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A Multi-Scale Model of Intergranular Fracture and Computer Simulation of Fracture Toughness of a Carburized Steel

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ABSTRACT

It is no doubt that material's atomic-electronic structure determines macroscale physical properties; however, broad varieties in microstructures and heterogeneities from different scales may significantly amplify or dilute the mechanical behavior of an alloy presented at quantum scale. Hence, breakthrough of computational alloy's design lies in the capabilities to quantitatively and completely integrate key-mechanisms from different scales at each processing step, so as to obtain a unified procedure to establish quantitative relationships between composition, process, structure, properties, and performance. For this purpose, a multi-scale hierarchical model of intergranular fracture has been developed for polycrystalline systems. As an application example, it has been applied to a carburized steel that is used for gears or other components in heavy power transmission system. Based on the computations at quantum, micro, and macro scales, the prediction of fracture toughness has been obtained in accord with experiments, which reveals that an integranular cracking is mainly triggered by the stress concentration at the junctions of grain boundaries when the adhesion between adjacent grains is low. An improved toughness can be achieved by reducing impurities grain boundary segregation while increasing grains' ductility.

Keywords

grain boundary, interfacial adhesion, first-principle, carburized steel, intergranular fracture, fracture toughness, multi-scale, hierarchical.

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1. INTRODUCTION

A practical issue in condense matter physics is to design atomistic structure and enhanced process treatment, so as to obtain optimized microscale structure with desirable macroscale properties. Quantum mechanics provides fundamental knowledge of the atomic-electronic structures of a material, which is the key to understand the corresponding micro and macro behaviors [1-4, 11, 12, 26]. To investigate properties in general, at sub-atomic scale a transition metal is usually treated as a bulk phase with periodic atomic array obeying Bloch's theorem [67]. However, in the perspective of fracture, a metal's failure is often triggered by lattice's defects. The micro-scale random-distributed heterogeneities such as grains, second phase particles, solution atoms and other point defects, essentially dominate the kinetics of the metal when environment and applied load change [5-9,19,20,23,24,43,75]. Nevertheless, at macro scale a structural evolution, for example, a plastic deformation that is an average of the accumulations of irreversible lattice's distortions and dislocations, is often modeled again under the approximations of uniform and continuous through constitutive relationship in the framework of continuum mechanics. This bottom-up, hierarchical variations in material's structural modeling reflect the challenges remaining for alloys' design, which also reveal the natural complexity of a material. An effective and quantitative analysis procedure to integrate sub-atomic physics with up-atomic mechanisms will open new avenue to explore new materials with desirable performance. In this paper an effort has been made to develop such a procedure to study the failure mechanisms from different scales which may affect the fracture toughness of carburized steels. The fundamental theories of solid plasticity and fracture mechanics can be found in [10,13,14,39]. The literatures regarding computational materials design can be found by, e.g. [4,5,17,20].

Directly quenched after carburizing, carburized steel is used for manufacturing machine components like gears in heavy power transmission systems [19,21,22]. The part near surface in this class of steels is termed "case", which has higher carbon content as compared with the core part. The case microstructure consists of low temperature tempered martensite and retained austenite, which is highly susceptible to intergranular fracture that often results in relatively low fracture toughness. Fig. 1 is a fractography of an AISI 8822 fully carburized specimen with multiple intergranular cracks. To achieve improved performance, it is vital important to clarify the mechanisms that induce intergranular cracking and to distinguish the effects of alloy additions and impurities which cause the transition between intergranular and transgranular failure modes.

Based on the literatures cited in this paper and the authors' previous works, a multi-scale model of the intergranular fracture is proposed and an associated computation-based procedure is developed. This approach focuses on the interaction between grain structure and grain boundary properties, especially the effects of phosphorous segregation [6]. The developed procedure hierarchically implements the quantum mechanical computation into a micro-scale polycrystalline system within plasticity finite elements and cohesive law, which is built into the computation of a laboratory-sized specimen to obtain fracture toughness. As an example of application, the AISI 8822 carburized steel case has been analyzed and the obtained results are compared with experiments.

Reviews and analysis of carburized steels can be found, e.g. in [19, 21-23]. The interactions between dislocation kinetics and intergranular fracture have been discussed, e.g. by [23] in the perspective of physical metallurgy. To link grain boundary with chemical composition and adhesion energies, an impurity segregation model has been developed in [6]. Thermodynamics analysis and more generalized investigations of grain boundaries chemistry can be found, e.g. in [7, 23, 74]. Based on the "universal feature" [26] of atomic bonding [11, 12, 67], interfacial debonding and cohesive models for solid mechanical computations have been proposed in [15, 29] with the applications of bulk phase fragmentation [38] and interfacial crack

propagation [36]. A systematical study of the elastoplastic constitutive model for polycrystalline system has been introduced in [9]. An ab initio computation of grain boundary and comparison with electron microscopy observation of Cu_3Bi grain boundary has been conducted in [74]. Regarding the general issues of grains and grain boundaries, researches and developments have been reported, e.g. in [18, 34, 35, 55, 66] of crystal plasticity, in [6, 31-33] of impurities grain boundary segregations, in [43] of grain boundary creep and sliding, in [6, 7, 23, 31] of intergranular fracture, in [68, 69] of the length scale in plasticity and in [39, 13, 14, 40-42] of fracture mechanics. Literatures of computation science can be found, e.g. in [44-49, 70, 71, 73] of the density function theory [2, 3] based quantum mechanics computation and in [50-53] of finite element analysis.

