximum fracture toughness. We define here the effective toughness as the maximum stress intensity along the crack tip at current time, and denoted as $K_e$. So the shielding
effect of these short dislocations has no contribution to $K_e$, we will disregard this shielding effect during the later calculation, and later when we mention a new loop is nucleated means a such kind of loop is generated.

It can be shown in FIG. 6.7 that $K_e$ is identical to the applied load $K_a$, until after around 9.0 sec when the first long dislocation covers the whole crack tip, there is a sharp drop of around 0.35 MPa√m in $K_e$, that means the dislocation has an effective shielding effect of 0.35 MPa√m. After that, $K_e$ increases with the increase of the applied load. After another 5 sec, the second long dislocation is generated, and there is a further drop in $K_e$ of 0.42 MPa√m. But the effective shielding of the two dislocation is only 0.71 MPa√m, that is because the first dislocation moves further away from the top as shown in FIG.6.6f. Due to the mutual interactions of the already nucleated dislocations, the time for forming a new one is longer and longer. It takes around around 7 and 13 seconds for the generation of the third and fourth dislocation respectively, as shown in FIG. 6.7. All these dislocations forming a pileup on the slip plane.

The lattice resistance to the movement of the dislocation loop, the Peierls stress, or friction stress, plays an important role in the motion of dislocation. The equation of motion EQN. 4.3 should be modified as:

$$v = \begin{cases} A \left( \frac{\tau - \tau_f}{\tau_0} \right)^{m(T)} \exp \left( -\frac{Q_{\text{dis}}}{k_BT} \right) & \tau > \tau_f \\ 0 & \tau \leq \tau_f \end{cases}$$

(6.6)

FIG. 6.8 shows the effect of the friction stress to the dislocation nucleation. With the increase of the friction stress, the speed of generation of new loops decreases. The maximum difference of the time for generation of the 1st loop is around 1.82 sec, the 3rd one is around 1.94 sec, while the difference of the fifth one is around 3.87 sec as shown in the figure. In this tendency, the time gap will increase more and more, after a longer time, there will be a huge difference of the
Figure 6.8: Time for nucleation of fresh new loops at different friction stress. Lines count from bottom to top stands for time for the nucleation of the 1st to 5th loop number of generated dislocations, and as a result there will be large difference of their totally shielding effect, which will lead to the brittle-to-ductile transition.

**6.4 Brittle to Ductile Transition**

Dislocation velocity varies significantly at different temperature under the same applied shear stress as shown in FIG. 6.9 (refer to EQN. 3.2). Under the applied shear stress $\tau=1$ and $5 \text{ GPa}$, the ratio of the highest and lowest velocity is about $10^6$. The huge difference of velocity at different temperature makes it possible for more dislocation generated at the same time interval for higher temperature, and thus have more shielding effects on the crack tip. On the other hand, the huge velocity difference makes the right selection of time step difficult, since the time step should be small enough to accommodate the high speed in higher
Figure 6.9: The variance of the dislocation velocity as a function of temperature under applied shear stress $\tau = 1$ & $5$ GPa respectively.

To obtain the brittle-to-ductile transition curve, FIG. 6.10 shows the flow chart of the simulation procedure, instead of traditional 2D dislocation dynamics method, FIG. 6.11 shows the detailed procedure of three dimensional simulation of dislocation dynamics and crack simulation method in the obtaining the ductile-to-brittle transition curve.

Follow the procedure in FIG. 6.11, FIG. 6.12 shows the rate of the dislocation generation as a function of temperature. The rate of dislocations generation is measured by counting the numbers of crystal dislocations generated within the same time slot. As shown in the figure, at lower temperature, dislocation generation is rather slow, the rate at 400K is only 1% of that at 1000K. Since at lower temperature, the motion of dislocation is rather slow (refer to FIG.6.9), at the same time interval, due to their mutual interactions, already nucleated
Figure 6.10: Flow chart of the simulation of Ductile-to-Brittle-transition curve. Dislocations move slowly, prohibits new dislocation generation. As shown in the figure, there is a sharp increase in the generation rate at $K \approx 800$ K.