In recent years great activities can be found regarding multiscale analysis and its application to materials science, for examples, these in [57-63, 20, 28, 72, 76]. In [63] a scheme has been proposed to compute the average heat conductivity of a cell containing aggregated heterogeneous composite inclusions through the cell's surface heat flow and temperature. This model has its counter part in solid mechanics analysis, i.e. the cell model originally developed in [64]. A quasicontinuum method, by which the basic unit is a finite element that contains atoms, has been developed in [60, 61]; the energy potential of the "embedded atomic method (EAM)" [48, 49] has been adopted as the strain energy in the finite element. In the "coarse-grain" method [58] an atomic segregate forms a finite element to represent a thermodynamic system, by which both mechanical deformation and statistic-based thermo-vibration-induced temperature are taken into account. A concurrent scheme between different physical domains in continuum theory has been developed in [62]. A formalism coupling density-function theory-based simulation in one domain to the continuum mechanics simulation in another domain has been developed in [57]. A variations-based scheme to compute many-body quantum system has been reported in [76]. A "Moving Particle Finite Element Method" (MPFEM) has been developed in [20, 25, 27, 30], which combines the salient features of finite element and particles to represent a solid. In order to accurately and efficiently represent atomistic behavior at micro scale, a "Particle Dynamics" (PD) method has been developed in [20, 28]. The MPFEM and PD are the computational methods applied in this study; detailed description of the "Particle Dynamics" and application to bcc iron are given in the sections 2.5-2.7 of [20]. The original concept for this class of methodologies can be found in the Lorentz's original work that establishes the connection between mean field electrostatic theory and microscopic theory, as described by the chapter 27 of [67].



Fig. 1 An experimental observation of intergranular fracture

2. MODEL AND PROCEDURE DEVELOPED

2.1 An Integranular Fracture Model

For an engineering material such as the case of a gear under cycling contacts, its hardness, strength and fracture toughness are the key-properties that determine performance. High case strength and hardness are usually produced by quench and tempering, whereby the tempered martensite is the major constituent that provides surface hardness with enhanced wear resistance. Fracture toughness, which also contributes to fatigue life, represents the resistances against micro-crack initiation and growth. Obviously, grain boundary properties have profound effects on the initial stage of intergranular separations – the phenomenon presented in Fig. 1. Considering a carburized steel case as a system, the strength and fracture toughness of this system are determined by the combinations of chemical composition, phase constituents, and the structural parameters associated with the micro- and nano-scale heterogeneities, such as precipitates, solute atoms and second phase particles in grains, lattice misorientation and impurities segregation at grain boundaries, grain size and morphology, and dislocations density. From the viewpoints of strength and fracture toughness, these parameters can be distinguished into two classes: the heterogeneities inside a grain which determine the mechanical properties of the grain; and the complexities around grain boundaries which determine the interaction between adjacent grains; by the latter grain boundary adhesion is the dominant factor among others.

Hence, an intergranular fracture model, illustrated in Fig. 2, has been developed, by which a macroscale crack propagation is an accumulation of the damage evolution within the small process zone in close vicinity around the crack tip. For an intergranular fracture in carburized steels, evolution of damage is an accumulation of the decohesion of grain boundaries since those martensite-dominant grains are mechanically stronger. This decohesion is governed by the traction-separation law between two adjacent grains, as plotted at the upper right corner of the figure, which essentially is a process to break the bonds between atoms pairs separated by a grain boundary. Therefore, this model schematically establishes the correlations among macroscale crack growth, microscopic damage evolution and atomistic debonding. Fracture toughness is material's resistance against crack growth. Accurate computations of grain boundary adhesion and grain's mechanical properties are the necessary pre-conditions for obtaining a prediction of fracture toughness. The associated idea for computational carburized steel's design is conceptualized by the flow chart in Fig. 3.



Fig. 2 A proposed multi-scale intergrannual fracture model



Fig. 3 The fundamental idea behind the proposed model in Fig. 2

2.2 Experiments

The uniaxial tension test and the compact tension test of fracture toughness for the AISI 8822 carburized steel have been conducted by Dana Corporation, following the ASTM Standard E8 and E399, respectively. The specimens have been fully carburized up to 0.91 pct carbon through entire thickness, representing the case part of a gear made of the steel. The tensile tests and fracture toughness tests were performed in normal laboratory environment at ambient temperature with no humidity control. Fracture toughness tests were conducted with the ASTM

Standard Compact Tension specimens (CT) of thickness B: 5.08 mm, length W: 50.8 mm, and height: 60.96 mm. The microscopic observation indicates that the tempered martensite is the dominant constituent (>80%). The grain size is in the range of the ASTM E-112 size 9 with the average diameter of 16 microns. The measured mechanical properties and the chemical composition are listed in Tables I and II.

С	Mn	Ni	Cr	Мо	Cu	S	Р	Si	Fe
0.91	1.01	0.51	0.57	0.30	0.15	0.0024	0.009	0.35	balance

Table I: Chemical Composition (wt %)

Table II: Mechanical Properties

Young's module (GPa)	Yielding Strength (MPa)	Ultimate Strength (MPa)	Engineering fracture strain	Section reduction (%)			
			(%)				
220	978	1369	13	2			



Fig. 4 An alternative expression of the multi-scale, multi-physic model of the intergranular fracture introduced by Fig. 2

2.3 A Bottom-Up Analysis Procedure

Fig. 4 details the procedure introduced by the model of Fig. 2. Starting at its right upper corner, a finite element model of the ASTM Standard CT specimen is plotted which is used to measure the conventional macroscale fracture parameters, such as CTOD (crack tip opening displacement), CMOD(crack mouth opening displacement), Stress Intensity Factor and J-integral. These parameters correlated to each other, defining a fracture toughness of the steel [39]. For

carburized steels, the evolution of the crack tip damage zone is mainly dominated by the decohesion of grain boundary, as illustrated by Fig. 2. A numerical procedure, which is termed "Moving Particle Finite Element" (MPFEM) [25, 27, 30], is employed to integrate the deformation of grains and separation of grain boundaries through the computation over the damage zone, as illustrated in the box left from the CT specimen. An intergranular decohesion can be mathematically described by the traction-separation relations between grain boundaries, which is similar to adhesion in bulk phase but with deducted adhesion energy due to localized concentration of heterogeneities and tilted angle between two grains. This traction-separation relation can be obtained through the sub-atomic quantum mechanical computations applying, e.g. the primitive cell plotted at the left lower corner of the figure 4.

The flow chart in Fig. 5 outlines the procedure; the details for each step will be explained in the following sections.



Fig. 5 An outline of the bottom-up analysis procedure for the intergranular fracture model introduced by Figs 2 and 4; where α , α' , γ , and

 β refer to ferrite, martensite, austenite, and bainite phase, respectively.

2.4 Sub-Atomic Computation

2.4.1 Interfacial adhesion and interatomic potential

Considering fracture as a motion to split an atomic array, the interfacial adhesion between the two separated surfaces governs this process. On other hand, it is a process to break bonded atoms pairs from adjacent grains. Obviously, the interfacial adhesion and interatomic potential are not identical. The relationship between them is crucial for establishing a hierarchical linkage in this multi-scale analysis, which has been derived as follows: Fig. 6 shows two chunks of atoms arrays, Ω_A and Ω_B , apart from each other with a distance λ_N . $T(\lambda_N)$, the attractive (or repulsive) force per unit area between the two paralleled surfaces S_A and S_B , can be expressed as the derivative of an interfacial cohesive potential $E^{coh}(\lambda_N)$ [26]:

$$T(\lambda_N) = \frac{\partial E^{coh}(\lambda_N)}{\partial \lambda_N},\tag{1}$$

On other hand, let f(r) be the interatomic force between two individual atoms A and B in Ω_A and Ω_B , respectively, with a distance r; and E(r) be the corresponding interatomic potential:

$$f(r) = \frac{\partial E(r)}{\partial r} \tag{2}$$

Assuming both Ω_A and Ω_B to be semi-infinite, the total force acting upon the single atom *B* from all atoms in Ω_A is:

$$f_B(\lambda) = n_A \int_{\Omega_A} \cos\theta \cdot f(r) d\Omega_A = n_A \int_0^{2\pi} r d\varphi \int_0^{\pi} 2r \cos\theta d\theta \int_{\frac{\lambda_N}{\cos\theta}}^{\infty} \frac{\partial E(r)}{\partial r} dr$$
(3)

where n_A is the number of atoms per unit volume of Ω_A . So the total traction/impulsion between the two bodies, represented by the adhesion $T(\lambda_N)$ between the surface pair S_A and S_B , yields

$$\frac{\partial E^{coh}(\lambda_N)}{\partial \lambda_N} = T(\lambda_N) = n_B \int_{unit \ area} dS \int_{\lambda_N}^{\infty} f_B(\lambda) d\lambda$$
(4)

where n_B is the number of atoms per unit volume in Ω_B and dS is an infinitesimal area element of S_B .



Fig. 7 The difference between interfacial adhesion and interatomic traction/repulsion: the normal traction/separation law between surfaces S_A of Ω_A and S_B of Ω_B is a function of the separation λ_N ; whereas the cohesion between atoms *A* and *B* is determined by the interatomic distance *r* when omitting the effects of spin-polarization.

The equations (3) and (4) establish the analytical relationship between interatomic cohesion and interfacial adhesion. For example, quantum mechanics computation usually gives the interfacial adhesion potential E^{coh} [26, 46]. When E^{coh} is written as a polynomial as following:

$$E^{coh}(\lambda_N) = \frac{a_2}{(\lambda_N)^2} + \frac{a_8}{(\lambda_N)^8} + \frac{a_n}{(\lambda_N)^n} \dots , \qquad n \neq 2,8$$
(5)

where a_i , i = 2,8,..., are constant. By substituting (5) into (4) and (3) one finds that the corresponding interatomic potential yields:

$$E(r) = 4\varepsilon_0 \left[\left(\frac{\sigma}{r}\right)^{12} - \left(\frac{\sigma}{r}\right)^6 \right]$$
(6)

if

$$a_2 = -\frac{n_A n_B \varepsilon_0 \pi^3 \sigma^6}{2}, \quad a_8 = \frac{7 n_A n_B \varepsilon_0 \pi^3 \sigma^{12}}{256} \quad \text{and} \quad a_n = 0$$
 (7)

Eq. (6) is the conventional Leonard-Jones Potential for a biatom system; where σ is the "collision diameter" that equals the separation when *E* is zero; ε_0 is related to the "well depth",

i.e. the minimum of E in the energy-separation (E, r) curve, representing the equilibrium position:

$$f(r) = \frac{\partial E(r)}{\partial r}\Big|_{r=r_0} = 0$$
 and $r_0 = \sqrt[6]{2}\sigma$

It should be noticed that the bi-atomic potential (6) is under the approximation to omit the effects of hyperfine structure of atoms.

Similarly to (1), the stress against sliding between two atomic surfaces, denoted as σ_T , is determined by the derivative of Peierls-Nabarro energy potential again stacking fault E_s [15, 20, 54]:

$$\sigma_T = \frac{\partial E_s}{\partial \lambda_T}$$
 and $E_s = \gamma_{US} \sin^4 \left(\frac{\pi \lambda_T}{b}\right)$ (8)

where λ_T is the relative sliding; γ_{US} is the Peierls-Nabarro energy barrier against dislocation.