As shown in FIG. 6.11, the most time-consuming part is updating the configuration of the crack dislocation at each time step, since the applied load increases and the position of crystal dislocation changed, to obtain correct crack tip stress field, the crack dislocation should be redistributed. Secondly, the time step for the motion of the half loops sliding along the crack tip direction is rather small, which also is a computational burden. Thirdly, as shown in FIG.6.6, the major difference between the 3D & 2D simulation is how the dislocation is generated, in 2D simulation, the infinite long dislocation is automatically generated, while in 3D, the dislocation is formed by coalescing short dislocation half loops nucleated from different site, but after a curtained distance away from the crack tip, the difference in 2D and 3D is very small, the long dislocation can be treated as straight 2-D dislocation as shown in FIG.6.6. The simulation in solely 3D is
Figure 6.11: Flow chart of the 3D simulation of PDD & crack dislocation simulation.
Figure 6.12: The rate of dislocation generation as a function of temperature at different applied load.

very time consuming and is beyond the current computation limitation unless the loading rate is artificially high, which will lead to the mistakenly shift of the ductile-to-brittle transition[73].

As a conclusion of this part, FIG.6.13 shows the ductile-to-brittle transition with 2D simulation method. The critical applied load $k_c$ when $K_e$ reaches the critical fracture toughness, increases as the temperature raise. As shown in FIG.6.13a, when the temperature raised from 300 K to 1100 K, $k_c$ is raised by a factor of 2.5. The decreasing of the loading rate also increase the overall $k_c$ as shown in the figure. When the loading rate $\dot{K}$ is decreased from 0.028 MPa$\sqrt{\text{m}/\text{s}}$ to 0.01 MPa$\sqrt{\text{m}/\text{s}}$, the overall $k_c$ is increased by an average of 6.0 %, which due to the more dislocation generation, and when the loading rate is further decreased to 0.001 MPa$\sqrt{\text{m}/\text{s}}$, there is another 10 % increase in $k_c$. Similar to FIG. 6.7, in 2D simulation, it is also a zigzag curve in the loading history curve as shown in
Figure 6.13: Relative fracture toughness as a function of temperature at different applied load. (a) Brittle-to-Ductile transition curve at different loading rate. (b) the loading history at T=1000 K.
CHAPTER 7

Conclusions and Discussion

The three dimensional nature of dislocation processes ahead of crack tip is quite complex and it is essential to understand the ductile-to-brittle transition in BCC metals.

In current work, several topics related to dislocation theory has been fully exploited.

7.1 Three Dimensional Parametric Dislocation Dynamics and its Convergence and Accuracy

Based on the Fast Sum Method, the present work attempts to present a comprehensive investigations of the the Parametric Dislocation Dynamics Method (PDD) as well as its fundamental mechanism in the simulation of plastic deformations.

Numerical studies of dislocation generation by the Frank-Read mechanism revealed a number of significant features. The shape accuracy of an simple equilibrium F-R source has been shown to be excellent with only a few parametric segments, with only two segments, a maximum relative accuracy of $10^{-3}$ can be achieved. For complicate unstable F-R source, it requires more segments, and six segments produce similar accuracy. The shapes of both equilibrium and unstable
F-R sources were shown to be absolutely converge with the increase of the number of segments. Adaptive node re-distribution and a higher nodal density on the dislocation line regions with high curvature is necessary for the annihilation event and the subsequent loop reconfiguration. The selection of implicit and explicit integration scheme of the system equation of motion EQN. 4.6 should take more consideration. The numerical stability for the implicit one can be always being assured, while for the same numerical stability, the time step should be reduced to less than less than 500 dimensionless units (on the order of 1 ps for a mobility of $10^4\text{Pa}^{-1}\text{s}^{-1}$).