2.4.2 Geometrical discontinuities and grain boundary

In Fig. 6 the two adjacent surfaces S_A and S_B are parallel to each other but no discussion about the details, for example, lattice's orientations of Ω_A and Ω_B . Grain boundary can be considered as accumulated discontinuities in a periodic atomic array, as categorized in Fig. 7. The discontinuities, in conjunction with impurities segregations, may reduce the grain boundary adhesion energies profoundly. To explain this concept, only the tilted grain boundary in polycrystalline bcc iron system, i.e. the case (a) of Fig. 7, will be discussed in this paper, focusing on the effects of phosphorous segregation on fracture toughness. Obviously, the grain boundary adhesion energy, denoted as E_{gb}^{coh} , is different from the E^{coh} , the adhesion energy in bulk phase:

$$E_{gb}^{coh} = E^{coh} - \Delta E^{tilt} - \Delta E^{segragate}$$
⁽⁹⁾

where ΔE^{tilt} and $\Delta E^{segragate}$ are the deductions due to tilt and impurities segregation, respectively. The similar expression also applies to the γ_{US} in (8), denoted as γ_{gb}^{US} .



Fig. 7 Geometrical heterogeneities caused by grain boundary

2.4.3 Impurities segregation

When chemical composition of steel is known, an issue is to find how many the hazard elements, such as phosphorous and sulfur, segregated around grain boundaries. Let the symbol Γ_P^G to represent the average weight percentage of phosphorous (P) segregated at grain boundaries, the Langmuir-McLean model [17] gives an estimate:

$$\Delta \Gamma_{p}^{G} = \frac{l}{l + x \exp\left(\frac{\Delta G^{0}}{RT}\right)}$$
(10)

where $\Delta \Gamma_p^G$ is the increment of Γ_p^G during a heat treatment at a thermodynamic equilibrium state with the temperature *T*, ΔG^0 is the chemical potential of the corresponding bulk phase which is about -78 kJ/mol⁻¹ (at 300K) [6], *R* is the universal gas constant and *x* is to be calibrated by test.

The equation (10) indicates phosphorous segregation can be controlled through adjusting heat treatment temperature, which also provides a way to estimate Γ_p^G according to heat treatment history. The experimental results of DANA Corp. have been used to calibrate the constant *x*. The total Γ_p^G for the steel analyzed is below 10%.

2.4.4 DFT [2, 3] Computation

In order to obtain E_{gb}^{coh} and γ_{gb}^{US} for establishing grain boundary traction-separation law, the Density Function Theory[2,3] based quantum mechanics computations have been conducted, applying the full potential all electrons linear augmented plane wave (FLAPW) numerical codes [4, 44-46,71]. The procedure to set up periodic atomic supercells for this class of computations has been introduced in [70].

The atomic cells with two tilted angles θ : $\Sigma 1 (\theta = 0^{\circ})$ and $\Sigma 5 (\theta = 51.13^{\circ})$ have been computed at ground state. The $\Sigma 1$ grain boundary is the case that two adjacent grains have the same lattice's orientation but with phosphorous segregation in-between, which is a degenerated case of low-angle grain boundaries. The primitive cell of $\Sigma 5$ grain boundary is given by the plot on the left hand side of Fig. 8, which is a typical high angle grain boundary. The corresponding distributions of electron charge density for the boundary cell and $\Sigma 5$ surface, respectively, are plotted on the right hand side of the figure. Applying the procedure introduced in [26, 32, 20], the differences between the cases with and without phosphorous segregation define the $\Delta E^{segragate}$, whereas the difference between non-segregation $\Sigma 1$ and $\Sigma 5$ cells defines the ΔE^{tilt} in (9). Grain boundaries with other tilted angles (less than $\theta = 51.13^{\circ}$) are interpolated between zero and the ΔE^{tilt} of $\Sigma 5$. The twin boundary is not taken into account for the intergranular fracture studied.

The computed adhesion energies for bulk bcc iron and for the $\Sigma 1$ grain boundary with phosphorous segregation are plotted in Fig. 9a, by which the discretized numerical data were

fitted into polynomials (5-7). As expected, a remarkable drop in adhesion energy can be seen when *P* segregate presents. For the Σ 5 boundary ΔE^{tilt} , the reduction of adhesion energy, is about 18% when $\Gamma_P^G = 0$. Considering grain boundary represents a "discontinuity" of periodic array of atoms, the reasons that cause the drop of adhesion energy can be: (i) the segregated interstitial phosphorous atoms weaken metallic Fe-Fe bonds; (ii) the tilted boundary is actually an array of empty sites that enlarges the interatomic distance between the atoms from adjacent grains; (iii) segregated *P* atoms or titled boundary breaks the periodicity in bulk phase, which destroys the ferromagnetic alignment of bcc iron and, thus, induces localized antiferromagnetic-like spin-polarization (see Fig. 9b); it is well-known that the iron fcc crystal is antiferromagnetic, which is with less stability as compared with bcc iron at ground state, see Appendix I. Table III lists the computed values of E^{coh} , ΔE^{tilt} and $\Delta E^{segragate}$. Results of other segregations or grain boundary tilt angles can be found, e.g. in [6, 10, 11, 31, 32]. According to the computation in [20]: $\gamma_{US} \approx 0.43$ (J/M²). At grain boundary the relation $\gamma_{US}^{gb} = \gamma_{US} \left(\frac{E_{gb}^{coh}}{E_{gb}} \right)$ is applied in this analysis.



Fig. 8 The supercell of $\Sigma 5$ bcc iron grain boundary with phosphorous segregate (a) and charge density in the middle layer of the cell (b, c); where the red ball in (a) and the small dark-gray ball in (b) and (c) are Phosphorous atoms whereas the others are iron atoms; a bcc cell is illustrated in the supercell of (a). The misorientation defined by Fig. 7 is zero in these computations.



(a) grain boundary adhesion energies (b) spin-charge density around $\Sigma 5$ grain boundary without phosphorous segregation; it demonstrates a semi-periodic distribution in vertical direction which is somewhat like the anti-ferromagnetic case in fcc; from the horizontal periodicity one sees that the computation is performed using the cell of Fig. 8(a).

Fig. 9. Computation of the grain boundary adhesion with phosphorous segregation

E^{coh}	$\Delta E^{tilt} \left(\sum 1 \right)$	$\Delta E^{tilt} \left(\sum 5\right)$	$\Delta E^{segragate} (\sum 1)$	$\Delta E^{segragate}$ ($\sum 1$)	
	$\Gamma_P^G = 0$	$\Gamma_P^G = 0$	$\Gamma_P^G = 4.6\%$	$\Gamma_P^G = 9.4\%$	
4.97	0	1.67	0.94	1.61	

Table III Grain boundary adhesion energy for BCC iron (J/M^2)

where *P*: phosphorous; Γ_P^G : weight percentage of *P* segregation

2.5 A Hierarchical Multi-Scale Procedure

A major challenge for the object studied is to bridge the sub-atomic computation with micro and macro analysis, which requires to hierarchically integrate the kinetics from angstroms to centimeters while highlighting the dominant mechanisms. This is somewhat different from many recently developed successful methods, for examples, [57-63]. In order to predict the steel's fracture toughness, the following two steps are vital important: (I) implement the results of quantum mechanical computation obtained into the polycrystalline system; (II) to embed the information of grain-sized analysis into the inch-sized fracture toughness specimen.

2.5.1 Sub-Atomic to Up-atomic

The step (I) requires bridging the sub-atomic quantum physics with up-atomic continuum analysis. As plotted in the flow chart of Fig. 5, the Moving Particle Finite Element Method (MPFEM) is employed for this purpose. It contains the methodologies in two perspectives: finite element and particle method. For sub-atomic to up-atomic bridging, the 'Particle Dynamics', introduced in [20, 28], is applied. The idea of this approach is to represent an atomic system as a particle system through lumping several atoms into a super-atom, termed "particle", while

preserving the essential properties of the atomic system via a proposed "equivalent stiffness rule". This rule requires that the particle system has the same periodic structure and stiffness as the atomic system but with less number of particles and a larger inter-particle spacing that is determined according to the scale of interest, see Figs. 10a,b. The sub-atomic physics, which may dominate the mechanical behavior at up-atomic scales, is preserved through transforming the inter-atomic potential into an inter-particle potential by the following way:

Assuming E^{atom} to be the interatomic potential, like (6), for the system in Fig. 10a; when it endures a deformation, for example, a_0 becomes a_1 ; accordingly $R \to R_1 = Na_1$ in the particle system of Fig. 10b. This deformation can be represented by a continuous strain field \mathcal{E}_{ij} for both systems. The corresponding stress tensor σ_{ij}^{atom} and stiffness tensor C_{ijkl}^{atom} of the atomic system yield [26, 56]:

$$\sigma_{ij}^{atom} = \rho \left(\frac{\partial E^{atom}}{\partial \varepsilon_{ij}} - 2\varepsilon_{ik} \frac{\partial E^{atom}}{\partial \varepsilon_{kj}} \right)$$
(11a)

and

$$C_{ijkl}^{atom} = \frac{\partial \sigma_{ij}^{atom}}{\partial \mathcal{E}_{kl}}$$
(11b)

where ρ is density and the dummy summation rule is applied. The second term on the right hand side of (11a) is corresponding to finite strain [56], which can be omitted when deformation is small.

Similarly, for the particle system in Fig. 10b with an inter-particle potential $E_{Particle}$:

$$\sigma_{ij}^{Particle} = \rho \left(\frac{\partial E_{Particle}}{\partial \varepsilon_{ij}} - 2\varepsilon_{ik} \frac{\partial E_{Particle}}{\partial \varepsilon_{kj}} \right)$$
(12a)

and

$$C_{ijkl}^{Particle} = \frac{\partial \sigma_{ij}^{Particle}}{\partial \varepsilon_{kl}}$$
(12b)

The "equivalent stiffness rule" requires:

$$C_{ijkl}^{Particle} = C_{ijkl}^{atom}$$
(13)

which leads to

$$\boldsymbol{\sigma}_{ij}^{Particle} = \boldsymbol{\sigma}_{ij}^{atom} \tag{14}$$

The equality (13) provides a group of conditions to determine $E_{Particle}$ that gives the same stresses for both systems, as confirmed by (14). However, the computational effort by Particle Dynamics is reduced to about $1/N^m$ of that by atomic system for a *m*-dimensional case; where *N* defines the size of the particle, see Fig. 10b, and *m*=1,2,3. An analysis of the bcc iron lattice with grain boundary is given in [20, 28].



(a) atomis system

(b) particle system

Fig. 10 The "Particle Dynamics" method [20] that transfers the atomic system (a) into the particle system (b) through segregating the atoms within the cell of dimension R into a particle whereas an "equal stiffness rule" is imposed to ensure the two systems having the same elastic properties.

According to the procedure introduced above, one can find that, as compared with other successful multiscale methods, e.g. the Quasicontinuum [60, 61], the novelty of the Particle Dynamics lies in the following two perspectives:

- The Particle Dynamics (PD) is a "hierarchically-structured" method to transfer an atomic system into a particle network that keeps the same lattice structure as the original; by contrast, in the methods of [58, 60, 61] a finite element contains a set of atoms and the finite element net work may have kinematically favorited slip system that differs from the particle network.

- The "equivalent stiffness rule" of Particle Dynamics provides a unique way to reproduce an inter-atomic interactions through an inter-particle interaction with the same "constitutive" representation.