In the case of a finite size dipole, with only 2 segments on each dislocation, the formation and destruction dynamics are resolved with a relative accuracy of less than $10^{-2}$. The formation dynamics reveals that the two interacting dislocations do not feel one another till their closest segments are within $\approx 100\,\text{a}$. It is found that the stress to unzip the dipole is smaller than the breakup stress of finite size dipoles, nearly by a factor of 2. On the other hand, to break up the dipole by pushing the two dislocations past one another (forward destruction) requires a stress that is nearly 30% higher than the infinite dipole.

Studies of the process of unstressed sessile junction formation with PDD, and its destruction by an externally applied stress has been consistent with similar recent studies with other methods. However, less number of required segments for each dislocation is required for spatial resolution of less than $10^{-3}$, and for temporal resolution of $10^{-6}$.

It's also shown that PDD is also a convenient tool for studies of elementary mechanisms of dislocation patterning caused by elastic interactions amongst defects. The method is accurate enough to deal with complex stress fields associated with the dynamic motion of dipolar dislocation loops and glide dislocations. The
present investigation has demonstrated a number of physical processes involved in the sweeping mechanism.

7.2 Three Dimensional Crack Simulation

Two dimensional continuous dislocation distribution method has been extensively studied analytically and numerically by previous work[88, 61, 50]. In the continuous method, the cracks are assumed to be a distribution of loops with infinitesimal length of Burgers vector, and by way of solving a complicated singular (in 2D) or hyper-singular (in 3D) integral equations, one can obtain its distribution function. In this work, we assume that the crack surface is filled with discrete dislocation loops with finite burgers vector. By way of minimization of system free Gibbs energy, PDD is utilized to obtain the final equilibrium distribution and configuration of these dislocation loops under the applied load. The stress field from these distributions is the right answer to the original crack tip field. The outer dislocation of the distribution is assumed to be the contour of the crack surface and being fixed, the rest dislocation loops can be obtained by a dynamic adding-deletion process. The crack opening displacement can be easily obtained from the distribution since each dislocation loop stands for one layer of displacement jump, and hence the crack opening shape can also be easily reconstructed. It is shown that high accuracy can be obtained in the mode I & II loading in penney-shaped and slit crack. The relative error of the stress field tends to be higher when the field point is near to the crack tip, and more nodal points and Gaussian Quadrature points are needed for the same accuracy. The Burgers vector is chosen as proportional to the applied load, such that the changes of the applied load will not changes the dislocation distribution significantly.
7.3 Crack-Dislocations Interactions as well as Ductile-to-Brittle Transition

It is easy to derive a closed formulation for the stress field of a dislocation ahead of a semi-infinite crack in two dimensional, but similar formulation for an arbitrary oriented dislocation loops ahead of the crack tip is beyond the state of the art. Three Dimensional Discrete Dislocation Distribution (DDD) provide us a useful tool to investigate the interactions of crack and dislocation in 3D space. In current work, the stress field of the crystal dislocation can be treated the same as the applied load. Thus modify the equation of Burgers vector, and add the contribution part from the stress field of the crystal dislocation, then rearrange the configuration of the dislocations, one can obtained the changes of the crack tip field due to the existence of crystal dislocations and change of the applied load. As a result of the redistribution, the shielding effect of the dislocation can be obtained.

The motion of dislocation ahead of crack tip is rather complicated. The motion of the dislocation changes its shielding effects to the crack tip, and on the other hand, the changed crack tip field will changes the motion speed of the dislocation. It is shown that different initial size has different behavior of motion, since the crack free surface has a different feedback to the motion of the dislocation. The concave configuration due to the dragging effect of the image stress can be easily recovered when the initial length is short. In the simulation, it is assumed that embryonic dislocation can be generated at separate nucleation site instantly and without any cost. The embryonic dislocation will bow out and once the neighboring dislocation loops meet, they will coalesce and form a longer one, until cover the whole crack front. We assume that only those dislocation covering the whole crack tip has a shielding effect to the crack,
since those short dislocation half loops only have local shielding effect. It is found that the major difference between the 3D and 2D simulation is how the dislocation is nucleated, but it is still important in the understanding of ductile-to-brittle transition behavior, especially when we need to take the friction stress, and other type of material defects into consideration, but we should pay the price for extensive computational time.
REFERENCES