Applying this approach, the quantum mechanical computation-based interfacial adhesions are applied to describe the grain boundary traction-separation relation for the polycrystalline system that plotted in the second column from right in Fig. 4, where each grain is a crystal made of "particles" which are segregations of atoms. The traction-separation relation has

been implemented into a group of "cohesive elements" that connect each grain boundary surfaces pair, e.g. the *AB* in Fig. 11a. On other hand, all grains are treated as bcc crystals with the same [001] direction perpendicular to the two-dimensional plane but with randomly pre-assigned inplane orientations. Each particle in a grain defines a "node", the connections among these nodes form small finite elements that partition the grain. The finite element is with the average size about one order smaller than average grain size, as illustrated in the middle box of the second column from right in Fig. 4. The finite elements obey the constitutive relation of crystal plasticity with Taylor's hardening (isotropic hardening) [18]; which is governed by the effective stress-strain relation that is calibrated according to the quantum mechanics analysis of bulk iron phase through the "Particle Dynamics" method, as described in [28]. The Moving Particle Finite Element method [25, 27, 30], which integrates all atomic segregates into a polycrystalline network that combines the "cohesive elements" and grain finite elements, has been applied for the micro-scale computation.



Fig. 11 MPFEM cohesive elements for the computations of grain boundary separation and sliding; (a) the cohesive element *AB* connects two adjacent grains through the nodes *A* and *B*; where S_A is a "master surface" enhanced to the node *A* in numerical contact analysis, defining the normal separation λ_N and relative sliding λ_S , respectively; thus, the grain boundary surface associated with the node *B* is defined as "slaver surface"; (b) at triple junction the three cohesive elements, i.e. *AB*, *AA'*, *A'B*, and corresponding master-slaver surfaces pair, are required in analysis. Moving Finite Element Method [25, 27, 30] is applied to integrate cohesive elements and grains.

2.5.2 Micro to Macro Scale

As mentioned in the previous subsection, the size of finite element inside a grain is in the order of micron; for a grain by grain micro-structured two-dimensional computation of an ASTM standard CT specimen, a model with about 6.25×10^{11} finite elements (10^{17} elements for 3D) is required. Obviously, more effective scheme will be preferred since the damage caused by intergranular cracking is limited within the close vicinity of the crack tip; the rest part of the specimen can be described by well-developed theory of isotropic and uniform plasticity; for the latter, finite element with the size of millimeter is sufficient.

In this analysis a scheme is developed by which the CT specimen is divided into two parts: a small chunk of material surround the crack tip which is modeled by the polycrystalline microstructure; an example of this crack tip zone, denoted as Ω_t , is plotted on the right upper corner of Fig. 12. The rest part of the specimen, denoted as Ω , is modeled by the continuum J_2 flow plasticity theory using regular finite element. The external load applied to the specimen is transferred into the crack tip through the continuity conditions of traction T and displacement \overline{u}_t on the shared boundary Σ . The detail of the three-dimensional meshes around Σ in Ω is given by Fig. 12, which shows the adoption from the relatively small finite elements near Σ to the coarse elements far away. T and \overline{u}_t at Σ on the middle section of the 3D block define the plane strain boundary condition on the Σ of Ω_t . This is similar to the "boundary layer method", e.g., in [37], by which the stresses of asymptotic analytical solution, such as mode-I K-field [39], is applied to the outer boundary of the crack tip zone.



Fig. 12 The developed scheme divides a standard specimen into two parts: the polycrystalline slab Ω_t containing, for example, a sharp crack tip, as plotted on the right upper corner; which "shakes" hands to the rest of the specimen through the continuities of displacement and J-integral on the boundary Σ

However, when massive microscale intergranular cracks occur within Ω_t , the stresses on Σ may decrease due to material's softening while \overline{u}_t keeps climbing up. In other word, the traction T alone does not uniquely determine the deformation field inside Ω_t . By contrast, the \overline{u}_t on Σ can be monotonic when an external load in the form of displacement increases monotonically. Generally it is very difficult to satisfy the both stress and displacement continuities simultaneously by boundary layer method. Hence, instead, a *J*-integral[13] based "shake hand" scheme is developed which requires the satisfaction of the continuous conditions of displacement and *J*-integral on Σ ; the latter represents the energy that flows into the crack tip.

This "shake hand" scheme includes three computations: beside the computation of the part Ω and that of the polycrystalline system Ω_t with intergranular cracking, respectively; the third computation of the entire specimen $(\Omega_t + \Omega)$ is conducted by which no intergranular cracking in Ω_t , so all material are obeying the simple J₂ plasticity law. The quantities associated with these three computations are distinguished in turn by the superscriptions Ω , Ω_t , and $\Omega_t + \Omega$. Then, for the two cases without intergranular cracking, the corresponding *J*-integral, applied load *P*, and CMOD (crack mouth opening displacement) *U* are J^{Ω} , P^{Ω} , U^{Ω} and $J^{\Omega_t+\Omega}$, $P^{\Omega_t+\Omega}$, $U^{\Omega_t+\Omega}$, respectively. The reason for the computation ($\Omega_t + \Omega$) is to find the external load ($P_{damage}^{\Omega_t}$) and CMOD ($U_{damage}^{\Omega_t}$) corresponding to the computation of Ω_t with intergranular cracking induced damage.

The displacement on Σ obtained from the third computation, denoted as $\overline{u}_{t}^{\Omega_{t}+\Omega}$, is used as the boundary condition imposed on Σ for the computation of the polycrystalline system Ω_{t} with intergranular cracking, which leads to the corresponding *J*-integrals $J^{\Omega_{t}}$ on Σ . On other hand, \overline{u}_{CTOD} , a reference crack tip opening displacements (CTOD), is recorded for all three computations; which is defined as an integral over Σ for the vertical component of \overline{u}_{t} with respect to dx and then divided by the length of the projection of Σ on the horizontal coordinate x_{t} . For a given \overline{u}_{CTOD} : $P^{\Omega_{t}+\Omega} > P^{\Omega}$, $U^{\Omega_{t}+\Omega} \cong U^{\Omega}$, and $J^{\Omega_{t}+\Omega} > J^{\Omega}$. So $P_{damage}^{\Omega_{t}}$, the corresponding external load for the computation of Ω_{t} with intergranular cracking, is obtained through the following interpolation:

$$P_{damage}^{\Omega_{t}} = P^{\Omega} + \left(P^{\Omega_{t}+\Omega} - P^{\Omega}\right) \frac{J^{\Omega_{t}} - J^{\Omega}}{J^{\Omega_{t}+\Omega} - J^{\Omega}}$$
(15)

which is the load actually applied to the specimen. This procedure is under the approximation that the displacement field in Ω changes monotonically when the CT specimen is under a monotonic load-line displacement. This is a reasonable approximation for small-scale yielding.

3. RESULTS AND DISSCUSION

Fig. 13 is a set of snap-shots of the evolutions of stress field in the damage zone Ω_t surrounding a blunted crack tip of the CT specimen, when applied load increases. Fig. 14 gives the comparison between the simulated load-CMOD relation and experimental results. Fig. 15 shows the computed fracture toughness according to ASTM E399 when the content of phosphorous varies. As expected, higher content of phosphorous leads to lower fracture toughness.



Fig. 13 Intergranular fracture – MPFEM microscale simulation for the case with blunted crack tip



Fig. 14 The computed Load-CMOD curves for the specimens with different weight percentages of phosphorous



Fig. 15 The relationship between fracture toughness and the content of phosphorous, computational results, by improving grain ductility, the curve may move up, as indicated by the arrow A.

It is well-known that phosphorous segregates will embrittle metals [6]. However, by comparing the drop in fracture toughness plotted in Fig. 15 and the deduction of adhesion energy plotted in Fig. 9a when phosphorous segregates at grain boundary, the changes in the former is much more significant than that in the latter. This can be explained by the simulations in Fig. 13, which shows a severe stress concentration at triple-junctions and the junctions of multi-grain boundaries when the average amplitude of the stress field is moderate. Thus, each junction becomes a source of microscale-cracking, which causes decohesion along grain boundaries. The accumulation of these microscale-cracks forms a damage zone ahead a macro-scaled crack tip. The evolution of this damage zone leads to the subsequent macroscale crack growth. Considering a steel be a system, an intergranular fracture essentially is the result of the interaction between grain boundary separation and deformation of grains. Therefore, removing the stress concentration at junctions of grain boundaries and increasing grain boundary adhesion are the two equally important goals for improving the mechanical properties of the steel. The study conducted inspires the ideas to achieve the first goal by increasing grain ductility through alloy additions, e.g. Ni, and phase constituents, e.g. with optimized ratio of bainite and retard austenite; while to improve grain boundary properties through the following three ways: (i) adding grain boundary clue elements such as B and Nb; which also have the function to pin grain boundary so reduce grain's size; (ii) alloying with appropriated processing to promote the formations of $(M, P)_x C_y$ and $(M, S)_x C_v$ compounds which extract P and S out from grain boundaries; (iii) adjusting heat treatment process to reduce P grain boundary concentration, as described (10).

4. Conclusions and Suggestions

From the respect of mechanical analysis, this paper reports an effect to understand the intergranular fracture phenomenon in polycrystalline system based on the underlying fundamental physics. For this purpose a multi-scale procedure has been developed, which hierarchicaly integrate the density function theory computations of $\Sigma 5$ and $\Sigma 1$ boundary within Phosphorous segregation into a molecule-like "particle dynamics" analysis of polycrystalline system; the

obtained model is built into a corresponding macro-scale particle-finite element analysis of laboratory-size fracture toughness specimen. A theoretical relation between the interatomic potential in a bi-atomic system and the interfacial adhesion in an atomic slab has been obtained; the difference between them is often being omitted in many analyses. The developed "particle dynamics" is able to efficiently reproduce the quantum physics-based lattice's mechanical behavior by a particle system of grains and grains boundaries at the scale of microns.

The developed procedure has been applied to the gear's case made of AISI 8822 carburized steel, which concludes that for the polycrystalline system consisting of the grains with high strength and hardness, an intergranular fracture is governed by the interaction of grain boundary adhesion and stress concentration at triple and other multiple grain boundaries' junctions. Hence, optimized mechanical properties for this class of steels can be achieved through the following conventional ways:

- Refine grain size: although finer grains may be detrimental to toughness, however, when the average size is not less than 10^2 nm, smaller grains may reduce the local resistance against dislocations motion around junctions of grain boundaries while increase the energy barriers against large scale yielding due to increased zigzags in slipping paths; the former may smear out the stress concentration at the junctions and the latter elevates the average strength according to conventional Hall-Petch relation. Also, finer grains enlarge the total area of grain boundaries which reduces average impurities segregation per unit area of grain boundary. Hence, the trade-off of these factors may finally lead to a positive effect to the system.

- Alloying and processing to simultaneously improve grain boundary adhesion and grain's ductility: high grain's ductility reduces the stress concentration at triple-junctions; a superposition of these two mechanisms may result in the transition from intergranular fracture to trangranular fracture.

- Adjusting heat treatment to reduce impurities grain boundary segregation, improve phase constituent distribution and grain size.

Appendix

In the following text boldface symbol denote tensor, the order of which is indicated by the context. Plain symbols denote scalars or a component of a tensor when a subscript is attached. Repeated indices are summed. For two order tensors \boldsymbol{a} and \boldsymbol{b} , $\mathbf{a} = [a_{ij}]$, $\mathbf{b} = [b_{ij}]$; then

 $\boldsymbol{a}^{T} = [a_{ji}], \ \mathbf{a} \cdot \mathbf{b} = [a_{ik}b_{kj}], \ \mathbf{a} \cdot \mathbf{b} = [a_{ij}b_{ij}], \ \mathrm{and} \ \mathbf{ab} = [a_{ij}b_{kl}].$

I Feromagnetic and Antiferromagnetic phases of Iron [65]



Fig. A1: (a) Comparison of the ground state ferromagnetic and antiferromagnetic system energies of unit atomic cell when it transforms from bcc to fcc crystal along "Bain path"; where η is the order parameter to characterize the lattice constant ($\eta = 0$: fcc, $\eta = 1$: bcc); so the fcc crystal has antiferromagnetic structure with higher ground state energy than bcc. (b) The magnetization moment per atom when the lattice structure varies along Bain path.



Fcc (100) density of spin-polarization bcc (110) density of spin-polarization Fig. A2: Spin-charge density distributions for bulk fcc (left) and bcc (right).

II: About the "Moving Particle Finite Element Method" (MPFEM) [25, 27, 30]

The network that connects all particles obtained from "Particle Dynamics" [20] forms a finite element mesh, which is the same as the mesh that represents a discretized body in solid

mechanics finite element analysis [50-53]. Fig. A3(a) shows an example of such a solid body Ω with the boundary segment $\partial \Omega_t$ under applied force (natural boundary condition) and the segment $\partial \Omega_u$ with given displacement (essential boundary condition). Ω is partitioned into 9 triangle elements in (b). Each corner of a triangle element is a particle that is termed "node" in finite element analysis. In the *i*th element, the strain incremental tensor inside a finite element, denoted as $\Delta \varepsilon$, is expressed as the function of the displacement increments vector Δu_i at all nodes associated with this element:

$$\Delta \boldsymbol{\varepsilon} = \boldsymbol{B}_i \cdot \Delta \boldsymbol{u}_i \tag{b1}$$

where the rule of dummy summation for the repeated index *i* applies; B_i is the "differential matrix", determined by displacement-strain relation (e.g. Cauchy geometric relation) and the interpolation scheme (shape function) of the finite element.



Fig. A3: The conception of MPFEM applied to grain boundary with cohesive element xx';

Assuming that the elements E_1, E_2, E_3 belong to one grain while E_4, E_5, E_6 belong to another grain, then the node between two grains actually is a "cohesive element" that connects the particles from two adjacent grains' surfaces; for example, the x and x' in Fig. A3(c). The "Moving Particle Finite Element Method" [25, 27, 30] is able to solve the equilibrium condition at nodes through the virtual work principle.

Considering the node x in the case of Fig. A3(b), the virtual work principle-based equilibrium condition at this node implies that the work done by a force ΔF_x imposed on the nodal x, which is the product of ΔF_x and a virtual nodal displacement Δu_x at this node, equals the summation of the strain energies caused by Δu_x in all elements shared the node x:

$$\Delta F_{x} \cdot \Delta u_{x} = \sum_{i=1}^{6} (\Delta \sigma : \Delta \varepsilon) V_{i} = \sum_{i=1}^{6} (\Delta \sigma : B_{i} \cdot \varDelta \overline{u}_{i}) V_{i}$$
(b2)

where V_i is the volume of the i^{th} element; $\Delta \overline{u}_i = [\Delta u_x, \theta]^T$ is a vector that contains the displacement increments at all nodes associated with the elements adjacent to the node x; but by the virtual work principle it is assumed all these nodal displacements are zero except Δu_x ; $\Delta \sigma$ is the stress increment tensor that is correlated to $\Delta \varepsilon$ through the constitutive law:

$$\Delta \boldsymbol{\sigma} = \boldsymbol{C}_t : \Delta \boldsymbol{\varepsilon} \tag{b3}$$

where C_t is the tangential stiffness matrix of the bulk phase, which is a fourth order tensor.

In MPFEM there is no restriction to the type of the elements in the nodal equilibrium condition (b2). For the case in Fig. A3(c) the node x is at a grain boundary formed by the upper edges of the triangle elements E_1 and E_3 . The developed cohesive element xx' establishes the connection between the two sides of the opposite grains. Similar to (b2), the equilibrium condition at the node x can be expressed as the following summation.

$$\Delta \boldsymbol{F}_{x} \cdot \Delta \boldsymbol{u}_{x} = \sum_{i=1}^{3} \left(\Delta \boldsymbol{\sigma} : \boldsymbol{B}_{i} \cdot \boldsymbol{\Delta} \overline{\boldsymbol{u}}_{i} \right) \boldsymbol{V} + \Delta \boldsymbol{F}_{x}^{\ decoh} \cdot \Delta \boldsymbol{u}_{x}^{\ decoh}$$
(b4)

where ΔF_x^{decoh} and Δu_x^{decoh} are the force and elongation vector of the cohesive element; when there is no displacement on the other end of the cohesive element:

$$\Delta \boldsymbol{u}_{x}^{decoh} = \Delta \boldsymbol{u}_{x} = \left[\boldsymbol{\lambda}_{N}, \boldsymbol{\lambda}_{S}\right]^{T}.$$
(b5)

where λ_N and λ_S are the normal and tangential separation in (1) and (8), respectively.

When all other nodes are fix but node x has the displacement Δu_x , one can remove Δu_x from the both sides of (b4), which becomes the equilibrium condition at this node in matrix form:

$$\Delta \boldsymbol{F}_{x} = \boldsymbol{K} \cdot \Delta \boldsymbol{u}_{x} \tag{b6}$$

where $\tilde{K} = K_t + K_{decoh}$; the grain matrix tangential stiffness K_t and decohesion stiffness K_{decoh} are defined as below:

$$\boldsymbol{K}_{t} = \sum_{i=1}^{3} \left(\boldsymbol{B}_{i}^{T} : \boldsymbol{C}_{t} : \boldsymbol{B}_{i} \right) \cdot \boldsymbol{V}$$
(b7)

and

$$\boldsymbol{K}_{decoh} = \frac{\partial \left(E^{coh}, E_{S} \right)}{\partial \left(\lambda_{N}, \lambda_{S} \right)}$$
(b8)

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